# **Financial Econometrics – 2023 Lecture Notes**

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# Lecture 1 – Review of Statistics and Linear Algebra (NOT Covered)

This lecture reviews basic probability concepts, from random variables to the Law of Large Numbers and the Central Limit Theory. In the Appendix, the lecture introduces Linear Algebra and its compact notation.

## **Random Variable**

In probability and statistics, a *random variable* (RV), or *stochastic variable*, is described informally as a variable whose values depend on outcomes of an *experiment* (or phenomenon). An experiment is an act or a process with an unknown outcome. For example, the CEO of Apple announces a new product, the effect on the price of Microsoft is unknown, thus, the price (and return) of Microsoft is a RV.

### **Examples:**

1. We throw two coins and count the number of heads.

2. We define X = 1 if the economy grows two consecutive quarters and X = 0, otherwise. (This is an example of a *Bernouille* (or *indicator*) RV.)

- 3. We read comments from IBM's CEO and compute IBM's daily return.
- 4. We count the days in a week that XOM has a positive return.

5. We look at a CEO and write his/her highest education degree.

6. We compute the weekly sign of stock returns of two unrelated firms: Positive (U: up) or

negative (D: down). We count the times at least one stock is up:  $\{D,U\}, \{U,D\}, \{U,U\}, \P$ 

For some RVs, it is easy to enumerate all possible outcomes. For instance, for the fourth (XOM) example above:  $\{0, 1, 2, 3, 4, 5\}$ . But, for some RV, it can be complicated. For example, for the third (IBM) example:  $\{-100\%, K\}$ , where K is a large positive number.

The set of all possible outcomes is called *sample space*, denoted by  $\Omega$ .

An event A is a set containing outcomes from the sample space. For example, for the IBM example, the returns are between 2% and 12.5% is an event.

The collection of all possible events is  $\Sigma$ . For example, for the IBM example, {(1.1%-1.2%), (-0.02%, -0001%), (2.0%, 12.5%), (-5%, 5%), (-100%, -13.95%), (0%, 350%), ... }

• In general, a RV is a *function* whose domain is the sample space,  $\Omega$ . It produces numbers. For instance, in example 6 above, instead of using {U, U} when both stocks go up, we use 2.

Mathematically, **X**:  $\Omega \rightarrow R$ .

<u>Remark</u>: The name "random variable" is confusing; it is just a function!

• We put some mathematical structure (pdf, pmf, CDF) to the concept of RV to describe what is more/less likely to happen to the (randomly determined) events.

For example, we would like to know which event is more/less likely for the IBM example: Is (1.1%-1.2%) more likely than (-0.02%, -0.001%)?

Definitions & Notation:

 $\Omega$ : The sample space –the set of possible outcomes from an experiment.

An event *A* is a set containing outcomes from the sample space.

 $\Sigma$ : The collection of all possible events involving outcomes chosen from  $\Omega$ . (Formally:  $\Sigma$  is a  $\sigma$  - algebra of subsets of the sample space.)

P is a probability measure over  $\Sigma$ . P assigns a number between [0,1] to each event in  $\Sigma$ .

#### Remarks:

- A random variable is a convenient way to express the elements of  $\Omega$  as numbers rather than abstract elements of sets.

- A random variable *X* is a function.

- It is a numerical quantity whose value is determined by a random experiment.

- It takes single elements in outcome set  $\Omega$ , which can be abstract elements, and maps them to points in R.

**Example:** We compute the weekly sign of stock returns of two unrelated firms: Positive (U: up) or negative (D: down).

The sample space is  $\Omega = \{DD; DU; UD; UU\}.$ 

Possible events (A):

- Both firms have the same signed return:  $\{U,U\}$  &  $\{D,D\}$ .

- At least one firm has positive returns:  $\{U,U\}$ ;  $\{D,U\}$  &  $\{U,D\}$ .

- The first firm has positive returns: {U,U} & {U,D}

Collection of all possible events:  $\Sigma = [\Phi, \{U,U\}, \{U,D\}, \{D,U\}, \{D,D\}, \{UU, UD\}, \{UU, DU\}, \{UU, DD\}, \{DD, DU\}, \{DD, UD\}, \{DU, DD\}, \{UU, DU, UD\}, \{UD, DU, DD\}, \{UU, UD, DU, DD\}]$ 

Define RV:  $\mathbf{X} =$  "*Number of Up cycles*." Recall,  $\mathbf{X}$  takes  $\Omega$  into  $\chi$ , & induces P<sub>X</sub> from P. Then,

$$\begin{split} \chi &= \{0; 1; 2\} \\ \Sigma_{\chi} &= \{\Phi; \{0\}; \{1\}; \{2\}; \{0;1\}; \{0;2\}; \{1;2\}; \{0;1;2\}\}. \end{split}$$



Assuming U and D have the same probability,  $P[U] = P[D] = \frac{1}{2}$ , we define  $P_X$ : Prob. of 0 Ups =  $P_X[0] = P[\{DD\}] = \frac{1}{4}$ Prob. of 1 Ups =  $P_X[1] = P[\{UD; DU\}] = \frac{1}{2}$ Prob. of 2 Ups =  $P_X[2] = P[\{UU\}] = \frac{1}{4}$ Prob. of 0 or 1 Ups =  $P_X[\{0; 1\}] = P[\{DD; UD; DU\}] = \frac{3}{4}$ Prob. of 0 or 2 Ups =  $P_X[\{0; 2\}] = P[\{DD; UU\}] = \frac{1}{2}$ Prob. of 1 or 2 Ups =  $P_X[\{1; 2\}] = P[\{DU; UD; DD\}] = \frac{3}{4}$ Prob. of 1, 2, or 3 Ups =  $P_X[\{0; 1; 2\}] = P[\{DD; UU; UD; UU\}] = 1$ Prob. of "nothing" =  $P_X[\Phi] = P[\Phi] = 0$ 

The empty set is simply needed to complete the  $\sigma$ -algebra (a technical point). Its interpretation is not important since P[ $\Phi$ ] = 0 for any reasonable P.

<u>Technical detail</u>: P is the probability measure over the sample space,  $\Omega$ , and P<sub>X</sub> is the probability measure over  $\chi$ , the range of the random variable.

#### **Example:** IBM Returns

We buy an IBM share at USD 120 today and plan to sell the share next week. The return of IBM next week,  $r_t$ , depends on how the market values IBM next week –this is the experiment.

The sample space is continuous, from -100% (worst case scenario) to potentially a huge undefined positive number. We set  $\Omega = \{r_t : r_t \in [-1, K], K > 0\}$ .

Possible events:

- IBM returns are positive.
- IBM returns are higher than 0.5%.
- IBM returns are lower than 10%.
- IBM returns are between -2% and 4%.

The collection of all possible events,  $\Sigma$ , is very, very large. We use a probability distribution, for example, the normal distribution, to describe the likelihood of possible events.

### **Probability Function & CDF**

**Definition** – The probability function, p(x), of a RV, X. For any random variable, *X*, and any real number, *x*, we define  $p(x) = P[X = x] = P[\{X = x\}],$ where  $\{X = x\}$  = the set of all outcomes (event) with X = x.

**Definition** – The *cumulative distribution function* (CDF), F(x), of a RV, X. For any random variable, *X*, and any real number, *x*, we define

 $F(x) = P[X \le x] = P[\{X \le x\}],$ where  $\{X \le x\}$  = the set of all outcomes (event) with  $X \le x$ .

**Example:** Let X be the number of days in a week that XOM has a positive return. Sample space  $S = \{0, 1, 2, 3, 4, 5\}$ . Assuming a binomial distribution with a probability of a daily positive return equal to 0.52, we have the following graph of the pdf:



Binomial Distribution (n=5, p=0.52)

The actual probabilities are given by:

p(x = 0) = 0.0255p(x = 1) = 0.1380p(x = 2) = 0.2990p(x = 3) = 0.3240p(x = 4) = 0.1755p(x = 5) = 0.0380

Note:  $\{X = x\} = \phi$  for all other *x*.

Below, we display the CDF of X:



CDF of Number of Days with positive retu

### PDF for a Continuous RV

**Definition**: Suppose that X is a random variable. Let f(x) denote a function defined for  $-\infty < x < \infty$  $\infty$  with the following properties:

1.  $f(x) \ge 0$ 

2. 
$$\int_{-\infty}^{\infty} f(x) dx =$$

2.  $\int_{-\infty}^{\infty} f(x) dx = 1.$ 3.  $P[a \le X \le b] = \int_{a}^{b} f(x) dx$ 

1.

Then, f(x) is called the *probability density function* (pdf) of X. The random variable X is called continuous.

• PDF



• If X is a continuous random variable with probability density function, f(x), the *cumulative distribution function* of X is given by:

$$F(x) = P[X \le x] = \int_{-\infty}^{x} f(t)dt$$



• Also because of the FTC (fundamental theorem of calculus):



## PDF for a Discrete RV

A random variable X is called *discrete* if

$$\sum_{x} p(x) = \sum_{i=1}^{\infty} p(x_i) = 1$$

All the probability is accounted for by values, *x*, such that p(x) > 0.

• For a discrete random variable X the probability distribution is described by the probability function p(x), which has the following properties:

- 1.  $0 \le p(x) \le 1$
- $2. \qquad \sum_{i=1}^{N} p(x_i) = 1$
- 3.  $P[a \le X \le b] = \sum_{a \le x \le b} p(x)$



#### **Bernouille and Binomial Distributions**

Suppose that we have a *Bernoulli trial* (an experiment) that has 2 results:

- 1. Success (S)
- 2. Failure (F)

Suppose that *p* is the probability of success (S) and q = 1 - p is the probability of failure (F). Then, the probability distribution with probability function:

$$p(x) = P[X = x] = \begin{cases} q & x = 0\\ p & x = 1 \end{cases}$$

is called the Bernoulli distribution.

• We observe an independent Bernoulli trial (**S**, **F**) n times. Let X be the number of successes in the n trials. Then, X has a *binomial distribution*:

$$p(x) = P[X = x] = {n \choose x} p^{x} q^{n-x} \quad x = 0, 1, 2, \dots, n$$

where

- 1. p = the probability of success (S), and
- 2. q = 1 p = the probability of failure (**F**)

**Example:** If a firm announces profits and they are "surprising," the chance of a stock price, P, increase is 85%. Assume there are n=20 (independent) announcements.

Let *X* be the number of increases in the stock price following *surprising announcements* in the n = 20 trials.

$$p(x) = P[X = x] = {n \choose x} p^{x} q^{n-x} \quad x = 0, 1, 2, ..., n$$
$$= {20 \choose x} (.85)^{x} (.15)^{20-x} \quad x = 0, 1, 2, ..., 20$$

x	0	1	2	3	4	5
p(x)	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
x	6	7	8	9	10	11
p(x)	0.0000	0.0000	0.0000	0.0000	0.0002	0.0011
x	12	13	14	15	16	17
p(x)	0.0046	0.0160	0.0454	0.1028	0.1821	0.2428
x	18	19	20			
p(x)	0.2293	0.1368	0.0388			



### **The Poisson Distribution**

Suppose events are occurring randomly and uniformly in time.

• The events occur with a known average.

• Let *X* be the number of events occurring (arrivals) in a fixed period of time (time-interval of given length).

• Typical example: X = Number of crime cases coming before a criminal court per year (original Poisson's application in 1838.)

• Then, *X* will have a *Poisson distribution* with parameter  $\lambda$ :

$$p(x) = \frac{\lambda^{x}}{x!} e^{-\lambda}$$
  $x = 0, 1, 2, 3, 4, ...$ 

• The parameter  $\lambda$  represents the expected number of occurrences in a fixed period of time. The parameter  $\lambda$  is a positive real number.

**Example**: On average, a trade occurs every 15 seconds. Suppose trades are independent. We are interested in the probability of observing 10 trades in a minute (X=10). A Poisson distribution can be used with  $\lambda = 4$  (4 trades per minute).

• Poisson probability function



#### **Poisson Distribution: Illustration**

Suppose a time interval is divided into n equal parts and that one event may or may not occur in each subinterval.



X = # of events is *Poisson*( $\lambda$ )

### **Poisson Distribution: Comments**

• The Poisson distribution arises in connection with Poisson processes - a stochastic process in which events occur continuously and independently of one another.

• It occurs most easily for time-events; such as the number of calls passing through a call center per minute, or the number of visitors passing through a turnstile per hour. However, it can apply to any process in which the mean can be shown to be constant.

• It is used in *finance* (number of jumps in an asset price in a given interval); *market microstructure* (number of trades per unit of time in a stock market); *sports economics* (number of goals in sports involving two competing teams); *insurance* (number of a given disaster - volcano eruptions/hurricanes/floods- per year); etc.

**Example**: The number of named storms over a period of a year in the Atlantic is known to have a Poisson distribution with  $\lambda = 13.1$ 

Determine the probability function of *X*.

Compute the probability that X is at most 8.

Compute the probability that *X* is at least 10.

Given that at least 10 hurricanes occur, what is the probability that *X* is at most 15? Solution:

$$p(x) = \frac{\lambda^{x}}{x!}e^{-\lambda}$$
  $x = 0, 1, 2, 3, 4, ...$ 

$$=\frac{13.1^{x}}{x!}e^{-13.1} \qquad x=0,1,2,3,4,\dots$$

x	p(x)	x	p(x)
0	0.000002	10	0.083887
1	0.000027	11	0.099901
2	0.000175	12	0.109059
3	0.000766	13	0.109898
4	0.002510	14	0.102833
5	0.006575	15	0.089807
6	0.014356	16	0.073530
7	0.026866	17	0.056661
8	0.043994	18	0.041237
9	0.064036	19	0.028432

 $P[\text{at most 8}] = P[X \le 8]$ 

$$= p(0) + p(1) + \dots + p(8) = .09527$$

 $P[\text{at least } 10] = P[X \ge 10] = 1 - P[X \le 9]$  $= 1 - (p(0) + p(1) + \dots + p(9)) = .8400$ 

$$= \frac{p(10) + p(11) + \dots + p(15)}{.8400} = 0.708$$

#### The Normal distribution

A random variable, X, is said to have a *normal distribution* with mean m and standard deviation s if X is a continuous random variable with probability density function f(x):



#### **Normal distribution: Properties**

**1.** Indexed by two parameters:  $\mu$  (central parameter) &  $\sigma$  (spread parameter).

**2.** Symmetric around  $\mu$ , which is the location of the maximum of f(x). Check:

$$f(x) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{(x-\mu)^2}{2\sigma^2}\right]$$
$$f'(x) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{(x-\mu)^2}{2\sigma^2}\right] * \left[-\frac{(x-\mu)}{\sigma^2}\right] = 0$$

The last equality holds when  $\mu = x$ . Thus,  $\mu$  is an extremum point of f(x). Since f(x) is a pdf, it is the mode.

**3.** The inflection points of f(x) are  $\mu - \sigma$ ,  $\mu + \sigma$ . (Check: set f''(x) = 0 and solve for x.)

#### Normal distribution: Comments

• The normal distribution is often used to describe or approximate any variable that tends to cluster around the mean. It is the most assumed distribution in economics and finance: rates of return, growth rates, IQ scores, observational errors, etc.

• The central limit theorem (CLT) provides a justification for the normality assumption when *n* is large.

<u>Notation</u>: PDF:  $x \sim N(\mu, \sigma^2)$ CDF:  $\Phi(x)$ 

## The Expectation of X: E(X)

The expectation operator defines the mean (or population average) of a random variable or expression.

### Definition

Let X denote a *discrete* RV with probability function p(x) (probability density function f(x) if X is *continuous*) then the expected value of X, E(X) is defined to be:

 $E(X) = \sum_{x} x p(x) = \sum_{i=1}^{\infty} x_i p(x_i)$ 

and if X is *continuous* with probability density function f(x)

 $E(X) = \int_{-\infty}^{\infty} x f(x) dx$ 

Sometimes we use E[.] as Ex[.] to indicate that the expectation is being taken over f x(x) dx

### **Interpretation of E(X)**

- 1. The expected value of X, E(X), is the center of gravity of the probability distribution of X.
- 2. The expected value of *X*, *E*(*X*), is the *long-run average value* of *X*. (To be discussed later: *Law of Large Numbers*)



#### **E[X]:** The Normal Distribution

Suppose *X* has a Normal distribution with parameters *m* and *s*. Then, E[X] = m. **Proof:** 

$$f(x) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{(x-\mu)^2}{2\sigma^2}\right]$$
$$E(X) = \int_{-\infty}^{\infty} x f(x) dx = \int_{-\infty}^{\infty} x \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{(x-\mu)^2}{2\sigma^2}\right] dx$$

Making the substitution:

$$z = \frac{x - \mu}{\sigma} \implies dz = \frac{1}{\sigma} dx$$
 and  $x = \mu + z\sigma$ 

Then,

$$E(X) = \int_{-\infty}^{\infty} \frac{1}{\sqrt{2\pi}} (\mu + z\sigma) e^{-\frac{z^2}{2}} dz$$
  
=  $\mu \int_{-\infty}^{\infty} \frac{1}{\sqrt{2\pi}} e^{-\frac{z^2}{2}} dz + \frac{\sigma}{\sqrt{2\pi}} \int_{-\infty}^{\infty} z e^{-\frac{z^2}{2}} dz$ 

Using the following results:

$$\int_{-\infty}^{\infty} \frac{1}{\sqrt{2\pi}} \exp\left[-\frac{z^2}{2}\right] dz = 1 \quad \text{and} \quad \int_{-\infty}^{\infty} z \, \exp\left[-\frac{z^2}{2}\right] dz = 0$$

Thus,  $E(X) = \mu$ 

### Expectation of a function of a RV

Let X denote a *discrete RV* with probability function p(x), then the expected value of g(X), E[g(X)], is defined to be:

$$E[g(X)] = \sum_{x} g(x) p(x) = \sum_{i=1}^{\infty} g(x_i) p(x_i)$$

and if *X* is *continuous* with probability density function f(x):  $E[g(X)] = \int_{-\infty}^{\infty} g(x) f(x) dx$ 

**Examples:**  $g(x) = (x - \mu)^2 \implies E[g(x)] = E[(x - \mu)^2]$  $g(x) = (x - \mu)^k \implies E[g(x)] = E[(x - \mu)^k]$ 

**Example**: Suppose *X* has a uniform distribution from 0 to *b*. Then:

$$f(x) = \begin{cases} 1/b & 0 \le x \le b \\ 0 & x \le 0, x > b \end{cases}$$

Find the *expected value* of  $A = X^2$ .

If X is the length of a side of a square (chosen at random from 0 to b) then A is the area of the square

$$E(X^{2}) = \int_{-\infty}^{\infty} x^{2} f(x) dx = \int_{a}^{b} x^{2} \frac{1}{b-a} dx = \left[\frac{1}{b} \frac{x^{3}}{3}\right]_{0}^{b} = \frac{b^{3} - 0^{3}}{3(b)} = \frac{b^{2}}{3}$$

= 1/3, the maximum area of the square

### Median: Another central measure

A median is the numeric value separating the higher half of a sample, a population, or a probability distribution, from the lower half.

### **Definition:** Median

The *median* of a random variable X is the unique number m that satisfies the following inequalities:

 $P(X \le m) \ge \frac{1}{2}$  and  $P(X \ge m) \ge \frac{1}{2}$ .

For a continuous distribution, we have that *m* solves:

$$\int_{-\infty}^{m} f_X(x) dx = \int_{m}^{\infty} f_X(x) dx = 1/2$$

<u>Note</u>: If the **mean** > **median** > **mode** (= most popular observation), the distribution will be skewed to the right. If the **mean** < **median** < **mode**, the distribution will be skewed to the left.

• Calculation of medians is a popular technique in summary statistics and summarizing statistical data, since it is simple to understand and easy to calculate, while also giving a measure that is more robust in the presence of outlier values than is the mean.

#### An optimality property

A median is also a central point which minimizes the average of the absolute deviations. That is, a value of c that minimizes

 $E(|\mathbf{X} - c|)$ 

is the median of the probability distribution of the random variable X.

**Example**: Let *X* have an exponential distribution with parameter  $\lambda$ . The probability density function of *X* is:

$$f(x) = \begin{cases} \lambda e^{-\lambda x} & x \ge 0\\ 0 & x < 0 \end{cases}$$

The median *m* solves the following integral of *X*:

$$\int_{m}^{\infty} f_X(x) dx = 1/2$$
  
$$\int_{m}^{\infty} \lambda e^{-\lambda x} dx = \lambda \int_{m}^{\infty} e^{-\lambda x} dx = -e^{-\lambda x} |_m^{\infty} = e^{-\lambda m} = 1/2$$

That is,  $m = \ln(2)/\lambda$ .

### **Moments of Random Variables**

The moments of a random variable *X* are used to describe the behavior of the RV (discrete or continuous).

#### **Definition**: *K*<sup>th</sup> Moment

Let X be a RV (discrete or continuous), then the  $k^{th}$  moment of X is:

$$\mu_{k} = E(X^{k}) = \begin{cases} \sum_{x} x^{k} p(x) & \text{if } X \text{ is discrete} \\ \int_{-\infty}^{\infty} x^{k} f(x) dx & \text{if } X \text{ is continuous} \end{cases}$$

#### **Definition:** Central Moments

Let X be a RV (discrete or continuous). Then, the  $k^{th}$  central moment of X is defined to be:

$$\mu_k^0 = E[(X - \mu)^k] = \begin{cases} \sum_x (x - \mu)^k p(x) & \text{if } X \text{ is discrete} \\ \int_{-\infty}^{\infty} (x - \mu)^k f(x) dx & \text{if } X \text{ is continuous} \end{cases}$$

where  $m = m_1 = E(X)$  = the first moment of *X*.

• The central moments describe how the probability distribution is distributed about the center of gravity, m.

• The first central moments is given by:

$$\mu_1^0 = E[X - \mu]$$

• The second central moment depends on the *spread* of the probability distribution of X about m. It is called the variance of X and is denoted by the symbol  $\sigma^2 = var(X)$ :

$$\mu_2^0 = E[(X - \mu)^2] = \operatorname{var}(X) = \sigma^2$$

The square root of var(X) is called the *standard deviation* of *X* and is denoted by the symbol s = SD(X). We also refer to it as *volatility*:

$$\sqrt{\mu_2^0} = \sqrt{E[(X-\mu)^2]} = \sigma$$

### **Moments of a RV: Skewness**

The third central moment:

$$\mu_3^0 = E[(X - \mu)^3]$$

 $\mu_3^0$  contains information about the *skewness* of a distribution.

• A popular measure of skewness:

$$\gamma_1 = \frac{\mu_3^0}{\sigma^3} = \frac{\mu_3^0}{(\mu_2^0)^2}$$

- Distribution according to skewness:
- 1) Symmetric distribution



2) Positively (right-) skewed distribution (with mode < median < mean)



3) Negatively (left-) skewed distribution (with mode > median > mean)



- Skewness and Economics
- Zero skew means symmetrical gains and losses.
- Positive skew suggests many small losses and few rich returns.
- Negative skew indicates lots of minor wins offset by rare major losses.

• In financial markets, stock returns at the firm level show positive skewness, but at the aggregate (index) level show negative skewness.

• From horse race betting and from U.S. state lotteries there is evidence supporting the contention that gamblers are not necessarily risk-lovers but skewness-lovers: Long shots are overbet (positive skewness loved!).

### Moments of a RV: Kurtosis

The fourth central moment:

$$\mu_4^0 = E[(X - \mu)^4]$$

It contains information about the *shape* of a distribution. The property of shape that is measured by this moment is called *kurtosis*, usually estimated by  $\kappa$ 

$$\kappa = \frac{\mu_4^0}{\sigma^4}.$$

• The measure of (excess) kurtosis:

$$\gamma_2 = \frac{\mu_4^0}{\sigma^4} - 3 = \frac{\mu_4^0}{(\mu_2^0)^2} - 3$$

<u>Note</u>: We subtract 3, because the kurtosis of the Normal distribution is  $\kappa=3$ .

• Distributions:

1) Mesokurtic distribution







3) Leptokurtic distribution (usual shape for asset returns)



## **Moments and Expected Values**

Note that moments are defined by expected values. We define the expected value of a function of a continuous RV X, g(X), as

$$E[g(X)] = \int_{-\infty}^{\infty} g(x)f(x)dx$$

• If *X* is *discrete* with probability function p(x)

$$E[g(X)] = \sum_{x} g(x)p(x) = \sum_{i} g(x_i)p(x_i)$$

**Examples:** 
$$g(x) = (x - \mu)^2$$
  $\Rightarrow E[g(x)] = E[(x - \mu)^2]$   
 $g(x) = (x - \mu)^k$   $\Rightarrow E[g(x)] = E[(x - \mu)^k]$ 

• We estimate expected values with sample averages. The Law of Large Numbers (LLN) tells us they are *consistent* estimators of expected values.

### **Estimating Moments**

We estimate expected values with sample averages. For example, the first moment, the mean, and the second central moment, the variance, are estimated by:

$$\bar{X} = \frac{\sum_{i=1}^{N} X_i}{N}$$

$$s^2 = \frac{\sum_{i=1}^{N} (X_i - \bar{X})^2}{N - 1}$$
(N - 1 adjustment needed for E[s<sup>2</sup>] =  $\sigma^2$ )

• Besides consistent, they are both are *unbiased* estimators of their respective population moments (unbiased = "on average, I get the population parameter"). That is,

 $E[\bar{X}] = \mu$  "population parameter"  $E[s^2] = \sigma^2$ 

### The Law of Large Numbers (LLN)

Long history: Gerolamo Cardano (1501-1576) stated it without proof. Jacob Bernoulli published a rigorous proof in 1713.

#### Theorem (Weak LLN)

Let  $x_1, x_2, ..., x_N$  be *N* mutually independent random variables each having mean  $\mu$  and a finite variance  $\sigma^2$ -i.e, the sequence  $\{x_N\}$  is *i.i.d*.

Let  $\bar{X} = \frac{\sum_{i=1}^{N} x_i}{N}$ . Then, for any  $\delta > 0$  (no matter how small)

 $P[|\bar{X} - \mu| < \delta] = P[|\mu - \delta < \bar{X} < \mu + \delta] \to 1, \qquad \text{as } N \to \infty$ 

• There are many variations of the LLN. It is a general result: A sample average as the sample size goes to infinite tends to its expected value. Also written as:

 $\bar{X}_N \xrightarrow{p} \mu$ . (convergence in probability)

### The Central Limit Theorem (CLT)

The Central Limit Theorem (CLT) states conditions for the sequence of RV  $\{x_N\}$  under which the mean or a sum of a sufficiently large number of  $x_i$ 's will be approximately normally distributed.

Let  $x_1, x_2, ..., x_N$  be a sequence of *i.i.d.* RVs with finite mean  $\mu$ , and finite variance  $\sigma^2$ . Then, as N increases,  $\overline{X}_N$ , the sample mean, approaches the normal distribution with mean  $\mu$  and variance  $\sigma^2/N$ .

This theorem is sometimes stated as:

$$\frac{\sqrt{N}(\bar{X}-\mu)}{\sigma} \stackrel{d}{\to} N(0,1)$$

where  $\stackrel{d}{\rightarrow}$  means "the limiting distribution (asymptotic distribution) is" (or *convergence in distribution*).

• Many version of the CLT. Two versions are commonly used in economics and finance:

- The one above is the *Lindeberg-Lévy CLT*, with  $\{x_N\}$  are *i.i.d.*, with finite  $\mu$  and finite  $\sigma^2$ . - The other one is the *Lindeberg-Feller CLT*. It requires  $\{x_N\}$  to be independent, with finite  $\mu_i$ ,  $\sigma_i^{2} < \infty$ ,  $S_n = \sum_i x_i$ ,  $s_n^{2} = \sum_i \sigma_i^{2}$  and for  $\varepsilon > 0$ ,

$$\lim_{n \to \infty} \frac{1}{s_n^2} \sum_{i=1}^n \int_{|x_i - \mu_i| > \delta S_n} (x_i - \mu_i)^2 f(x_i) dx = 0$$

Note:

Lindeberg-Levy assumes random sampling –observations are *i.i.d.*, with the same mean and same variance.

Lindeberg-Feller allows for heterogeneity in the drawing of the observations --through different variances. The cost of this more general case: More assumptions about how the  $\{x_N\}$  vary.

• The CLT gives only an asymptotic distribution. We usually take it as an approximation for a finite number of observations. In these cases, the notation goes from  $\stackrel{d}{\rightarrow}$  to  $\stackrel{a}{\rightarrow}$ .

<u>Technical Note</u>: The *Berry–Esseen theorem (Berry–Esseen inequality)* attempts to quantify the rate at which the convergence to normality takes place.

$$|F_n(x) - \Phi(x)| \le \frac{C\rho}{\sigma^3 n^{1/2}}$$

where  $\rho = E(|X|) < \infty$  and C is a constant (best current C=0.7056).

#### **Asymptotic Distribution**

An asymptotic distribution is a hypothetical distribution that is the *limiting* distribution of a sequence of distributions.

We will use the asymptotic distribution as a finite sample *approximation* to the true distribution of a RV when *N*-i.e., the sample size- is *large*.

Practical question: When is N large?

#### **Sampling Distributions**

All statistics, T(X), are functions of RVs and, thus, they have a distribution. Depending on the sample, we can observe different values for T(X), thus, the finite sample distribution of T(X) is called the *sampling distribution*.

For the sample mean,  $\overline{X}$ , if the  $X_i$ 's are normally distributed, then the sampling distribution is normal with mean  $\mu$  and variance  $\sigma^2/N$ . Or

$$\overline{X} \sim N(\mu, \sigma^2/N).$$

Then,  $E[\overline{X}] = \mu$  $Var[\overline{X}] = \sigma^2/N \implies variance of sample mean decreases as N increases!$ 

The SD of the sampling distribution is called the *standard error* (SE). Then,  $SE(\bar{X}) = \sigma/sqrt(N)$ .

We usually associate the standard error with the precision of the estimate. That is, the precision of the estimation of the mean increases as *N* increases.

• Below, we show the sampling distribution for the sample mean of a normal population for different sample sizes (N).



<u>Note</u>: As  $N \to \infty$ ,  $\overline{X} \to \mu$  —i.e., the distribution becomes a spike at  $\mu$ !

<u>Note</u>: If the data is not normal, the CLT is used to approximate the sampling distribution by the asymptotic one, usually, after some manipulations. Again, in those cases, the notation goes from  $\stackrel{d}{\rightarrow}$  to  $\stackrel{a}{\rightarrow}$ .

• For the sample variance  $\sigma^2$ , if the  $X_i$ 's are normally distributed, then the sampling distribution is derived from this result:

$$(N-1) s^2/\sigma^2 \sim \chi^2_{N-1}.$$

It can be shown that a random variable that follows a  $\chi_{\nu}^2$  distribution has a variance equal to 2 times the degrees of freedom (=2\* $\nu$ ). Then,

 $\operatorname{Var}[(N-1) s^2/\sigma^2] = 2 * (N-1) \implies \operatorname{Var}[s^2] = 2 * \sigma^4 / (N-1)$ 

Then,  $SE(s^2) = SD(s^2) = \sigma^2 * \sqrt{2/(N-1)}].$ 

<u>Note</u>: If the data is not normal (& N is large), the CLT can be used to approximate the sampling distribution by the asymptotic one:

 $s^2 \xrightarrow{a} N(\sigma^2, \sigma^4 * (\kappa - 1)/N)$ where  $\kappa = \frac{\mu_4^0}{\sigma^4}$  (recall when data is normal,  $\kappa = 3$ ).

<u>Remark</u>: The precision of the estimation increases as N increases.

This remark is especially relevant in Finance, where we derive relations between expected returns and risk factors, like market risk or volatility. As we gather more data, expected returns and the volatility of returns will be more precisely estimated.

## **Hypothesis Testing**

A *statistical hypothesis test* is a method of making decisions using experimental data. A result is called *statistically significant* if it is unlikely to have occurred by chance.

• These decisions are made using (null) hypothesis tests. A hypothesis can specify a particular value for a population parameter, say  $q=q_0$ . Then, the test can be used to answer a question like:

Assuming  $q_0$  is true, what is the probability of observing a value for the (test) statistic used that is at least as big as the value that was actually observed?

• Uses of hypothesis testing:

- Check the validity of theories or models.
- Check if new data can cast doubt on established facts.

• In general, there are two kinds of hypotheses:

(1) About the form of the probability distribution

**Example**: Is the random variable normally distributed?

(2) About the parameters of a distribution function **Example**: Is the mean of a distribution equal to 0?

• The second class is the traditional material of econometrics. We may test whether the effect of income on consumption is greater than one, or whether the size coefficient on a CAPM regression is equal to zero.

• Hypothesis testing involves the comparison between two competing hypothesis (sometimes, they represent partitions of the world).

- The null hypothesis, denoted  $H_0$ , is sometimes referred to as the maintained hypothesis.
- The alternative hypothesis, denoted *H*<sub>1</sub>, is the hypothesis that will be considered if the null hypothesis is "rejected."

<u>Idea</u>: We collect a sample of data  $X = \{X_1, ..., X_N\}$ . We construct a statistic T(X) = f(X), called the *test statistic*. Now we have a decision rule:

- If T(X) is contained in space *R*, we reject  $H_0$  (& we learn).
- If T(X) is in the complement of  $R(R^{C})$ , we fail to reject  $H_0$ .

<u>Note</u>: T(X), like any other statistic, is a RV. It has a distribution.

**Example:** Suppose we want to test if the mean of IBM annual returns,  $\mu_{IBM}$ , is 10%. That is,  $H_0$ :  $\mu_{IBM} = 10\%$ .

From the population, we get a sample: { $X_{1962}$ ,  $X_{1963}$ , ...,  $X_{N=2024}$ }, with N=63. We use  $T(X) = \overline{X}$ , which is unbiased, consistent, and, assuming X is normally distributed, we know its distribution,  $\overline{X} \sim N(\mu, \sigma^2/N)$ .



Now, we need to determine the rejection region, *R*, such that if

$$T(X) = \overline{X} \notin [T_{LB}, T_{UB}] \implies \text{Reject H}_0: \mu_{\text{IBM}} = 10\%$$

That is,

$$\mathbf{R} = [\bar{X} < T_{LB}, T_{UB} > \bar{X}]$$



• Q: How do we determine  $T_{LB}$  and  $T_{UB}$  and, thus, make a decision? We use the distribution of  $\overline{X}$ , which is derived under H<sub>0</sub>. As we will se below, we determine R in such a way that the probability of rejecting a true H<sub>0</sub> is "small."

<u>Note</u>: In the Graph above, the blue area gives us the associated probability with R. That is, the probability, under H<sub>0</sub>, that the observed  $T(X) = \overline{X}$  falls in the rejection region. The blue are is called the *significance level*.

## **Hypothesis Testing: Steps**

We present the *classical approach*, a synthesized approach, known as *significance testing*. It relies on Fisher's *p*-value: the probability, of observing a result at least as extreme as the test statistic, under  $H_0$ .

We follow these steps:

**Step 1.** Identify H<sub>0</sub> & decide on a *significance level* ( $\alpha$ %) to compare your test results.

**Step 2.** Determine the appropriate test statistic T(X) and its distribution under the assumption that H<sub>0</sub> is true.

**Step 3.** Calculate T(X) from the data.

Step 4. Decision Rule:

Reject H<sub>0</sub> if the *p*-value is sufficiently small, that is, we consider T(X) in *R* (we learn). Otherwise, we reach no conclusion (no learning).

• Q: What *p*-value is "sufficiently small" as to warrant rejection of H<sub>0</sub>? <u>Rule</u>: If *p*-value <  $\alpha$  (say, 5%)  $\Rightarrow$  test result is *significant*: Reject H<sub>0</sub>. If the results are "*not significant*," no conclusions are reached (no learning here). Go back gather more data or modify model.

• The father of this approach, Ronald Fisher, favored 5% or 1%.

**Example:** From the U.S. Jury System  $H_0$ : The defendant is not guilty  $H_1$ : The defendant is guilty

In statistics we learn when we reject. In this case, we learn a defendant is guilty when the jury finds the defendant guilty, by rejecting  $H_0$ .

Example: From the U.S. Jury System
Step 1. Identify H<sub>0</sub> & decide on a *significance level* (α%)
H<sub>0</sub>: The defendant is not guilty
H<sub>1</sub>: The defendant is guilty
Significance level α = "beyond reasonable doubt," presumably small level.

**Step 2.** After judge instructions, each juror forms an "innocent index"  $T(X)_i$ .

**Step 3.** Through deliberations, jury reaches a conclusion  $T(X) = \sum_{i=1}^{12} T(X)_i$ .

**Step 4.** Decision Rule:

If *p*-value of  $T(X) < \alpha \Rightarrow$  Reject  $H_0$ . That is, guilty! If *p*-value of  $T(X) > \alpha \Rightarrow$  Fail to reject  $H_0$ . That is, non-guilty. Alternatively, we build a rejection region, R, around  $H_0$ . Then, if  $T(X) \in \mathbb{R}$ , we reject  $H_0$ .

Note: Mistakes are made. We want to quantify these mistakes.

• Failure to reject  $H_0$  does not necessarily mean that the defendant is not guilty, or rejecting  $H_0$  does not mean necessarily the defendant is guilty. *Type I error* and *Type II error* give us an idea of both mistakes.

**Definition**: Type I and Type II errors

A *Type I error* is the error of rejecting  $H_0$  when it is true. A *Type II error* is the error of "accepting"  $H_0$  when it is false (that is, when  $H_1$  is true).

<u>Notation</u>: Probability of Type I error:  $\alpha = P[X \in R | H_0 \text{ is true}]$ Probability of Type II error:  $\beta = P[X \in R^C | H_1 \text{ is true}]$ 

	State of World	
Decision	H <sub>0</sub> true	H <sub>1</sub> true (H <sub>0</sub> false)
Cannot reject ("accept") H <sub>0</sub>	Correct decision	Type II error
Reject H <sub>0</sub>	Type I error	Correct decision

Need to control both types of error:

 $\alpha = P[rejecting H_0 | H_0 is true]$ 

 $\beta = P[\text{not rejecting } H_0 | H_1 \text{ is true}]$ 

**Example:** From the U.S. Jury System

*Type I error* is the error of finding an innocent defendant guilty. *Type II error* is the error of finding a guilty defendant not guilty.

• In general, we think *Type I error* is the worst of the two errors, we try to minimize the error of sending to jail an innocent person.

Actually, we would like *Type I error* to be zero. However, the only way to do this (100% of innocent defendants are found not guilty) is to never reject H<sub>0</sub>. Then, we maximize *Type II error*.

• There is a clear trade-off between both errors. Traditional view: Set *Type I error* equal to a small number (defined in the U.S. court system as "*beyond reasonable doubt*") and design a test that minimizes *Type II error*.

The usual tests (t-tests, F-tests, Likelihood Ratio tests) incorporate this traditional view.

### Hypothesis Testing: z-test & t-test

For inferences about the population mean, the usual test statistic is the t-test. It is a modification of the z-test statistic.

• **z-test.** Assuming  $\{X_1, X_2, X_3, ..., X_N\}$  is generated by a N( $\mu$ ,  $\sigma^2$ ), then, the sampling distribution of the sample mean is:

$$\overline{X} \sim N(\mu, \sigma^2/N).$$
  
Using the CLT, the distribution of the standardize sample mean, z, is:  
 $z = \frac{\overline{X} - \mu}{\sigma/\sqrt{N}} \sim N(0, 1)$ 

• **t-test.** In practice,  $\sigma$  is unknown. We need to estimate it, which we do with use *s*. Then, keeping the assumption  $\{X_i\} \sim N(\mu, \sigma^2)$ :

$$\mathbf{t} = \frac{\bar{x} - \mu}{s_{/\sqrt{N}}} \sim t_{N-1} \qquad -\text{when } N > 30, t_N \sim \mathcal{N}(0, 1).$$

Below, we plot a simulated t-distribution with  $\nu = 5$  (in red), along a normal distribution (in blue). It has thicker tails. As  $\nu$  increases,  $t_{\nu}$  converges to a N(0, 1) distribution.



<u>Technical Note 2</u>: The distribution of t is exact if  $\{X_i\} \sim N(\mu, \sigma^2)$ , otherwise, the distribution is asymptotic (for large *N*). That is,

$$t = \frac{\bar{x} - \mu}{s / \sqrt{N}} \xrightarrow{d} N(0, 1).$$

**Example:** We want to test if the mean is equal to  $\mu_0$ . Then,

1. 
$$H_0: \mu = \mu_0.$$

H<sub>1</sub>:  $\mu \neq \mu_0$ .

2. Appropriate T(X): *t-test* (based on  $\sigma$  unknown and estimated by *s*).

Determine distribution of T(X) under H<sub>0</sub>. Sampling distribution of  $\overline{X}$ , under H<sub>0</sub>:  $\overline{X} \sim N(\mu_0, \sigma^2/N)$ .

Then, distribution of T(X) under H<sub>0</sub>:

$$=\frac{\bar{X} - \mu_0}{s_{/\sqrt{N}}} \sim t_{N-1} \qquad - \text{ when } N > 30, t \sim N(0, 1).$$

3. Compute t,  $\hat{t}$ , using  $\bar{X}$ ,  $\mu_0$ , s, and N. Get *p*-value( $\hat{t}$ ).

t

4. <u>Rule</u>: Set an  $\alpha$  level. If *p*-value( $\hat{\mathbf{t}}$ ) <  $\alpha$   $\Rightarrow$  Reject H<sub>0</sub>:  $\mu = \mu_0$ . Alternatively, if  $|\hat{\mathbf{t}}| > t_{N-1,\alpha/2}$  (=1.96, if  $\alpha$ =.05)  $\Rightarrow$  Reject H<sub>0</sub>:  $\mu = \mu_0$ .

<u>Technical Note 2</u>: In step 2, we determine the distribution of t, by using the sampling distribution of  $\overline{X}$  under H<sub>0</sub>. If H<sub>0</sub> is not true, suppose  $\mu = \mu_I$ , then

$$\overline{X} \sim N(\mu_1, \sigma^2/N),$$

and, thus, t is distributed N(0, 1) only under H<sub>0</sub>, since only under H<sub>0</sub> the E[ $\overline{X} - \mu_0$ ] = 0.

## Lecture 1 – Appendix: Review of Linear Algebra

## A Matrix

A matrix is a set of elements, organized into rows and columns



- *a* and *d* are the diagonal elements.
- *b* and *c* are the off-diagonal elements.

• Matrices are like plain numbers in many ways: they can be added, subtracted, and, in some cases, multiplied and inverted (divided).

### **Example:**

$$\boldsymbol{A} = \begin{bmatrix} a_{11} & a_{21} \\ a_{12} & a_{22} \end{bmatrix}; \quad \boldsymbol{b} = \begin{bmatrix} b_1 & b_2 & b_3 \end{bmatrix}. \P$$

Dimensions of a matrix: numbers of rows by numbers of columns. The Matrix A is a  $2x^2$  matrix, **b** is a  $1x^3$  matrix.

A matrix with only 1 column or only 1 row is called a vector.

If a matrix has an equal numbers of rows and columns, it is called a *square* matrix. Matrix A, above, is a square matrix.

Usual Notation:	Upper case letters	$\Rightarrow$ matrices	
	Lower case	$\Rightarrow$ vectors	

### **Matrices - Information**

Information is described by data. A tool to organize the data is a list, which we call a vector. Lists of lists are called matrices. That is, we organize the data using matrices.

We think of the elements of **X** as data points ("data entries", "observations"), in economics, we usually have numerical data.

We store the data in rows. In a Txk matrix, **X**, over time we build a database:

$$X = \begin{bmatrix} x_{11} & \cdots & x_{k1} \\ \vdots & \ddots & \vdots \\ x_{1T} & \cdots & x_{kT} \end{bmatrix}$$

• Once the data is organized in matrices it can be easily manipulated: multiplied, added, etc. (this is what Excel does).

• In econometrics, we have a model  $y = f(x_1, x_2, ..., x_k)$ , which we want to estimate. We collect data, say T (or N) observations, on a dependent variable, y, and on k explanatory variables, X.

• Under the usual notation, vectors will be column vectors: y and  $x_k$  are Tx1 vectors:

$$\mathbf{y} = \begin{bmatrix} y_1 \\ \vdots \\ y_T \end{bmatrix} & \& \quad \mathbf{x}_j = \begin{bmatrix} x_{j1} \\ \vdots \\ x_{jT} \end{bmatrix} \quad j = 1, \dots, k$$
  
$$\mathbf{X} \text{ is a } Txk \text{ matrix:} \quad \mathbf{X} = \begin{bmatrix} x_{11} & \cdots & x_{k1} \\ \vdots & \ddots & \vdots \\ x_{1T} & \cdots & x_{kT} \end{bmatrix}$$

Its columns are the k Tx1 vectors  $x_i$ . It is common to treat  $x_1$  as vector of ones, *i*.

#### **Special Matrices – Identity and Null**

• *Identity Matrix:* A square matrix with 1's along the diagonal and 0's everywhere else. Similar to scalar "1."

$$I = \begin{bmatrix} 1 & \cdots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \cdots & 1 \end{bmatrix}$$

• Null matrix: A matrix in which all elements are 0's. Similar to scalar "0."

$$\mathbf{0} = \begin{bmatrix} 0 & \cdots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \cdots & 0 \end{bmatrix}$$

• Both are *diagonal* matrices  $\Rightarrow$  off-diagonal elements are zero.

Note: Both are examples of symmetric and idempotent matrices. As we will see later:

- Symmetric:  $A = A^T$ - Idempotent:  $A = A^2 = A^3 = \dots$ 

### **Elementary Row Operations**

Elementary row operations:

- Switching: Swap the positions of two rows
- Multiplication: Multiply a row by a non-zero scalar
- Addition: Add to one row a scalar multiple of another.

• An *elementary matrix* is a matrix which differs from the identity matrix by one single elementary row operation.

• If the matrix subject to elementary row operations is associated to a system of linear equations, then these operations do not change the solution set. Row operations can make the problem easier.

• Elementary row operations are used in Gaussian elimination to reduce a matrix to row echelon form.

#### **Matrix multiplication: Details**

Multiplication of matrices requires a conformability condition

• The <u>conformability condition</u> for multiplication is that the <u>column</u> dimensions of the <u>lead</u> matrix **A** must be equal to the <u>row</u> dimension of the <u>lag</u> matrix **B**.

• If **A** is an  $(m \ge n)$  and **B** an  $(n \ge p)$  matrix (**A** has the same number of columns as **B** has rows), then we define the product of **AB**. **AB** is  $(m \ge p)$  matrix with its  $ik^{\text{th}}$  element is  $c_{ik} = \sum_{j=1}^{n} a_{ij} b_{jk}$ .

**Example:** Suppose we have a 1x2 vector a, and a 2x3 matrix B. What are the dimensions of the product:  $a^*B$ ?

$$aB = \begin{bmatrix} a_{11}a_{12} \end{bmatrix} \begin{bmatrix} b_{11} & b_{12} & b_{13} \\ b_{21} & b_{22} & b_{23} \end{bmatrix} = c = \begin{bmatrix} c_{11} & c_{12} & c_{13} \end{bmatrix}$$
$$= \begin{bmatrix} a_{11}b_{11} + a_{12}b_{21} & a_{11}b_{12} + a_{12}b_{22} & a_{11}b_{13} + a_{12}b_{23} \end{bmatrix}$$

<u>Dimensions</u>:  $a(1x^2)$ , B(2x3)  $\Rightarrow$  c(1x3).

**Example**: We want to multiply A (2x2) and B (2x2), where A has elements  $a_{ij}$  and B has elements  $b_{jk}$ . Recall the ik element is

$$c_{ik} = \sum_{j=1}^{n=2} a_{ij} b_{jk}$$

$$\mathbf{A} = \begin{bmatrix} 2 & 1 \\ 7 & 9 \end{bmatrix}$$

$$\mathbf{B} = \begin{bmatrix} 1 & 0 \\ 2 & 3 \end{bmatrix}$$

$$\mathbf{C} = \begin{bmatrix} 2 & 1 \\ 7 & 9 \end{bmatrix} * \begin{bmatrix} 1 & 0 \\ 2 & 3 \end{bmatrix} = \begin{bmatrix} \mathbf{4} = \mathbf{2} * \mathbf{1} + \mathbf{1} * \mathbf{2} & \mathbf{3} = \mathbf{2} * \mathbf{0} + \mathbf{1} * \mathbf{3} \\ \mathbf{25} = \mathbf{7} * \mathbf{1} + \mathbf{9} * \mathbf{2} & \mathbf{27} = \mathbf{7} * \mathbf{0} + \mathbf{9} * \mathbf{3} \end{bmatrix}$$

$$C_{2x2} = A_{2x2} * B_{2x2}$$

<u>Dimensions</u>: A(2x2),  $B(2x2) \Rightarrow C(2x2)$ , a square matrix. ¶

**Example**: We want to multiply X (2x2) and  $\beta$  (2x1), where X has elements  $x_{ij}$  and **b** has elements  $\beta_j$ :

$$\mathbf{X} = \begin{bmatrix} x_{11} & x_{21} \\ x_{12} & x_{22} \end{bmatrix} \qquad \& \qquad \mathbf{\beta} = \begin{bmatrix} \beta_1 \\ \beta_2 \end{bmatrix}$$

We compute

$$y = X \beta$$

Recall the  $i^{th}$  element is

$$y_i = \sum_{j=1}^{n=2} x_{ij} \beta_j$$

Then,

$$\mathbf{y} = \begin{bmatrix} y_1 \\ y_2 \end{bmatrix} = \begin{bmatrix} x_{11} & x_{21} \\ x_{12} & x_{22} \end{bmatrix} * \begin{bmatrix} \beta_1 \\ \beta_2 \end{bmatrix} = \begin{bmatrix} x_{11} & \beta_1 + x_{21} & \beta_2 \\ x_{12} & \beta_1 + x_{22} & \beta_2 \end{bmatrix}$$

<u>Dimensions</u>:  $X(2x^2)$ ,  $\beta(2x^1) \Rightarrow y(2x^1)$ , a row vector. ¶

### **Transpose Matrix**

The transpose of a matrix A is another matrix  $A^{T}$  (also written A') created by any one of the following equivalent actions:

.

-write the rows (columns) of A as the columns (rows) of  $A^{T}$ 

-reflect A by its main diagonal to obtain  $A^{T}$ 

Formally, the (i,j) element of  $\mathbf{A}^{\mathrm{T}}$  is the (j,i) element of  $\mathbf{A}$ :  $[\mathbf{A}^{\mathrm{T}}]_{ij} = [\mathbf{A}]_{ji}$ 

**Example**: 
$$A = \begin{bmatrix} 3 & 8 & -9 \\ 1 & 0 & 4 \end{bmatrix} \Rightarrow A' = \begin{bmatrix} 3 & 1 \\ 8 & 0 \\ -9 & 4 \end{bmatrix} \cdot \P$$

• Results:

- If **A** is a 
$$m \times n$$
 matrix  $\Rightarrow \mathbf{A}^{\mathrm{T}}$  is a  $n \times m$  matrix.

- Conformability changes unless the matrix is square.
- $-(\mathbf{AB})' = \mathbf{B'A'}$

**Example:** In econometrics, an important matrix is **X'X**. Recall **X**:

$$X = \begin{bmatrix} x_{11} & \cdots & x_{k1} \\ \vdots & \ddots & \vdots \\ x_{1T} & \cdots & x_{kT} \end{bmatrix}$$
 a (*Txk*) matrix

Then,

$$X' = \begin{bmatrix} x_{11} & \cdots & x_{1T} \\ \vdots & \ddots & \vdots \\ x_{k1} & \cdots & x_{kT} \end{bmatrix}$$
 a (*kxT*) matrix. ¶

## **Basic Operations**

Addition, Subtraction, Multiplication

$$\begin{bmatrix} a & b \\ c & d \end{bmatrix} + \begin{bmatrix} e & f \\ g & h \end{bmatrix} = \begin{bmatrix} a+e & b+f \\ c+g & d+h \end{bmatrix}$$
$$\begin{bmatrix} a & b \\ c & d \end{bmatrix} - \begin{bmatrix} e & f \\ g & h \end{bmatrix} = \begin{bmatrix} a-e & b-f \\ c-g & d-h \end{bmatrix}$$
$$\begin{bmatrix} a & b \\ c & d \end{bmatrix} \begin{bmatrix} e & f \\ g & h \end{bmatrix} = \begin{bmatrix} ae+bg & af+bh \\ ce+dg & cf+dh \end{bmatrix}$$
$$k \begin{bmatrix} a & b \\ c & d \end{bmatrix} = \begin{bmatrix} ka & kb \\ kc & kd \end{bmatrix}$$

Just add elements

Just subtract elements

Multiply each row by each column and add

Multiply each row by each column and add

### **Example:**

$\begin{bmatrix} 2 & 1 \\ 7 & 9 \end{bmatrix} + \begin{bmatrix} 3 & 1 \\ 0 & 2 \end{bmatrix} = \begin{bmatrix} 5 & 2 \\ 7 & 11 \end{bmatrix}$ $A_{2x2} + B_{2x2} = C_{2x2}$	Addition
$\begin{bmatrix} 2 & 1 \\ 7 & 9 \end{bmatrix} - \begin{bmatrix} 1 & 0 \\ 2 & 3 \end{bmatrix} = \begin{bmatrix} 1 & 1 \\ 5 & 6 \end{bmatrix}$ $A_{2x2} - B_{2x2} = C_{2x2}$	Subraction
$\begin{bmatrix} 2 & 1 \\ 7 & 9 \end{bmatrix} * \begin{bmatrix} 1 & 0 \\ 2 & 3 \end{bmatrix} = \begin{bmatrix} 4 & 3 \\ 25 & 27 \end{bmatrix}$ $A_{2x2} * B_{2x2} = C_{2x2}$	Multiplication
$\frac{1}{8} \begin{bmatrix} 2 & 4 \\ 6 & 1 \end{bmatrix} = \begin{bmatrix} 1/4 & 1/2 \\ 3/4 & 1/8 \end{bmatrix}$	Scalar multiplication.

## **Basic Matrix Operations:** ε'ε

In Least Squares (LS) estimation, we minimize a sum of square errors ( $\varepsilon_i$  for = 1, 2, ..., T):

$$S(x_i, \beta) = \sum_{i=1}^T \varepsilon_i^2$$

Let  $\varepsilon$  be the *T*x1 vector of errors. We use linear algebra to write the sum of squares of its elements as (dot product of 2 *T*x1 vectors):

$$S(x_i, \beta) = \sum_{i=1}^T \varepsilon_i^2 = \varepsilon' \varepsilon$$

Check:

$$\boldsymbol{\varepsilon}' \, \boldsymbol{\varepsilon} = \left[\varepsilon_1 \, \varepsilon_2 \dots \, \varepsilon_T\right] * \begin{bmatrix} \varepsilon_1 \\ \vdots \\ \varepsilon_T \end{bmatrix} = \left[\varepsilon_1^2 + \varepsilon_2^2 + \dots + \varepsilon_T^2\right] = \sum_{i=1}^T \varepsilon_i^2$$

Thus, if we define  $y = X\beta + \varepsilon$ , LS estimation picks  $\beta$  to minimize:

$$S(x_i, \beta) = \varepsilon'\varepsilon = (\mathbf{y} - \mathbf{X}\beta)' (\mathbf{y} - \mathbf{X}\beta).$$

### **Basic Matrix Operations: X'X**

A special matrix in econometrics, **X'X** (a *kxk* matrix):

Recall X (Txk): 
$$X = \begin{bmatrix} x_{11} & \cdots & x_{k1} \\ \vdots & \ddots & \vdots \\ x_{1T} & \cdots & x_{kT} \end{bmatrix} \& X' = \begin{bmatrix} x_{11} & \cdots & x_{1T} \\ \vdots & \ddots & \vdots \\ x_{k1} & \cdots & x_{kT} \end{bmatrix}$$
$$X' X = \begin{bmatrix} \sum_{i=1}^{T} x_{1i}^2 & \cdots & \sum_{i=1}^{T} x_{1i} x_{ki} \\ \vdots & \ddots & \vdots \\ \sum_{i=1}^{T} x_{ki} x_{1i} & \cdots & \sum_{i=1}^{T} x_{ki}^2 \end{bmatrix} =$$
$$= \sum_{i=1}^{T} \begin{bmatrix} x_{1i}^2 & \cdots & x_{1i} x_{ki} \\ \vdots & \ddots & \vdots \\ x_{ki} x_{1i} & \cdots & x_{ki}^2 \end{bmatrix} =$$
$$= \sum_{i=1}^{T} \begin{bmatrix} x_{1i} \\ \vdots \\ x_{ki} \end{bmatrix} [x_{1i} & \cdots & x_{ki}] = \sum_{i=1}^{T} x_i x_i'$$

## **Basic Matrix Operations:** *i***'X**

Recall i is a column vector of ones (in this case, a Tx1 vector):

$$i = \begin{bmatrix} 1 \\ 1 \\ \dots \\ 1 \end{bmatrix}$$

Given X (Txk), then **i**' X is a 1xk vector:

$$\mathbf{i}'X = \begin{bmatrix} \mathbf{1} & \dots & \mathbf{1} \end{bmatrix} \begin{bmatrix} x_{11} & \cdots & x_{k1} \\ \vdots & \ddots & \vdots \\ x_{1T} & \cdots & x_{kT} \end{bmatrix} = \begin{bmatrix} \sum_{t=1}^{T} x_{1t} & \dots & \sum_{t=1}^{T} x_{kt} \end{bmatrix}$$

<u>Note</u>: If  $\mathbf{x}_1$  is a vector of ones (representing a constant in the linear classical model), then:  $\mathbf{\hat{i}} \cdot \mathbf{x}_1 = \sum_{t=1}^T \mathbf{x}_{1t} = \sum_{t=1}^T \mathbf{1} = T$  ("dot product")

**Inverse of a Matrix** Identity matrix: **AI** = **A** 

$$I_3 = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

<u>Notation</u>:  $I_j$  is a *j*x*j* identity matrix.

Given A (mxn), the matrix B (nxm) is a *right-inverse* for A iff AB = I<sub>m</sub>

Given A (mxn), the matrix C (mxn) is a *left-inverse* for A iff CA = I<sub>n</sub>

**Theorem:** If A (mxn), has both a *right-inverse* B and a *left-inverse* C, then C = B. <u>Proof</u>:

We have  $AB = I_m$  and  $CA = I_n$ .

Thus,

$$C(AB) = C I_m = C$$
 and  $C(AB) = (CA)B = I_nB = B$   
 $\Rightarrow C(nxm) = B(mxn)$ 

Note:

- This matrix is unique. (Suppose there is another left-inverse **D**, then  $\mathbf{D} = \mathbf{B}$  by the theorem, so  $\mathbf{D} = \mathbf{C}$ ).

- If **A** has both a right and a left inverse, it is a square matrix. It is usually called *invertible*. We say "the matrix **A** is *non-singular*."

• Inversion is tricky:  $(ABC)^{-1} = C^{-1} B^{-1} A^{-1}$ 

**Theorem:** If **A** (*mxn*) and **B** (*nxp*) have inverses, then **AB** is invertible and  $(AB)^{-1} = B^{-1}A^{-1}$ <u>Proof</u>:

We have  $\mathbf{A}\mathbf{A}^{-1} = \mathbf{I}_{\mathbf{m}}$  and  $\mathbf{A}^{-1}\mathbf{A} = \mathbf{I}_{\mathbf{n}}$   $\mathbf{B}\mathbf{B}^{-1} = \mathbf{I}_{\mathbf{n}}$  and  $\mathbf{B}^{-1}\mathbf{B} = \mathbf{I}_{\mathbf{p}}$ Thus,  $\mathbf{B}^{-1}\mathbf{A}^{-1}(\mathbf{A}\mathbf{B}) = \mathbf{B}^{-1}(\mathbf{A}^{-1}\mathbf{A}) \mathbf{B} = \mathbf{B}^{-1} \mathbf{I}_{\mathbf{n}} \mathbf{B} = \mathbf{B}^{-1} \mathbf{B} = \mathbf{I}_{\mathbf{p}}$   $(\mathbf{A}\mathbf{B}) \mathbf{B}^{-1}\mathbf{A}^{-1} = \mathbf{A} (\mathbf{B}\mathbf{B}^{-1}) \mathbf{A}^{-1} = \mathbf{A} \mathbf{I}_{\mathbf{n}} \mathbf{A}^{-1} = \mathbf{A} \mathbf{A}^{-1} = \mathbf{I}_{\mathbf{m}}$  $\Rightarrow \mathbf{A}\mathbf{B}$  is invertible and  $(\mathbf{A}\mathbf{B})^{-1} = \mathbf{B}^{-1}\mathbf{A}^{-1}$ 

<u>Note</u>: It is not possible to divide one matrix by another. That is, we can not write A/B. For two matrices **A** and **B**, the quotient can be written as  $AB^{-1}$  or  $B^{-1}A$ .

• In general, in matrix algebra  $AB^{-1} \neq B^{-1}A$ .

Thus, writing A/B does not clearly identify whether it represents AB-1 or B-1A.

We'll say  $B^{-1}$  post-multiplies A (for  $AB^{-1}$ ) and  $B^{-1}$  pre-multiplies A (for  $B^{-1}A$ )

### **Transpose and Inverse Matrix**

 $(\mathbf{A} + \mathbf{B})' = \mathbf{A}' + \mathbf{B}'$ 

If  $\mathbf{A}' = \mathbf{A}$ , then  $\mathbf{A}$  is called a *symmetric* matrix.

#### **Theorems**:

- Given two conformable matrices A and B, then (AB)' = B'A'
- If A is invertible, then  $(A^{-1})' = (A')^{-1}$  (and A' is also invertible).

### **Properties of Symmetric Matrices**

#### **Definition:**

If  $\mathbf{A}' = \mathbf{A}$ , then  $\mathbf{A}$  is called a *symmetric* matrix.

• In many applications, matrices are often symmetric. For example, in statistics the *correlation matrix* and *the variance covariance matrix*.

• Symmetric matrices play the same role as real numbers do among the complex numbers.

• We can do calculations with symmetric matrices like with numbers: for example, we can solve  $B^2 = A$  for B if A is symmetric matrix (& B is square root of A.) This is not possible in general.

#### **Theorems**:

- If **A** and **B** are  $n \times n$  symmetric matrices, then (AB)' = BA
- If **A** and **B** are nxn symmetric matrices, then (A+B)' = B+A
- If C is any n x n matrix, then  $\mathbf{B} = \mathbf{C}'\mathbf{C}$  is symmetric.
- (*Spectral decomposition*) If **A** is *n*x*n* symmetric matrix, then it can be diagonalized as  $\mathbf{B} = \mathbf{X}^{-1}\mathbf{A}\mathbf{X}$ , with an orthogonal **X**.

• Useful symmetric matrices:

V = X'X  $P = X(X'X)^{-1}X'$   $M = I - P = I - X(X'X)^{-1}X'$  $Var[b] = \sigma^{2} (X'X)^{-1}$ 

**P**: Projection matrix **M**: Residual maker OLS Variance of **b** 

### **Application 1: Linear System**

There is a functional form relating a dependent variable, y, and k explanatory variables,  $\mathbf{X}$ . The functional form is linear, but it depends on k unknown parameters,  $\boldsymbol{\beta}$ . The relation between y and  $\mathbf{X}$  is not exact. There is an error,  $\boldsymbol{\varepsilon}$ . We have T observations of y and  $\mathbf{X}$ .

Then, the data is generated according to:

$$y_i = \sum_{j=1}^k x_{ji} \beta_j + \varepsilon_i$$
  $i = 1, 2, ..., T.$ 

Or using matrix notation:
$\mathbf{y} = \mathbf{X} \boldsymbol{\beta} + \boldsymbol{\varepsilon}$ where  $\mathbf{y} \& \boldsymbol{\varepsilon}$  are (*T*x1);  $\mathbf{X}$  is (*T*x*k*); and  $\boldsymbol{\beta}$  is (*k*x1).

We will call this relation *data generating process* (DGP).

The goal of econometrics is to estimate the unknown vector  $\beta$ .

- Assume an economic model as system of linear equations with:
  - $a_{ij}$  parameters, where i = 1, ..., m rows, j = 1, ..., n columns
  - $x_i$  endogenous variables (*n*),
  - $d_i$  exogenous variables and constants (*m*).

 $\begin{cases} a_{11} x_1 + a_{12} x_2 + \dots + a_{1n} x_n = d_1 \\ a_{21} x_1 + a_{22} x_2 + \dots + a_{2n} x_n = d_2 \\ \vdots & \vdots & \vdots \\ a_{m1} x_1 + a_{m2} x_2 + \dots + a_{mn} x_n = d_m \end{cases}$ 

We can write this system using linear algebra notation:  $\mathbf{A} \mathbf{x} = \mathbf{d}$ 

$$\begin{bmatrix} a_{11} & \cdots & a_{1n} \\ \vdots & \ddots & \vdots \\ a_{m1} & \cdots & a_{mn} \end{bmatrix} \begin{bmatrix} x_1 \\ \vdots \\ x_n \end{bmatrix} = \begin{bmatrix} \mathbf{d}_1 \\ \vdots \\ \mathbf{d}_m \end{bmatrix} \mathbf{d} = \text{column vector}$$
  
$$\mathbf{A} = (mxn) \text{ matrix} \mathbf{A} = \text{column vector}$$

• Summary: System of linear equations:

 $\mathbf{A}\mathbf{x} = d$ 

where

 $\mathbf{A} = (m \mathbf{x} n)$  matrix of parameters

 $\mathbf{x}$  = column vector of endogenous variables (*n*x1)

d = column vector of exogenous variables and constants (mx1)

Solve for x\*.

• Questions:

- For what combinations of **A** and *d* there will zero, one, many or an infinite number of solutions?

- How do we compute (characterize) those sets of solutions?

**Theorem:** Given A (mxn) invertible. Then, the equation Ax = d has one and only one solution for every d(mx1). That is, there is a unique  $x^*$ .

$$\Rightarrow \mathbf{x}^* = \mathbf{A}^{-1} d$$

### Linear dependence and Rank: Example

A set of vectors is *linearly dependent* if any one of them can be expressed as a linear combination of the remaining vectors; otherwise, it is linearly independent.

• Formal definition: Linear independence (LI)

The set  $\{u_1, u_2, ..., u_k\}$  is called a *linearly independent* set of vectors iff

 $c_1 u_1 + c_2 u_2 + \dots + c_k u_k = 0 \implies c_1 = c_2 = \dots = c_k = 0.$ 

Notes:

- Dependence prevents solving a system of equations. More unknowns than independent equations.

- The number of linearly independent rows or columns in a matrix is the *rank* of a matrix  $(rank(\mathbf{A}))$ .

### **Examples:**

(1)  $v'_{1} = \begin{bmatrix} 5 & 12 \end{bmatrix}$   $v'_{2} = \begin{bmatrix} 10 & 24 \end{bmatrix}$   $A = \begin{bmatrix} 5 & 10 \\ 12 & 24 \end{bmatrix} = \begin{bmatrix} v'_{1} \\ v'_{2} \end{bmatrix}$  (a 2x2 matrix)  $2v'_{1} - v'_{2} = 0 \Rightarrow rank(A) = 1 \Rightarrow cannot invert A$ (2)  $v_{1} = \begin{bmatrix} 2 \\ 7 \end{bmatrix}; v_{2} = \begin{bmatrix} 1 \\ 8 \end{bmatrix}; v_{3} = \begin{bmatrix} 4 \\ 5 \end{bmatrix};$   $A = \begin{bmatrix} 2 & 1 & 4 \\ 7 & 8 & 5 \end{bmatrix}$   $3v'_{1} - 2v'_{2} = \begin{bmatrix} 6 & 21 \end{bmatrix} - \begin{bmatrix} 2 & 16 \end{bmatrix}$   $= \begin{bmatrix} 4 & 5 \end{bmatrix} = v'_{3}$  $3v'_{1} - 2v'_{2} - v'_{3} = \mathbf{0} \Rightarrow rank(A) = 2. \P$ 

A matrix **A** has *full row* rank when each of the rows of the matrix are linearly independent and *full column rank* when each of the columns of the matrix are linearly independent. For a square matrix these two concepts are equivalent and we say matrix **A** has full rank.

# **Determinant Test**

We can check if a matrix square matrix A has full rank, that is, all its rows/columns are linearly independent by computing the determinant. If a square matrix A has full rank, it is invertible. That is, the *determinant* of a square matrix A detects whether A is invertible:

If  $det(\mathbf{A}) = 0$  then  $\mathbf{A}$  is not invertible (equivalently, the rows/columns of  $\mathbf{A}$  are linearly dependent).

# Lecture 2 - Introduction: Review, Returns and Data

• All the information and material is on my webpage: https://www.bauer.uh.edu/rsusmel/4397/4397.htm

• Textbook:

Required: Introductory Econometrics for Finance, Cambridge University Press; 4th edition or older, by Chris Brooks.

Recommended: **R Guide for Introductory Econometrics for Finance,** written by Chris Brooks. You can download it from my homepage (pdf format). It's also available for free through Amazon (kindle format).

• Install R in your machine. Many students strongly prefer R Studio. Both will do fine. We will run programs and do some simple programing.

• Two midterms and a final (optional paper for MBA/MS class). There is a project in between midterms.

• Three homework: Two before first Midterm and one before second Midterm.

### **This Class**

This is an applied technical class, with some econometric theory and many stats concepts, followed by related financial applications.

• We will review many math and statistical topics.

• Some technical material may be new to you, for example Linear Algebra. The new material is introduced to simplify the exposition of the main concepts. You will not be required to have a deep understanding of the new material, but you should be able to follow the intuition behind it.

• This is not a programming class, but we will use R to do computation and to estimate models. I will cover some of the basics in class. But, the more you know, the more comfortable you will be running the programs.

• For some students, the class will be dry ("He fried my brain," a student recently wrote.)

### • Main Topics

We will go over a lot of statistics and math tools: tests of hypothesis, bootstrap, linear regression, time series modeling, etc. But keep in mind that the goal of this class is to use statistical tools to understand financial issues. In this class we will try to answer the following questions:

- How do we measure returns and risks of financial assets?

- Can we estimate expected returns with precision? What about the variance of returns?
- Is the equity risk premium (excess returns of stocks over bonds) too high?

- Can we explain asset returns?
- How can one explain variations in stock returns across various stocks?
- Is the CAPM a good model? What about the Fama-French factor models?
- Do we need normality to test financial economics hypothesis or models?
- How do we incorporate structural breaks in our models? Do we have to do it?
- Are asset returns predictable? In the short run? In the long run?
- How do we select a model to forecast asset returns?
- Are markets efficient?

- Does the risk of an asset vary with time? What are the implications? How can one model time-varying risk?

### • Topics Not Covered

This course provides an introduction to the basics of financial econometrics, focusing on estimation of linear models and analysis of time series. There are many more topics in financial econometrics that we will not cover, among them:

- Credit risk management and probability of default
- Interest rate models and term structure models
- Analyzing high-frequency data and modeling market microstructure
- Estimating models for options
- Multivariate time series models
- Technical methods such as state-space models and the Kalman filter, Markov processes, copulae, nonparametric methods, etc.

### What is Econometrics?

Ragnar Frisch, Econometrica Vol.1 No. 1 (1933), defined the field:

"Experience has shown that each of these three view-points, that of *statistics*, *economic theory*, and *mathematics*, is a necessary, but not by itself a sufficient, condition for a real understanding of the quantitative relations in modern economic life.

It is the unification of all three aspects that is powerful. And it is this unification that constitutes econometrics."



**Examples:** We want to estimate the annual expected excess return for Exxon,  $E[r_{XOM} - r_f]$ . - Simple approach: We get monthly XOM return data to compute the average annualized return of XOM, since 1973. We use this average to estimate the annual expected return. Then, we get an annualized **2.81%** estimate. Good, we use data & statistics (the average estimates the expectation).

- More sophisticated approach: Add economic theory. That is, use econometrics. For example, we can use the Capital Asset Pricing Model (CAPM) that states a linear relation, in equilibrium, between excess market returns,  $r_M - r_f$ , & excess returns,  $r_i - r_f$ , for any asset *i*:

$$\mathbb{E}[r_i - r_f] = \beta_i \mathbb{E}[(r_M - r_f)].$$

We get data on  $r_i$ ,  $r_f$ , and  $r_M$ . Then, we use a linear regression to estimate  $\beta_i$ .

• Steps:

(1) Economic Theory: The CAPM:  $E[r_{i=XOM} - r_f] = \beta_i E[(r_M - r_f)]$ 

(2) Data: Collect data, 1973-now for  $r_{XOM}$ ,  $r_f$ , &  $r_M$ .

- (3) Mathematical Statistics: Use a linear regression to estimate  $\beta_i$ :  $r_{XOM} - r_f = \alpha_{XOM} + \beta_{XOM} (r_M - r_f) + \varepsilon_{XOM}$  $\Rightarrow$  Compute **b**<sub>XOM</sub> (the regression estimator of  $\beta_{XOM}$ ), say **0.665**.
- Now, we are ready to compute the expected excess return for XOM: Expected excess XOM return: b<sub>XOM</sub> \* Average(r<sub>M</sub> - r<sub>f</sub>).
   : 0.665 \* 0.0727 = 0.0483 (= 4.83%). ¶

Financial Econometrics is applied econometrics to financial data. That is, we study the statistical tools that are needed to analyze and address the specific types of questions and modeling challenges that appear in analyzing financial data.

Always keep in mind that almost in all cases, financial data is not "experimental data." We have no control over the data. We have to learn how to deal with the usual problems in financial data.

Typical applications of econometric tools to finance:

- Describe data. For example, expected returns & volatility.
- Test hypothesis. For example, are stocks riskier than bonds?
- Build and test models. For example, the different Fama-French factor Models.

In general, in finance we deal with trade-off. The usual trade-off: Risk & Return. Then:

- How do we measure risk and return?
- Can we predict them?
- How do we measure the trade-off?
- How much should I be compensated for taking a given risk?

Thus, we will be concerned with quantifying rewards and risks associated with uncertain outcomes.

• Trade-off application: Fund Management

A fund manager has to allocate money across potentially many different investment alternatives to form portfolios.

At the time of the investment, the fund manager does not know what the return will be on each investment opportunity. (As we will see soon, returns are random variables.)

However, the fund manager can still make good investment decisions.

Q: How? By quantifying the uncertainty associated with all the investment alternatives. For this purpose, the fund manager needs a **model** for the returns of all the different investment alternatives.

From the model, the fund manager gets expected returns, variances & covariances. Using these pieces of information, the fund manager builds a portfolio.

### **This Lecture**

In the first part of the lecture, we review some of the concepts discussed in Lecture 1 (sample statistics, distributions, random variables, descriptive statistics, etc.). In the second part, we go over returns, yields and, then, we start to apply statistical concepts to financial data. We also start to introduce R concepts and to write some R programs.

# **Review – Population and Sample**

#### **Definition:** Population

A population is the totality of the elements under study. We are interested in learning something about this population.

**Examples:** Number of alligators in Texas, percentage of unemployed workers in cities in the U.S., the total return of all stocks in the U.S., the 10-year Japanese government bond yield from 1960-2023.

A Random Variable (RV) X defined over a population is called the population RV. The population RV generates the data. We call the population RV the "*Data Generating Process*," or DGP.

Usually, the population is large, making a complete enumeration of all the values in the population impractical or impossible. Thus, the descriptive statistics describing the population – i.e., the *population parameters*– will be considered unknown.

Typical situation in statistics: we want to make inferences about an unknown population parameter  $\theta$  using a sample –i.e., a small collection of observations from the general population  $\{X_1, X_2, ..., X_N\}$ .

We summarize the information in the sample with a *statistic*, which is a function of the sample. That is, any statistic summarizes the data, or reduces the information in the sample to a single number. To make inferences, we use the information in the statistic instead of the entire sample.



Get a statistic and Make inferences ("learn")

#### **Definition:** Sample

The sample is a (manageable) subset of elements of the population.

**Example:** The total returns of the stocks on the S&P 500 index.

Samples are collected to learn about the population. The process of collecting information from a sample is referred to as *sampling*.

### **Definition:** Random Sample

A *random sample* is a sample where the probability that any individual member from the population being selected as part of the sample is exactly the same as any other individual member of the population.

**Example:** The total returns of the stocks on the S&P 500 index is *not* a random sample.

In mathematical terms, given a random variable X with distribution F, a *random sample* of length N is a set of N independent, identically distributed (*i.i.d.*) random variables with distribution F.

We will estimate population parameters using sample analogues: mean, sample mean; variance, sample variance;  $\beta$ , **b**; etc.

• In general, in finance and economics, we do not deal with random samples. The collected observations will have issues that make the sample not a truly random sample.

# **Review – Samples and Types of Data**

The samples we collect to learn about the population by computing sample statistics are classified in three groups:

- *Time Series Data*: Collected over time on one or more variables, with a particular *frequency* of observation. For example, we record for 10 years the monthly S&P 500 returns, or 10' IBM returns.

<u>Usual notation</u>:  $x_t$ , t = 1, 2, ..., T.

- *Cross-sectional Data*: Collected on one or more variables collected at a single point in time. For example, today we record all closing returns for the members of the S&P 500 index. <u>Usual notation</u>:  $x_i$ , i = 1, 2, ..., N.

- *Panel Data*: Cross-sectional Data collected over time. For example, the CRSP database collects daily prices of all U.S. traded stocks since 1962. <u>Usual notation</u>:  $x_{i,t}$ , i = 1, 2, ..., N & t = 1, 2, ..., T.

The different types of data will present different problems; for example, autocorrelated data is a common problem in time series.

# **Review – Sample Statistic**

A *statistic* (singular) is a single measure of some attribute of a sample (for example, its arithmetic mean value). It is calculated by applying a function (statistical algorithm) to the values of the items comprising the sample, which are known together as a set of data.

Definition: Statistic

A *statistic* is a function of the observable random variable(s), which does not contain any unknown parameters.

**Examples:** Sample mean  $(\overline{X})$ , sample variance  $(s^2)$ , minimum, median,  $(x_1 + x_N)/2$ , etc. ¶

<u>Note</u>: A statistic is distinct from a population parameter. A statistic will be used to estimate a population parameter. In this case, the statistic is called an *estimator*.

### **Review – Population and Sample**

Sample Statistics are used to estimate population parameters.

**Example:**  $\overline{X}$  is an estimate of the population mean,  $\mu$ .

Notation: Population parameters: Greek letters ( $\mu$ ,  $\sigma$ ,  $\theta$ , etc.) Estimators: A hat over the Greek letter ( $\hat{\theta}$ ).

Suppose we want to learn about the mean of IBM annual returns,  $\mu_{\text{IBM}}$ . From the population, we get a sample:  $\{X_{1962}, X_{1963}, \dots, X_{N=2023}\}$ . Then, we compute a statistic,  $\bar{X}$ . As we will see later, on average  $\bar{X}$  is a good estimator of  $\mu$ .



The definition of a sample statistic is very general. For example,  $(x_1 + x_N)/2$  is by definition a statistic; we could claim that it estimates the population mean of the variable X. However, this is probably not a good estimate.

We would like our estimators,  $\hat{\theta}$ , to have certain desirable properties, for example, low bias and low variance, where bias and varianc are defined below:

- Bias[
$$\hat{\theta}$$
] = E[ $\hat{\theta}$ ] -  $\theta$   
- Var[ $\hat{\theta}$ ] = E[( $\hat{\theta}$  - E[ $\theta$ ])<sup>2</sup>]

Ideally, we would like to have  $\hat{\theta}$  with both low bias and low variance, but as we would see later, in general, we have a trade-off between these two properties.

# **Review – Sample Statistic**

Some simple properties for estimators:

- An estimator  $\hat{\theta}$  is *unbiased* estimator of  $\theta$  if  $E[\hat{\theta}] = \theta$ .

- An estimator is *most efficient* if the variance of the estimator is minimized.

- An estimator is BUE, or Best Unbiased Estimate, if it is the estimator with the smallest variance among all unbiased estimates.

- An estimator is *consistent* if as the sample size, *n*, increases to  $\infty$ ,  $\hat{\theta}_n$  converges to  $\theta$ . We write  $\widehat{\theta}_n \xrightarrow{p} \theta$ 

(A LLN is behind this result.)

- An estimator is *asymptotically normal* if as the sample size, *n*, increases to  $\infty$ ,  $\hat{\theta}_n$ , often standardized or transformed, converges in distribution to a Normal distribution. We write

 $\hat{\theta}_n \xrightarrow{d} N(\theta, \operatorname{Var}(\hat{\theta}_n)).$ (A CLT is behind this result.)

The first two properties for estimators hold for samples of any size, not just large samples –i.e., when  $N \rightarrow \infty$ . We associate bias with lack of accuracy and efficiency/variance with uncertainty.

It is common to evaluate an estimator using the Mean Squared Error (MSE), which combines the bias and the variance:

$$MSE[\hat{\theta}] = E[(\hat{\theta} - \theta)^{2}] = Bias[\hat{\theta}]^{2} + Var[\hat{\theta}].$$

### **Review – PDF for a Discrete RV**

**Definition**: Let X be a discrete RV. Let p(x) be a function with the following properties:

1. 
$$0 \le p(x) \le 1$$
  
2.  $\sum_{i=1}^{\infty} p(x_i) = 1$   
3.  $P[a \le X \le b] = \sum_{a \le x \le b} p(x)$ 

Then, p(x) is called the *probability function* or *probability mass function* (pmf) of X. We use p(x) to describe the behavior of a discrete RV.

**Example:** Suppose the discrete RV *X* is the number of days in a week that XOM has a positive return. Using Property 3, we can compute the probability that XOM's has a positive return in 3 or more days in a week:

$$P[a = 3 \le X \le b = 5] = p(x = 3) + p(x = 4) + p(x = 5).$$

#### **Review – PDF for a Continuous RV**

Analogous definition applies for a continuous RV, where the notation uses f(x) instead and the summation sign is replaced by the integral.

**Definition**: Suppose that *X* is a random variable. Let f(x) denote a function defined for  $-\infty < x < \infty$  with the following properties:

1.  $f(x) \ge 0$ 

2. 
$$\int_{-\infty}^{\infty} f(x) dx = 1$$

3. 
$$P[a \le X \le b] = \int_a^b f(x) dx$$

Then, f(x) is called the *probability density function* (pdf) of X. The RV X is called *continuous*.

The pdf is non-negative and integrates to  $\int_{-\infty}^{\infty} f(x) dx = 1$ . The probability that



<u>Remark</u>: We use the pdf to describe the behavior of a continuous RV.

**Example:** Suppose the continuous RV X is IBM's daily stock returns and we know the pdf. Then, using Property 3, we can compute the probability that IBM's daily return is between a = -1.64% and *b*=1.64%:

 $P[-1.64\% \le X \le 1.64\%] = \int_{a=-1.64}^{b=1.64} f(x) dx$  (the red area in the above graph).

### **Review – Popular PDFs: Normal Distribution**

A RV X is said to have a normal distribution with parameters  $\mu$  (mean) and  $\sigma^2$  (variance) if X is a continuous RV with pdf f(x):

$$f(x) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{(x-\mu)^2}{2\sigma^2}\right]$$

Note: Described by two parameters:  $\mu$  and  $\sigma^2$ . We write  $X \sim N(\mu, \sigma^2)$ 

When  $\mu = 0$  and  $\sigma^2 = 1$ , we call the distribution *standard normal*. We write  $X \sim N(0, 1)$ . This is the distribution that is tabulated.

The normal distribution is often used to describe or approximate any variable that tends to cluster around the mean. It is the most assumed distribution in economics and finance: rates of return, growth rates, IQ scores, observational errors, etc.

The central limit theorem (CLT) provides a justification for the normality assumption when the sample size, *n*, is large.

PDF:  $X \sim N(\mu, \sigma^2)$ Notation: CDF:  $\Phi(x)$ 

# **Review – Popular PDFs: Gamma Distribution**

Let the continuous RV *X* have density function):

$$f(x) = \begin{cases} \frac{\lambda^{\alpha}}{\Gamma(\alpha)} x^{\alpha-1} e^{-\lambda x} & x \ge 0\\ 0 & x < 0 \end{cases}$$

where  $\alpha$ ,  $\lambda > 0$  and  $\Gamma(\alpha)$  is the gamma function evaluated at  $\alpha$ .

Then, X is said to have a *Gamma distribution* with parameters  $\alpha$  and  $\lambda$ , denoted as  $X \sim$  Gamma( $\alpha$ ,  $\lambda$ ) or  $\Gamma(\alpha, \lambda)$ .

It is a family of distributions, with special cases:

- Exponential Distribution, or  $Exp(\lambda)$ :  $\alpha = 1$ .
- Chi-square Distribution, or  $\chi_{\nu}^2$ :  $\alpha = \nu/2$  and  $\lambda = \frac{1}{2}$ .

The Chi-square distribution,  $\chi_{\nu}^2$ , will appear a lot in this class, since it is derived from a sum of independent square standard normals. It is the distribution of many popular test statistics. Below we plot the Chi-square distribution with parameter v, which we refer as degrees of freedom:



### **Review – Popular PDFs: Other Distributions**

Other distributions that we will use in this class: the t-distribution and the F-distribution.

#### (1) t-distribution

The t-distribution is the ratio of a standard normal and the square root of a Chi-squared distribution, divided by its degrees of freedom. That is, let  $Y \sim N(0, 1)$  and  $W \sim \chi_{\nu}^2$ , then  $t = \frac{Y}{\sqrt{W/\nu}} \sim t_{\nu}$ .

The t-distribution is indexed by its degrees of freedom. Below, we plot a simulated t-distribution with  $\nu = 5$  (in red), along a normal distribution (in blue). It looks like a normal distribution, but with thicker tails. As  $\nu$  increases, the t-distribution converges to a standard normal distribution.



#### (2) F-distribution

The F-distribution is the ratio of two independent Chi-squared distributions, divided by their degrees of freedom. That is, let  $Z_1 \sim \chi^2_{\nu_1}$  and  $Z_2 \sim \chi^2_{\nu_2}$ , then

$$\mathbf{F} = \frac{Z_1 / \nu_1}{Z_2 / \nu_2} \sim F_{\nu_1, \nu_2}$$

The F distribution is indexed by two degrees of freedom, informally referred as "numerator and denominator degrees of freedom."



We will use both distributions in the context of testing null hypothesis.

<u>Note</u>:  $t^2 \sim F_{1,\nu}$ .

#### **Review – CDF for a Continuous RV**

If X is a continuous random variable with probability density function, f(x), the *cumulative distribution function* (CDF) of *X* is given by:



The plot of F(x) is:





**Example**: Let *X* above, in the previous graph, be the daily IBM's returns, then, the probability that IBM's daily returns are 1.645% or less is 95% (red area in the first graph).

### **Review – The Empirical Distribution**

The empirical distribution (ED) of a dataset is simply the distribution that we observe in the data.

The ED is a discrete distribution that gives equal weight to each data point, assigning a 1/N probability to each of the original N observations.

We form a cumulative distribution function,  $F^*$ , that is a step function that jumps up by 1/N at each of the N data point:

 $F^*(x) = 1/N \sum_{i=1}^N I(x_i \le x),$ where I(.) is the indicator function:  $I(x_i \le x) = 1,$  if  $x_i \le x$ 

 $I(x_i \le x) = 0, \qquad \text{if } x_i \ge x$   $I(x_i \le x) = 0, \qquad \text{if } x_i > x$ 

**Example:** We throw 100 times two dice and sum the results. The CDF is given below:



In general, we use a histogram to describe the ED of a dataset.

<u>Important result</u>: Let *F* be the true distribution of the data and  $F^*$  be the ED of the data. As  $N \rightarrow \infty$ , the Law of large numbers (LLN) tells us that  $F^*$  becomes a good approximation of *F*.

### **Review – Histogram of a RV**

Recall that a *histogram* is an approximate representation of the distribution of numerical data.

**Example:** We use a histogram to estimate the distribution of a RV. Below we display two histograms. For the first histogram, let X = Percentage changes in the CHF/USD exchange rate  $= e_f$ ; while for the second one, let X = MSCI USA Index returns.

Data: For the CHF/USD exchange rate, we have monthly data from January 1971 to March 2024 (N= 6155 observations) and for the MSCI USA returns we have monthly data from January 1970 to June 2020 (N=607).





Distribution - Histogram for MSCI USA Returns



Note: We overlay a Normal density (blue line) over the histograms.

# **Review – Moments of Random Variables**

The moments of a random variable *X* are used to describe the behavior of the RV (discrete or continuous).

**Definition**:  $K^{th}$  Moment Let X be a RV (discrete or continuous), then the  $k^{th}$  moment of X is:

$$\mu_{k} = E(X^{k}) = \begin{cases} \sum_{x} x^{k} p(x) & \text{if } X \text{ is discrete} \\ \int_{-\infty}^{\infty} x^{k} f(x) dx & \text{if } X \text{ is continuous} \end{cases}$$

**Example:** Suppose *X* is the number of days in a week that XOM has a positive return. We want to know the first moment, the mean, of the distribution. That is,

 $\mu_1 = \sum_x x \, p(x) = 1 * p(x = 1) + 2 * p(x = 2) + 3 * p(x = 3) + 4 * p(x = 4) + 5 * p(x = 5)$ 

Suppose we can describe X with a Binomial distribution, with p=0.52, that is, we sssume that XOM has a 52% probability of having a positive return. Then,

 $\mu_1 = 0 * 0.0255 + 1 * 0.1380 + 2 * 0.2990 + 3 * 0.3240 + 4 * 0.1755 + 5 * 0.0380 = 2.60$ 

Interpretation: The expected number of days in week with positive returns for XOM is 2.6 days.

Note: For a continuous RV, we need to integrate to get moments.

#### **Definition:** Central Moments

Let X be a RV (discrete or continuous). Then, the  $k^{th}$  central moment of X is defined to be:

$$\mu_k^0 = E[(X - \mu)^k] = \begin{cases} \sum_x (x - \mu)^k p(x) & \text{if } X \text{ is discrete} \\ \int_{-\infty}^{\infty} (x - \mu)^k f(x) dx & \text{if } X \text{ is continuous} \end{cases}$$

where  $\mu = \mu_1 = E(X)$  = the first moment of *X*.

The central moments describe how the probability distribution is distributed about the center of gravity, m.

The first central moments is given by:

$$\mu_1^0 = E[X - \mu]$$

The second central moment depends on the *spread* of the probability distribution of *X* about *m*. It is called the variance of *X* and is denoted by the symbol  $\sigma^2 = var(X)$ :

$$\mu_2^0 = E[(X - \mu)^2] = \operatorname{var}(X) = \sigma^2$$

The square root of var(X) is called the *standard deviation* of *X* and is denoted by the symbol s = SD(X). We also refer to it as *volatility*:

$$\sqrt{\mu_2^0} = \sqrt{E[(X-\mu)^2]} = \sigma$$

**Example:** Suppose X is the number of days in a week that XOM has a positive return. We want to know the second central moment,  $\mu_2^0 = \sigma^2$  (& volatility), and the third central moment,  $\mu_3^0$ . (Recall that  $\mu_1 = \mu = 2.6$  days)

Then, the second central moment is given by:

 $\mu_2^0 = \sigma^2 = \sum_x (x - \mu)^2 p(x) = (0 - 2.6)^2 * p(x = 0) + (1 - 2.6)^2 * p(x = 1)$  $+ (2 - 2.6)^2 * p(x = 2) + (3 - 2.6)^2 * p(x = 3)$  $+ (4 - 2.6)^2 * p(x = 4) + (5 - 2.6)^2 * p(x = 5)$ 

Again, assume X follows a Binomial distribution, with p=0.52. Then,  $\sigma^{2} = (0 - 2.6)^{2} * 0.0255 + (1 - 2.6)^{2} * 0.1380 + (2 - 2.6)^{2} * 0.2990 + (3 - 2.6)^{2} * 0.3240 + (4 - 2.6)^{2} * 0.1755 + (5 - 2.6)^{2} * 0.0380 = 1.24802 \implies \sigma = \operatorname{sqrt}(1.24802) = 1.117148$ 

Interpretation: The volatility of *X* is 1.12 days.

Then, the second central moment is given by:  $\mu_3^0 = (0 - 2.6)^3 * 0.0255 + (1 - 2.6)^3 * 0.1380 + (2 - 2.6)^3 * 0.2990 + (3 - 2.6)^3 * 0.3240 + (4 - 2.6)^3 * 0.1755 + (5 - 2.6)^3 * 0.0380 = -0.04989$ 

Note: Again, for a continuous RV, we need to integrate to get central moments.

#### **Review – Moments of a RV: Skewness**

The third central moment:

 $\mu_3^0 = E[(X - \mu)^3]$ 

 $\mu_3^0$  contains information about the *skewness* of a distribution. We use skewness as a gauge of symmetry. If  $\mu_3^0 = 0$  the distribution is symmetric; otherwise, asymmetric.

A popular measure of skewness:

$$\gamma_1 = \frac{\mu_3^0}{\sigma^3} = \frac{\mu_3^0}{(\mu_2^0)^{\frac{3}{2}}}$$

• Distribution according to skewness:

1) Symmetric distribution



2) Positively (right-) skewed distribution (with mode < median < mean)



3) Negatively (left-) skewed distribution (with mode > median > mean)





**Example (continuation):** Now, we can compute  $\gamma_1$  for *X*, the number of days in a week that XOM has a positive return. Recall  $\mu_2^0 = 1.24802$  and  $\mu_3^0 = -0.04989$ , then,

$$\gamma_1 = \frac{\mu_3^0}{\sigma^3} = \frac{-0.04989}{(1.11715)^3} = -0.03578522$$

Interpretation: *X* has a small, but negative skewness. The left tail is a little bit longer.

• Skewness and Economics

For changes in asset prices:

- Zero skew means symmetrical gains and losses –i.e., extreme values tend to occur on both sides of the curve on similar proportions.

- Positive skew suggests many small losses and few rich returns –i.e., extreme values tend to occur in the right tail

- Negative skew indicates a lot of minor wins offset by rare major losses –i.e., extreme values tend to occur in the left tail.

In financial markets, stock returns at the firm level show positive skewness, but at the aggregate (index) level show negative skewness.

From horse race betting and from U.S. state lotteries there is evidence supporting the contention that gamblers are not necessarily risk-lovers but skewness-lovers: Long shots are overbet (positive skewness loved!).

### **Review – Moments of a RV: Kurtosis**

The fourth central moment:

 $\mu_4^0 = E[(X - \mu)^4]$ 

It contains information about the *shape* of a distribution. The property of shape that is measured by this moment is called *kurtosis*, usually estimated by  $\kappa$ :

$$\kappa = \frac{\mu_4^0}{\sigma^4}.$$

Kurtosis measures how much weight there is in the tails of the distribution relative to the middle (we call this a measure of the "*fatness*" of the tails). We usually compare the kurtosis of a series relative to the kurtosis of a normal distribution, which is equal to 3. We measure the "excess" fatness of the tail over the normal curve. That is, the *measure of (excess) kurtosis*:

$$\gamma_2 = \frac{\mu_4^0}{\sigma^4} - 3 = \frac{\mu_4^0}{(\mu_2^0)^2} - 3$$

• Distributions:

1) Mesokurtic distribution



2) Platykurtic distribution



3) Leptokurtic distribution (usual shape for asset returns)



• Positive excess kurtosis,  $\gamma_2 > 0$ , is the norm for financial returns. Below I simulate a series with  $\mu=0$ ,  $\sigma=1$ , zero skewness & kurtosis = 6 ( $\gamma_2=3$ ), overlaid with a standard normal distribution. Fat tails are seen on both sides of the distribution.



#### **Review – Moments and Expected Values**

Note that moments are defined by expected values. We define the expected value of a function of a continuous RV X, g(X), as

$$E[g(X)] = \int_{-\infty}^{\infty} g(x) f(x) dx$$

If *X* is *discrete* with probability function p(x) $E[g(X)] = \sum_{x} g(x)p(x) = \sum_{i} g(x_{i})p(x_{i})$ 

**Examples:**  $g(x) = (x - \mu)^2 \implies E[g(x)] = E[(x - \mu)^2]$  $g(x) = (x - \mu)^k \implies E[g(x)] = E[(x - \mu)^k]. \P$ 

We estimate expected values with sample averages. As we will see below, the Law of Large Numbers (LLN) tells us they are *consistent* estimators of expected values.

### **Review – Estimating Moments**

We estimate expected values with sample averages. For example, the first moment, the mean, and the second central moment, the variance, are estimated by:

$$\bar{X} = \frac{\sum_{i=1}^{N} X_i}{N}$$

$$s^2 = \frac{\sum_{i=1}^{N} (X_i - \bar{X})^2}{N - 1}$$
(N - 1 adjustment needed for E[s<sup>2</sup>] =  $\sigma^2$ )

They are both *unbiased* estimators of their respective population moments (unbiased = "on average, I get the population parameter"). That is,

 $E[\overline{X}] = \mu$  "\mu is the population parameter of interest"  $E[s^2] = \sigma^2$  "\sigma^2 is population parameter of interest"

### **Review – Law of Large Numbers (LLN)**

Long history: Gerolamo Cardano (1501-1576) stated it without proof. Jacob Bernoulli published a rigorous proof in 1713.

#### Theorem (Weak LLN)

Let  $X_1, \ldots, X_N$  be N mutually independent random variables each having mean m and a finite s - i.e, the sequence  $\{X_N\}$  is *i.i.d*.

Let 
$$\overline{X} = \frac{\sum_{i=1}^{N} X_i}{N}$$
.

Then, for any  $\delta > 0$  (no matter how small)

 $P[|\overline{X} - \mu| < \delta] = P[|\mu - |\delta| < \overline{X} < \mu + \delta] \to 1, \qquad \text{as } N \to \infty$ 

There are many versions of the LLN. It is a general result: A sample average as the sample size goes to infinite tends to its expected value. Also written as:

 $\bar{X}_N \xrightarrow{p} \mu$ . (convergence in probability)

As an illustration, below, we randomly generate 10,000 observations from a N(0, 1) and compute the sample mean as a function of N. As expected, as N increases, the sample mean gets closer to the population mean (m = 0).



#### Simulated LLN - Mean as a function of N

#### **Review – Central Limit Theorem (CLT)**

Let  $X_1, X_2, ..., X_N$  be a sequence of *i.i.d.* RVs with finite mean *m*, and finite variance  $s^2$ . Then as *N* increases,  $\overline{X}_N$ , the sample mean, approaches the normal distribution with mean  $\mu$  and variance  $s^2/N$ .

This theorem is sometimes stated as

$$\frac{\sqrt{N(X-\mu)}}{\sigma} \stackrel{d}{\to} N(0,1)$$

where  $\stackrel{a}{\rightarrow}$  means "the limiting distribution (asymptotic distribution) is" (or *convergence in distribution*).

Many versions of the CLT. This one is the Lindeberg-Lévy CLT.

The CLT gives only an asymptotic distribution. We usually take it as an approximation for a finite number of observations. In these cases, the notation goes from  $\stackrel{d}{\rightarrow}$  to  $\stackrel{a}{\rightarrow}$ .

#### **Review – Sampling Distributions**

All statistics, T(X), are functions of RVs and, thus, they have a distribution. Depending on the sample, we can observe different values for T(X), thus, the finite sample distribution of T(X) is called the *sampling distribution*.

• For the sample mean,  $\overline{X}$ , if the  $X_i$ 's are normally distributed, then the sampling distribution is normal with mean  $\mu$  and variance  $\sigma^2/N$ . Or

$$\overline{X} \sim N(\mu, \sigma^2/N).$$

<u>Note</u>: If the data is not normal, the CLT is used to approximate the sampling distribution by the asymptotic one, usually after some manipulations. Again, in those cases, the notation goes from  $\stackrel{d}{\rightarrow}$  to  $\stackrel{a}{\rightarrow}$ .

The SD of the sampling distribution is called the *standard error* (SE). Then,  $SE(\overline{X}) = \sigma/sqrt(N)$ .

**Example:** We plot a Sampling Distribution for the sample mean,  $\overline{X}$ , of a normal population, as a function of the sample size (*N*). For this purpose, we generate 10,000 samples from a N(2, 4) population. We plot the distribution of  $\overline{X}$  for three sizes of N = 10, 60 & 200:



<u>Note</u>: As  $N \to \infty$ ,  $\overline{X} \to \mu$  –i.e., the distribution becomes a spike at  $\mu=2!$ 

• For the sample variance  $\sigma^2$ , if the  $X_i$ 's are normally distributed, then the sampling distribution is derived from this result:

$$(N-1) s^2/\sigma^2 \sim \chi^2_{N-1}$$

We use the properties of a  $\chi_k^2$  to derive the mean & variance of  $s^2$ :

Property 1. Let  $Z \sim \chi_k^2$ . Then, E[Z] = k. Property 2. Let  $Z \sim \chi_k^2$ . Then, Var[Z] = 2 \* k. Application:  $(N - 1) s^2/\sigma^2 \sim \chi_{N-1}^2$ From Property 1:  $E[(N - 1) s^2/\sigma^2] = N - 1$   $\Rightarrow E[s^2] = \sigma^2$ From Property 2:  $Var[(N - 1) s^2/\sigma^2] = 2 * (N - 1)$  $\Rightarrow Var[s^2] = 2 * \sigma^4/(N - 1)$ .

$$\Rightarrow SE(s^2) = SD(s^2) = \sigma^2 * sqrt[2/(N-1)].$$

Summary for  $s^2$ :

Sampling distribution: $(N-1) s^2/\sigma^2 \sim \chi^2_{N-1}$ .Mean: $E[s^2] = \sigma^2$ Variance: $Var[s^2] = 2 * \sigma^4/(N-1)$ .

<u>Note</u>: If the data is not normal (& N is large), the CLT can be used to approximate the sampling distribution by the asymptotic one:

 $s^2 \xrightarrow{a} N(\sigma^2, \sigma^4 * (\kappa - 1)/N)$ where  $\kappa = \frac{\mu_4^0}{\sigma^4}$  (recall when data is normal,  $\kappa = 3$ ).

**Example:** We plot a Sampling Distribution for the sample variance,  $s^2$ , of a normal population, as a function of the sample size (*N*). Above, we generated 10,000 samples from a N(2, 4) population. Now, we plot the distribution of  $s^2$  for three sizes of N = 10, 60 & 200:



<u>Note</u>: As  $N \to \infty$ , the distribution of  $s^2$  looks more Normal – the CLT at work!

### **Review – Estimating Moments in R**

First, we need to import the data. In R, we use the **read** function, usually followed by the type of data we are importing. Below, we import a *comma separated values* (csv) file with monthly data for the S&P Composite Index (P), Dividends (D), Earnings (E), CPI, Long interest rates (Long\_i), and some transformations of the data (Real Prices, Real Dividends, Real Returns, etc). We use the **read.csv** function:

Sh\_da <- **read.csv**("https://www.bauer.uh.edu/rsusmel/4397/Shiller\_2021\_m\_data.csv", head=TRUE, sep=",")

To check the names of the variables we imported, we use the **names()** function. It describes the headers of the file imported (41 headers):

> names(Sh_da	a)					
[1] "Date"	"P"	"D"	"E"	"CPI"	"Fraction"	"Long_i"
[8] "Real_P"	"Real	_D" "]	Real_Pd"	"Real_E"	"Scaled_Real_	P"

The **summary()** function provides some stats of variables imported: > summary(Sh\_da)

2 mining (2 m_				
Date	Р	D	E	CPI
Min. :1871	Min. : 2.73	Min. : 0.180	Min. : 0.16	Min. : 6.28
1st Qu.:1908	1st Qu.: 7.89	1st Qu.: 0.420	1st Qu.: 0.56	1st Qu.: 10.19
Median :1946	Median : 17.35	Median : 0.870	Median : 1.45	Median : 20.30
Mean :1946	Mean : 321.51	Mean : 6.732	Mean : 15.15	Mean : 62.05
3rd Qu.:1983	3rd Qu.: 163.25	3rd Qu.: 7.053	3rd Qu.: 14.71	3rd Qu.:101.25
Max. :2021	Max. :4358.13	Max. :59.680	Max. :139.47	Max. :270.80

• Second, we extract from the imported data, Sh\_da, the column corresponding to the i\_10 and for, later use, the S&P 500 Index (SP):That is, we extract from Sh\_da, the column corresponding to the 10-year interest rate (Long\_i) and, for later use, the S&P Composite Index (P):

SP <- Sh_da\$P	# Extract P = S&P500 series
i_10 <- Sh_da\$Long_i	<pre># Extract Long_i = Interest rates</pre>
N <- length(SP)	# Length of data

• Then, we estimate the sample moments for.

x <- i_10	# Series to be analyzed
$n \leq - length(x)$	# Number of observations
$m1 \leq sum(x)/n$	# Mean $(\overline{X})$
$m2 \le sum((x-m1)^2)/n$	# Used in denominator of both
$m3 \le sum((x-m1)^3)/n$	# For numerator of S
$m4 <- sum((x-m1)^4)/n$	# For numerator of K
$b1 <- m3/m2^{(3/2)}$	# Sample Skewness (γ1)
$b2 \le (m4/m2^2)$	# Sample Kurtosis ( $\gamma_2$ )
$s2 \le sum((x-m1)^2)/(n-1)$	# Sample Variance $(s^2)$
sd_s <- sqrt(s2)	# Sample SD (s)
• R output:	
> m1	# Sample mean (4.51% annual)
[1] 4.509972	
> s2	# Sample Variance
[1] 5.306247	
> sd_s	# Sample SD (2.30% annual)
[1] 2.303529	
> b1	# Sample Skewness
[1] 1.795057	
> b2	# Sample Kurtosis

#### [1] 6.751023

• Table 2.A presents a summary of the moments:

Statistic	e <sub>f</sub>
Mean	4.51
Median	3.82
Maximum	15.32
Minimum	0.62
Std. Dev.	2.30
Skewness	1.7951
Kurtosis	6.7510

### Table 2.A – 10-year Bond Rate ("Long interest Rate"): (1871: February – 2021: September)

Interest rates are right skewed and have kurtosis greater than 3, pointing out to non-normality of data ("*fatter tails*"):

$$\Rightarrow \gamma_2 = \frac{\mu_4^0}{\sigma^4} - 3 = 3.7510. \P$$

#### Returns

Returns have better statistical properties than prices, as we will mention below, returns have a well-defined (long-run) mean and variance, while asset prices, in general, do not. Thus, financial models tend to focus on returns. The return is the profit rate of holding an asset from time t - 1 to t.

We define net or simple (total) return,  $R_t$ , as:

$$R_t = \frac{(P_t - P_{t-1}) + D_t}{P_{t-1}} = \text{capital gain} + \text{dividend yield}$$

where  $P_t$  = Stock price or Value of investment at time t  $D_t$  = Dividend or payout of investment at time t

<u>Note</u>: This is the return from time t-1 to time t. To be very explicit we can write this as  $R_{t-1,t}$ .

Then, the gross (total) return is given by:

$$R_t + 1 = \frac{P_t + D_t}{P_{t-1}}$$

In general, when the word "total return" is used in the definition, it means "*returns including dividends*." Sometimes, total returns are also called "*overall returns*."

If  $D_t = 0$ , the total return is just the capital gain. In this situation, it is common to just use the word *returns*.

• There is another commonly used definition of return, the *log return*,  $r_t$ , defined as the log of the gross return:

 $r_t = \log(1 + R_t) = \log(P_t + D_t) - \log(P_{t-1})$ 

<u>Note</u>: When the values are small (-0.1 to +0.1), the two returns are approximately the same:  $r_t \approx R_t$ . In general –i.e., when returns are not small,  $r_t < R_t$ .

Derivation:

Recall:  $\ln(1) = 0$ , &  $\frac{\delta \ln(x)}{\delta x} = \frac{1}{x}$ .

Now do a 1<sup>st</sup>-order Taylor expansion around  $x_0$  to get

$$\log(x) \approx \log(x_0) + \frac{\delta \log(x)}{\delta x} |_{x_0} * (x - x_0) = \log(x_0) + \frac{1}{x_0} (x - x_0)$$

Thus, expanding around  $x_0 = 1$ , we have for  $x \approx 1$ :

$$\log(x) \approx 0 + \frac{1}{1}(x - 1) = x - 1$$

Set  $x = (1 + R_t)$  to get the result.

The log return is also called *continuously compounded return*.

When returns are small, say for daily or weekly data, the numerical differences between simple and compounded returns are very small.

In this class, we will use log returns.

**Example:** We estimate sample averages for  $e_f = \log$  returns for the CHF/USD. Note that there is no dividends or payouts for holding currency. That is, in this case, returns = capital gains.

PPP\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/ppp\_2020\_m.csv",head=TRUE,sep=",")

x\_chf <- PPP\_da\$CHF\_USD

• Now, we define  $e_f = \log returns$  ( $\approx \%$  changes) for the CHF/USD.

$T \leq -length(x_chf)$	# Size of series read ( $T$ or $N$ notation is OK)
$e_chf \le log(x_chf[-1]/x_chf[-T])$	# Log returns

• Then, we estimate the sample moments for.

$x \le e_chf$	# Series to be analyzed
$N \le -length(x)$	# Number of observations
$m1 \leq sum(x)/N$	# Mean $(\overline{X})$
$m2 \le sum((x - m1)^2)/N$	# Used in denominator of both
$m3 \le sum((x - m1)^3)/N$	# For numerator of S
$m4 \le sum((x - m1)^4)/N$	# For numerator of K
$b1 \le m3/m2^{(3/2)}$	# Sample Skewness (γ <sub>1</sub> )
$b2 \le (m4/m2^2)$	# Sample Kurtosis ( $\gamma_2$ )
$s2 \le sum((x - m1)^2)/(N-1)$	# Sample Variance $(s^2)$
sd_s <- sqrt(s2)	# Sample SD ( <i>s</i> )
• R output:	
> m1	
[1] -0.002550636	
> s2	
[1] 0.001115257	
> sd_s	
[1] 0.03339546	
> b1	# Sample Skewness (y1)
[1] -0.06733514	
> b2	# Sample Kurtosis ( $\kappa$ )
[1] 4.621602	

• Summary of moments of  $e_f = \%$  changes in the CHF/USD exchange rate (1971:Jan – 2020:Jun):

Statistic	ef	
Mean	-0.002551	
Median	-0.001431	
Maximum	0.145542	
Minimum	-0.145639	
Std. Dev.	0.033395	
Skewness	-0.067335	
Kurtosis	4.621602	

Small mean (-0.25%), slight negative skewness, kurtosis greater than 3, pointing out to *fatter tails*:

$$\Rightarrow \gamma_2 = \frac{\mu_4^0}{\sigma^4} - 3 = 1.62. \P$$

#### **Portfolio Returns**

For portfolios, the simple rate of return has an advantage: The rate of return on a portfolio is the portfolio of the rates of return.

Let  $V_{P,t}$  be the value of a portfolio at time t.

$$V_{P,t} = \sum_{i=1}^{N} N_i P_{i,t}$$

where  $N_i$  is the investment in asset *i*, which has a value  $P_{i,t}$  at time t. Then, the return is:

$$R_{P,t} = \frac{V_{P,t} - V_{P,t-1}}{V_{P,t-1}} = \frac{\sum_{i=1}^{N} N_i P_{i,t} - \sum_{i=1}^{N} N_i P_{i,t-1}}{\sum_{i=1}^{N} N_i P_{i,t-1}} = \sum_{i=1}^{N} w_i r_{i,t}$$

where  $w_i$  is the portfolio weight in asset *i*.

This relationship does not hold for log returns because the log of a sum is not the sum of the logs.

#### **Multi-period holding return**

To simplify notation, include dividends into prices. That is,  $P_{d,t+1} = P_{t+1} + D_t$ 

The two-period holding return is:

$$R_{t,t+2} = \frac{P_{d,t+2}}{P_{d,t}} - 1 = \frac{P_{d,t+2}}{P_{d,t+1}} * \frac{P_{d,t+1}}{P_{d,t}} - 1 = (1 + R_{t+1,t+2}) * (1 + R_{t,t+1}) - 1$$

Or

$$1 + R_{t,t+2} = (1 + R_{t+1,t+2}) (1 + R_{t,t+1})$$

For small returns, we can use the log approximation:

$$r_{t,t+2} \approx r_{t,t+1} + r_{t+1,t+2}$$

The *k*-period gross holding return  $(1 + R_{t,t+k}) = \prod_{j=0}^{k-1} (1 + R_{t+j,t+j+1})$ 

Or with the log approximation:  $r_{t,t+k} = \sum_{j=0}^{k-1} r_{t+j,t+j+1}$ 

If the (expected) returns are equal, that is,  $R_{t+j,t+j+1} = R$ . Then, the log approximation produces:

$$r_{t,t+k} = \sum_{j=0}^{k-1} r_{t+j,t+j+1} = k * r$$

If the returns,  $R_{t+j,t+j+1}$ , are independent (covariance is 0) and with a constant variance equal to  $\sigma_r^2$  (a constant), then under the log approximation

$$\operatorname{Var}(r_{t,t+k}) = \sum_{j=0}^{k-1} \operatorname{var}(r_{t+j,t+j+1}) = k * \sigma_r^2$$

Then, the SD is equal to

$$SD(r_{t,t+k}) = \sqrt{k * \sigma_r^2} = \sqrt{k} * \sigma_r$$

# **Real returns**

We will deflate values by a Price Index, for example the CPI. Then,

Real Price<sub>t</sub> = 
$$P_t^{real} = \frac{P_t}{CPI_t}$$

Then, the real return becomes:

$$R_t^{real} = \frac{P_t^{real}}{P_{t-1}^{real}} - 1 = \frac{\frac{P_t}{CPI_t}}{\frac{P_{t-1}}{CPI_{t-1}}} - 1 = \frac{P_t}{P_{t-1}} * \frac{CPI_{t-1}}{CPI_t} - 1 = \frac{(1+R_t)}{(1+\pi_t)} - 1$$

where  $\pi_t$  is the inflation rate at time t.

The log approximation (for small returns) produces  $r_t^{real} \approx r_t - \pi_t$ 

**Example:** Below, we plot the **long-run S&P Composite Index** monthly data, nominal and real. Data taken from Robert Shiller's website (1871:Jan - 2021: Sep).

Sh_da <- read.csv("http://www.bauer.uh.edu	u/rsusmel/4397/Shiller_2021_m_data.csv",
head=TRUE, sep=",")	
SP <- Sh_da\$P	# Extract SP = S&P Composite series
D <- Sh_da\$D	# Extract D = S&P Dividends series
CPI <- Sh_da\$CPI	# Extract CPI = Price Index series
R_SP <- Sh_da\$Real_P	# Extract R_SP = Real S&P500 series
i_10 <- Sh_da\$Long_i	# Extract Long_i = Interest rates
$lr \le log(SP[-1]/SP[-T])$	# Define log returns
$lr_R \le log(R_SP[-1]/R_SP[-T])$	# Define log Real returns
plot(SP, col="blue",ylab ="S&P Index", typ lines(R_SP, col="red") title("S&P Composite Index: Nominal & Re legend("topleft", legend = c("Nominal", "Re lwd = 3, col = c("blue", "red"))	e="l", xlab ="Date") eal (1871-2021)") eal"),



<u>R Note</u>: A more elegant plot, with dates, can be done with the package ggplot2. You need to install it first with install.packages("ggplot2"). Then, you need to call it, with:

```
library(ggplot2)
ggplot(data = Sh_da, aes(x = Date)) +
geom_line(aes(y = SP, color = "Nominal")) +
geom_line(aes(y = R_SP, color = "Real")) +
scale_colour_manual("",
            breaks = c("Nominal", "Real"),
            values = c("Nominal"="blue", "Real"="red")) +
scale_y_continuous("S&P Index", limits = c(0,4500)) +
labs(title = "S&P Composite Index: Nominal and Real",
        subtitle = "Period: January 1871: September 2021")
```



**Long-run S&P 500 monthly log returns**, from Robert Shiller's website (1871:Jan -2020: Sep; T = N = 1805).

 $T \leq - length(SP)$ 



Prices have a clear trend, returns do not. In statistics, we prefer to work with data with no trends, like returns; they have better properties, for example, a well defined *long-run mean* (expected value).



Distribution of S&P 500 monthly log returns



<u>Note</u>: We observe slight negative skewness –left tail events are more common (steeper) than right tail events and fatter tails than the normal distribution. These two features are part of the *"stylized facts"* for stock (index) returns.

**Example:** Table 2.B reports univariate statistics for Shiller's monthly **S&P 500 returns (total, capital gains & real)** and **U.S. inflation**.

			-	-
	Total Return	Capital Gains	Inflation	Real Total Return
Mean	0.007378	0.003801	0.001707	0.005615
Median	0.009928	0.006756	0.001432	0.009367
Maximum	0.414151	0.407459	0.068054	0.421504
Minimum	-0.30365	-0.30753	-0.06805	-0.30364
Std. Dev.	0.040455	0.040650	0.010344	0.040726
Skewness	-0.47050	-0.51512	-0.14404	-0.36439
Kurtosis	14.61046	14.35809	9.81997	14.33933
Jarque-Bera	10205.0	9782.2	3504.3	9719.3
P-value (JB)	2.2e-16	2.2e-16	2.2e-16	2.2e-16

Table 2.B – S&P Composite Returns (Total,	<b>Capital Gains and Real) and Inflation Rate:</b>
(1871: February –	2021: September)

Monthly total returns are slightly (left-) skewed (& median > mean), and with "*fat tails*"–i.e., kurtosis is higher than 3.

• Check some results from log approximation:

(1)  $r_t^{real} \approx r_t - \pi_t$  (for small % changes)  $0.005615 \approx 0.007378 - 0.001707 = 0.005671$ 

(2) Multiperiod return: k = 12 –i.e., from monthly to annual,

- annual return = 0.007378 \* 12 = 0.088531 (8.85%)

- annual SD = 
$$0.040455 * \text{sqrt}(12) = 0.14014$$
 (14.01%)

<u>Note</u>: Compounding the return:  $(1 + 0.007378)^{12} - 1 = 0.0922$ .

According to the annualized numbers from the above Table, since 1871, the average total stock market return has been 8.85% per year.

#### **Returns: Sample Moments – Changing Frequency**

Assuming independence of returns and constant moments, we can use the log returns to easily change frequencies for the mean and variance of returns.

Suppose we have compounded data in base frequency b (say, monthly), but we are interested in compounded data in frequency q (say, annual). The approximation formulas for mean and standard deviation (SD) are:

*q*-frequency mean = *b*-freq mean \* q/n*q*-freq SD = *b*-freq SD \*  $\sqrt{(q/b)} \implies$  *q*-freq Variance = sqrt(*q*-freq SD)

**Example:** Using the data from the previous table we calculate the weekly mean and standard deviation for returns (b=30, q=7).

- weekly return = 0.007378 \* (7/30) = 0.00172 (0.172%)

- weekly SD = 0.040455 \* sqrt(7/30) = 0.01954 (1.95%)

<u>Note</u>: de-compounding the return:  $(1 + 0.007378)^{(7/30)} = 0.00172$ .

### **Returns: Sampling Distribution**

Recall that the sampling distribution of the sample mean is:  $\overline{X} \sim N(\mu, \sigma^2/N)$ 

**Example:** Before, using monthly S&P 500 log returns (N = 1805), we got:

Estimated Monthly mean return =  $\overline{X} = 0.007378$ 

Estimated Var( $\overline{X}$ ) =  $s^2/N$  = 0.040455<sup>2</sup> /1805 = 9.067075e-07

The SD of the monthly mean (also called the Standard Error, SE):

S.E. $(\bar{X}) = \text{sqrt}(9.067075\text{e-}07) = 0.000952 \text{ (or } 0.095\%\text{)}.$  $\Rightarrow \bar{X} \sim \text{N}(0.007378, .00095^2\text{)}$ 

<u>Note</u>: Compared to returns, expected returns, estimated by the sample mean, are more precisely estimated (0.1% vs 4.05%). Not surprised, the sampling distribution of the mean shrinks towards the population mean as N increases.

#### **Yields**

Consider an *n*-period discount bond. Time is measured in years. Today is *t*. Bond (asset) pays  $F_{t+n}$  dollars *n* years from now, at t + n.

 $F_{t+n}$  = Face value (value at time t + n).

 $P_t$  = Market price of the bond.

 $r_{n,t}$  = Yield to maturity (YTM) at time t for a maturity of n years.

n = Maturity of bond

$$P_t = \frac{F_{t+n}}{(1+r_{n,t})^n}$$

Interpretation: If our initial capital,  $P_t$  dollars, is invested today at the interest rate  $r_{n,t}$  for *n* years *compounded* annually, then, at time t + n, the payoff is  $F_{t+n}$ .

YTM,  $r_{n,t}$ , is a raw number: 4% at an annual rate is 0.04.

### **Continuous Compounding**

More generally, suppose an investment is compounded *m* times per *n* years; where *m* is number of times return (yield) is compounded, for example, m = 4 for quarterly, m = 12 for monthly,  $m = \infty$  for continuous compounding. Then, the market price of the bond is:

$$P_t = \frac{F_{t+n}}{(1 + \frac{r_t}{m})^{mn}}$$

As  $m \to \infty$ ,  $F_{t+n} = P_t (1 + \frac{r_t}{m})^{mn} \to P_t e^{rn}$ 

where we used

$$\lim_{m\to\infty} \left(1+\frac{x}{m}\right)^{mn} = e^{xn}$$

Then,

$$P_t = e^{-r_t n} F_{t+n}$$

• Suppose the continuously compounded bond at maturity pays  $1 (=F_{t+n})$  and the remaining duration is D units of time. Then,

$$P_t = e^{-r_t D}$$
\$1

Then, the log return per year is:

$$\log(P_{t+1}) - \log(P_t) = D(r_t - r_{t+1})$$
where we ignore that D has one unit of time less at time t+1. That is, the daily return of a bond is the change of yields multiplied by its duration.

• Now, suppose we invest  $P_0$  in a bond with continuous compounding at an annual rate r. Then, the value of the investment at year t is:

$$V_t = P_0 \ e^{rt}$$

The log return (log of gross return) per year is r:  $log(V_{t+1}) - log(V_t) = r$ 

The simple annual interest rate r quoted in the market is the annual log return if the interest is compounded continuously.

The *effective annual interest rate*,  $r^a$ , is simple the annual rate of return:

$$(1+r^a)^n = \left(1+\frac{r}{m}\right)^{mn} \qquad \Rightarrow \qquad r^a = \left(1+\frac{r}{m}\right)^m - 1$$

**Example:** Table 2.C reports descriptive statistics for monthly **long-run S&P 500 (blue) returns** and **10-year bond (red) rates**, from Robert Shiller's website. Interest Returns are reported annually.

	Total Return	10-year Interest
		Rates
Mean	0.007378	0.04511
Median	0.009928	0.0382
Maximum	0.414151	0.1532
Minimum	-0.30365	0.0062
Std. Dev.	0.040455	0.02304
Skewness	-0.47050	1.79664
Kurtosis	14.61046	6.75173
Jarque-Bera	10205.0	2029.7
P-value (JB)	2.2e-16	2.2e-16

### Table 2.C – S&P Composit Returns & 10-year Bond Rates (1871-2021)

We plot long-run S&P returns (blue) and 10-year Bond interest rates (red) data.



<u>Note</u>: Interest rates have a lower monthly mean (0.003759 = 0.04511/12), & lower volatility and kurtosis than stock returns.

We plot the histogram for **bond rates**, with a normal curve (in blue) for comparison purposes



Distribution - Histogram with Normal Curve

# **Returns: Expected Returns & The Equity Risk Premium**

As mentioned above, returns are not very precisely estimated. They have a large variance. Things get better for expected returns, which are estimated by the sample mean, since the S.E. of the sample mean gets smaller with N. That expected returns are better estimated that returns is a good property for financial models, since expected returns are a key component of every valuation model.

The expected return on any investment can be written as the sum of a *risk-free rate* and a *risk premium* to compensate for the risk of the investment. A key element in equity valuation models

is the risk premium that investors demand to hold "average (equity) risk" or *market (equity) risk,* which in turn affects the prices of all risky investments.

The difference between the expected risky market return and a risk-free rate is called the *equity risk premium* or ERP:

$$\mathrm{ERP} = \mathrm{E}[(r_m - r_f)],$$

where  $E[r_m]$  is the expected return on a well-diversified market portfolio and  $r_f$  is the risk-free rate. It is common to find that the ERP changes over time. In this case, we write the ERP at time t + k,  $ERP_{t+k}$ , as:

$$ERP_{t+k} = E_t[(r_m - r_f)_{t+k}]$$

where the subscript t means the expectations for period t + k are taken at time t.

In standard macroeconomic-finance models, the ERP is determined by the aggregate risk aversion of investors and the volatility of equity markets. The ERP drives expectations of future equity market returns. It is an input in equilibrium asset pricing models like the capital asset pricing model (CAPM) and multi-factor models, like the Fama-French 3-factor model.

# **Returns: Expected Returns & The ERP - Components**

To calculate the ERP we need  $r_f$  and  $E[r_m]$ .

(1) The risk-free rate. The risk-free rate,  $r_f$ , is approximated by the mean yield of government securities, typical examples, the 3-month U.S. Treasury bill or the 10-year U.S. Treasury bond. In general, given the upward sloping terms structure, using the T-bill rate will give result in a higher  $E[(r_m - r_f)]$ .

The risk-free rate used to compute the ERP has to be consistent with the risk-free rate used to compute expected returns. If we are estimating the cost of capital for a project that is expected to last 10 years, r<sub>f</sub> is approximated by the yield of 10-year government securities, for example, 10-year Treasury bonds.

(2) Expected Market Return. To determine  $E[r_{m,t}]$  we need to determine the Market Portfolio.

Q: What is the Market Portfolio? In theory, it represents the universe of risky assets. In practice it represents the universe of traded equities, not just domestic, but in the world. Returns on this equity market portfolio should be measured free of survivor bias.

In general, we approximate ("proxy") the Market Portfolio, with a well-diversified equity index. Also, in general, market-weighted indexes are preferred. For example, the S&P 500 Index, the MSCI World Index, or the Weighted Average of CRSP returns.

Q: How do we calculate  $E[r_{m,t}]$ ? There are three different ways to compute  $E[r_{m,t}]$ : 1) **Surveys**. Usually an average of ERPs provided by individual investors, institutional investors, managers and, even, academics.

2) **Historical data**. Expectations are computed using past data. This is the most popular approach. For example, compute  $E[r_{m,t}]$  with sample averages of market returns,  $\overline{X}$ . As we have seen above, if we use this approach, it pays to use as much data as possible –more data, lower S.E. We think of  $E[r_{m,t}]$  as a *long-run* average of market returns. Robert Shiller's website has U.S. market equity returns from 1871, but if we think that the ERP is time varying, for example we consider the stock market in 1871 and in 2021 are very different, it may be better to use a shorter period, for example, 50 years or, may be shorter period, 20 years to compute the ERP.

3) **Forward-looking data**. We derive an (implied) ERP using market prices, for example, market indexes, options & futures on market indexes, etc. Of course, we also need a model (a formula) that extracts the ERP from market prices.

# **Returns: Expected Returns & The ERP – The Equity Risk Premium Puzzle**

Once we determine  $E[r_{m,t}]$  and the risk-free rate, we are ready to calculate the ERP. But, keep in mind that we make decisions along the way.

For example, using Shiller's monthly data, with 150 years of data, we produce an estimate of the ERP =  $E[(r_m - r_f)]$ . Decisions made: computation of returns (log returns); method of computing ERP (Historical data); sample period (1871-2021); market portfolio (S&P Composite Index); risk-free rate (10-year U.S. bond rate). Then,

Annualized Market return = 0.007378 \* 12 = 0.088536Annualized risk-free rate = 0.04511ERP = 0.088536 - 0.04511 = 0.043426 (4.34%)

Many economists would consider this estimate of the ERP "*too high*." Why is **4.34%** too high? Using standard macroeconomic (neoclassical) models, the *degree of risk aversion* to justify it is unreasonable high, between 30 and 50, while a reasonable degree of risk aversion is no larger than 10.

A *too high* (for economic models) ERP was first reported by Mehra and Prescott (1985), which they estimated around 6%. According to their calibration of their "standard" model, the ERP should be, at most, 1%. Mehra and Prescott (1985) labeled the incompatibility of theory & observed data the *equity risk premium puzzle*.

There have been many attempts to explain the puzzle: statistical artifact (survivor bias); disaster insurance (peso problem/sample period), transaction costs & taxation, model's preferences, behavioral issues (mainly, myopic loss aversion & overreactions).

Damodaran (2021), who produces an annual update of the literature and the numbers, said in overview of the possible explanations:

"It is true that historical risk premiums are higher than could be justified using conventional utility models for wealth. However, that may tell us more about the dangers of using historical data and the failures of classic utility models than they do about equity risk premiums."

# **Returns: Expected Returns & The ERP – Wide Range**

Is it 4% or 6%? It turns out that even with 100+ years of data for developed markets there is no consensus on an ERP. Different choices in how to compute the ERP produce different estimates. For example, for the U.S. market, considered one of the best in terms of quality of data, Duarte and Rosa (2015) list over 20 approaches ("models") to estimate the ERP in the U.S.

With 1960-2013 data, D&R (2015) report estimates from -0.4% to 13.1%, with a 5.7% average for all models. A wide range! This wide range is consistent with the reported averages for developed markets that we present below in Tables 2.D and 2.E.

In Table 2.D, using **50 years of monthly return data** of log returns for some developed markets, we report mean equity annualized returns (in USD), the annualized standard deviation of returns (in USD) and the annualized ERP, which, again, is defined as the expected market excess return over the risk-free rate.

There is a wide range of ERP estimates from 0.88% (Italy) to 11.56% (HK), using the sample mean return for the MSCI country index (in USD) and the average U.S. T-Bill rate for the period ( $\approx 4.50\%$ ). The World Index, a weighted average of all equity markets, has an ERP equal to 3.17%, where the EAFE (Europe, Australia and the Far East) Index has a slightly lower ERP, 3.06%.

<b>Market</b> (N = 620)	Equity Return	Standard Deviation	ERP
U.S.	8.31	15.01	3.82
Canada	7.95	19.21	3.46
France	8.80	21.95	4.31
Germany	8.80	21.48	4.31
Italy	5.37	25.25	0.88
Switzerland	10.34	17.64	5.85
U.K.	7.37	21.20	2.88
Japan	9.56	20.46	5.06
Hong Kong	16.06	33.23	11.56
Singapore	11.71	27.48	7.22
Australia	7.35	23.42	2.73
World	7.66	14.54	3.17
EAFE	7.69	16.64	3.06

# Table 2.D - MSCI Index USD Equity Returns and ERP: (1970-2021)

In Table 2.E, using **34 years of monthly return data or less**, we report ERP (annualized) estimates for some emerging markets (in USD), where we observe a big dispersion of estimates, with higher mean returns, but also, higher standard deviations.

# Table 2.E - MSCI Index USD Equity Returns and ERP: (1987\* - 2021)

Market (N)	Equity Return	Standard Deviation	ERP
Argentina (404)	24.21	51.49	19.72
Brazil (404)	22.23	47.67	17.74
<b>Mexico</b> (404)	17.67	29.26	13.18
Poland (344)	15.88	43.24	11.39
Russia (320)	21.09	47.54	16.60

<b>India</b> (344)	12.10	28.35	7.60
<b>China</b> (344)	4.90	31.94	0.41
Korea (404)	11.75	34.08	0.726
Thailand (404)	11.58	32.24	6.06
Egypt (320)	11.61	31.69	8.62
South Africa (344)	9.47	26.31	4.98
World (620)	7.66	14.54	3.17
EM Asia	8.85	23.13	4.36

• We use the SE as a measure of precision of an estimate. For the sample mean,  $\overline{X}$ , we have: S.E. $(\overline{X}) = \frac{S}{\sqrt{N}}$ 

Using the previous data, we calculate the S.E.( $\overline{X}$ ) for several markets:

U.S. : 15.01/sqrt(620/12) = 2.0882% Germany: 21.48/sqrt(620/12) = 2.9883% Singapore: 27.48/sqrt(620/12) = 3.8231% Hong Kong: **33.23**/sqrt(620/12) = 4.6230 %

Brazil: 47.67/sqrt(404/12) = 8.2157 % Russia: 47.54/sqrt(320/12) = 9.2061% India: 28.35/sqrt(344/12) = 5.2950% China: 31.94/sqrt(344/12) = 5.9654%

A big difference in precision between Developed and Emerging Markets.

<u>Note</u>: Notice the effect of *N*. Hong Kong has a larger SD than India and China, but more observations make the mean estimate more precise.

### **Returns: Risk-Return – The Sharpe Ratio**

The most commonly cited statistics that provides a measure of the risk-return trade-off for an asset is the Sharpe ratio (SR), the ratio of the excess expected return of an asset to its risk, measured by its return volatility (SD). We estimate the SR of asset i with

$$\widehat{SR} = \frac{\widehat{\mu}_{i} - r_{f}}{s_{i}},$$

where  $\hat{\mu}_i$  is the sample mean  $(\bar{X})$  return of asset *i*,  $r_f$  is the risk-free rate and  $s_i$  is the SD of the return of asset *i*.

Interpretation: A 1% change in risk, increases excess returns by SR%.

The higher the SR, the better the risk-return trade-off. That is, if we compare assets, the asset with the higher SR provides the better trade-off.

Using the previous data, we calculate the SR for several markets:

U.S.:	<b>0.0382</b> /. <b>1501</b> = 0.254497	
Switzerland:	0.0585/ <b>17.64</b> = 0.3316	
Hong Kong:	<b>0.1156</b> /. <b>3323</b> = 0.3479	
Russia:	<b>0.1660</b> /.4754 = 0.3492	$\Leftarrow$ Best trade-off!
India:	<b>0.0760</b> /. <b>2835</b> = 0.2681.	

# **Review – Hypothesis Testing**

A *statistical hypothesis test* is a method of making decisions using experimental data. A result is called *statistically significant* if it is unlikely to have occurred by chance.

These decisions are made using (null) hypothesis tests. A hypothesis can specify a particular value for a population parameter, say  $q=q_0$ . Then, the test can be used to answer a question like:

Assuming  $q_0$  is true, what is the probability of observing a value for the (test) statistic used that is at least as big as the value that was actually observed?

Uses of hypothesis testing:

- Check the validity of theories or models.
- Check if new data can cast doubt on established facts.

Testing involves the comparison between two competing hypothesis (sometimes, they represent partitions of the world).

- The null hypothesis, denoted  $H_0$ , is sometimes referred to as the maintained hypothesis.
- The alternative hypothesis, denoted *H*<sub>1</sub>, is the hypothesis that will be considered if the null hypothesis is "rejected."

<u>Idea</u>: We collect a sample of data  $X = \{X_1, X_2, X_3, \dots, X_N\}$ . We construct a statistic T(X) = f(X), called the *test statistic*. Now we have a decision rule:

- If T(X) is contained in space R, we reject  $H_0$  (& we learn).

- If T(X) is in the complement of  $R(R^{C})$ , we fail to reject  $H_0$ .

<u>Note</u>: T(X), like any other statistic, is a RV. It has a distribution. We will use the distribution of T(X) under  $H_0$  to determine R.

**Example:** Suppose we want to test  $H_0: \mu = \mu_0$ . We collect data,  $\{X_1, X_2, X_3, \dots, X_N\}$ , and decide to use the statistic T(X). We know the distribution that T(X) follows under  $H_0$ . Suppose this distribution is the Normal distribution.

Then, we build a symmetric rejection region, *R*, around  $\mu = \mu_0$ , in such a way that *R* contains  $\alpha$ % of the Normal distribution. Note that by deriving the distribution of *T*(*X*) under  $\mu = \mu_0$ , we assume  $H_0$  to be the true. Then, *R* is:



We call the blue area "significance level" ( $\alpha$ %). If  $H_0$  is true, the blue area represents the probability of rejecting a true  $H_0$  or, just, P[ $R|H_0$ ]. ¶

<u>Remark</u>: We determine  $T_{LB}$  and  $T_{UB}$  in such a way that the probability of rejecting  $H_0$  when it is true –i.e., when  $\mu = \mu_0$ - is equal to  $\alpha$ . In practice, since we try to avoid rejecting a true  $H_0$ , we usually set  $\alpha$  (= P[ $R|H_0$ ]) equal to a small number.

**Example:** Following the above example, we now test if the mean of IBM annual returns,  $\mu_{IBM}$ , is 10%. That is,  $H_0$ :  $\mu_{IBM} = 10\%$ .

From the population, we get a sample:  $\{X_{1962}, X_{1963}, \ldots, X_{N=2024}\}$ , with N = 63. We use  $\overline{X}$ , which is unbiased, consistent, and, assuming X is normally distributed, we know its distribution,  $\overline{X} \sim N(\mu, .15^2/N)$ , where  $\sigma$  is .15.



Now, we need to determine the rejection region, *R*. We define  $T(X) = (\overline{X} - .10)/(.15/\sqrt{63})$ , which follows a N(0,1). Then, setting  $\alpha$  equal to 5%,





R is represented in the above graph by the blue area. That is, if T(X) falls in the "blue region," we reject H<sub>0</sub>:  $\mu_{IBM} = 10\%$ .

# **Review – Hypothesis Testing:** *p-value* and steps

We present the *classical approach*, a synthesized approach, known as *significance testing*. It relies on Fisher's *p-value*:

*p-value* is the probability of observing a result at least as extreme as the test statistic, under  $H_0$ .

**Example:** Suppose  $T(X) \sim \chi_2^2$ . We compute  $\widehat{T(X)} = 7.378$ . Then,

p-value( $\widehat{T(X)} = 7.378$ ) = 1 - Prob[T(X) < 7.378] = 0.025





<u>**R** Note</u>: We compute the *p*-value using pchisq(q, df), which computes the CDF at value q of a Chi-square distribution with df degrees of freedom. Then,

> pchisq(q = 7.378, df = 2) # Prob[T(X) < 7.378] 0.975003 p\_val <- 1 - pchisq(q = 7.378, df = 2) # p-value( $\widehat{T(X)} = 7.378$ ) = 1 - Prob[T(X) < 7.378] > p\_val [1] 0.02499699. ¶

Using the distribution of the test statistic T(X) under the null hypothesis, Fisher's *significance testing* approach determines a rejection region, based on the significance level ( $\alpha$ %).

We follow these steps:

**1.** Identify H<sub>0</sub> & decide on a *significance level* ( $\alpha$ % = P[ $R|H_0$ ]) to compare your test results.

**2.** Determine the appropriate test statistic T(X) and its distribution under the assumption that  $H_0$  is true.

**3.** Calculate T(X) from the data.

**4.** <u>Rule</u>: Reject H<sub>0</sub> if the *p*-value is "sufficiently small," then, we consider T(X) in *R* (we learn). Otherwise, we reach no conclusion (no learning).

<u>Note</u>: In Step 4, setting  $\alpha$ % is equivalent to setting *R*.

Q: What *p-value* is "sufficiently small" as to warrant rejection of H<sub>0</sub>?
 <u>Rule</u>: If *p-value* < α (say, 5%) ⇒ test result is *significant*: Reject H<sub>0</sub>.
 If *p-value* > α ⇒ test results are "*not significant*." No conclusions are reached (no learning here). Go back and gather more data or modify model.

The father of this approach, Ronald Fisher, favored 5% or 1%.

**Example:** From the U.S. Jury System  $H_0$ : The defendant is not guilty.  $H_1$ : The defendant is guilty. ¶

In statistics, we learn when we reject. In this case, we learn a defendant is guilty when the jury finds the defendant guilty, by rejecting  $H_0$ .

Example: From the U.S. Jury System
1. Identify H<sub>0</sub> & decide on a *significance level* (α%) H<sub>0</sub>: The defendant is not guilty H<sub>1</sub>: The defendant is guilty
Significance level α = "beyond reasonable doubt," presumably, a small level.

**2.** After judge instructions, each juror forms an "innocent index"  $T(X)_i$ .

**3.** Through deliberations, jury reaches a conclusion  $T(X) = \sum_{i=1}^{12} T(X)_i$ .

<b>4.</b> Rule:	If <i>p</i> -value of $T(X) < \alpha$	$\Rightarrow$ Reject H <sub>0</sub> . That is, guilty!
	If <i>p</i> -value of $T(X) > \alpha$	$\Rightarrow$ Fail to reject H <sub>0</sub> . That is, non-guilty.
A	Alternatively, jury builds a re	ejection region around $H_0$ . If $T(X) \in \mathbb{R}$ , jury rejects $H_0$ .

Note: Mistakes are made. We want to quantify these mistakes.

Failure to reject  $H_0$  does not necessarily mean that the defendant is not guilty, or rejecting  $H_0$  does not mean necessarily the defendant is guilty. *Type I error* and *Type II error* give us an idea of both mistakes.

Definition: Type I and Type II errors

A *Type I error* is the error of rejecting  $H_0$  when it is true. A *Type II error* is the error of "accepting"  $H_0$  when it is false (that is, when  $H_1$  is true).

Notation:Probability of Type I error:  $\alpha = P[X \in R | H_0]$ <br/>Probability of Type II error:  $\beta = P[X \in R^C | H_1]$ We call  $1 - \beta$  the power of the test –i.e., the probability of rejecting a false null hypothesis.

**Example:** From the U.S. Jury System

*Type I error* is the error of finding an innocent defendant guilty. *Type II error* is the error of finding a guilty defendant not guilty.

	State of World		
Decision	<i>H</i> <sub>0</sub> true ("not guilty")	<i>H</i> <sup>1</sup> true ("guilty")	
Cannot reject H <sub>0</sub>	Correct decision	Type II error	
Reject H <sub>0</sub>	Type I error	Correct decision	

<u>Note</u>: We usually think that we learn when we reject H<sub>0</sub>. Note that some "learning" comes from Type I error –i.e., from *false positives*. ¶

In general, we think *Type I error* is the worst of the two errors: We try to minimize the error of sending to jail an innocent person.

Actually, we would like *Type I error* to be zero. However, the only way to do this (100% of innocent defendants are found not guilty) is to never reject  $H_0$ . Then, we maximize *Type II error*.

There is a clear trade-off between both errors. Traditional view: Set *Type I error* equal to a small number (defined in the U.S. court system as "*beyond reasonable doubt*") and design a test that minimizes *Type II error*.

The usual tests (*t-tests*, *F-tests*, Likelihood Ratio tests) incorporate this traditional view.

**Example:** We want to test if the mean is equal to  $\mu_0$ , against the alternative hypothesis of different than  $\mu_0$ . Then,

1. H<sub>0</sub>:  $\mu = \mu_0$ . H<sub>1</sub>:  $\mu \neq \mu_0$ .

Notice that we have a double sided alternative, which creates a rejection region on both sides of the distribution of T(X).

2. Appropriate T(X): *t-test* (based on  $\sigma$  unknown and estimated by *s*). Determine distribution of T(X) under H<sub>0</sub>. Sampling distribution of  $\overline{X}$ , under H<sub>0</sub>:  $\overline{X} \sim N(\mu_0, \sigma^2/N)$ .

Then, distribution of T(X) under H<sub>0</sub>:

$$t = \frac{\bar{X} - \mu_0}{s_{1/\sqrt{N}}} \sim t_{N-1}$$
 -when  $N > 30, t \sim N(0, 1).$ 

**3.** Compute t,  $\hat{t}$ , using  $\bar{X}$ ,  $\mu_0$ , s, and N. Since it is a double sided t-test, we look at  $|\hat{t}|$ . Then, get *p*-value( $|\hat{t}|$ ).

4. <u>Rule</u>: Set  $\alpha$  level. If *p*-value( $|\hat{t}|$ ) <  $\alpha$   $\Rightarrow$  Reject H<sub>0</sub>:  $\mu = \mu_0$ . Alternatively, if  $|\hat{t}| > t_{N-1,1-\alpha/2}$  (=1.96, if  $\alpha$ =.05)  $\Rightarrow$  Reject H<sub>0</sub>:  $\mu = \mu_0$ .

Notice the alternative Rule; it sets a Rejection region:



If  $\alpha = 5\%$  and N > 30, then  $t_{N>30,.025} = -1.96 \& t_{N>30,.975} = 1.96 (\approx 2)$ . (The distribution is symmetric, that is,  $-t_{N>30,.025} = t_{N>30,.975} = 1.96$ ).

<u>Remark</u>: Under  $H_0$ , the blue area represents the Type I error –i.e., the probability of rejecting a true  $H_0$ , which is by design equal to .

<u>Technical Note 1</u>: In step 2, the distribution of the t-statistic, t, is exact if  $\{X\}$  follows a normal distribution, otherwise, the distribution is asymptotic (for this we need a large *N*); that is

$$t = \frac{\bar{X} - \mu_0}{s/\sqrt{N}} \xrightarrow{d} N(0, 1)$$

<u>Technical Note 2</u>: In step 2, we determine the distribution of t, by using the sampling distribution of  $\overline{X}$  under H<sub>0</sub>. If H<sub>0</sub> is not true, say  $\mu = \mu_1$ . then

$$\overline{X} \sim \mathrm{N}(\mu_1, \, \sigma^2/N),$$

thus, t is distributed N(0, 1) only under H<sub>0</sub>, since only under H<sub>0</sub> the E[ $\overline{X} - \mu_0$ ] = 0.

# **Review – Hypothesis Testing: Examples**

**Example 1:** We want to test if the monthly mean total return of the S&P 500 is equal to zero using  $\alpha = .05$ . We use the **S&P 500 monthly returns** (1871-2021) with the following mean and variance:  $\overline{X} = 0.007378$ , s = 0.04046, N = 1805.

1. H<sub>0</sub>: 
$$\mu = 0$$
.  $(\mu_0 = 0)$   
H<sub>1</sub>:  $\mu \neq 0$ . (a two-sided alternative.)

$$2. t = \frac{\bar{X} - \mu_0}{s/\sqrt{N}}$$

3.  $\hat{t} = \frac{0.007378}{0.04046/\sqrt{1805}} = 7.7478 \& p-value(|\hat{t}|) = 9.325873e-15$  -multiply by 2 p-value( $\hat{t}$ ).

4. Rule: p-value( $|\hat{t}|$ ) = 9.325873e-15 <  $\alpha$  = .05  $\Rightarrow$  Reject H<sub>0</sub>:  $\mu$  = 0. Alternatively,  $|\hat{t} = 7.7478| > t_{1789,.025} = 1.96$   $\Rightarrow$  Reject H<sub>0</sub>:  $\mu$  = 0.

### Conclusion: S&P 500 monthly mean total returns are not equal to zero.

<u>**R** Note</u>: We find the  $t_{1789,025}$  using qt (p, df), which gives the quantile of the t-distribution with df degrees of freedom. That is,

> qt(.975, 1789)  $\# = (-1)^*$ qt(.025, 1789) by symmetry. [1] **1.961291** # Check result with pt (q, df) > pt (q=**1.96**, df=1789)

Q: How do we calculate the *p*-value? Recall, it is the probability of observing a result at least as extreme as the test statistic, under  $H_0$ .

 $\Rightarrow$  1 - pt (q=1.96, df=1789)  $\approx$  0.025. ¶

In this case, we know that under H<sub>0</sub>:  $\mu = 0$ , the t-stat is well approximated by a N(0,1) distribution (since *N*>30). Then, we use the R function *pnorm* to calculate the cumulative standard normal value up to **7.7478**, and then subtract it from 1:

 $p_val_1 <-1 - pnorm(7.7478)$ # p-value of (one-sided) t\_test $> p_val_1 * 2$ # Multiply by 2 since it is a double sided test[1] 9.325873e-15

The observed t ( $\hat{t} = 7.7478$ ) is outside the non-rejection region ( $R^C$ ) built around  $H_0$ : (-1.96, 1.96).

 $\Rightarrow$  Rejection region = R = ([- $\infty$ , -1.96] U [1.96,  $\infty$ ]). ¶

[1] 0.9749246.



| î = 7.7478 |

<u>Note</u>: : p-value( $|\hat{t}|$ ) = p-value( **3.7121**) \* **2** 

**Example 2:** We want to test if **monthly S&P 500 total returns** (1871-2021) follow a normal distribution using  $\alpha = .05$ . If the distribution is normal, skewness is zero and kurtosis is equal to 3 (or excess kurtosis equals 0). The estimated moments are:  $\hat{\gamma}_1 = -0.4705$ ,  $\hat{\gamma}_2 = (14.6105 - 3) = 11.6105$ , & N = 1805.

- 1. H<sub>0</sub> (Data is normal):  $\gamma_1 = \frac{\mu_3^0}{\sigma^3} = 0$  and  $\gamma_2 = \frac{\mu_4^0}{\sigma^4} 3 = 0$ . H<sub>1</sub> (Data is not normal):  $\gamma_1 \neq 0$  and/or  $\gamma_2 \neq 0$ .
- 2. Appropriate T(X): the Jarque-Bera test (JB),  $JB = \frac{N}{6} * (\gamma_1^2 + \frac{\gamma_2^2}{4})$ Under H<sub>0</sub>, JB  $\xrightarrow{d} \chi_2^2$  (chi-square distribution with 2 degrees of freedom)
- 3.  $\widehat{\text{JB}} = \frac{1805}{6} * \left[ (-0.4705)^2 + \frac{(11.6105)^2}{4} \right] = 10,204.89$
- 4. <u>Rule</u>: p-value( $\widehat{JB} = 10,204.9$ )  $\approx 0 < \alpha = .05 \implies$  Reject H<sub>0</sub>. Alternatively, compare  $\widehat{JB}$  to the  $\chi^2_{2,95}$  value ( $\chi^2_{2,95} = 5.991$ ). That is,

$$\widehat{JB} > \chi^2_{2,95}$$
  $\Rightarrow$  Reject H<sub>0</sub>. (A strong rejection!)

#### Chi-square Distribution (df=2): One-sided 95% C.I.



<u>Conclusion</u>: We strongly reject H<sub>0</sub>. That is, monthly S&P 500 returns are not normally distributed. ¶

# **Review – Confidence Intervals (C.I.)**

When we estimate parameters with an estimator,  $\hat{\theta}$ , we get a point estimate for  $\theta$ , meaning that  $\hat{\theta}$  is a single value in  $R^k$ . For example, in the previous example, we get  $\bar{X} = 0.003571$ .

Broader concept: Estimate a set  $C_n$ , a collection of values in  $\mathbb{R}^k$ . For example,  $\mu \in \{0.00155, 0.00554\}$ .

It is common to focus on intervals  $C_n = [L_n; U_n]$ , called an *interval estimate* for  $\theta$ . The goal of  $C_n$  is to contain the true population value,  $\theta$ . We want to see  $\theta \in C_n$ , with high probability.

<u>Technical detail</u>: Since  $C_n$  is a function of the data, it is a RV and, thus, it has a pdf associated with it. The *coverage probability* of the interval  $C_n = [L_n; U_n]$  is  $Prob[\theta \in C_n]$ .

Intervals estimates  $C_n$  provide an idea of the uncertainty in the estimation of  $\theta$ : The wider the interval  $C_n$ , the more uncertain we are about our estimate,  $\hat{\theta}$ .

Interval estimates  $C_n$  are called *confidence intervals* (C.I.) as the goal is to set the coverage probability to equal a pre-specified target, usually 90% or 95%.  $C_n$  is called a  $(1 - \alpha)$ % C.I.

When we know the distribution for the point estimate, it is straightforward to construct a C.I. For example, if  $\hat{\theta} \sim N(\theta, Var[\hat{\theta}])$ , then a  $(1 - \alpha)\%$  C.I. is given by:  $C_n = [\hat{\theta} + z_{\alpha/2} * \text{Estimated SE}(\hat{\theta}), \hat{\theta} + z_{1-\alpha/2} * \text{Estimated SE}(\hat{\theta})]$ 

This C.I. is symmetric around  $\hat{\theta}$ . Its length is proportional to SE( $\hat{\theta}$ ).

The *z* values are taken from the standard normal distribution, which is symmetric around 0. That is,  $z_{(1-\alpha/2)} = -z_{\alpha/2} = |z_{\alpha/2}|$ .

hus, we can write the above  $(1 - \alpha)$ % C.I. as:  $C_n = [\hat{\theta} - z_{(1 - \alpha/2)} * \text{Estimated SE}(\hat{\theta}), \hat{\theta} + z_{(1 - \alpha/2)} * \text{Estimated SE}(\hat{\theta})]$ 

Popular values for  $\alpha$  and z:

$\alpha = .10 \implies z_{.95} = 1.645$	$(z_{.05} = -1.645)$
$\alpha = .05 \implies z_{.975} = 1.96$	$(z_{.025} = -1.96)$
$\alpha = .02 \implies z_{.99} = 2.33$	$(z_{.01} = -2.33)$

If the data follows a Normal distribution, then for the sample mean a  $(1 - \alpha)$ % C.I. is given by:  $C_n = [\overline{X} - z_{1-\alpha/2} * SD(\overline{X}), \overline{X} + z_{1-\alpha/2} * SD(\overline{X})]$ 

The size of the symmetric C.I. depends on the SD (=SE). The higher SD, the wider the C.I.

**Example:** Two 95% C.I. for the mean, with  $\overline{X} = 0$ , with two different SD (=1, 2), are plotted below. (Recall:  $z_{1-.05/2} = 1.96$ ).



Normals with Different Volatilities

**Example:** We estimate a 95% C.I. for the **monthly total mean return of the S&P 500**. The sampling distribution of the sample mean (assuming normality) is  $\overline{X} \sim N(\mu, \sigma^2/N)$ , then, a  $(1 - \alpha)$ % C.I. is given by:

 $C_n = [\overline{X} - z_{1-\alpha/2} * SD(\overline{X}), \overline{X} + z_{1-\alpha/2} * SD(\overline{X})] \Rightarrow$  The higher SD, the wider the C.I. Then,

$$\begin{split} C_n &= [ \underbrace{0.00738 - 1.96 * (0.04046 / \sqrt{1805}), 0.00738 + 1.96 * ((0.04046 / \sqrt{1805})] \\ &= [ 0.005511, 0.00924 ] = [ 0.55\%, 0.92\% ]. \end{split}$$



By looking at the 95% C.I., we can reject the null hypothesis that monthly S&P 500 total returns are 0, since 0% is outside the 95% C.I. But, the C.I is wide, even after 150 years of data.

<u>Conclusion</u>: Reject H<sub>0</sub>:  $\mu = 0$ , since 0 is outside the observed 95% C.I.

<u>Note</u>: Using the above confidence interval, we can also reject that monthly excess returns are equal to 0.0833% (= 1%/12). Recall that Mehra & Prescott (1983) reported that the ERP is too high since in their calculation the annualized equilibrium ERP is equal to 1%. ¶

### **Confidence Interval (C.I.) for the Variance**

We want to estimate a  $(1 - \alpha)$ % C.I. for the variance. Assuming normality, the sample variance, once scaled, is distributed:

$$(N - 1) s^2 / \sigma^2 \sim \chi^2_{N-1}.$$

To derive a  $(1 - \alpha)$ % C.I. for the variance, we rewrite the standard confidence interval for a chisquared variable:

$$P(\chi_{v,\alpha/2}^2 < \chi_v^2 < \chi_{v,1-\alpha/2}^2) = P(\chi_{v,\alpha/2}^2 < (N-1) s^2/\sigma^2 < \chi_{v,1-\alpha/2}^2) = 1 - \alpha$$

For example, for N=11 (v = 10), a 95% C.I. for a  $\chi^2_{v=10}$  RV can be built as:

$$P[\chi^{2}_{10,.025} = 3.246973 < (10) s^{2}/\sigma < \chi^{2}_{10,.975} = 20.48318] = .95\%$$



Then, after some algebra (recall inversion changes inequality signs), we derive:  $P[(N-1) s^2/\chi^2_{\nu,1-\alpha/2} < \sigma^2 < (N-1) s^2/\chi^2_{\nu,\alpha/2}] = 1 - \alpha.$ 

<u>Note</u>: This C.I. is not symmetric. But, as the degrees of freedom get large, the  $\chi^2_{N-1}$  starts to look like the normal distribution and, thus, CIs will look more symmetric.

**Example:** We estimate a 95% C.I. for the variance of **monthly S&P 500 mean total return** (*N*=1805). Then, from the  $\chi^2_{1804}$  distribution, we get:  $\chi^2_{1804,025} = 1688.2 \& \chi^2_{1804,975} = 1923.7$ . (You get these values in R with qchisq(.025, df=N-1) & qchisq(.975, df=N-1), respectively.)



 $\begin{array}{l} P[(1804)*(0.04046)^2/(1923.6) < \sigma^2 < (1804)*(0.04046)^2/(1688.2)] = .95 \\ P[0.001535 < \sigma^2 < 0.001749] = .95 \end{array}$ 

Taking square root above delivers a 95% C.I. for  $\sigma$ :  $\Rightarrow$  95% C.I. for  $\sigma$  is given by (3.918%, 4.182%).

The C.I. is quite compact around the sample point estimate. Compared to the mean,  $\sigma$  is measured with accuracy.

<u>Note</u>: Usually *N* is large (N > 30). We can use the normal approximation to calculate CIs for the population  $\sigma$  (check the pdf above!). For the S&P data, we estimate the S.E. for the sample SD:

 $SE(s) = s/\sqrt{2 * (N-1)} = 0.04046/sqrt(2*1804) = 0.000673$  (or .067%).

A 95% CI for  $\sigma$  is given by

 $(4.046\% \pm 1.96 * .067\%) = (3.914\%, 4.178\%).$  (Very close!)

### C.I. Application: Using the ED – The Bootstrap

In the previous examples, we assumed that we knew the distribution of the data: Stock returns follow a normal distribution. What happens when the data follows an unknown distribution, *F*?

We still can use the sample mean,  $\overline{X}$ , or the sample variance,  $s^2$ , as estimates of  $\mu$  and  $\sigma^2$ , since the LLN tell us that they are both consistent estimators. If we have a "large" dataset –i.e., large N– we can use the CLT to justify a C.I based on a normal distribution.

But, when we have an unknown distribution F and we do not have a large enough N or we suspect the normal approximation is not a good one, we still can build a C.I. for any statistic using a new method: a *bootstrap*.

*Bootstrapping* is the practice of estimating the properties of an estimator -say, its variance- by measuring those properties when sampling from an approximating distribution (the *bootstrap DGP*).

That is, it is necessary to estimate a bootstrap DGP from which to draw the simulated samples. The DGP that generated the original data is unknown, and so it cannot be used to generate simulated data.

 $\Rightarrow$  The bootstrap DGP estimates the unknown true DGP.

<u>Idea</u>: We use the data at hand -the empirical distribution (ED)- to estimate the variation of statistics that are themselves computed from the same data. Recall that, for large samples, the ED approximates the CDF very well.

The *empirical bootstrap* is a statistical technique, easy to implement, that takes advantage of today's modern computers, by resampling from the ED. Bootstrapping uses the ED –i.e., sample-as if it were the true CDF.



• Suppose we have a sample with *N* observations drawn from F(x):

$$\{x_1, x_2, \dots, x_N\}$$

From the ED,  $F^*$ , we sample ("*resample*") with replacement N observations:  $\{x_1^* = x_2, x_2^* = x_4, x_3^* = x_4, \dots, x_N^* = x_{N-8}\}$ 

This is an *empirical bootstrap sample*, which is a resample of the same size N as the original data, drawn from  $F^*$ .

For any statistic  $\theta$  computed from the original sample data, we can define a statistic  $\theta^*$  by the same formula, but computed instead using the resampled data. Then,

$$\{x_1^* = x_2, x_2^* = x_4, x_3^* = x_4, \dots, x_N^* = x_{N-8}\} \quad \Rightarrow \hat{\theta}_1^*$$

 $\theta^*$  is computed by resampling the original data; we can compute many  $\theta^*$  by resampling many times from  $F^*$ . Say, we resample  $\theta^* B$  times.

We have a collection of estimated  $\theta^*$ :

$$\{\hat{\theta}_1^*, \hat{\theta}_2^*, \hat{\theta}_3^*, \dots, \hat{\theta}_B^*\}.$$

From this collection of  $\hat{\theta}^*$ 's, we can compute the mean, the variance, skewness, draw a histogram, etc., and confidence intervals. From this collection of  $\hat{\theta}^*$ 's, we learn about the behavior of statistic  $\theta$ .

**Example:** We are interested in estimating the variance of monthly S&P 500 returns. We have already estimated it, using Shiller's data:  $(0.04046)^2$ . We also estimated a 95% C.I. based on the normal distribution, but, we are not sure it is a reliable C.I. since we already know that monthly returns are not normally distributed.

We use a bootstrap to study the distribution of the sample variance.

• Randomly construct a sequence of *B* samples (all with N=1,871). Say,

$$B_{1} = \{x_{1}, x_{2}, x_{6}, x_{6}, x_{6}, x_{6}, x_{16}, \dots, x_{1458}, x_{1758}, x_{1859}\} \Rightarrow \theta_{1} = s_{1}^{2}$$

$$B_{2} = \{x_{5}, x_{7}, x_{8}, x_{9}, x_{21}, x_{21}, x_{26}, \dots, x_{1661}, x_{1663}, x_{1870}\} \Rightarrow \hat{\theta}_{2}^{*} = s_{2}^{2}$$

$$B_{B} = \{x_{2}, x_{3}, x_{8}, x_{11}, x_{21}, x_{21}, x_{22}, \dots, x_{1805}, x_{1805}, x_{1806}\} \Rightarrow \hat{\theta}_{B}^{*} = s_{B}^{2}$$

We have a collection of estimated *sample variances:*  $\{s_1^2, s_2^2, s_3^2, \dots, s_B^2\}.$ 

From this collection of  $s^2$  's, we can compute the mean, the variance, skewness, draw a histogram, etc., and confidence intervals. From this collection of  $s^2$  's, we learn about  $\sigma^2$ .

- Bootstrap Steps:
- 1. From the original sample, draw a random sample with size N.
- 2. Compute statistic  $\theta$  from the resample in 1:  $\hat{\theta}_1^*$ .
- 3. Repeat steps 1 & 2 *B* times  $\Rightarrow$  Get *B* statistics:  $\{\hat{\theta}_1^*, \hat{\theta}_2^*, \hat{\theta}_3^*, \dots, \hat{\theta}_B^*\}$
- 4. Compute moments; draw histograms; etc. for these *B* statistics.

Using the histogram or the sorted  $\{\hat{\theta}_1^*, \hat{\theta}_2^*, \hat{\theta}_3^*, \dots, \hat{\theta}_B^*\}$ , we can build a  $(1 - \alpha)$ % C.I. Using the histogram, the lower bound leaves  $\alpha/2$ % of the  $\hat{\theta}^*$  to the right and  $(1 - \alpha/2)$ % of the  $\hat{\theta}^*$  to the left.

**Example**: We construct a 95% C.I. for the variance of S&P 500 returns. (You need to install R package *resample*, using the **install.packages()** function.)

```
Sh da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/Shiller 2020data.csv", head=TRUE,
sep=",")
SP \leq Sh da P
T \leq - length(SP)
lr \leq log(SP[-1]/SP[-T])
\ln var < var(\ln)
T s <- length (lr)
sim size <- 1000
                                                              \# B = size of bootstrap
                                                              # call library resample
library(resample)
data_star <- sample(lr, T_s*sim_size, replace=TRUE)
                                                              # create B resamples of size T_s
boot_sample <- matrix(data_star, nrow= T s, ncol=sim_size) # organize resamples in matrix
boots_vars <- colVars(boot_sample)</pre>
                                              # compute the variance for each bootsrap sample
q \leq -quantile(boots_vars, c(0.025, 0.975))
                                              # Find the 0.025 and 0.975 quantile for q
c_i <- lr_var - c(q[1], q[2])
                                              # Calculate the 95% C.I. for the variance.
cat("Confidence interval: ",ci, "\n")
                                              # Print C.I using cat
```

```
> lr_var
```

[1] 0.001637
> ci
97.5% 2.5%
0.001376664 0.001909769
> cat("Confidence interval: ", ci, "\n")
Confidence interval: 0.001376664 0.001909769
> Or, taking square roots above, we can get a 95% CI for σ: (3.71%, 4.37%). ¶

• Results (bootstrap principle):

**1.** With a large enough *B*, the LLN allows us to use the  $\hat{\theta}^*$ 's to estimate the distribution of  $\hat{\theta}$ ,  $F(\hat{\theta})$ .

**2.** The variation in  $\hat{\theta}$  is well approximated by the variation in  $\hat{\theta}^*$ .

Result 2 is the one that we use to estimate the size of a C.I.

Technical Note: The bootstrap delivers consistent results only.

# C.I. Application: The Bootstrap Percentile Method

There are many ways to construct a C.I. using bootstrapping. The easier one is the one described above. Just use the distribution of the  $\hat{\theta}^*$ 's to compute directly a C.I. This is the *bootstrap percentile method*.

The percentile method uses the distribution of  $\hat{\theta}^*$  as an approximation to the distribution of  $\hat{\theta}$ . It is very simple, but not as appealing, since comparing differences tends to work better.

**Example**: (Continuation of previous example.) We construct a 95% C.I. for the variance of S&P 500 returns. Using the boot.ci function, with type=**perc**, from *boot* package (install boot first, using the **install.packages()** function and then call library(boot) before you use it):

library(**boot**) boot.samps <- boot(data=lr, statistic=var\_p, R=sim\_size) # resampling and  $\theta^*$  estimation boot.ci(boot.samps, type = "**perc**")

> boot.ci(boot.samps, type = "perc")
BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
Based on 1000 bootstrap replicates

CALL : boot.ci(boot.out = boot.samps, type = "**perc**")

Intervals : Level Percentile 95% (0.0014, 0.0020) Calculations and Intervals on Original Scale • Draw a Histogram to Check Distribution of  $\hat{\theta}^*$  (=boot.samps\$t).

> hist(boot.samps\$t, breaks=12, main="Histogram for Boostrapped Variances", xlab="Bootstrapped Variances")



Histogram for Boostrapped Variances

• Check results by sorting boot.samps\$t.	
<pre>&gt; new &lt;- sort(boot.samps\$t)</pre>	
> new[25]	# CI's Lower Bound
[1] 0.001398215	
> new[975]	# CI's Upper Bound
[1] 0.001955096	
On fan - talving gewone na et af the ale ave have de th	$\sim 0.50/$ CI is given by (2.740/

Or for  $\sigma$ , taking square root of the above bounds, the 95% CI is given by (3.74%, 4.42%).

> new <- sort(boot.sampst) > new[25] # CI's Lower Bound [1] 0.001398215 > new[975] # CI's Upper Bound [1] 0.001955096 Or for  $\sigma$ , the 95% CI is given by (3.74%, 4.42%). ¶

# C.I. Application: The Empirical Bootstrap

The percentile method uses the distribution of  $\hat{\theta}^*$  as an approximation to the distribution of  $\hat{\theta}$ . It is very simple, but there are more appealing methods. In general, a bootstrap based on comparing differences is sounder. This is the key to the *empirical bootstrap*.

To build a C.I. for  $\theta$ , we use  $\hat{\theta}$ , computed from the original sample. As in the previous C.I.'s, we want to know how far is  $\hat{\theta}$  from  $\theta$ . For this, we would like to know the distribution of

$$q = \hat{\theta} - \theta.$$

If we knew the distribution of  $q = \hat{\theta} - \theta$ , we build a  $(1 - \alpha)$ % C.I., by finding the critical values  $q_{\alpha/2} \& q_{(1-\alpha/2)}$  to have:

$$\Pr\left(q_{\alpha/2} \leq \widehat{\theta} - \theta \leq q_{(1-\alpha/2)} | \theta\right) = 1 - \alpha$$

Or, after some manipulations:

 $\Pr\left(\widehat{\theta} - q_{\alpha/2} \geq \theta \geq \widehat{\theta} - q_{(1-\alpha/2)} | \theta\right) = 1 - \alpha,$ 

which gives a  $(1 - \alpha)$ % C.I.:

$$\mathbf{C}_{\mathrm{n}} = [\hat{\theta} - q_{(1-\alpha/2)}, \ \hat{\theta} - q_{\alpha/2}]$$

We do not know the distribution of q, but we can use the bootstrap to estimate it with  $q^* = \hat{\theta}^* - \hat{\theta}$ .

and then to get  $q_{\alpha/2}^* \& q_{(1-\alpha)/2}^*$ :

$$\mathbf{\hat{C}}_{n} = [\hat{\theta} - q^{*}_{(1-\alpha)/2}, \ \hat{\theta} - q^{*}_{\alpha/2}]$$

This C.I. is called the *pivotal* C.I.

<u>Intuition</u>: The distribution of  $\hat{\theta}$  is 'centered' at  $\theta$ , while the distribution of  $\theta^*$  is centered at  $\hat{\theta}$ . If there is a significant separation between  $\hat{\theta}$  and  $\theta$ , these two distributions will also differ significantly.

On the other hand, the distribution of  $q = \hat{\theta} - \theta$  describes the variation of  $\hat{\theta}$  about its center. Similarly, the distribution of  $q^* = \theta^* - \hat{\theta}$  describes the variation of  $\theta^*$  about its center.

Then, even if the centers are quite different, the two variations about the centers can be approximately equal.

**Example**: (Continuation of previous example.) We want to estimate a 95% C.I. for the variance of monthly returns of the S&P 500. (You need to install R package *resample*, using the **install.packages()** function.)

#### sim size <- 1000

# B = size of bootstrap

library(resample)# call library resampledata\_star <- sample(lr, T\_s \* sim\_size, replace=TRUE)</td># create B resamples of size T\_sboot\_sample <- matrix(data\_star, nrow=T\_s, ncol=sim\_size)</td># organize resamples in matrix

boots\_vars <- colVars(boot\_sample) q\_star <- boots\_vars - lr\_var q <- quantile(q\_star, c(0.025, 0.975)) ci <- lr\_var -c(q[2], q[1]) cat("Confidence interval: ",ci, "\n")

> lr\_var
[1] 0.001637
> ci
97.5% 2.5%
0.001376664 0.001909769
> cat("Confidence interval: ",ci, "\n")

# variance for each bootstrap sample
# Compute q\* for each bootstrap sample
# Find the 0.025 & 0.975 quantile for q\*
# Calculate the 95% C.I. for the variance.
# Print C.I using cat

Confidence interval: **0.001376664 0.001909769** > Or for  $\sigma$ , the 95% CI is given by (**3.71%**, **4.37%**).

Note: We got very similar results to the percentile bootstrap.

**Example**: Now, we construct the same 95% C.I. for the variance of monthly S&P 500 returns but using the R package *boot*. You need to install package first, using the **install.packages()** function.

```
library(boot)
# function to obtain the variance from the data
var p <- function(data, i) {</pre>
       d \leq -data[i]
return(var(d))
}
boot.samps <- boot(data=lr, statistic=var p, R=sim size) # resampling and \theta^* estimation
boot.ci(boot.samps, type = "basic")
                                                    # boot computes the CI with type=basic.
> boot.ci(boot.samps, type = "basic")
BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
Based on 1000 bootstrap replicates
CALL:
boot.ci(boot.out = boot.samps, type = "basic")
Intervals :
Level
       Basic
95% (0.0014, 0.0019)
Calculations and Intervals on Original Scale
• Check results using previous step-by-step process:
q star <- boot.samps$t - lr var
                                                           \# q^* = \theta^* - \hat{\theta}
q ad <- sort(q star)
                                                           # sort q^*
> lr var - q ad[975]
                                                           # CI's Lower Bound
[1] 0.001357793
> lr var - q ad[25]
                                                           # CI's Upper Bound
[1] 0.001914674
We can transform this CI for the variance into a CI for the SD:
```

> sqrt(lr\_var - q\_ad[975])
[1] 0.03684825
> sqrt(lr\_var - q\_ad[25])
[1] 0.04375699

A 95% CI for  $\sigma$  is given by (3.68%, 4.38%), wider than the CI assuming a Normal distribution for returns.

Note that we can also gauge the uncertainty of the estimation of  $\theta$  by computing the sample standard error, SE( $\hat{\theta}^*$ ). (Recall we call the standard deviation of an estimator its standard error.):

#### • Steps

1. Computing the sample variance:

$$\operatorname{Var}(\widehat{\theta}^*) = \frac{1}{B-1} \sum_{i=1}^{B} (\widehat{\theta}_i^* - \overline{\theta}^*)^2,$$

where  $\bar{\theta}^* = \frac{1}{B} \sum_{i=i}^{B} \hat{\theta}_i^*$ . 2. Estimate the S.E. of  $\hat{\theta}^*$ : SE $(\hat{\theta}^*) = \operatorname{sqrt}[\operatorname{Var}(\hat{\theta}^*)]$ .

### C.I. Application: The Parametric Bootstrap Method

If we assume the data is from a parametric model (say, from a Normal or a Gamma distribution), we can use the parametric bootstrap to access the uncertainty (variance, C.I.) of the estimated parameter. In a parametric bootstrap, we generate bootstrap samples from the assumed distribution, based on moments computed from the sample. We do not use the ED.

Suppose we have a sample with *N* observations drawn from  $F(x; \theta)$ :

$$\{x_1, x_2, ..., x_N\}$$

In the parametric bootstrap, we know  $F(x; \theta)$ , the distribution of *x*, but we do not know its parameters. Suppose there is only one unknown parameter,  $\theta$  (say, the variance). From the sample, we compute  $\hat{\theta}$ , the estimator of  $\theta$ . Then, we bootstrap from  $F(x; \hat{\theta})$  and proceed as before to form a C.I..

• Steps:

- 1. Draw *B* samples of size *N* from  $F(x; \hat{\theta})$ .
- 2. For each bootstrap sample,  $\{x_1^*, x_2^*, x_3^*, ..., x_N^*\}$ , calculate  $\hat{\theta}^*$ .  $\Rightarrow$  Get  $B \hat{\theta}^*$ .

3. Estimate a C.I. using the previous methods.

**Example**: Suppose S&P 500 monthly returns follow a N(0,  $\sigma^2$ ). We estimate  $\sigma^2$  with  $s^2 = 0.04046^2$ . > lr var

/ Ir\_var
[1] 0.001637

 $\begin{array}{l} x <- \operatorname{rnorm}(\mathbf{T}_{s}*\operatorname{sim}_{size}, \operatorname{mean}=0, \operatorname{sd}=\operatorname{lr}_{sd}) \\ \text{boot}\_\operatorname{sample} <- \operatorname{matrix}(x, \operatorname{nrow}=\mathbf{T}_{s}, \operatorname{ncol}=\operatorname{sim}_{size}) \\ \text{boots}\_\operatorname{vars} <- \operatorname{colVars}(\operatorname{boot}\_\operatorname{sample}) \\ q\_\operatorname{star} <- \operatorname{boots}\_\operatorname{vars} - \operatorname{lr}\_\operatorname{var} \\ q <- \operatorname{quantile}(q\_\operatorname{star}, \operatorname{c}(0.025, 0.975)) \\ \operatorname{ci} <- \operatorname{lr}\_\operatorname{var} - \operatorname{c}(q[2], q[1]) \\ > \operatorname{ci} \\ 97.5\% \qquad 2.5\% \end{array}$ 

# generate normal data
# organize simulated data
# compute variances

#### 0.001547382 0.001760286

Or for  $\sigma$ , the 95% CI is given by (3.94%, 4.20%). Very close to the C.I.'s we obtained before assuming a Normal distribution for returns. Not a surprise! ¶

<u>Note</u>: In the previous example, to gauge the uncertainty of the estimation of  $s^2$ , we can also compute the sample standard error, SE( $s^2$ ).

• Steps

- 1. Draw *B* samples of size *N* from a N(0,  $s^2$ )  $\Rightarrow$  Get  $B s^{2^*}$ .
- 2. Estimate the variability of  $s^2$  by computing the sample variance

$$\operatorname{Var}(s^{2}) = \frac{1}{B-1} \sum_{i=1}^{B} (s_{i}^{2*} - s_{B}^{2})^{2}$$

where  $s_B^2 = \frac{1}{B} \sum_{i=1}^{B} s_i^{2*}$ .

3. Estimate the S.E. of  $s^2$ : SE $(s^2)$  = sqrt[Var $(s^2)$ ].

<u>Remark</u>: An important difference between the nonparametric and parametric bootstrap procedures is that in the nonparametric procedure, only values of the original sample appear in the bootstrap samples. In the parametric bootstrap, the range of values in the bootstrap sample is the entire support of  $F(x; \theta)$ . In the parametric bootstrap of the above example, the values in the bootstrap sample could be any value between negative and positive infinity.

# C.I. Application: Bootstrapping – Why?

Question: Why do we need a bootstrap?

- Sample sizes are "small" and asymptotic assumptions do not apply
- DGP assumptions are violated.
- Distributions are complicated.

Usually, we would not use a bootstrap to compute C.I.'s for the mean; in general, the normal distribution works well, as long as N is large enough. The bootstrap is used to generate standard errors for estimates of other statistics where the normal distribution is not a good approximation. A typical example is the median, where for non-normal underlying distributions the SE of the median is complicated to compute.

Efron (1979) is the seminal paper. But, the related literature is older. It became popular in the 1980's due to the explosion of computer power.

Disadvantages and Advantages:

- Disadvantage: Only consistent results, no finite sample results.
- Advantage: Simplicity.

# C.I. Application: Value-at-Risk

What is the most an investor can lose with a particular investment over a given time framework? Or, what is the worst case scenario? *Value-at-Risk* (VaR) provides one answer to this question: It gives a (lower) bound with a probability attached to it.

So far, we have measured risk of an asset/investment with its volatility.

Volatility is calculated including positive (right tail) and negative (left tail) returns. Investors, however, love the right tail of the returns distribution, but dislike the other tail. *Value-at-Risk* (VaR) focuses on the left tail.

VaR gives a formal definition of "worst case scenario" for an asset over a period of time.

VaR: *Maximum expected amount (loss)* in a given time interval within a (*one-sided*)  $(1 - \alpha)$ % C.I.:

 $VaR(1 - \alpha) = Amount exposed * (1 + worst % change scenario in C.I.)$ 

It is common to express the "*expected loss*" relative to today's expected value of asset/investment:

VaR-mean $(1 - \alpha) = VaR - E[Amount exposed]$ 

There are different ways to compute the worst case scenario within a time interval. We go over two approaches:

- Assuming a probability distribution (normal, in our case).
- Using the empirical distribution (a bootstrap, using the past).

**Example**: Let  $\alpha = .025$ 

VaR = Amount exposed \* (1 + worst change scenario in 97.5% C.I.).VaR-mean(97.5%) = VaR - E[Amount exposed].



### VaR: One-sided 97.5% C.I.

# C.I. Application: Range Estimates & VaR for Transaction Exposure - Normal

When a company is involved with transactions denominated in foreign currency (FC), it is exposed to *currency risk*. *Transaction exposure* (TE) provides a simple measure of this exposure:

 $TE_t$  = Value of a fixed future transaction in FC \*  $S_t$ 

where  $S_t$  is the exchange rate expressed as units of domestic currency (USD for us) per unit of FC (say, EUR).

**Example**: A Swiss company, Swiss Cruises, sells packages in USD. Amount = USD 1 million. Payment: 30 days.  $S_t = 0.92$  CHF/USD

### $\Rightarrow$ *TE*<sub>t</sub> = USD 1M \* 0.92 CHF/USD = CHF 0.92M.

If  $S_t$  is described by a Random Walk ( $E[S_{t+T}]=S_t$ ), then  $TE_t$  is a forecast of the value of the transaction in 30 days ( $TE_{t+30}$ ).

Swiss Cruises wants a measure of the uncertainty related to the amount to receive in CHF in 30 days, since  $S_{t+30}$  is unknown.

We can use a range to quantify this uncertainty; we want to say

 $TE_{t+30} \in [\text{TE}_{\text{LB}}, \text{TE}_{\text{UB}}].$  with high probability. To determine this range for TE, we assume that (log) changes in  $S_t$ ,  $e_{f,t}$ , are normally distributed:  $e_{f,t} \sim N(\mu, \sigma^2)$ .

Then, we build a  $(1-\alpha)$ % interval around the mean:  $[\mu \pm z_{1-\alpha/2} * \sigma]$ .

Usual  $\alpha$ 's in interval calculations:  $\alpha = .05 \implies |z_{.025}| = 1.96 \ (\approx 2)$  $\alpha = .02 \implies |z_{.01}| = 2.33$ 

As usual, we estimate  $(\mu, \sigma)$  using  $(\overline{X}, s)$ .

**Example**: Range estimate based on a Normal distribution.

Assume Swiss Cruises believes that CHF/USD monthly changes  $(e_{f,t})$  follow a normal distribution. Swiss Cruises estimates the mean and variance using the last 15 years of data:  $\bar{X} =$  Monthly mean = -0.00152  $\approx$  -0.15% s<sup>2</sup> = Monthly variance = 0.001014 ( $\Rightarrow$  s = 0.03184, or 3.18%)

 $e_{f,t} \sim N(-0.00152, 0.03184^2),$   $e_{f,t} = CHF/USD$  monthly log changes.

Swiss Cruises constructs a 95% CI for CHF/USD monthly changes.

Recall that a 95% C.I. for  $e_{f,t+30}$  (which applies to any t) is given by:  $e_{f,t} \in [-0.00152 \pm 1.96 * 0.03184] = [-0.06393; 0.06089].$  Based on this range for  $e_{f,t}$ , we build a 95% C.I. for  $S_{t+30}$  and, then, for  $TE_{t+30}$  (= **USD 1M** \*  $S_{t+30}$ ).





Now, we derive a range for  $S_{t+30}$ : (A) Upper bound

 $S_{t+30,UB} = S_t^* (1 + e_{f,t,UB}) = 0.92 \text{ CHF/USD} * (1 + 0.06089) = 0.97602 \text{ CHF/USD}$ 

(B) Lower bound  $S_{t+30,LB} = S_t^* (1 + e_{f,t,LB})) = 0.92 \text{ CHF/USD} * (1 + (-0.06393))] = 0.86118 \text{ CHF/USD}$  $\Rightarrow S_{t+30} \in [= 0.86118 \text{ CHF/USD}; 0.97602 \text{ CHF/USD}].$ 

Finally, we derive the bounds for the TE:

(A) Upper bound (with  $S_{t+30,UB} = 0.97602$  CHF/USD) TE<sub>UB</sub>: USD 1M \* [0.97602 CHF/USD] = CHF 976,019.

(B) Lower bound (with  $S_{t+30,LB} = 0.86118 \text{ CHF/USD}$ ) TELB: USD 1M \* [0.86118 CHF/USD] = CHF 861,184.  $\Rightarrow$  TE<sub>t+30</sub>  $\in$  [CHF 0.861M; CHF 0.976M]. ¶

• The lower bound, for a receivable, represents the worst case scenario within the interval in 30 days. That is, this is the VaR:

VaR = Amount exposed \* (1+ worst % change scenario in C.I.) =  $TE_t * (1 + e_{f,t,LB})$ 



VaR(97.5%): Minimum revenue within a 97.5% C.I.

It is common to express the "*expected loss*" relative to today's expected value of transaction (or asset):

VaR-mean = VaR - 
$$TE_t$$
 =  $TE_t^* (1 + e_{f,t,LB}) - TE_t$   
=  $TE_t^* e_{f,t,LB}$ 

Or just

VaR-mean = Amount exposed \* worst case scenario

The minimum revenue to be received by SC in the next 30 days, within a 97.5% CI. VaR(97.5%) = CHF 0.92M \* [1+ (-0.06393)]= CHF 0.8612M.

Interpretation of VaR: If SC expects to cover expenses with this USD 1M inflow, the maximum amount in CHF to cover, within a 97.5% one-sided CI, should be CHF 0.8612M.

Relative to today's valuation (or *expected valuation*, according to RWM), the maximum expected loss in 30 days within a 97.5% one-sided C.I. is:

VaR-mean(.975) = CHF 0.8612M - CHF 0.92M = CHF -0.0927M.

Note that we can also compute the VaR-mean as: VaR-mean(.975)= CHF 0.92M \* (-0.06393)= CHF -0.0588M. ¶

Technically speaking, the VaR is a *quantile*, where a quantile is the fraction of observations that lie below a given value (in this case the VaR).

**Example**: In the previous example, the 0.025 quantile (or 2.5% quantile) for expected loses is **CHF -0.0588M**.



<u>Note</u>: We could have used a different quantile –i.e. a different significant level- to calculate the VaR, for example  $1\% \implies z_{.01} = 2.33$ ). Then,

VaR(99%) = CHF 0.92M \* [1+ (-0.00152 - 2.33 \* 0.03184)]= CHF 0.92M \* [1 + (-0.0757072)]= CHF 0.8503M (A more*conservative*bound.)

 $\Rightarrow$  VaR-mean (.99) = CHF 0.92M \* (-0.0757072) = CHF -0.0697M.

<u>Interpretation of VaR-mean</u>: Relative to today's valuation (or *expected valuation*, according to RWM), the maximum *expected loss* with a 99% "chance" is **CHF -0.0697M**.

<u>Note</u>: As the C.I. gets wider, Swiss Cruises can spend less CHF on account of the **USD 1M** receivable.

### C.I. Application: Range Estimates & VaR for TE - Bootstrap

VaR is a statistic –a function of the data, in our case, a function of  $e_{f,t}$ . We can do an empirical bootstrap to calculate the mean, SE (=SD), C.I., etc.

**Example:** We want to calculate the average VaR(97.5%) and its S.E., using all CHF/USD data from 1990:Jan - 2020:Sep. Then,

```
T \le \text{length}(S)

Tstart \le 229

SP \le S[Tstart: T]

T \le \text{length}(SP)

Val \le 1000000

S_0 \le S[T]

e_f \le \log(SP[-1]/SP[-T])

T_s \le -\log(e_f)

alpha \le .05

T_s \ low \le -round(T_s*alpha/2)

TE_o \le -Val*S_0*(1+e_f)

STE_o \le -sort(TE_o)

VaR_o \le STE_o[T_s_low]

> VaR_o

[1] 860293
```

# Check total T (1971:1 to 2020:9)
# Start of sample period: 1990:1
# FX rate during relevant period (1990:1 on)
# Value of transaction in FC (in M)

# FX Rate at T (Today's S\_t) # Log changes in FX Rate

# Specify alpha level for VaR
# Obs corresponding to alpha/2\*T\_s
# calculate Original TE values
# sort Original TE
# Original VaR

# calculate R TE values
# sort TE

library(boot) sim\_size <- 1000 boot.samps <- boot(data=e\_f, statistic=varisk, R=sim\_size) > boot.samps

ORDINARY NONPARAMETRIC BOOTSTRAP

Call: boot(data = e\_f, statistic = varisk, R = sim\_size)

Bootstrap Statistics : original bias std. error t1\* 860293 1929.305 4870.733

> boot.ci(boot.samps, type = "basic") # boot computes the CI. BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS Based on 1000 bootstrap replicates

CALL : boot.ci(boot.out = boot.samps, type = "basic")

Intervals :

```
Level Basic
95% (849352, 867587)
```

> mean(boot.samps\$t)
[1] 862222.3
> sd(boot.samps\$t)
[1] 4870.733

hist(boot.samps\$t, xlab="VaR (in CHF)", breaks=30)

#### Histogram of boot.samps\$t



Bootstrap estimated VaR(97.5%) = CHF 0.8622M.

# **C.I. Application: Performance Evaluation**

In the 1990s, Bankers Trust evaluated traders based on a risk-adjusted performance measure called RAROC: Risk-adjusted return on capital.

RAROC = Profits/Capital-at-Risk

RAROC adjusts profits taking into account the exposure of the bank, called *capital-at-risk*. BT defined this exposure as the amount of capital needed to cover 99% of the maximum expected loss over a year.

That is, capital-at-risk is the worst loss within one-sided 99% C.I. We called this VaR-mean(99%).

The rationale for this measure: BT needs to hold enough cash to cover 99% of possible losses.

**Example:** Ranking two traders I and II, dealing in different markets.

	Segment	Profits (in USD, annualized)	Position (in USD)	Volatility (annualized)
Trader I	Futures stock indices	3.3 M	45 M	21%
Trader II	FX Market	3.0 M	58 M	14%

To calculate RAROC, we calculate the VaR-mean(99%) -i.e., worst possible loss in a 99% CI.

Assuming normality for profits with mean equal to zero (not important, since all traders are evaluated using the same mean). Then, (since  $\alpha = .01 \Rightarrow z_{.01} = 2.33$ ):

VaR-mean(99%) = Amount exposed \* worst case scenario

= Position \* **z**.01 \* Volatility

Since  $\alpha = .01 \Rightarrow \mathbf{z}_{.01} = \mathbf{2.33}$ .

(1) Calculate VaR-mean (99%) for each trader (under normal distribution) **Trader I:** USD 45M \* 2.33 \* 0.21 = USD 22,018,500. **Trader II:** USD 58M \* 2.33 \* 0.14 = USD 18,919,600.

(2) Calculate RAROC:

**Trader I**: RAROC = **USD 3.3M** /USD 22,018,500 = .1499.

**Trader II**: RAROC = **USD 3.0M** /USD 18,919,600 = .1586.

Conclusion: Once adjusted for risk, Trader II provided a better return.
## Lecture 3 – Least Squares

So far, we have focused on one RV at a time, say stock returns, and learning about its distribution, for example, using descriptive statistics. In econometrics, we are more interested in describing or measuring the expected effect of a tax on consumption or the expected effect of education on an employee's salary or a CEO's compensation.

That is, we usually care about a functional relation between *y*, the *dependent variable*, and *x*, a set of *explanatory variables*.

In this lecture, we **linearly** relate *y* to x & an *error term*,  $\varepsilon$ :

 $y_i = \alpha + \beta x_i + \varepsilon_i,$  i = 1, 2, ..., N,

where  $\alpha \& \beta$  are parameters to be estimated and  $\varepsilon$ , the error term or *disturbance*, has zero mean and constant variance,  $\sigma^2$ . That is,  $\varepsilon$  is a RV with  $E[\varepsilon_i] = 0 \& Var[\varepsilon_i] = \sigma^2$ .

We think of  $\varepsilon$  as the effect of individual variation that have not been "controlled for" with *x*. The disturbance  $\varepsilon_i$  is part of the model. Even if we know that the relation between *y* and *x* is linear and we also know  $\alpha \& \beta$  with certainty –i.e, no need to estimate them- we still would not be able to compute *y* with 100% accuracy.

We call the above equation the *Data Generating Process* (**DGP**), that is, the data we observe  $(y_i \text{ and } x_i)$  are generated following this equation.

Given the always present uncertainty, we focus on expected values. Then, under the assumption  $E[\varepsilon_i] = 0$ , we have:

 $\mathbf{E}[y_i] = \alpha + \beta \mathbf{E}[x_i].$ 

We have a linear relation between the expected value of y and the expected value of x.

**Example:** The CAPM posits a relation between the excess return of asset *i*,  $y_i = r_{i,t} - r_f$ , and the excess return of the market,  $x_i = r_{m,t} - r_f$ . In equilibrium, the CAPM states:

$$E[(r_{i,t} - r_f)] = \beta_i E[(r_{m,t} - r_f)],$$

where  $\beta_i$  is the sensitivity of asset *i* to market risk.

**CAPM DGP:**  $y_i = \alpha + \beta_i x_i + \varepsilon_i$ , i = 1, 2, ..., N. where  $\alpha \& \beta_i$  are parameters to be estimated. Once we estimate  $\alpha \& \beta$ , we can test the CAPM for IBM, since according to the CAPM  $\alpha = 0$ .

The previous mathematical structure allows us to estimate  $\beta_{\text{IBM}}$  and also test the CAPM for asset i = IBM. Define y = excess returns for IBM

x = excess returns for the market (the "Market").

Using time series data, we express the underlying relation behind the CAPM as:

 $y_t = \alpha + \beta x_t + \varepsilon_t, \qquad t = 1, 2, ..., T,$ 

where  $\alpha \& \beta$  are parameters to be estimated and  $\varepsilon$  is the *error term* with  $E[\varepsilon] = 0 \& Var[\varepsilon] = \sigma^2$ .

Taking expectations:

 $\mathbf{E}[y_t] = \alpha + \beta \mathbf{E}[x_t].$ 

Then, once we estimate  $\alpha \& \beta$ , we can compute the expected excess return for IBM. We can also test the CAPM for IBM, since according to the CAPM  $\alpha = 0$ . That is, we test: H<sub>0</sub>:  $\alpha = 0$  vs H<sub>1</sub>:  $\alpha \neq 0$ . ¶

To gain intuition and easy interpretation of the model, it is useful to think of x as given or *predetermined* (realized before y) variable. Then, we can express the relation between y & x, in terms of the conditional expectation of y, conditioning on the predetermined value of x:

 $E[y_i|x_i] = \alpha + \beta x_i.$  ("Regression equation")

The conditional expectation of y is what we model; in general, based on finance theory or the experience of the practitioner. To be technically precise, for the regression equation we require  $E[\varepsilon_i|x_i] = 0$ .

Note: We started with:	$y_i = \alpha + \beta x_i + \varepsilon_i$
which can be converted to:	$y_i = \mathrm{E}[y_i   x_i] + \varepsilon_i$

That is,  $y_i$  is what we model plus something unexpected, a surprise.

• In the CAPM example above, we have that IBM excess returns are only related to ("explained by") the market excess returns. This is a *one variable model*.

But, we could have used a DGP with more explanatory variables, for example the 3 factors in the standard Fama-French (1993) model: Excess market returns (market factor), SMB (size factor), and HML (book-to-market factor). This represents a *multivariate model* for IBM returns: **Fama-French DGP**:  $y_i = \alpha + \beta_1 x_{1,i} + \beta_2 x_{2,i} + \beta_3 x_{3,i} + \varepsilon_i$ 

Though not necessary correct, we usually think of y as the *endogenous* variable and x as the *exogenous* variable, determined "outside" the model. (If x is not endogenous, we have a lot of issues that will force us to change the model or the way we estimate the model.)

• The goal of this lecture is to learn how to estimate the population parameters  $\alpha \& \beta$  and, at the same time, learn the properties of estimators.

<u>Technical Note</u>: We can study the joint distribution of y & x, f(x, y) and describe the joint behavior in terms of expectations, conditional expectations, correlations, etc. For example, assuming joint normality for y & x, we can derive the conditional expectation of y, given x:  $E[y_i|x_i] = \alpha + \beta x_i,$ which gives a functional (& linear) relation between y & x. Thus, in a joint normality context, we can study the effect of a change in x on y. Moreover, after a lot of manipulations and applying statistical definitions, we get a formula to estimate  $\alpha \& \beta$  in terms of moments of y & x.

Then, why do we need other methods to estimate  $\alpha \& \beta$ ? Two things to consider:

- 1) In general, assuming joint normality is not realistic in economics and finance.
- 2) In many situations, we think of the explanatory variable, x, as control, not necessarily as RV.

<u>Remark</u>: Without making any reference to a joint distribution, we will derive the formulas to estimate parameters in a linear relation.

#### **LS Estimation – OLS**

Old method: Gauss (1795, 1801) used it in astronomy.

<u>Idea</u>: We relate a dependent variable y to a set of k explanatory variables x. This function depends on unknown parameters,  $\theta$ , which we want to estimate. The relation between y and x is not exact. There is an error,  $\varepsilon$ . We have T observations of y and x.

The model is:

$$y_i = f(x_{1,i}, x_{2,i}, ..., x_{k,i}; \theta) + \varepsilon_i, \quad i = 1, 2, ..., T.$$

If the functional form is known, we estimate the parameters  $\theta$  by minimizing a sum of squared errors:

$$\min_{\theta} \{ S(x_{1,i}, x_{2,i}, \dots, x_{k,i}; \theta) = \sum_{i}^{T} \varepsilon_{i}^{2} = \sum_{i}^{T} (y_{i} - f(x_{1,i}, x_{2,i}, \dots, x_{k,i}; \theta))^{2} \}$$

The estimator obtained is called the Least Squares (LS) estimator.

LS is a general estimation method. It can be applied to almost any function.

The functional form,  $f(x_i, \theta)$ , is dictated by theory or experience. In this class, we mainly work with the **linear** case:

$$f(x_i, \theta) = \beta_l \ x_{1,i} + \beta_2 \ x_{2,i} + \beta_3 \ x_{3,i} + \dots + \beta_k \ x_{k,i}$$

Now, we estimate the vector  $\mathbf{\theta} = \{\beta_1, \beta_2, \dots, \beta_k\}$  by minimizing

 $S(\mathbf{x}, \boldsymbol{\theta}) = \sum_{i}^{T} \varepsilon_{i}^{2} = \sum_{i}^{T} (y_{i} - \beta 1 x_{1,i} - \beta 2 x_{2,i} - \dots - \beta k x_{k,i})^{2}$ In this case, we call this estimator the *Ordinary Least Squares* (OLS) estimator. (Ordinary Linear functional form.)

<u>Notation</u>: In lecture 2, we used  $\wedge$  over the estimator of the parameter of interest. For example,  $\hat{\theta}$  is the estimator of the parameter  $\theta$ . Sometimes, to emphasize the method of estimation, we add to the estimated parameter the initials of the method used, say  $\hat{\theta}_{LS}$ .

For historical reasons, in the linear model, **b** is popularly used to denote the OLS estimator of  $\beta$ .

**Example 1**: We want to study the effect of the tech boom (x) on the San Francisco housing market (y). We rely on a simple linear model, with only one explanatory variable, the tech boom variable. That is,

$$y_i = \alpha + \beta x_i + \varepsilon_i.$$

In this model, we are interested in estimating  $\beta$ , our parameter of interest.  $\beta$  measures the *marginal effect* of *x* on *y*. We can use the estimate of  $\beta$  to check if the high tech boom has a positive effect on SF housing prices. In this case we test:

H<sub>0</sub> (No or Negative effect):  $\beta \leq 0$ .

H<sub>1</sub> (Positive effect):  $\beta > 0$ .

We have monthly data on SF Housing Prices and a Tech Indicator, developed by the Federal Reserve. We transform the data in percentage changes. Below, we plot the data: SF House Prices vs Tech Indicator (both in % changes).



SF House Prices & Tech Indicator

**Example 2**: We want to study the effect of a CEO's education (x) on a firm's CEO's compensation (y). We build a CEO's compensation model including a CEO's education (x) and other "*control variables*" (*W*: experience, gender, etc.), controlling for other features that make one CEO's compensation different from another. That is,

$$y_i = f(x_i, W_i, \theta) + \varepsilon_i,$$
  $i = 1, 2, ..., T.$ 

The term  $\varepsilon_i$  represents the effects of individual variation that have not been controlled for with  $W_i$ , or  $x_i$  and  $\theta$  is a vector of parameters.

Usually,  $f(x, \theta)$  is linear. Then, the compensation model becomes:

 $y_i = \alpha + \beta x_i + \gamma_1 W_{1,i} + \gamma_2 W_{2,i} + \dots + \varepsilon_i$ 

Again, in this model, we are interested in the estimation of  $\beta$ , our parameter of interest, which measures the effect of a CEO's education on a CEO's compensation. We can use the estimate of  $\beta$  to check if CEO's education has a positive effect on CEO's compensation. In this case we test:

H<sub>0</sub> (No or Negative effect):  $\beta \le 0$ .

H<sub>1</sub> (Positive effect):  $\beta > 0$ . ¶

#### LS Estimation – General Functional Form

We start with a general functional form,  $f(x_i, \theta)$ , where  $\theta$  is a vector of k parameters.

The general model:

$$y_i = f(x_i, \theta) + \varepsilon_i$$

We want to estimate *k* parameters.

Objective function:

$$S(x_{i}, \boldsymbol{\theta}) = \sum_{i}^{T} \varepsilon_{i}^{2} = \sum_{i}^{T} \{y_{i} - f(x_{i}, \boldsymbol{\theta})\}^{2}$$
  
=  $(y_{1} - f(x_{1}, \boldsymbol{\theta}))^{2} + (y_{2} - f(x_{2}, \boldsymbol{\theta}))^{2} + \dots + (y_{T} - f(x_{T}, \boldsymbol{\theta}))^{2}$ 

• We minimize  $S(x_i, \theta)$  with respect to  $\theta$ . That is,

min<sub>$$\theta$$</sub> { S(x<sub>i</sub>,  $\boldsymbol{\theta}$ ) =  $\sum_{i}^{T} \varepsilon_{i}^{2} = \sum_{i}^{T} \{y_{i} - f(x_{i}, \boldsymbol{\theta})\}^{2}$ }

$$\Rightarrow \frac{\partial S(x_i,\theta)}{\partial \theta} = 2 \{ y_1 - f(x_1, \theta) \} (-f'(x_1, \theta)) + \dots + 2 \{ y_T - f(x_T, \theta) \} (-f'(x_T, \theta)) \\ = -2 \sum_i^T \{ y_i - f(x_i, \theta) \} f'(x_i, \theta) \}$$

f.o.c. 
$$\Rightarrow -2\sum_{i}^{T} \{y_{i} - f(x_{i}, \hat{\theta}_{LS})\} f'(x_{i}, \hat{\theta}_{LS})\} = 0$$
$$\Rightarrow \sum_{i}^{T} \{y_{i} - f(x_{i}, \hat{\theta}_{LS})\} f'(x_{i}, \hat{\theta}_{LS})\} = 0$$

Suppose we have q elements in  $\theta$ , the f.o.c.'s have set up a qxq system of equations. This system of equations is called the *normal equations*. The solution to the normal equation,  $\hat{\theta}_{LS}$ , is the Least Squares estimator.

We do not always can solve analytically the normal equations. Two cases:

- When  $f(x_i,\theta)$  is linear, we have an analytic, explicit solution, the OLS estimator,  $\hat{\theta}_{OLS} = \mathbf{b}$ . - When  $f(x_i,\theta)$  is *non-linear*, we **do not** have an explicit solution for  $\hat{\theta}_{LS}$ . The system, however, can be solved numerically. In this case, the estimator is usually referred as Non-linear Least Squares estimator,  $\hat{\theta}_{NLLS}$ .

The estimator  $\hat{\theta}_{LS}$  is a function of the data (y<sub>i</sub>, x<sub>i</sub>).

#### **OLS Estimation – One Variable Model**

One explanatory variable in a linear model:

$$f(x_i, \theta) = \beta_1 + \beta_2 x_i$$

Linear Model:  $y_i = \beta_1 + \beta_2 x_i + \varepsilon_i$ 

We have two parameters to estimate.

Objective function:

S(**x**; 
$$\beta_1$$
,  $\beta_2$ ) =  $\sum_i^T \varepsilon_i^2 = \sum_i^T (y_i - \beta_1 - \beta_2 x_i)^2$   
=  $\{y_1 - \beta_1 - \beta_2 x_1)^2 + \{y_2 - \beta_1 - \beta_2 x_2)^2 + \dots + \{y_T - \beta_1 - \beta_2 x_T)^2$ 

First, we take first derivatives:

( $\beta_1$ ):  $2 \sum_i^T (y_i - \beta_1 - \beta_2 x_i)$  (-1) ( $\beta_2$ ):  $2 \sum_i^T (y_i - \beta_1 - \beta_2 x_i)$  (- $x_i$ )

Second, we set the f.o.c. and get the normal equations (2 equations, 2 unknowns):

$$\begin{array}{ll} (\beta_1): & 2\sum_i^T (y_i - b_1 - b_2 x_i) \ (-1) = 0 \\ (\beta_2): & 2\sum_i^T (y_i - b_1 - b_2 x_i) \ (-x_i) = 0 \end{array} \qquad \Rightarrow \sum_i^T (y_i - b_1 - b_2 x_i) = 0 \qquad (1) \\ \Rightarrow \sum_i^T (y_i x_i - b_1 x_i - b_2 x_i^2) = 0 \qquad (2) \end{array}$$

Now, we solve for  $b_1 \& b_2$ , the OLS estimators:

From (1):  

$$\sum_{i}^{T} y_{i} - \sum_{i}^{T} b_{1} - b_{2} \sum_{i}^{T} x_{i} = 0$$

$$\Rightarrow \sum_{i}^{T} y_{i} - T b_{1} - b_{2} \sum_{i}^{T} x_{i} = 0$$

$$\Rightarrow b_{1} = \overline{y} - b_{2} \overline{x}$$
From (2):  

$$\sum_{i}^{T} y_{i} x_{i} - (\overline{y} - b_{2} \overline{x}) \sum_{i}^{T} x_{i} - b_{2} \sum_{i}^{T} x_{i}^{2} = 0$$

$$\Rightarrow \sum_{i}^{T} y_{i} x_{i} - \overline{y} \sum_{i}^{T} x_{i} - b_{2} (\sum_{i}^{T} x_{i}^{2} - \overline{x} \sum_{i}^{T} x_{i}) = 0$$

$$\Rightarrow b_{2} = \frac{\sum_{i}^{T} (y_{i} - \overline{y}) x_{i}}{\sum_{i}^{T} (x_{i} - \overline{x}) x_{i}}$$

or, more elegantly,

$$b_{2} = \frac{\sum_{i}^{T} (y_{i} - \bar{y})(x_{i} - \bar{x})}{\sum_{i}^{T} (x_{i} - \bar{x})^{2}} = \frac{cov(y_{i}, x_{i})}{var(x_{i})}$$

Note that we need  $var(x_i) \neq 0$  to get b<sub>2</sub>.

• Interpretation of coefficients

-  $b_1$  estimates the *constant* of the regression, the value of  $y_i$ , when  $x_i$  equals to 0.

- b<sub>2</sub> estimates the *slope* of the regression, the marginal effect –i.e., the first derivative of  $y_i$  with respect to  $x_i$ :

$$\frac{\delta y_i}{\delta x_i} = \beta_2$$

That is, if x increases by one unit (say, 1%), then, y is estimated to increase by  $b_2$  units (say,  $b_2$ %).

• Conditional Prediction

Suppose analysts estimate that  $x_i$  will be z%, then, you estimate (or predict, given the z% value of  $x_i$ ) y<sub>i</sub>:

Predicted[ $y_i | x_i = \mathbf{z}\%$ ] =  $\mathbf{b}_1 + \mathbf{b}_2 * \mathbf{z}$ .

## **OLS Estimation – One Variable Model: CAPM**

As mentioned in the introduction, a typical finance application of a one variable linear model is the CAPM. Recall that the (Sharpe-Litner) CAPM, in equilibrium, implies:

 $E[(r_{i,t} - r_f)] = \beta_i E[(r_{m,t} - r_f)],$ 

where

 $r_{i,t}$  = return on asset *i* at time *t*.  $r_f$  = return of riskless asset at time *t*.

 $r_{m,t}$  = return on the market portfolio at time *t*.

 $\beta_i$  = asset *i*'s sensitivity to market (systematic) risk.

<u>Note</u>: The market portfolio in the CAPM represents wealth. All wealth. We need to include not only all stocks, but all bonds, real estate, privately held capital, publicly held capital (roads, universities, etc.), and human capital in the world. (Easy to state, but complicated to form.) In general, we proxy the Market Portfolio, with a well-diversified index, that only includes equities, like the S&P 500 Index, or the MSCI World Index.

The CAPM is a particular case of what in financial theory we call "*factor models*." Factors represent the systematic component that drives the cross-section of returns over time; they can be observed or unobserved. For example, a *k-factor model* for returns is given by:

 $r_{i,t} = \alpha + \beta_1 f_{1,t} + \beta_2 f_{2,t} + ... + \beta_k f_{k,t} + \varepsilon_{i,t}$ where  $f_{j,t}$  is the *j* (common) factor at time *t*, and constant over *i*, and  $\varepsilon_{i,t}$  represents the idiosyncratic component of asset *i*.

Thus, we think of returns as driven by common or systematic factors (undiversifiable) and idiosyncratic factors (diversifiable in large portfolios.) Thus, in equilibrium, investors get compensated only for the systematic risk they take.

The CAPM has only one factor: market excess returns ("*the market*"). The higher the exposure to this factor –i.e.,  $\beta_i$ -, the higher the expected compensation.

• A linear data generating process (DGP) consistent with the CAPM is:

 $(r_{i,t} - r_f) = \alpha_i + \beta_i (r_{m,t} - r_f) + \varepsilon_{i,t}, \qquad i = 1, ..., N \& t = 1, ..., T$ where

 $\alpha_i$  and  $\beta_i$  are the coefficients to be estimated by LS.  $\operatorname{Cov}(r_{m,t}, \varepsilon_{i,t}) = 0$  -i.e., market returns are *exogenous*.

If  $\beta_i = 0$ , asset *i* is not exposed to market risk. Thus, the investor is not compensated with a higher return than  $r_f$ .

If  $\beta_i > 0$ , asset *i* is exposed to market risk &  $r_{i,t} \ge r_f$ , provided that  $E[(r_{m,t} - r_f)] > 0$ .

If  $\beta_i > 1$  ( $\beta_i < 1$ ), asset *i* is "*riskier*" ("*safer*") than the market. That is, the expected return for asset *i* is higher (lower) than the expected return for the market. For  $\beta_i > 1$ , we have  $E[(r_{i,t} - r_f)] > \beta_i E[(r_{m,t} - r_f)]$  -higher compensation for higher risk.

If  $\alpha_i > 0$ , then asset *i* has higher expected returns than what is expected in equilibrium –i.e., what the CAPM implies.

• Then, in our linear model let  $y_i$  represent IBM excess returns  $(r_{IBM,t} - r_f)$  at time *t* and let  $x_i$  represent Market excess returns (say,  $r_{m,t} - r_f$ ) at time *t*. Then, b<sub>2</sub> estimates IBM's beta in the CAPM.

Then, b<sub>2</sub> (=  $\hat{\beta}_{CAPM}$ ) estimates the stock's beta in the CAPM; in the IBM case, we have :

$$\mathbf{b}_2 = \widehat{\boldsymbol{\beta}}_{IBM} = \frac{cov(r_{IBM,t} - r_f, r_{m,t} - r_f)}{var(r_{m,t} - r_f)}$$

That is, the CAPM  $\beta$  is the ratio of a covariance over a variance.

Recall that the CAPM  $\beta$  measures a stock's risk in relation to the risk (volatility) of the market. That is, we think of  $\beta$  as a measure of the relative risk exposure of holding a particular stock (IBM, in this case) in relation to the market.

• Interpretation of coefficients

-  $b_1$  estimates the *constant* of the regression: IBM excess returns in excess of Market excess returns. In the CAPM, it should be  $0 (= \alpha_i)$ .

- b<sub>2</sub> estimates the *slope* of the regression. In the CAPM:  $\beta_{i=IBM}$ 

$$\frac{\delta y_i}{\delta x_i} = \beta_{i=IBM} \Rightarrow \text{estimated by } b_2.$$

That is, if Market excess returns increases by one 1% (unit), then we estimate that IBM excess returns are expected to increase by b<sub>2</sub>% (b<sub>2</sub> units). The  $\beta_{\text{IBM}}$  also tells us if IBM is riskier ( $\beta_{\text{IBM}} > 1$ ) or safer ( $\beta_{\text{IBM}} < 1$ ) than the market.

Conditional Prediction

Suppose analysts estimate that Market excess returns will be 10%, then, we estimate (or predict, given the 10% value for Market excess returns):

Predicted 
$$[(r_{IBM,t} - r_f) | (r_{m,t} - r_f) = 10\%] = b_1 + b_2 * .10.$$

We will call the Predicted  $y_i = \hat{y}_i =$ fitted value.

#### **Example:** We estimate the CAPM for **IBM returns** using *lm* R function.

• Import data with read function

SFX\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/Stocks\_FX\_1973.csv", head=TRUE, sep=",")

• Extract variables from imported data	
x_ibm <- SFX_da\$IBM	# extract IBM price data
x_Mkt_RF <- SFX_da\$Mkt_RF	# extract Market excess returns (in %)
x RF <- SFX da\$RF	# extract Risk-free rate (in %)
	1 1 1 1

• Define log returns & adjust size of varia	ables accordingly
$T \leq length(x_ibm)$	# sample size
$lr_ibm \le log(x_ibm[-1]/x_ibm[-T])$	# create IBM log returns (in decimal returns)
$Mkt_RF \leq x_Mkt_RF[-1]/100$	# Adjust sample size to (T-1) by removing 1 <sup>st</sup> obs
RF <- x_RF[-1]/100	# Adjust sample size and use decimal returns.

Define excess returns and estimate CAPM with lm function. Then, print results to screen with summary function:
 ibm\_x <- lr\_ibm - RF</li>
 fit ibm\_capm <- lm(ibm\_x ~ Mkt\_RF)</li>
 # IBM excess returns
 # lm (=linear model) package

summary(fit\_ibm\_capm) # min(-unear mode # print lm results

> summary(fit\_ibm\_capm)
Call:
lm(formula = ibm x ~ Mkt RF)

Residuals: Min 1Q Median 3Q Max -0.314401 -0.031692 -0.000537 0.031447 0.248201

Coefficients:

Estimate Std. Error t value Pr(>|t|)(Intercept) -0.005791 0.002487 -2.329 0.0202 \*  $b_1 = -0.005791$ xMkt\_RF 0.895774 0.053867 16.629 <2e-16 \*\*\*  $b_2 = 0.895774$ ---Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 0.05887 on 567 degrees of freedom Multiple R-squared: 0.3278, Adjusted R-squared: 0.3266 F-statistic: 276.5 on 1 and 567 DF, p-value: < 2.2e-16

Interpretation of b<sub>1</sub> and b<sub>2</sub>:

 $b_1$  = constant. The additional IBM return, after excess market returns are incorporated, is 0.58% (under the CAPM,  $b_1$  should be close to 0).

**b**<sub>2</sub> = slope. It is the marginal effect. If market excess returns increase by 1%, IBM excess returns increase by .90%. The estimate of the CAPM  $\beta < 1$ , implying that IBM is less volatile ("safer") than the market.

Unconditional (average) Expected IBM excess returns

We observed an average excess market monthly return in the sample: 0.0056489. Then, the expected IBM excess monthly return in the sample was:

-0.005791 + 0.895774 \* 0.0056489 = -0.000731 (0.07%).

Conditional prediction of IBM excess returns:

Suppose market excess returns are expected to be 1% next month, then we predict next month IBM excess returns:

-0.005791 + 0.895774 \* .1 = 0.00317 (0.32%).

Note: According to the CAPM, IBM excess monthly returns should have been:

- IBM excess returns (IBM) = 0.895774 \* mean(Mkt RF)

= 0.895774 \* 0.0056489 = 0.0050601

But, in the sample, we observed

- IBM excess returns = mean(ibm x) = -0.00073141.

That is, IBM underperformed relative to the CAPM.

## LS Estimation – Application 1: The CAPM & The Cost of Equity

As mentioned in Chapter 2, the ERP is central to many financial theories, for example, as illustrated above the CAPM uses the ERP as an input to price assets. The CAPM states that, in equilibrium, the expected excess return for asset *i* is proportional to the expected market excess return or *expected market risk premium (ERP)*, given by  $E[(r_{m,t} - r_f)]$ . That is:

$$\mathbf{E}[(r_{i,t}-r_f)] = \beta_i \mathbf{E}[(r_{m,t}-r_f)].$$

In equilibrium, the cost of equity, ke, is equal to the required (expected) rate of return a firm has to pay to investors/shareholders. Firms need to calculate the cost of equity to estimate the *cost of capital, kc*, which is used to discount the cash flows of a firm or a firm's project. According to the weighted average cost of capital (WACC) method,  $k_c$  is given by:

$$k_c = \frac{D}{D+E}k_d(1-t) + \frac{E}{D+E}k_e$$

where E represents total equity, D is total Debt, t is the effective tax rate and  $k_d$  is the cost of debt.

Firms routinely use expected returns to calculate the cost of equity. For example, using the CAPM we have:

$$k_e = r_f + \beta_i \operatorname{E}[(r_{m,t} - r_f)].$$

Thus, in this case, in order to compute  $k_e$ , a firm needs to determine  $\beta_i$ , the risk-free rate,  $r_f$ , and the Market Portfolio, usually, a local market index, like the S&P 500 or the Nikkei 225, or a global index like the MSCI World Index.

**Example**: Suppose IBM wants to determine its cost of equity. IBM decides to use the CAPM, with a U.S. ERP and U.S  $r_f$ .

Data:

Estimated  $\beta_{\text{IBM}} = 0.895774 \approx 0.90$ Risk-free rate,  $r_f = 4.50\%$ ERP = E[ $(r_{m,t} - r_f)$ ] = 0.0382  $k_{e,IBM} = r_f + \beta_{\text{iBM}}$  E[ $(r_{m,t} - r_f)$ ] = 0.045 + 0.90 \* 0.0382 = 0.07938

The required (or expected) rate of return for IBM investors is **7.938%**. This is what IBM will use as  $k_e$  to determine its cost of capital and, therefore, discount the cash flows associated with new projects.

<u>Note</u>: If IBM decides to use the MSCI World Index as the benchmark for the Market Porfolio, then,

 $k_{e,IBM} = r_f + \beta_{iBM} E[(r_{m,t} - r_f)] = 0.045 + 0.90 * 0.0317 = 0.07353$ 

a smaller number, which would produce a smaller cost of capital and, thus, increase the NPV of IBM or an IBM's project! ¶

Q: Which one should a firm use: a Domestic-based ERP or a World-based ERP? It depends on the view that a company has regarding capital markets. If capital markets are integrated (or if the shareholders are world-wide diversified) the appropriate equity risk premium should reflect a world benchmark (say, MSCI World Index),  $(r_{m,t} - r_f)$ w. But, if markets are segmented (or if the shareholders hold domestic portfolios), then the appropriate equity risk premium should be based on a domestic benchmark (say, the MSCI US Index for U.S. companies),  $(r_{m,t} - r_f)$ D. The risk-free rate should also be adjusted accordingly. Then, using the CAPM we have:

- World CAPM:	$k_{\rm e} = k_{\rm e,W} = r_f, w + \beta w E[(r_{m,t} - r_f)w]$
- Domestic CAPM:	$k_{\rm e} = k_{\rm e,D} = r_{f,\rm D} + \beta_{\rm D}  {\rm E}[(r_{m,t} - r_{f})_{\rm D}]$

The difference between these two models can be considerable. In our previous example we have a 0.585% difference. According to Bruner et al. (2008), on average, there is a 5.55% absolute difference for emerging markets and a 3.58% absolute difference for developed markets.

#### LS Estimation – Application 2: Hedging

In the linear model, we can estimate the optimal hedge ratio using a regression. To see this, we derive the optimal hedge ratio for a position in foreign currency (FC).

Notation:

 $S_t$ : Exchange rate at time t. We use direct quotations, that is, DC units per unit of FC, say  $S_t = 1.30$  USD/GBP.

Ft,T: Forward/Futures price at time t with a T maturity.

*n*<sub>s</sub>: Number of units of foreign currency held.

*n*f: Number of futures foreign exchange units held (opposite position).

 $\pi_{h,t}$ : (Uncertain) profit of the hedger at time t.

 $\Delta X$ : Change in X (= X<sub>t</sub> - X<sub>t-1</sub>)

**h** = hedge ratio =( $n_f/n_s$ )= Number of futures per spot in position.

We want to calculate  $h^*$  (optimal h): We minimize the variability of  $\pi_{h,t}$ .

 $\pi_{\rm h,T} = \Delta S_{\rm t} \, n_{\rm s} + \Delta F_{\rm t,T} \, n_{\rm f} \qquad ({\rm Or}, \, \pi_{\rm h,T}/n_{\rm s} = \Delta S_{\rm t} + \frac{\hbar}{\hbar} \, \Delta F_{\rm t,T}.)$ 

We want to select *h* to minimize:

 $\operatorname{Var}(\pi_{h,T}/n_{s}) = \operatorname{Var}(\Delta S_{T}) + h^{2} \operatorname{Var}(\Delta F_{t,T}) + 2 h \operatorname{Covar}(\Delta S_{T}, \Delta F_{t,T})$  $= \sigma_{s}^{2} + h^{2} \sigma_{F}^{2} + 2 h \sigma_{sF}$ 

f.o.c.

$$2 \mathbf{h}^* \sigma_{\mathrm{F}}^2 + 2 * \sigma_{\mathrm{SF}} = 0$$
$$\implies \mathbf{h}^* = -\sigma_{\mathrm{SF}}/\sigma_{\mathrm{F}}^2$$

<u>Note</u>: A covariance over a variance. It can be estimated by LS:  $\Delta S_t = \beta_1 + \beta_2 \Delta F_{t,T} + \varepsilon_t \implies b_2 \text{ estimates } h^*.$ 

**Example:** In March, we are long a GBP 1M position. We are uncertain about St in the next 90

**Example:** In March, we are long a GBP IM position. We are uncertain about S<sub>t</sub> in the next 90 days. We hedge this position using June GBP futures (size of contract = GBP 62,500). We want to determine  $h^*$ .

Get Data (St and Ft,90-days , for say 10 years). Do a regression.  $\Delta S_t = \beta_1 + \beta_2 \, \Delta F_{t,T} + \epsilon_t$ 

Suppose we estimate this regression:

 $\Delta S_t = .001 + .82 \Delta F_{t,T},$  $\Rightarrow h^* = -.82.$ 

Now, we determine the number of June GBP futures contracts:

 $\Rightarrow n_{\rm f}/{\rm size of the contract} = \frac{h * n_{\rm s}}{1,000,000} / \frac{62,500}{62,500} = -13.12 \approx 13 \text{ contracts sold! } \P$ 

## LS Estimation – Multivariate OLS

The CAPM is a particular case of what in financial theory is called "*factor models*." Factors represent the systematic component that drives the cross-section of returns over time; they can be observed or unobserved. For example, a *k*-factor model for returns is given by:

$$f_{i,t} = \alpha + \beta_1 f_{1,t} + \beta_2 f_{2,t} + \ldots + \beta_k f_{k,t} + \varepsilon_i,$$

where  $f_{j,t}$  is the *j* (common) factor at time *t*, and constant over *i*, and  $\varepsilon_{i,t}$  represents the idiosyncratic component of asset *i*.

Thus, we think of returns as driven by common or systematic factors (undiversifiable) and idiosyncratic factors (diversifiable in large portfolios.) Thus, in equilibrium, investors get compensated only for the systematic risk they take.

The CAPM has only one factor: market excess returns ("*the market*"). The higher the exposure to this factor -i.e.,  $\beta_i$ -, the higher the expected compensation.

LS is a general estimation method. It allows any functional form for the relation between  $y_i$  and  $x_i$ , and it allows  $y_i$  to be related to many explanatory variables, like the above mentioned multi-factor models for excess returns.

In this lecture, we cover the case where  $f(x_i, \theta)$  is **linear**. We assume a linear system with k independent variables and T observations. That is,

$$y_i = \beta_1 x_{1,i} + \beta_2 x_{2,i} + ... + \beta_k x_{k,i} + \varepsilon_i,$$
  $i = 1, 2, ..., T$ 

The whole system (for all *i*) is:

$$y_{1} = \beta_{1} x_{11} + \beta_{2} x_{21} + \dots + \beta_{k} x_{k1} + \varepsilon_{1}$$
  

$$y_{2} = \beta_{1} x_{12} + \beta_{2} x_{22} + \dots + \beta_{k} x_{k2} + \varepsilon_{2}$$
  
....  

$$y_{T} = \beta_{1} x_{1T} + \beta_{2} x_{2T} + \dots + \beta_{k} x_{kT} + \varepsilon_{T}$$

It is cumbersome and complicated to write the whole system. Using linear algebra, we can rewrite the system in a more compact and simplify derivations.

For example, after some definitions, we can write the whole system as:

$$y = X \beta + \varepsilon$$

#### **Linear Algebra: Brief Review – Matrix**

Life (& notation) becomes easier with linear Algebra. Concepts:

• Matrix.

A matrix is a set of elements, organized into rows and columns

Columns

Rows	ſa	b
Rowb	$L_{C}$	d

• *a* & *d* are the diagonal elements.

• *b* & *c* are the off-diagonal elements.

Matrices are like plain numbers in many ways: they can be added, subtracted, and, in some cases, multiplied and inverted (divided).

# Linear Algebra: Matrices and Vectors Examples:

$$A = \begin{bmatrix} a_{11} & a_{21} \\ a_{12} & a_{22} \end{bmatrix}; \quad b = \begin{bmatrix} b_1 & b_2 & b_3 \end{bmatrix}.$$

<u>Dimensions of a matrix</u>: numbers of rows by numbers of columns. The Matrix  $\mathbf{A}$  is a 2x2 matrix,  $\mathbf{b}$  is a 1x3 matrix.

A matrix with only 1 column or only 1 row is called a *vector*.

If a matrix has an equal numbers of rows and columns, it is called a *square* matrix. Matrix **A**, above, is a square matrix.

Usual Notation:	Upper case letters	$\Rightarrow$ matrices
	Lower case	$\Rightarrow$ vectors

## **Linear Algebra: Matrices – Information**

Information is described by data. A tool to organize the data is a list, which we call a vector. Lists of lists are called matrices. That is, we organize the data using matrices.

We think of the elements of **X** as data points ("data entries", "observations"), in economics, we usually have numerical data.

We store the data in rows. In a Txk matrix, **X**, over time we build a database:

 $X = \begin{bmatrix} x_{11} & x_{21} & \cdots & x_{k1} \\ \vdots & \vdots & \ddots & \vdots \\ x_{1T} & x_{2T} & \cdots & x_{kT} \end{bmatrix}$ 

Once the data is organized in matrices it can be easily manipulated: multiplied, added, etc. (This is what Excel does).

## **Linear Algebra: Matrices in Econometrics**

In econometrics, we have a model  $y = f(x_1, x_2, ..., x_k)$ , which we want to estimate. We collect data, say *T* (or *N*) observations, on a dependent variable, y, and on *k* explanatory variables, X.

Under the usual notation, vectors will be column vectors:  $\mathbf{y}$  and  $\mathbf{x}_k$  are Tx1 vectors:

$$\mathbf{y} = \begin{bmatrix} y_1 \\ \vdots \\ y_T \end{bmatrix} \qquad \& \qquad x_j = \begin{bmatrix} x_{j1} \\ \vdots \\ x_{jT} \end{bmatrix} \qquad j = 1, \dots, k$$
  
is a *Txk* matrix: 
$$\mathbf{X} = \begin{bmatrix} x_{11} & x_{21} & \cdots & x_{k1} \\ \vdots & \vdots & \ddots & \vdots \\ x_{1T} & x_{2T} & \cdots & x_{kT} \end{bmatrix}$$

Х

Its columns are the k Tx1 vectors  $\mathbf{x}_i$ . It is common to treat  $\mathbf{x}_1$  as vector of ones,  $\mathbf{i}$ .

In general, we import matrices (information) to our programs.

**Example:** In R, we use the **read** function, usually followed by the type of data we are importing. Below, we import a comma separated values (csv) file with monthly CPIs and exchange rates for 20 different countries, then we use the **read.csv** function: PPP da <-

read.csv("https://www.bauer.uh.edu/rsusmel/4397/ppp\_2020\_m.csv",head=TRUE,sep=",")

The **names()** function describes the headers of the file imported (41 headers): > names(PPP\_da) [1] "Date" "BG\_CPI" "IT\_CPI" "GER\_CPI" "UK\_CPI" [6] "SWED\_CPI" "DEN\_CPI" "NOR\_CPI" "IND\_CPI" "JAP\_CPI" [11] "KOR CPI" "THAI CPI" "SING CPI" "MAL CPI" "KUW CPI"

[16] "SUAD CPI" "CAN CPI" "MEX CPI" "US CPI" "EGY CPI" [...]

The **summary()** function provides some stats of variables imported: > summary(PPP da)

Date	BG_CPI IT_CPI GER_CPI
1/15/1971: 1	Min. : 19.77 Min. : 5.90 Min. : 31.20
1/15/1972: 1	1st Qu.: 49.32 1st Qu.: 32.25 1st Qu.: 57.17
1/15/1973: 1	Median : 69.91 Median : 67.30 Median : 75.30
1/15/1974: 1	Mean : 67.92 Mean : 60.14 Mean : 72.29
1/15/1975: 1	3rd Qu.: 89.40 3rd Qu.: 89.65 3rd Qu.: 91.17
1/15/1976: 1	Max. :109.71 Max. :103.50 Max. :106.60
(Other) :588	

We extract a variable from the matrix by the name of file followed by **\$** and the header of variable:

> x\_chf <- PPP\_da\$CHF\_USD # extract CHF/USD exchange rate data</pre>

We can transform the vector x chf. For example, for % changes:

$T \leq - length(x_chf)$	# length of CHF/USD exchange rate data
$lr_chf <- log(x_chf[-1]/x_chf[-T])$	# create log returns (changes) for the CHF/USD.

#### **Linear Algebra: Special Matrices**

• *Identity Matrix,* I: A square matrix with 1's along the diagonal and 0's everywhere else. Similar to scalar "1." A\*I = A

[1	0	[0
0	1	0
Lo	0	1

• *Null matrix*, **0**: A matrix in which all elements are 0's. Similar to scalar "0." A\*0 = 0

[0]	0	[0
0	0	0
Lo	0	0]

Both are *diagonal* matrices  $\Rightarrow$  off-diagonal elements are zero.

Both are examples of *symmetric* matrices. That is, element  $a_{ij}$  is equal to element  $a_{ji}$ . (Later, we'll see  $\mathbf{A} = \mathbf{A}^{T}$ ). For example:

$$\mathbf{A} = \begin{bmatrix} 2 & 5 & 9 \\ 5 & -1 & 0 \\ 9 & 0 & 1 \end{bmatrix}$$
 is a symmetric

#### **Linear Algebra: Multiplication**

We want to multiply two matrices: **A\*B**. But, multiplication of matrices requires a *conformability condition*.

<u>Conformability condition</u>: The <u>column</u> dimensions of the <u>lead</u> matrix  $\mathbf{A}$  must be equal to the <u>row</u> dimension of the <u>lag</u> matrix  $\mathbf{B}$ .

If **A** is an  $(m \ge n)$  and **B** an  $(n \ge p)$  matrix (**A** has the same number of columns as **B** has rows), then we define the product of **AB**. **AB** = **C** is  $(m \ge p)$  matrix with its *ik*-th element is  $c_{ik} = \sum_{j=1}^{n} a_{ij} b_{jk}$ 

Q: What are the dimensions of the vector, matrix, and result?

$$aB = \begin{bmatrix} a_{11}a_{12} \end{bmatrix} \begin{bmatrix} b_{11} & b_{12} & b_{13} \\ b_{21} & b_{22} & b_{23} \end{bmatrix} = c = \begin{bmatrix} c_{11} & c_{12} & c_{13} \end{bmatrix}$$
$$= \begin{bmatrix} a_{11}b_{11} + a_{12}b_{21} & a_{11}b_{12} + a_{12}b_{22} & a_{11}b_{13} + a_{12}b_{23} \end{bmatrix}$$
$$(2) B(2x3) \rightarrow c(1x3)$$

Dimensions: a(1x2),  $B(2x3) \Rightarrow c(1x3)$ 

**Example 1**: We want to multiply A (2x2) and B (2x2), where A has elements  $a_{ij}$  and B has elements  $b_{jk}$ . Recall the  $ik^{ih}$  element is  $\sum_{j=1}^{n=2} a_{ij} b_{jk}$ 

$$\mathbf{A} = \begin{bmatrix} 2 & 1 \\ 7 & 9 \end{bmatrix}$$
$$\mathbf{B} = \begin{bmatrix} 1 & 0 \\ 2 & 3 \end{bmatrix}$$

 $\mathbf{C} = \begin{bmatrix} 2 & 1 \\ 7 & 9 \end{bmatrix} * \begin{bmatrix} 1 & 0 \\ 2 & 3 \end{bmatrix} = \begin{bmatrix} \mathbf{4} = \mathbf{2} * \mathbf{1} + \mathbf{1} * \mathbf{2} & \mathbf{3} = \mathbf{2} * \mathbf{0} + \mathbf{1} * \mathbf{3} \\ \mathbf{26} = \mathbf{7} * \mathbf{1} + \mathbf{9} * \mathbf{2} & \mathbf{27} = \mathbf{7} * \mathbf{0} + \mathbf{9} * \mathbf{3} \end{bmatrix}$  $C_{2x2} = A_{2x2} * B_{2x2}$ 

Dimensions:  $A(2x^2)$ ,  $B(2x^2) \Rightarrow C(2x^2)$ , a square matrix.

**Example 2**: We want to multiply **X** (2x2) and **b** (2x1), where **X** has elements  $x_{ij}$  and  $\beta$  has elements  $\beta_j$ :

$$\mathbf{X} = \begin{bmatrix} x_{11} & x_{21} \\ x_{12} & x_{22} \end{bmatrix} \qquad \& \qquad \mathbf{\beta} = \begin{bmatrix} \beta_1 \\ \beta_2 \end{bmatrix}$$

We compute

$$y = X \beta$$

Recall the *i*-th element is

$$y_i = \sum_{j=1}^{n=2} x_{ij} \beta_j$$

Then,

$$\mathbf{y} = \begin{bmatrix} y_1 \\ y_2 \end{bmatrix} = \begin{bmatrix} x_{11} & x_{21} \\ x_{12} & x_{22} \end{bmatrix} * \begin{bmatrix} \beta_1 \\ \beta_2 \end{bmatrix} = \begin{bmatrix} x_{11} & \beta_1 + x_{21} & \beta_2 \\ x_{12} & \beta_1 + x_{22} & \beta_2 \end{bmatrix}$$

Dimensions: **X** (2x**2**),  $\beta$  (2x1)  $\Rightarrow$  **y** (2x1), a row vector. ¶

#### Linear Algebra: Transpose

The transpose of a matrix  $\mathbf{A}$  is another matrix  $\mathbf{A}^{T}$  (also written  $\mathbf{A}'$ ) created by any one of the following equivalent actions:

-write the rows (columns) of A as the columns (rows) of  $A^{T}$ 

-reflect  $\mathbf{A}$  by its main diagonal to obtain  $\mathbf{A}^{\mathrm{T}}$ 

**Example**: 
$$A = \begin{bmatrix} 3 & 8 & -9 \\ 1 & 0 & 4 \end{bmatrix} \Rightarrow A' = \begin{bmatrix} 3 & 1 \\ 8 & 0 \\ -9 & 4 \end{bmatrix}$$
.

Some transpose results:

- If **A** is a  $m \times n$  matrix  $\Rightarrow \mathbf{A}^{\mathrm{T}}$  is a  $n \times m$  matrix.
- $(\mathbf{A'})' = \mathbf{A}$
- Conformability changes unless the matrix is square.
- (AB)' = B'A'

**Example**: In econometrics, an important matrix is X'X. Recall X (usually, the matrix of k independent variables):

$$X = \begin{bmatrix} x_{11} & x_{21} & \cdots & x_{k1} \\ x_{12} & x_{22} & \cdots & x_{k2} \\ \vdots & \vdots & \ddots & \vdots \\ x_{1T} & x_{2T} & \cdots & x_{kT} \end{bmatrix}$$
 a (*Txk*) matrix

Then,

$$X' = \begin{bmatrix} x_{11} & x_{12} & \cdots & x_{1T} \\ x_{21} & x_{22} & \cdots & x_{2T} \\ \vdots & \vdots & \ddots & \vdots \\ x_{k1} & x_{k2} & \cdots & x_{kT} \end{bmatrix}$$
 a (*kxT*) matrix

## **Linear Algebra: Math Operations**

Addition, Subtraction, Multiplication

- Addition: Just add elements

$$\begin{bmatrix} a & b \\ c & d \end{bmatrix} + \begin{bmatrix} e & f \\ g & h \end{bmatrix} = \begin{bmatrix} a+e & b+f \\ c+g & d+h \end{bmatrix}$$

- Subtraction: Just subtract element

$$\begin{bmatrix} a & b \\ c & d \end{bmatrix} - \begin{bmatrix} e & f \\ g & h \end{bmatrix} = \begin{bmatrix} a - e & b - f \\ c - g & d - h \end{bmatrix}$$

- Multiplication: Multiply each row by each column and add

$$\begin{bmatrix} a & b \\ c & d \end{bmatrix} \begin{bmatrix} e & f \\ g & h \end{bmatrix} = \begin{bmatrix} ae + bg & af + bh \\ ce + dg & cf + dh \end{bmatrix}$$

- Scalar Multiplication: Multiply each element by the scalar, k

$$k \begin{bmatrix} a & b \\ c & d \end{bmatrix} = \begin{bmatrix} ka & kb \\ kc & kd \end{bmatrix}$$

**Examples**:

Addition:	$\begin{bmatrix} 2 \\ 7 \\ A_{2x} \end{bmatrix}$	1 9]- 2 +	$+\begin{bmatrix}3\\0\\B_{2x2}\end{bmatrix}$	$\begin{bmatrix} 1\\2\\=\\=\\($	$5 \\ 7 \\ -2x^2$	2 11
Subtraction:	$[{}^2_7$	1 9] -	$- \begin{bmatrix} 1 \\ 2 \end{bmatrix}$	${0 \atop 3} =$	[1  5	$\binom{1}{6}$
Multiplication:	$\begin{bmatrix} 2\\7\\A_{2x} \end{bmatrix}$	1 9]x .2 x	$\begin{bmatrix} 1\\ 2\\ B_{2x2} \end{bmatrix}$	$\begin{bmatrix} 0\\3 \end{bmatrix} = \begin{bmatrix} \\ \\ \\ \\ \\ \\ \\ \end{bmatrix} = C_2$	4 26 2x2	3 27
Scalar Multiplication:	$\frac{1}{8}$	[2 [6	$\binom{4}{1} =$	[1/4 [3/4]	1/ 1/	2 8].¶

## Linear Algebra: Math Operations – X'X

A special matrix in econometrics, **X'X** (a kxk matrix). First, we look at this matrix for the simple case, with k = 2:

$$X(Tx2) = \begin{bmatrix} x_{11} & x_{21} \\ x_{12} & x_{22} \\ \vdots & \vdots \\ x_{1T} & x_{2T} \end{bmatrix} \qquad \& \qquad X' = \begin{bmatrix} x_{11} & x_{12} & \cdots & x_{1T} \\ x_{21} & x_{22} & \cdots & x_{2T} \end{bmatrix}$$
$$X''X(2x2) = \begin{bmatrix} \sum_{i=1}^{T} x_{1i}^2 & \sum_{i=1}^{T} x_{2i} x_{1i} \\ \sum_{i=1}^{T} x_{1i} x_{2i} & \sum_{i=1}^{T} x_{2i}^2 \end{bmatrix} = \sum_{i=1}^{T} \begin{bmatrix} x_{1i}^2 & x_{2i} x_{1i} \\ x_{2i} x_{1i} & x_{2i}^2 \end{bmatrix}$$
$$= \sum_{i=1}^{T} \begin{bmatrix} x_{1i} \\ x_{2i} \end{bmatrix} [x_{1i} & x_{2i} \end{bmatrix}$$
$$= \sum_{i=1}^{T} x_{i} x_{i}'$$

For the general case, with k explanatory variables, we have  $\mathbf{X}'\mathbf{X}$  (a kxk matrix):

$$X(Txk) = \begin{bmatrix} x_{11} & x_{21} & \cdots & x_{k1} \\ x_{12} & x_{22} & \cdots & x_{k2} \\ \vdots & \vdots & \ddots & \vdots \\ x_{1T} & x_{2T} & \cdots & x_{kT} \end{bmatrix} \& X' = \begin{bmatrix} x_{11} & x_{12} & \cdots & x_{1T} \\ x_{21} & x_{22} & \cdots & x_{2T} \\ \vdots & \vdots & \ddots & \vdots \\ x_{k1} & x_{k2} & \cdots & x_{kT} \end{bmatrix}$$
$$X''X(kxk) = \begin{bmatrix} \sum_{i=1}^{T} x_{1i}^2 & \sum_{i=1}^{T} x_{1i} x_{2i} & \cdots & \sum_{i=1}^{T} x_{1i} x_{ki} \\ \sum_{i=1}^{T} x_{2i} x_{1i} & \sum_{i=1}^{T} x_{2i}^2 & \cdots & \sum_{i=1}^{T} x_{2i} x_{ki} \\ \vdots & \vdots & \ddots & \vdots \\ \sum_{i=1}^{T} x_{ki} x_{1i} & \sum_{i=1}^{T} x_{ki} x_{2i} & \cdots & \sum_{i=1}^{T} x_{ki}^2 \end{bmatrix} =$$

$$= \sum_{i=1}^{T} \begin{bmatrix} x_{1i}^2 & \cdots & x_{1i}x_{ki} \\ \vdots & \ddots & \vdots \\ x_{ki}x_{1i} & \cdots & x_{ki}^2 \end{bmatrix} = \sum_{i=1}^{T} \begin{bmatrix} x_{1i} \\ \vdots \\ x_{ki} \end{bmatrix} \begin{bmatrix} x_{1i} & \cdots & x_{ki} \end{bmatrix}$$
$$= \sum_{i=1}^{T} x_i x_i'$$

## Linear Algebra: Math Operations – í'X

Recall  $\mathbf{i}$  is a column vector of ones (in this case, a Tx1 vector):

$$\mathbf{i} = \begin{bmatrix} 1\\1\\\vdots\\1 \end{bmatrix}$$

Given **X** (Txk), then **i**' **X** is a 1xk vector:

$$\mathbf{i}'X = \begin{bmatrix} 1 & \dots & 1 \end{bmatrix} \begin{bmatrix} x_{11} & \cdots & x_{k1} \\ \vdots & \ddots & \vdots \\ x_{1T} & \cdots & x_{kT} \end{bmatrix} = \begin{bmatrix} \sum_{t=1}^{T} x_{1t} & \dots & \sum_{t=1}^{T} x_{kt} \end{bmatrix}$$

Note: If  $\mathbf{x}_1$  is a vector of ones (representing a constant in the linear classical model), then:  $\mathbf{i}' \mathbf{x}_1 = \sum_{t=1}^T x_{1t} = \sum_{t=1}^T 1 = T$  (dot product, "•")

# Linear Algebra: Inverse of a Matrix

Identity matrix: $\mathbf{AI} = \mathbf{A}$ ,		11	0	•••	0	
	where $I_j =$	0	1	•••	0	
		-   :	÷	•••	:	
		LO	0	•••	1	

<u>Notation</u>:  $I_j$  is a *j*x*j* identity matrix.

• Given A (mxn), the matrix B (nxm) is a *right-inverse* for A iff AB = I<sub>m</sub>

• Given A (mxn), the matrix C (nxm) is a *left-inverse* for A iff CA = I<sub>n</sub>

• Theorem: If A (mxn), has both a right-inverse B and a left-inverse C, then  $C = B = A^{-1}$ 

Note:

- If **A** has both a right and a left inverse, it is a square matrix (*m*=*n*). It is usually called *invertible*. We say "the matrix **A** is *non-singular*."
- This matrix,  $A^{-1}$ , is unique.
- If det(A)  $\neq 0 \Rightarrow$  A is non-singular.

## Linear Algebra: Symmetric Matrices

Definition:

If  $\mathbf{A}' = \mathbf{A}$ , then  $\mathbf{A}$  is called a *symmetric* matrix.

In many applications, matrices are often symmetric. For example, in statistics the *correlation matrix* and *the variance covariance matrix*.

Symmetric matrices play the same role as real numbers do among the complex numbers.

We can do calculations with symmetric matrices like with numbers: for example, we can solve  $B^2 = A$  for B if A is symmetric matrix (& B is square root of A.) This is not possible in general. X'X is symmetric. It plays a very important role in econometrics.

## Linear Algebra: Operations in R

```
Many ways to create a vector (c, 2:7, seq, rep, etc) or a matrix (c, cbind, rbind). We use c(), the
combine function:
v1 \le c(1, 3, 8)
                             \# a (3x1) vector (vectors are usually treated as a column list)
> v1
[1] 1 3 8
A <- matrix (c(1, 2, 3, 7, 8, 9), ncol = 3) # a (2x3) matrix
> A
  [,1] [,2] [,3]
[1,] 1 3 8
[2,] 2 7 9
B \le matrix(c(1, 3, 1, 1, 2, 0), nrow = 3)
> B
       [,1] [,2]
[1,] 1 1
[2,] 3 2
[3,] 1 0
• Now, we use rbind to create A and cbind to create B
v1 \le c(1, 3, 8)
                                     \# a (3x1) vector
v2 \le c(2, 7, 9)
A \leq rbind(v1, v2)
> A
                                     \# a (2x3) matrix
 [,1] [,2] [,3]
v1 1 3 8
v2 2 7 9
v3 \le c(1, 3, 1)
v4 \le c(1, 2, 0)
B \leq cbind(v3,v4)
>B
                                     \# a (3x2) matrix
   v3 v4
[1,] 1 1
[2,] 3 2
```

[3,] 1 0

• Matrix addition/subtraction: +/-• Matrix multiplication: %\*%C <- A%\*%B > C [,1] [,2] [1,] 18 7 [2,] 32 16 • Scalar multiplication: \* > 2\*C [,1] [,2] [1,] 36 14 [2,] 64 32 -element by element. #A is 2x3; B is  $3x2 \implies C$  is 2x2#A is 2x3; B is  $3x2 \implies C$  is 2x2# elementwise multiplication of C by scalar 2

Note: Usually, matrices will be data -i.e., read as input.

• Dot product "•" is a function that takes pairs of vectors, with same length, and produces a number. For vectors **c** & **z**, it is defined as:

 $c \cdot z = c_1 * z_1 + c_2 * z_2 + \dots + c_n * z_n = \sum_{i=1}^n c_i z_i$ 

• Dot product with 2 vectors: v1 • v2: sum of the elementwise multiplied elements of both vectors > t(v1) % \*% v2# v1 <- c(1, 3, 8) & v2 <- c(2, 7, 9) [,1] [1,] 95 • Dot product with a vector itself: v1 • v1: Sum of the square elements of vector > t(v1) %\*% v1 [,1] [1,] 74 • Dot product with *i* (a vector of ones): sum of elements of vector  $i \le c(1,1,1)$ # vector of ones (iota) > t(i) % \* % v1# v1 <- c(1, 3, 8) [,1] [1,] 12

Product of 2 vectors: v1 & t(v2): A (3x3) matrix.
> v1%\*%t(v2) # v1 <- c(1, 3, 8) --a (3x1) vector x (1x3) vector [,1] [,2] [,3]</li>
[1,] 2 7 9
[2,] 6 21 27
[3,] 16 56 72

<u>Property of dot product</u>: If the dot product of two vectors is equal to zero, then the vectors are *orthogonal* (perpendicular *or* " $\perp$ ") vectors. We interpret this result as "the vectors are *uncorrelated*."

• Matrix transpose: t > t(B)#B is  $3x2 \Rightarrow t(B)$  is 2x3[,1] [,2] [,3] [1,] 1 3 1 [2,] 1 2 0 • X'X (a symmetric matrix) > t(B)%\*%B# command crossprod(B) is more efficient [,1] [,2] [1,] 11 7 [2,] 7 5 (a symmetric matrix) • Determinant: **det** > det(t(B)%\*%B)# Matrix has to be square. If  $det(A)=0 \Rightarrow A$  non-invertible [1] 6 • (X'X)<sup>-1</sup>: Inverse: **solve** > solve(t(B)%\*%B) # Matrix inside solve() has to be square [,1] [,2] [1,] 0.8333333 -1.166667 [2,]-1.1666667 1.833333 • Take the diagonal elements of a matrix A: diag() > diag(solve(t(B)%\*%B)) [1] 0.8333333 1.833333 • Square root of (positive) elements of a matrix A: sqrt() > sqrt(diag(solve(t(B)%\*%B))) v3 v40.9128709 1.3540064

## Linear Algebra: Examples

### **Example 1 – Linear DGP**

There is a functional form relating a dependent variable, y, and k explanatory variables, X. The functional form is linear, but it depends on k unknown parameters,  $\beta$ . The relation between y and X is not exact. There is an error,  $\varepsilon$ . We have T observations of y and X.

• Then, the data is generated according to:

 $y_i = \sum_{j=1}^k x_{k,i} \beta_k + \varepsilon_i$  i = 1, 2, ..., T.Or using matrix notation:

 $y = X \beta + \varepsilon$ 

where  $\mathbf{y} \& \mathbf{\varepsilon}$  are (*T*x1);  $\mathbf{X}$  is (*T*x*k*); and  $\boldsymbol{\beta}$  is (*k*x1).

- We will call this relation data generating process (DGP).
- The goal of econometrics is to estimate the unknown vector  $\beta$ .

#### **Example 2 – Linear System**

Assume an economic model as system of linear equations with:

 $a_{ij}$  parameters, where i = 1,..., m rows, j = 1,..., n columns

- $x_i$  endogenous variables (*n*),
- $d_i$  exogenous variables and constants (m).

$$\begin{cases} a_{11}x_1 + a_{12} x_2 + \dots + a_{1n} x_n = d_1 \\ a_{21}x_1 + a_{22} x_2 + \dots + a_{2n} x_n = d_2 \\ \dots & \dots & \dots \\ a_{m1}x_1 + a_{m2} x_2 + \dots + a_{mn} x_n = d_n \end{cases}$$

• Using linear algebra, we have a system of linear equations: Ax = d

 $\begin{bmatrix} a_{11} & \cdots & a_{1n} \\ \vdots & \ddots & \vdots \\ a_{m1} & \cdots & a_{mn} \end{bmatrix} \begin{bmatrix} x_1 \\ \cdots \\ x_n \end{bmatrix} = \begin{bmatrix} d_1 \\ \cdots \\ d_m \end{bmatrix}$ 

where

A = (mxn) matrix of parameters x = column vector of endogenous variables (*nx*1) d = column vector of exogenous variables and constants (*mx*1). ¶

We want to solve for the solution of Ax = d,  $\Rightarrow x^*$ .

**Theorem:** Given A (mxn) invertible. Then, the equation Ax = d has one and only one solution for every d(mx1). That is, there is a unique  $x^*$ .

$$\Rightarrow$$
 **x**<sup>\*</sup> = **A**<sup>-1</sup> *d*

Example: In practice, we avoid computing A<sup>-1</sup>, we solve a system. A <- matrix(c(1, 1, 5, 7, 9, 11, 10, 10, 14), ncol = 3) # check det(A) for singularity (det(A)=-72) d <- c(2, 5, 2) > solve(A,d) [1] -0.7222222 1.5000000 -0.7777778 ¶

## Linear Algebra: Linear Dependence and Rank

A set of vectors is *linearly dependent* if any one of them can be expressed as a linear combination of the remaining vectors; otherwise, it is linearly independent.

Formal definition: Linear independence (LI) The set  $\{u_1, u_2, ..., u_k\}$  is called a *linearly independent* set of vectors iff

$$c_1 u_1 + c_2 u_2 + \dots + c_k u_k = 0 \implies c_1 = c_2 = \dots = c_k = 0.$$

**Example**: We want to check if  $\{v_1, v_2, v_3\}$  are linearly independent:

$$v_1 = \begin{bmatrix} 2 \\ 7 \end{bmatrix}; v_2 = \begin{bmatrix} 1 \\ 8 \end{bmatrix}; v_3 = \begin{bmatrix} 4 \\ 5 \end{bmatrix}$$

Then,

$$3v_1 - 2v_2 = \begin{bmatrix} 6\\21 \end{bmatrix} - \begin{bmatrix} 2\\16 \end{bmatrix} = \begin{bmatrix} 4\\5 \end{bmatrix} = v_3$$
  

$$\Rightarrow \quad 3v_1 - 2v_2 - v_3\mathbf{0} \quad (c_1 = 3, c_2 = -2, c_3 = -1)$$
  

$$\Rightarrow \text{ The set } \{v_1, v_2, v_3\} \text{ is linearly dependent. } \P$$

Notes:

- Dependence prevents solving a system of equations (A is not invertible). More unknowns than independent equations.

- The number of linearly independent rows or columns in a matrix is the *rank* of a matrix  $(rank(\mathbf{A}))$ .

- If A, a  $(k \ge k)$  square matrix, has rank(A) = k, then A is invertible.

**Examples**:

(1) 
$$v_1 = \begin{bmatrix} 5\\12 \end{bmatrix}; v_2 = \begin{bmatrix} 10\\24 \end{bmatrix}$$
  
 $A = \begin{bmatrix} 5&10\\12&24 \end{bmatrix} = \begin{bmatrix} v_1 & v_1 \end{bmatrix}$   
 $2v_1 = v_2 \qquad \Rightarrow rank(A) = 1$ 

 $(2) \qquad \boldsymbol{B} = \begin{bmatrix} 2 & 1 & 6 \\ 2 & 4 & 0 \end{bmatrix}$ 

 $4v_1 - 2v_2 = v_3 \qquad \Rightarrow rank(\mathbf{B}) = 2. \P$ 

**Least Squares Estimation with Linear Algebra – Rules for Vector Derivatives** Below we present the tules for vector differentiation of linear functions and quadratic forms (for derivation of the rules, see Appendix at end of Lecture 4):

(1) Linear function:  $y = f(x) = x' \gamma + \omega$ where x and  $\gamma$  are k-dimensional vectors and  $\omega$  is a constant. Then,  $\nabla f(x) = \gamma$ 

(2) Quadratic form:  $\mathbf{q} = f(\mathbf{x}) = \mathbf{x}' \mathbf{A} \mathbf{x}$ where  $\mathbf{x}$  is  $k \ge 1$  vector and  $\mathbf{A}$  is a  $k \ge k$  matrix, with  $a_{ji}$  elements. Then,  $\nabla f(\mathbf{x}) = \mathbf{A}' \mathbf{x} + \mathbf{A} \mathbf{x} = (\mathbf{A}' + \mathbf{A}) \mathbf{x}$ If  $\mathbf{A}$  is symmetric, then  $\nabla f(\mathbf{x}) = 2 \mathbf{A} \mathbf{x}$ . In the next section, we apply these rules to

$$S(\mathbf{x}; \boldsymbol{\beta}) = \sum_{i=1}^{T} \varepsilon_i^2 = \varepsilon' \varepsilon = (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})' (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})$$
$$= (\mathbf{y}' - \boldsymbol{\beta}'\mathbf{X}') (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})$$
$$= (\mathbf{y}'\mathbf{y} - \boldsymbol{\beta}'\mathbf{X}'\mathbf{y} - \mathbf{y}'\mathbf{X}\boldsymbol{\beta} + \boldsymbol{\beta}'\mathbf{X}'\mathbf{X}\boldsymbol{\beta})$$
$$= (\mathbf{y}'\mathbf{y} - 2 \ \boldsymbol{\beta}'\mathbf{X}'\mathbf{y} + \boldsymbol{\beta}'\mathbf{X}'\mathbf{X}\boldsymbol{\beta})$$

where we take derivatives with respect to the  $k \ge 1$  vector  $\beta$ . Note that since you take k derivatives, the first derivative vector will be a  $k \ge 1$  vector, like  $\beta$ .

## Least Squares Estimation with Linear Algebra

Let's assume a linear system with k independent variables and T observations. That is,

$$y_i = \beta_1 x_{1i} + \beta_2 x_{2i} + ... + \beta_k x_{ki} + \varepsilon_i,$$
  $i = 1, 2, ..., T$ 

The whole system (for all *i*) is:

$$y_{1} = \beta_{1} x_{11} + \beta_{2} x_{21} + \dots + \beta_{k} x_{k1} + \varepsilon_{1}$$
  

$$y_{2} = \beta_{1} x_{12} + \beta_{2} x_{22} + \dots + \beta_{k} x_{k2} + \varepsilon_{2}$$
  

$$\dots \qquad \dots \qquad \dots \qquad \dots$$
  

$$y_{T} = \beta_{1} x_{1T} + \beta_{2} x_{2T} + \dots + \beta_{k} x_{kT} + \varepsilon_{T}$$

Using linear algebra we can rewrite the system as:

$$y = X \beta + \varepsilon$$

Vectors will be column vectors: y,  $x_i$ , and  $\varepsilon$  are Tx1 vectors:

$$\mathbf{y} = \begin{bmatrix} y_1 \\ \vdots \\ y_T \end{bmatrix} \qquad \Rightarrow \qquad \mathbf{y}' = \begin{bmatrix} y_1 \ y_2 \ \dots \ y_T \end{bmatrix}$$
$$\mathbf{x}_j = \begin{bmatrix} x_{j1} \\ \vdots \\ x_{jT} \end{bmatrix} \qquad \Rightarrow \qquad \mathbf{x}_j '= \begin{bmatrix} x_{j1} \ x_{j2} \ \dots \ x_{jT} \end{bmatrix}$$
$$\mathbf{\varepsilon} = \begin{bmatrix} \varepsilon_1 \\ \vdots \\ \varepsilon_T \end{bmatrix} \qquad \Rightarrow \qquad \mathbf{\varepsilon}' = \begin{bmatrix} \varepsilon_1 \ \varepsilon_2 \ \dots \ \varepsilon_T \end{bmatrix}$$

**X** is a  $T\mathbf{x}k$  matrix.  $\Rightarrow$  **X** = [ $\mathbf{x}_1 \ \mathbf{x}_2 \ \dots \ \mathbf{x}_k$ ]

$$\Rightarrow \qquad X = \begin{bmatrix} x_{11} & x_{21} & \cdots & x_{k1} \\ \vdots & \vdots & \ddots & \vdots \\ x_{1T} & x_{2T} & \cdots & x_{kT} \end{bmatrix}$$
$$\beta = \begin{bmatrix} \beta_1 \\ \vdots \\ \beta_k \end{bmatrix}$$

• With the linear assumption:  $f(\mathbf{X}, \theta) = \mathbf{X} \boldsymbol{\beta}$ , we can write the objective function as:  $S(\boldsymbol{x}; \theta) = \sum_{i=1}^{T} \varepsilon_i^2 = \varepsilon' \varepsilon = (\boldsymbol{y} - \mathbf{X} \boldsymbol{\beta})' (\boldsymbol{y} - \mathbf{X} \boldsymbol{\beta}) = (\boldsymbol{y}' - \boldsymbol{\beta}' \mathbf{X}') (\boldsymbol{y} - \mathbf{X} \boldsymbol{\beta})$ 

We want to minimize  $S(x_i, \theta)$ . After some simple algebra we have:

$$S(\mathbf{x}; \boldsymbol{\beta}) = (\mathbf{y}'\mathbf{y} - \boldsymbol{\beta}'\mathbf{X}'\mathbf{y} - \mathbf{y}'\mathbf{X}\boldsymbol{\beta} + \boldsymbol{\beta}'\mathbf{X}'\mathbf{X}\boldsymbol{\beta})$$
  
=  $(\mathbf{y}'\mathbf{y} - 2 \ \boldsymbol{\beta}'\mathbf{X}'\mathbf{y} + \boldsymbol{\beta}'\mathbf{X}'\mathbf{X}\boldsymbol{\beta})$   
=  $(\mathbf{c} - 2 \ \boldsymbol{\beta}'\mathbf{d} + \boldsymbol{\beta}'\mathbf{A}\boldsymbol{\beta})$  ( $\mathbf{d} = \mathbf{X}'\mathbf{y} \& \mathbf{A} = \mathbf{X}'\mathbf{X}$  is symmetric)

First derivative w.r.t.  $\beta'$ :  $\nabla S(\mathbf{x}; \theta) = (-2 \mathbf{d} + 2 \mathbf{A} \beta)$  (kx1 vector) = -2 ( $\mathbf{X}'\mathbf{y} - \mathbf{X}'\mathbf{X} \beta$ )

F.o.c. (normal equations): (X'y - X'X b) = 0

Simple algebra  $(X'X) \mathbf{b} = X'y$ 

Assuming (X'X) is non-singular –i.e., invertible-, we solve for **b**:  $\Rightarrow$  **b** = (X'X)<sup>-1</sup> X'y

<u>Note</u>: **b** is called the Ordinary Least Squares (**OLS**) estimator. (*Ordinary* =  $f(\mathbf{X}, \theta)$  is linear.)

<u>Remark</u>: Technically, we still need to check the Second Order condition, we need the 2nd derivative to be positive for a minimum:

$$\frac{\partial^2 \mathbf{S}(x_i, \boldsymbol{\beta})}{\partial \boldsymbol{\beta} \partial \boldsymbol{\beta}'} = 2 \mathbf{X}' \mathbf{X},$$

which is a positive definite (**pd**) matrix, the counterpart to positive numbers for matrices.  $\Rightarrow$  **b** is a minimum!

 $\Rightarrow$  **D** is a minimum!

X is a Txk matrix. Its columns are the k Tx1 vectors  $x_j$ . It is common to treat  $x_1$  as vector of ones:  $\begin{bmatrix} x_{11} \end{bmatrix} \begin{bmatrix} 1 \end{bmatrix}$ 

$$x_1 = \begin{bmatrix} 1 \\ \vdots \\ x_{1T} \end{bmatrix} = \begin{bmatrix} 1 \\ \vdots \\ 1 \end{bmatrix} \Rightarrow x'_1 = \begin{bmatrix} 1 & 1 & \dots & 1 \end{bmatrix} = i'$$

This vector of ones represent the usual constant in the model. Then,

$$\mathbf{X} = \begin{bmatrix} 1 & x_{21} & \cdots & x_{k1} \\ 1 & x_{22} & \cdots & x_{k1} \\ \vdots & \vdots & \ddots & \vdots \\ 1 & x_{2T} & \cdots & x_{kT} \end{bmatrix}$$

<u>Note</u>: Recall the dot product: Post-multiplying a vector  $(1 \times T) \mathbf{x}_k$  by  $\mathbf{i}$  (or  $\mathbf{i}' \mathbf{x}_k$ ) produces a scalar, the sum of all the elements of vector  $\mathbf{x}_k$ :

$$x_{k}' i = i' x_{k} = x_{k1} + x_{k2} + \dots + x_{kT} = \sum_{i}^{T} x_{ki}.$$

#### Least Squares Estimation with Linear Algebra: The Fama-French Model

The CAPM is routinely rejected. A popular alternative model is the empirically derived 3-factor Fama-French (1993) model, which adds two risk factors to the market factor: a size factor,

measured as the returns of a portfolio of small firms minus the returns of a portfolio of big firms (SMB), and a book-to-market (BM) factor (or value factor), measured as the returns of a portfolio of high BM firms minus the returns of a portfolio of low BM firms (HML). SMB accounts for companies with small market caps that generate higher returns, while HML accounts for value stocks with high book-to-market ratios that generate higher returns in comparison to the market.

Then, a linear DGP generating this model is:

 $r_{i,t} - r_f = \alpha_i + \beta_1 (r_{m,t} - r_f) + \beta_2 SMB_t + \beta_3 HML_t + \varepsilon_{i,t}.$ 

The interpretation of the coeffcientes is the usual interpretation, they measure the sensitivity of excess returns to the risk factors. For example,  $\beta_2$  measures the exposure of asset *i* to the size factor (in general,  $\beta_2 > 0$  means that returns of asset *i* behaves like small stocks).

Like the CAPM, the 3-factor FF model produces expected excess returns:  $E[r_{i,t} - r_f] = \beta_1 E[r_{m,t} - r_f] + \beta_2 E[SMB_t] + \beta_3 E[HML_t].$ 

A significant constant would be evidence against this model: something is missing.

Below, we present in more detail this 3-factor model.

**Example**: Fama-French 3-factor Model for IBM returns: SFX da <read.csv("https://www.bauer.uh.edu/rsusmel/4397/Stocks FX 1973.csv",head=TRUE,sep=",") x ibm <- SFX da\$IBM # Read IBM price data (Mkt RF Factor) x Mkt RF <- SFX da\$Mkt RF # Read Factor data -Mkt RF Factor (in %) x SMB <- SFX da\$SMB # Read Factor data -SMB Factor (in %) x\_HML <- SFX\_da\$HML # Read Factor data -HML Factor (in %) x RF <- SFX\_da\$RF # Read Factor data – Risk free rate (in %)  $T \leq - length(x ibm)$ # Sample size  $\ln ibm \le \log(x ibm[-1]/x ibm[-T])$ # Log returns for IBM (lost one observation) Mkt RF  $\leq x$  Mkt RF[-1]/100 # Adjust size (take one observation out ) SMB <- x SMB[-1]/100  $HML \le x HML[-1]/100$  $RF \le x RF[-1]/100$ # Define y (IBM excess returns)  $y \le ibm x$ # Regressor 1 (Mkt RF)  $x1 \le Mkt RF$ x2 <- SMB # Regressor 2 (SMB) x3 <- HML # Regressor 3 (HML) # New sample size (Original - 1 observation)  $T \leq - length(x1)$ x0 <- matrix(1,T,1)# Define vector of ones (the constant in X)  $x \le cbind(x0,x1,x2,x3)$ # Matrix X # Number of regressors (=rank(X)=k)  $k \leq -ncol(x)$  $b \le solve(t(x)\%*\% x)\%*\% t(x)\%*\%y$  $\# \mathbf{b} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \mathbf{y} \text{ (OLS regression)}$ 

> t(b)

#### [1,] -0.005088944 0.9082989 -0.2124596 -0.1715002

<u>R Note</u>: You should get the same numbers using R's *lm* (use *summary*(.) to print results):

```
fit_ibm_ff3 <- lm(ibm x \sim Mkt RF + SMB + HML)
> summary(fit ibm ff3)
                                                           # print lm results
Call:
lm(formula = ibm x \sim Mkt RF + SMB + HML)
Residuals:
   Min
           10
                      Median
                                 3Q
                                        Max
-0.307488 -0.030388 -0.000861 0.034350 0.252667
Coefficients:
            Estimate
                        Std. Error t value Pr(>|t|)
            -0.005089 0.002488 -2.046 0.0412 *
(Intercept)
Mkt RF
            0.908299 0.056722
                                 16.013 <2e-16 ***
SMB
            -0.212460 0.084112
                                  -2.526 0.0118 *
HML
            -0.171500 0.084682 -2.025 0.0433 *
____
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Residual standard error: 0.05848 on 565 degrees of freedom Multiple R-squared: 0.3389, Adjusted R-squared: 0.3354 F-statistic: 96.55 on 3 and 565 DF, p-value: < 2.2e-16.

<u>Conclusion</u>: IBM has a positive exposure to the market, and negative exposure to the size and value factors.  $\P$ 

#### **OLS** – Assumptions

Typical OLS Assumptions (1) DGP:  $y_i = \beta_1 x_{1i} + \beta_2 x_{2i} + ... + \beta_k x_{ki} + \varepsilon_i, \qquad i = 1, 2, ..., T$  $\Rightarrow$  functional form known, but  $\beta$  is unknown.

(2)  $E[\varepsilon_i] = 0.$   $\Rightarrow$  expected value of the errors is 0.

(3) Explanatory variables  $X_1, X_2, ..., X_k$ , are given (& non random)  $\Rightarrow$  no correlation with  $\varepsilon$  (Cov( $\varepsilon_i, X_{ki}$ ) = 0.) for all *k*.

(4) The k explanatory variables are independent.

(5) Var[ $\varepsilon_i$ ] = E[ $\varepsilon_i^2$ ] =  $\sigma^2 < \infty$  (homoscedasticity = same variance for all *i*)

(6) Cov( $\varepsilon_i, \varepsilon_j$ ) = E[ $\varepsilon_i \varepsilon_j$ ] = 0. (no serial/cross correlation for all  $i \neq j$ )

• These are the assumptions behind the *classical linear regression model* (CLM).

#### Least Squares – Assumptions with Linear Algebra Notation

We can rewrite the assumptions, conditioning on **X**, which allows **X** to be a random variable (though, once we condition, **X** becomes a matrix of numbers). Using linear algebra:

(A1) DGP:  $\mathbf{y} = f(\mathbf{X}, \mathbf{\theta}) + \mathbf{\varepsilon}$  is correctly specified. (A2)  $\mathrm{E}[\mathbf{\varepsilon}|\mathbf{X}] = 0$ (A3)  $\mathrm{Var}[\mathbf{\varepsilon}|\mathbf{X}] = \sigma^2 \mathbf{I}_{\mathrm{T}}$ (A4) X has full column rank  $-\mathrm{rank}(\mathbf{X})=k-$ , where  $\mathrm{T} \ge k$ .

• Assumption (A1) is called *correct specification*. We know how the data is generated. We call  $y = f(\mathbf{X}, \mathbf{\theta}) + \mathbf{\epsilon}$  the Data Generating Process (DGP).

<u>Note</u>: The errors,  $\varepsilon$ , are called *disturbances*. They are not something we add to  $f(\mathbf{X}, \boldsymbol{\theta})$  because we don't know precisely  $f(\mathbf{X}, \boldsymbol{\theta})$ . No. The errors are part of the DGP.

• Assumption (A2) is called *regression*.

From Assumption (A2) we get:

(i)  $E[\boldsymbol{\varepsilon}|\mathbf{X}] = 0 \implies E[\boldsymbol{y}|\mathbf{X}] = E[f(\mathbf{X}, \theta)|\mathbf{X}] + E[\boldsymbol{\varepsilon}|\mathbf{X}] = f(\mathbf{X}, \theta)$ That is, the observed **y** will equal  $E[\boldsymbol{y}|\mathbf{X}]$  + random variation.

(ii) Using rules of expectations and the law of iterated expectations (LIE), we get two results: (1)  $E[\mathbf{\epsilon}|\mathbf{X}] = 0 \implies E[\mathbf{\epsilon}] = 0$  $\implies$  The conditional expectation = unconditional expectation

(2) 
$$\operatorname{Cov}(\varepsilon, \mathbf{X}) = \operatorname{E}[(\varepsilon - 0)(\mathbf{X} - \mu_{\mathbf{X}})] = \operatorname{E}[\varepsilon \mathbf{X} - \varepsilon \mu_{\mathbf{X}}]$$
  

$$= \operatorname{E}[\varepsilon \mathbf{X}] - \mu_{\mathbf{X}} \operatorname{E}[\varepsilon] = \operatorname{E}[\varepsilon \mathbf{X}] = 0 \qquad \text{(by LIE, E}[\varepsilon \mathbf{X}] = \operatorname{E}_{\mathbf{X}}[\mathbf{X} \operatorname{E}[\varepsilon |\mathbf{X}]] = 0.)$$

$$\Rightarrow \operatorname{That is,} \quad \operatorname{E}[\varepsilon \mathbf{X}] = 0 \qquad \Rightarrow \varepsilon \perp \mathbf{X}.$$
There is no information about  $\varepsilon$  in  $\mathbf{X}$  and viceversa.

• Assumption (A3) gives the model a constant variance for all errors and no relation between the errors at different measurements/times. That is, we have a diagonal variance-covariance matrix:

$$\operatorname{Var}[\boldsymbol{\varepsilon}|\mathbf{X}] = \boldsymbol{\Sigma} = \begin{bmatrix} \sigma^2 & 0 & \cdots & 0\\ 0 & \sigma^2 & \cdots & 0\\ \vdots & \vdots & \vdots & \vdots\\ 0 & 0 & \cdots & \sigma^2 \end{bmatrix} = \sigma^2 \mathbf{I}_{\mathrm{T}} \quad (kxk) \text{ matrix}$$

This assumption implies

(i) homoscedasticity	$\Rightarrow \mathrm{E}[\varepsilon_i^2   \mathbf{X}] = \sigma^2$	for all <i>i</i> .
(ii) no serial/cross correlation	$\Rightarrow \mathrm{E}[\varepsilon_i \varepsilon_j   \mathbf{X}] = 0$	for $i \neq j$ .

It can be shown using the law of total variance that

 $\operatorname{Var}[\boldsymbol{\varepsilon}|\mathbf{X}] = \sigma^2 \mathbf{I}_{\mathrm{T}} \qquad \qquad \Rightarrow \operatorname{Var}[\boldsymbol{\varepsilon}] = \sigma^2 \mathbf{I}_{\mathrm{T}}$ 

$$\underbrace{Note:} \operatorname{Var}[\boldsymbol{\varepsilon}|\mathbf{X}] = \operatorname{E}[(\boldsymbol{\varepsilon} - \operatorname{E}[\boldsymbol{\varepsilon}]) (\boldsymbol{\varepsilon} - \operatorname{E}[\boldsymbol{\varepsilon}])'|\mathbf{X}] \\ = \operatorname{E}[(\boldsymbol{\varepsilon} - \mathbf{0}) (\boldsymbol{\varepsilon} - \mathbf{0})'|\mathbf{X}] \\ = \operatorname{E}[\boldsymbol{\varepsilon} \, \boldsymbol{\varepsilon}'|\mathbf{X}] \\ = \begin{bmatrix} \operatorname{E}[\boldsymbol{\varepsilon}_{1}^{2}|\mathbf{X}] & \operatorname{E}[\boldsymbol{\varepsilon}_{2} \, \boldsymbol{\varepsilon}_{1}|\mathbf{X}] & \cdots & \operatorname{E}[\boldsymbol{\varepsilon}_{T} \, \boldsymbol{\varepsilon}_{1}|\mathbf{X}] \\ \operatorname{E}[\boldsymbol{\varepsilon}_{1} \, \boldsymbol{\varepsilon}_{2}|\mathbf{X}] & \operatorname{E}[\boldsymbol{\varepsilon}_{2}^{2}|\mathbf{X}] & \cdots & \operatorname{E}[\boldsymbol{\varepsilon}_{T} \, \boldsymbol{\varepsilon}_{2}|\mathbf{X}] \\ \vdots & \vdots & \vdots & \vdots \\ \operatorname{E}[\boldsymbol{\varepsilon}_{1} \, \boldsymbol{\varepsilon}_{T}|\mathbf{X}] & \operatorname{E}[\boldsymbol{\varepsilon}_{2} \, \boldsymbol{\varepsilon}_{T}|\mathbf{X}] & \cdots & \operatorname{E}[\boldsymbol{\varepsilon}_{T}^{2}|\mathbf{X}] \end{bmatrix} \\ = \begin{bmatrix} \sigma^{2} & 0 & \cdots & 0 \\ 0 & \sigma^{2} & \cdots & 0 \\ \vdots & \vdots & \vdots & \vdots \\ 0 & 0 & \cdots & \sigma^{2} \end{bmatrix} = \sigma^{2} \mathbf{I}_{T}$$

• From Assumption (A4)  $\Rightarrow$  the *k* independent variables in X are linearly independent. Then, the *kxk* matrix X'X will also have full rank –i.e., rank(X'X) = *k*.

Thus,  $\mathbf{X}'\mathbf{X}$  is invertible. We need this result to solve a system of equations given by the 1<sup>st</sup>-order conditions of Least Squares Estimation (normal equations):

$$\mathbf{X}'\mathbf{y} - \mathbf{X}'\mathbf{X} \mathbf{b} = 0$$

Note: To get asymptotic results we will need more assumptions about X.

• We assume a linear functional form for  $f(x, \theta) = X \beta$ : (A1') DGP:  $y = X \beta + \varepsilon$ 

#### **CLM: OLS – Summary**

Classical linear regression model (CLM) - Assumptions: (A1) DGP:  $y = X \beta + \varepsilon$  is correctly specified. (A2)  $E[\varepsilon|X] = 0$ (A3)  $Var[\varepsilon|X] = \sigma^2 I_T$ (A4) X has full column rank -rank(X) = k, where  $T \ge k$ .

Objective function:  $S(x_i, \beta) = \sum_{i=1}^T x_i^2 = \varepsilon' \varepsilon = (y - X\beta)' (y - X\beta)$ =  $(y'y - 2\beta'X'y + \beta'X'X\beta)$ First order conditions: -2(X'y - X'Xb) = 0Solving for b:  $b = (X'X)^{-1}X'y$  (kx1) vector

## **OLS Estimation: Second Order Condition**

$$\frac{\partial^2 S(x_i, \boldsymbol{\beta})}{\partial b \partial b'} = 2\mathbf{X}'\mathbf{X} = 2 \begin{bmatrix} \Sigma_{i=1}^T x_{i1}^2 & \Sigma_{i=1}^T x_{i1} x_{i2} & \dots & \Sigma_{i=1}^T x_{i1} x_{iK} \\ \Sigma_{i=1}^T x_{i2} x_{i1} & \Sigma_{i=1}^T x_{i2}^2 & \dots & \Sigma_{i=1}^T x_{i2} x_{iK} \\ \dots & \dots & \dots & \dots \\ \Sigma_{i=1}^T x_{iK} x_{i1} & \Sigma_{i=1}^T x_{iK} x_{i2} & \dots & \Sigma_{i=1}^T x_{iK}^2 \end{bmatrix}$$

If there were a single **b**, we would require this to be positive, which it would be:  $2 \mathbf{x}' \mathbf{x} = 2 \Sigma_{i=1}^{T} x_i^2 > 0.$ 

The matrix counterpart of a positive number is a positive definite (pd) matrix. We need X'X to be pd.

A square matrix (mxm) A "takes the sign" of the quadratic form, z'A z, where z is a mx1 vector. Then, z'A z is a scalar.

A form is a polynomial expression in which each component term has a uniform degree. A quadratic form has a uniform  $2^{nd}$  degree.

#### **Examples**:

9 x + 3 y + 2 z6  $x^2 + 2 x y + 2 y^2$   $d^2z = f_{xx} dx^2 + 2 f_{xy} dx dy + f_{yy} dy^2$ - 1st degree form. - 2nd degree (quadratic) form. - quadratic form. ¶

A quadratic form can be written in matrix notation as  $\mathbf{z'A} \mathbf{z}$ , where A is  $(mxm) \mathbf{A}$  and  $\mathbf{z}$  is an mx1 vector. Then,  $\mathbf{z'A} \mathbf{z}$  is a scalar.

**Example**: The fiear quadratic form from the previous example can be written as

$$q = [x \ y] * \begin{bmatrix} 6 & 1 \\ 1 & 2 \end{bmatrix} * \begin{bmatrix} x \\ y \end{bmatrix} = 6 \ x^2 + 2 \ x \ y + 2 \ y^2$$
  
we know x & y o is a number

Once we know x & y, q is a number.

• Let q be a quadratic form. We say q is:

- Positive definite if q is invariably positive (q > 0)
- Positive semi-definite if q is invariably non-negative  $(q \ge 0)$
- Negative semi-definite if q is invariably non-positive (q  $\leq 0$ )
- Negative definite if q is invariably negative (q < 0)

#### **Definition**: Positive definite matrix

A matrix **A** is *positive definite* (pd) if  $\mathbf{z'A} \mathbf{z} > 0$  for *any*  $\mathbf{z}$  (a *k*x1 vector).

For some matrices, it is easy to check. Let  $\mathbf{A} = \mathbf{X'X}$  (a *kxk* matrix). Then,  $\mathbf{z'A} \mathbf{z} = \mathbf{z'X'X} \mathbf{z} = \mathbf{v'v} = \sum_{i=1}^{N} v_i^2 > 0.$  (**v=Xz** is an *N*x1 vector)  $\Rightarrow \mathbf{X'X}$  is pd  $\Rightarrow \mathbf{b}$  is a min!

#### Technical notes:

1) In general, we need eigenvalues of A to check this. If all the eigenvalues are positive, then A is pd.

2) If  $\mathbf{A}$  is pd, then  $\mathbf{A}^{-1}$  is also pd.

3) In optimization problems in multivariate calculus, the second order condition requires the evaluation of the matrix of second derivatives, the Hessian. If all the leading principal minors are positive, then the critical point obtained is a minimum. In our case, this means that the Hessian is pd.

Loosely speaking, a matrix is positive definite if the diagonal elements are positive and the offdiagonal elements are not too large in absolute value relative to the diagonal elements. This is a very informal way of looking at a pd matrix, but, keep in mind for later, that the diagonal elements are positive.

#### **OLS Estimation – Properties of b**

The OLS estimator of  $\beta$  in the CLM is

 $\begin{aligned} \mathbf{b} &= (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \ \mathbf{y} \implies \mathbf{b} \text{ is a (linear) function of the data } (\mathbf{y}_{i}, \mathbf{x}_{i}). \\ \mathbf{b} &= (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \ \mathbf{y} = (\mathbf{X}'\mathbf{X})^{-1} \ \mathbf{X}' (\mathbf{X}\boldsymbol{\beta} + \boldsymbol{\varepsilon}) = \boldsymbol{\beta} + (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \boldsymbol{\varepsilon} \\ \implies \mathbf{b} - \boldsymbol{\beta} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \boldsymbol{\varepsilon} \end{aligned}$ 

Under the typical assumptions, we can establish properties for **b**.

1)  $E[\mathbf{b}|\mathbf{X}] = E[\boldsymbol{\beta}|\mathbf{X}] + E[(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\boldsymbol{\epsilon}|\mathbf{X}]$ =  $\boldsymbol{\beta} + (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' E[\boldsymbol{\epsilon}|\mathbf{X}] = \boldsymbol{\beta}$  (b is unbiased.)

2)  $\operatorname{Var}[\mathbf{b}|\mathbf{X}] = \operatorname{E}[(\mathbf{b} - \boldsymbol{\beta}) (\mathbf{b} - \boldsymbol{\beta})' | \mathbf{X}] = \operatorname{E}[(\mathbf{X}'\mathbf{X})^{-1} \mathbf{X}' \boldsymbol{\epsilon} \boldsymbol{\epsilon}' \mathbf{X} (\mathbf{X}'\mathbf{X})^{-1}]$ =  $(\mathbf{X}'\mathbf{X})^{-1} \mathbf{X}' \operatorname{E}[\boldsymbol{\epsilon} \boldsymbol{\epsilon}' | \mathbf{X}] \mathbf{X} (\mathbf{X}'\mathbf{X})^{-1}$ =  $(\mathbf{X}'\mathbf{X})^{-1} \mathbf{X}' \{\sigma^2 \mathbf{I}_{\mathrm{T}}\} \mathbf{X} (\mathbf{X}'\mathbf{X})^{-1} = \sigma^2 (\mathbf{X}'\mathbf{X})^{-1} \mathbf{X}'\mathbf{X} (\mathbf{X}'\mathbf{X})^{-1}$ =  $\sigma^2 (\mathbf{X}'\mathbf{X})^{-1} (kxk)$  matrix

3) Gauss-Markov Theorem: b is BLUE (*Best Linear Unbiased Estimator*). No other linear & unbiased estimator has a lower variance.

Proof:

Let  $\mathbf{b}^* = \mathbf{C}\mathbf{y}$  (linear in  $\mathbf{y}$ )  $E[\mathbf{b}^*|\mathbf{X}] = E[\mathbf{C}\mathbf{y}|\mathbf{X}] = E[\mathbf{C}(\mathbf{X}\boldsymbol{\beta} + \boldsymbol{\varepsilon})|\mathbf{X}] = \boldsymbol{\beta}$  (unbiased if  $\mathbf{C}\mathbf{X}=\mathbf{I}$ )  $\operatorname{Var}[\mathbf{b}^*|\mathbf{X}] = E[(\mathbf{b}^* - \boldsymbol{\beta})(\mathbf{b}^* - \boldsymbol{\beta})'|\mathbf{X}] = E[\mathbf{C}\boldsymbol{\varepsilon} \boldsymbol{\varepsilon}'\mathbf{C}'|\mathbf{X}] = \sigma^2 \mathbf{C}\mathbf{C}'$ 

Now, we relate  $Var[\mathbf{b}|\mathbf{X}]$  to  $Var[\mathbf{b}^*|\mathbf{X}]$ .

Let  $\mathbf{D} = \mathbf{C} - (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'$  (note  $\mathbf{D}\mathbf{X} = 0$ )

Then,

 $Var[\mathbf{b}^*|\mathbf{X}] = \sigma^2 (\mathbf{D} + (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}') (\mathbf{D}' + \mathbf{X}(\mathbf{X}'\mathbf{X})^{-1})$ =  $\sigma^2 \mathbf{D}\mathbf{D}' + \sigma^2 (\mathbf{X}'\mathbf{X})^{-1} = Var[\mathbf{b}|\mathbf{X}] + \sigma^2 \mathbf{D}\mathbf{D}'.$ 

Since **DD**' is positive definite  $\Rightarrow$  Var[**b**\*|**X**] > Var[**b**|**X**] •

4) If we make an additional assumption:

(A5)  $\boldsymbol{\varepsilon} | \mathbf{X} \sim i.i.d. \ \mathrm{N}(\mathbf{0}, \sigma^2 \mathbf{I}_{\mathrm{T}})$ we can derive the distribution of **b**. Since  $\mathbf{b} = \mathbf{\beta} + (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{\epsilon}$ , we have that **b** is a linear combination of normal variables and, thus, follows a normal distribution:

**b**| $\mathbf{X} \sim i.i.d.$  N( $\boldsymbol{\beta}, \sigma^2 (\mathbf{X}'\mathbf{X})^{-1}$ )

then,  $SD[\mathbf{b}|\mathbf{X}] = sqrt(diagonal elements of \sigma^2 (\mathbf{X}'\mathbf{X})^{-1})$ 

<u>Note</u>: The marginal distribution of a multivariate normal distribution is also normal, then  $b_k | \mathbf{X} \sim N(\beta_k, v_k^2)$ Std Dev  $[b_k | \mathbf{X}] = sqrt \{ [\sigma^2 (\mathbf{X}' \mathbf{X})^{-1}]_{kk} \} = v_k$ 

<u>Remark</u>: With (A5) we can do tests of hypothesis.

5) If (A5) is not assumed, we still can obtain a (limiting) distribution for **b**. Under additional assumptions –mainly, the matrix X'X does not explode as T becomes large–, as  $T \rightarrow \infty$ 

(i) $\mathbf{b} \xrightarrow{p} \mathbf{\beta}$	( <b>b</b> is consistent)
(ii) $\mathbf{b} \stackrel{a}{\rightarrow} N(\boldsymbol{\beta}, \sigma^2 (\mathbf{X}'\mathbf{X})^{-1})$	( <b>b</b> is asymptotically normal)

• Properties (1)-(4) are called *finite* (or *small*) sample properties, they hold for every sample size.

• Properties (5.i) and (5.ii) in (5) are called *asymptotic* properties, they only hold when *T* is large (actually, as *T* tends to  $\infty$ ). Property (5.ii) is very important: When the errors are not normally distributed we still can do testing about  $\beta$ , but we rely on an "approximate distribution."

## **OLS Estimation – Fitted Values and Residuals**

OLS estimates  $\beta$  with **b.** Now, we define *fitted values* as:

 $\widehat{y} = X b$ 

Now we define the estimated error, *e*:

$$e = y - \hat{y}$$

e represents the unexplained part of y, what the regression cannot explain. They are usually called *residuals*.

Note that **e** is uncorrelated (orthogonal) with  $\mathbf{X} \implies \mathbf{\epsilon} \perp \mathbf{X}$  $\mathbf{e} = \mathbf{y} - \mathbf{X}\mathbf{b} \implies \mathbf{X}'\mathbf{e} = \mathbf{X}' (\mathbf{y} - \mathbf{X}\mathbf{b}) = \mathbf{X}'\mathbf{y} - \mathbf{X}'\mathbf{X} (\mathbf{X}'\mathbf{X})^{-1} \mathbf{X}'\mathbf{y} = \mathbf{0}$ 

Using **e**, we can define a measure of unexplained variation: Residual Sum of Squares (RSS) =  $e'e = \sum_i e_i^2$ 

## **OLS Estimation** – Var[b|X]

We use the variance to measure precision of estimates. For OLS:  $Var[\mathbf{b}|\mathbf{X}] = \sigma^2 (\mathbf{X}'\mathbf{X})^{-1}$ 

**Example**: One explanatory variable model.

$$(\mathbf{A1'}) \text{ DGP: } \mathbf{y} = \beta_1 + \beta_2 \mathbf{x} + \mathbf{\epsilon}$$
  

$$\operatorname{Var}[\mathbf{b}|\mathbf{X}] = \sigma^2 (\mathbf{X'X})^{-1} = \sigma^2 \begin{bmatrix} \sum_i 1 & \sum_i 1x_i \\ \sum_i 1x_i & \sum_i x_i^2 \end{bmatrix}^{-1} = \sigma^2 \begin{bmatrix} T & T\bar{x} \\ T\bar{x} & \sum_i x_i^2 \end{bmatrix}^{-1}$$
  

$$= \sigma^2 \frac{1}{T(\sum_i x_i^2 - T\bar{x}^2)} \begin{bmatrix} \sum_i x_i^2 & -T\bar{x} \\ -T\bar{x} & T \end{bmatrix}$$
  

$$\operatorname{Var}[\mathbf{b}_1|\mathbf{X}] = \sigma^2 \frac{\sum_i x_i^2}{T(\sum_i x_i^2 - T\bar{x}^2)} = \sigma^2 \frac{\sum_i x_i^2/T}{\sum_i (x_i - \bar{x})^2}$$
  

$$\operatorname{Var}[\mathbf{b}_2|\mathbf{X}] = \sigma^2 \frac{1}{(\sum_i x_i^2 - T\bar{x}^2)} = \sigma^2 \frac{1}{\sum_i (x_i - \bar{x})^2}$$
  

$$\operatorname{Covar}[\mathbf{b}_1, \mathbf{b}_2|\mathbf{X}] = \sigma^2 \frac{-\bar{x}}{\sum_i (x_i - \bar{x})^2}.$$

• In general, we do not know  $\sigma^2$ . It needs to be estimated. We estimate  $\sigma^2$  using the residual sum of squares (RSS):

 $RSS = \sum_i e_i^2$ The natural estimator of  $\sigma^2$  is  $\hat{\sigma}^2 = RSS/T$ . Given the LLN, this is a consistent estimator of  $\sigma^2$ . However, this not unbiased.

• The unbiased estimator  $\sigma^2$  is  $s^2$  $s^2 = \text{RSS}/(T-k) = \sum_i e_i^2/(T-k) = e'e/(T-k)$ 

To get E[ $s^2$ ], we use a property of a RV with a  $\chi_v^2$  distribution: E[ $\chi_v^2$ ] = v

Given that

$$(T-k) s^2/\sigma^2 \sim \chi^2_{T-k}.$$

Then,

$$E[(T - k) s^{2}/\sigma^{2} | \mathbf{X}] = (T - k)$$
  

$$E[\mathbf{e}'\mathbf{e}/\sigma^{2} | \mathbf{X}] = (T - k)$$
  

$$E[\mathbf{e}'\mathbf{e}/(T - k) | \mathbf{X}] = E[s^{2}|\mathbf{X}] = \sigma^{2} \qquad \Rightarrow E[s^{2}|\mathbf{X}] = \sigma^{2}$$

Note: (*T-k*) is referred as a *degrees of freedom* correction.

• Then, the estimator of  $Var[\mathbf{b}|\mathbf{X}] = s^2 (\mathbf{X}'\mathbf{X})^{-1}$  (a *kxk* matrix)

This estimator gives us the *standard errors* (SE) of the individual coefficients. For example, for the  $b_k$  coefficient:

$$\operatorname{SE}[b_k|\mathbf{X}] = \operatorname{sqrt}[s^2(\mathbf{X}'\mathbf{X})^{-1}]_{kk} = s_{b,k}$$

#### **OLS Estimation – Testing Only One Parameter**

We are interested in testing a hypothesis about one parameter in our linear model:  $y = X \beta + \epsilon$ 

1. Set H<sub>0</sub> and H<sub>1</sub> (about only one parameter): H<sub>0</sub>:  $\beta_k = \beta_k^0$ H<sub>1</sub>:  $\beta_k \neq \beta_k^0$ . **2.** Appropriate T(X): *t-statistic*. To derive the distribution of the test under H<sub>0</sub>, we will rely on assumption (A5)  $\varepsilon | X \sim N(0, \sigma^2 I_T)$  (otherwise, results are only asymptotic).

Let  $b_k = \text{OLS estimator of } \beta_k$  $\text{SE}[b_k|\mathbf{X}] = \text{sqrt}\{[s^2(\mathbf{X}^*\mathbf{X})^{-1}]_{kk}\} = s_{b,k}$ 

From assumption (A5), we know that

$$(b_k | \mathbf{X} \sim \mathrm{N}(\beta_k, \mathrm{v}_k^2) \qquad \Rightarrow \text{Under H}_0: \quad b_k | \mathbf{X} \sim \mathrm{N}(\beta_k^0, s_{b,k}^2).$$
$$\Rightarrow \text{Under H}_0: \quad t_k = \frac{b_k - \beta_k^0}{s_{b,k}} | \mathbf{X} \sim t_{T-k}.$$

• We measure distance in standard error units:

$$t_k = \frac{b_k - \beta_k^0}{s_{b,k}}$$

<u>Note</u>:  $t_k$  is an example of the *Wald* (normalized) *distance measure*. Most tests statistics in econometrics will use this measure.

**3.** Compute  $t_k$ ,  $\hat{t}$ , using  $b_k$ ,  $\beta_k^0$ , s, and  $(\mathbf{X}'\mathbf{X})^{-1}$ . Get *p*-value( $\hat{t}$ ).

<b>4.</b> <u>Rule</u> : Set an $\alpha$ level. If <i>p</i> -value( $\hat{t}$ ) < $\alpha$	$\Rightarrow$ Reject H <sub>0</sub> : $\beta_k = \beta_k^0$
Alternatively, if $ \hat{\mathbf{t}}  > t_{T-k.1-\alpha/2}$	$\Rightarrow$ Reject H <sub>0</sub> : $\beta_k = \beta_k^0$ .

• Special case: H<sub>0</sub>:  $\beta_k = 0$ 

H<sub>1</sub>: 
$$\beta_k \neq 0$$
.

Then,

$$t_k = \frac{b_k}{\operatorname{sqrt}\{s^2(\mathbf{X}'\mathbf{X})^{-1}\}_{kk}} = \frac{b_k}{\operatorname{SE}[b_k]} \qquad \Rightarrow t_k \sim t_{T-k}.$$

This special case of  $t_k$  is called the *t-value* or *t-ratio* (also refer as the "t-stats"). That is, the t-value is the ratio of the estimated coefficient and its SE.

• The t-value is routinely reported in all regression packages. In the lm() function, it is reported in the third column of numbers.

• Usually,  $\alpha = 5\%$ , then if  $|t_k| > 1.96 \approx 2$ , we say the coefficient b<sub>k</sub> is "significant."

**Example:** Using the 1-factor CAPM for IBM returns, we test if IBM's market  $\beta = 1$ , that is, if IBM bears the same market risk as the market. WE use R *lm* function.

 $\begin{array}{ll} T <- \mbox{ length}(x\_ibm) & \# \mbox{ Sample size} \\ \mbox{lr\_ibm} <- \mbox{ log}(x\_ibm[-1]/x\_ibm[-T]) & \# \mbox{ Log returns for IBM (lost one observation)} \\ \mbox{Mkt}_RF <- x\_Mkt_RF[-1]/100 & \# \mbox{ Adjust size (take one observation out )} \\ \mbox{RF} <- x\_RF[-1]/100 & \# \mbox{ Adjust size (take one observation out )} \\ \mbox{ibm } x <- \mbox{ lr ibm} - \mbox{RF} & \# \mbox{ Define excess returns for IBM} \end{array}$ 

fit\_ibm\_capm <- lm(ibm\_x ~ Mkt\_RF) # OLS estimation with lm package in R
> summary(fit\_ibm\_capm)

Coefficients:

Estimate Std. Error t value Pr(>|t|) (Intercept) -0.005791 0.002487 -2.329 0.0202 \* xMkt\_RF 0.895774 0.053867 16.629 <2e-16 \*\*\* ---Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 b ibm <- fit ibm capm\$coefficients # Extract from Im function OLS coefficients

SE\_ibm <- sqrt(diag(vcov(fit\_ibm\_ff3))) # SE from fit\_ibm (also a kx1 vector) t beta1 <- (b ibm[2] - 1)/SE ibm[2] # t-stat for H<sub>0</sub>:  $\beta_1$  - 1

> t\_beta1

[1] -1.934877  $\Rightarrow |\hat{t}_1 = -1.934877| < 1.96 \Rightarrow \text{Cannot reject H}_0 \text{ at 5\% level}$ 

p\_val <- (1- pnorm(abs(t\_beta1))) \* 2 # pvalue for t\_beta (adjusted b/c two sided test) > p\_val [1] 0.0530  $\Rightarrow$  cannot reject H<sub>0</sub>:  $\beta_{IBM} = 1$  at 5% level, but a borderline case!

<u>Conclusions</u>: Cannot reject H<sub>0</sub>:  $\beta_{IBM} \Rightarrow$  IBM has a one-to-one risk relation with the market, but borderline test decision!

Note: You can get same results using linear algebra. From last class:

$b \le solve(t(x)\%*\% x)\%*\% t(x)\%*\%$	у	$\# \mathbf{b} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \mathbf{y} \text{ (OLS regression)}$
SE <- sqrt(diag(vcov( <b>fit_ibm_capm</b> )	))	# SE from fit_ibm (also a <i>kx</i> 1 vector)
$t_{beta1} <- (b[2] - 1)/SE[2]$	# t-stat	for $H_0$ : Beta <sub>1</sub> - 1

```
> t(b)
```

```
Mkt_RF
[1,] -0.005791039 0.895773564
> t(SE_b)
Mkt_RF
[1,] 0.002487 0.053867
> t_beta1
```
[1] **-1.934877**). ¶

# **OLS Estimation – Testing Only One Parameter: The CAPM**

Recall that the CAPM states:

 $E[r_{i,t} - r_{f,t}] = \beta_i E[(r_{M,t} - r_{f,t})].$ 

A linear data generating process (DGP) consistent the CAPM is:

 $r_{i,t} - r_{f,t} = \alpha_i + \beta_i (r_{M,t} - r_{f,t}) + \varepsilon_{i,t}, \qquad i = 1, ..., N \& t = 1, ..., T$ 

Then, using the time series of stock returns, we test the CAPM for asset *i* by testing:

H<sub>0</sub> (CAPM holds):  $\alpha_i = 0$ 

H<sub>1</sub> (CAPM rejected):  $\alpha_i \neq 0$ 

**Example**: Testing the CAPM for IBM returns with time series.

For IBM, we test the CAPM by testing:	H <sub>0</sub> (CAPM holds): $\alpha_{i=IBM} = 0$
	H <sub>1</sub> (CAPM rejected): $\alpha_{i=IBM} \neq 0$
SFX da <-	
read.csv("https://www.bauer.uh.edu/rsusme	el/4397/Stocks_FX_1973.csv",head=TRUE,sep=",")
x_ibm <- SFX_da\$IBM	# Read IBM price data
x_Mkt_RF <- SFX_da\$Mkt_RF	# Read Market excess returns (in %)
x_RF <- SFX_da\$RF	
$T \le length(x_ibm)$	# Sample size
$lr_ibm \le log(x_ibm[-1]/x_ibm[-T])$	# Log returns for IBM (lost one observation)
$Mkt_RF \le x_Mkt_RF[-1]/100$	# Adjust size (take one observation out )
$RF <- x_RF[-1]/100$	
ibm_x <- lr_ibm – RF	# Define excess returns for IBM
<pre>fit_ibm_capm &lt;- lm(ibm_x ~ Mkt_RF)</pre>	# OLS estimation with <i>lm</i> package
<pre>&gt; summary(fit_ibm_capm)</pre>	
Coefficients:	
Estimate Std. Error t value	Pr(> t )
(Intercept) -0.005791 0.002487 -2.329 0	.0202 *
xMkt_RF 0.895774 0.053867 16.629 <	2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0	0.05 `.' 0.1 ` ' 1

• Q: Is the intercept ( $\alpha_{\text{IBM}}$ ) equal to 0? We use the t-value:  $t_{\alpha_i} = \frac{\alpha_{i=IBM}}{\text{SE}[\alpha_{i=IBM}]} = \frac{-0.005791}{0.002487} = -2.329 \quad \text{(the t-value)}$   $\Rightarrow |\hat{t}_{\alpha} = -2.329| > 1.96 \quad \Rightarrow \text{Reject H}_0 \text{ at 5\% level}$  Conclusion: The CAPM is rejected for IBM at the 5% level.

<u>Note</u>: You can also reject H<sub>0</sub> by looking at the *p*-value of intercept (0.0202 < .05)

<u>Interpretation</u>: Given that the intercept is significant (& negative), IBM underperformed relative to what the CAPM expected:

- IBM excess returns: mean(ibm\_x) = -0.00073141

- IBM excess returns (CAPM) = 0.895774 \* mean(Mkt RF)

= 0.895774 \* 0.0056489 = 0.0050601

- Ex-post difference: -0.00073141 - 0.0050601 = -0.00579151 ( $\approx \alpha_{IBM}$ ).

<u>Remark</u>: Above we tested (& rejected) the CAPM for one asset only, IBM. But, the CAPM should apply to all assets, that is, in the cross-section. Suppose we have N assets. Then, a test for the CAPM involves testing  $\alpha_i$ 's:

H<sub>0</sub>:  $\alpha_1 = \alpha_2 = \dots = \alpha_N = 0$ H<sub>0</sub>: at least one  $\alpha_i \neq 0$ .

This test is a **joint** test. It requires a simultaneous estimation of *N* CAPM DGPs (with a constant). There are different ways to approach testing the CAPM in a cross-sectional setting. The popular approaches use a **two-step estimation**, for example, the popular Fama-MacBeth (1973) two-step estimation.

#### **OLS Estimation – Testing The CAPM (Cross-Section)**

The CAPM tells also a cross-section story for asset returns: Assets with higher  $\beta_i$  should get, on average, higher compensation.

CAPM (cross-section):  $E[r_{i,t} - r_f] = \beta_i \lambda$ where  $\lambda$ , in equilibrium, is the market excess return (or factor return). It is sometimes referred as the *price of risk*.

If we have  $\beta_i$ 's for *N* assets, we can estimate the *security market line* (SML), where we show the effect of  $\beta_i$  on  $E[r_{i,t} - r_f]$ . Below we show the SML in red. All stocks on the SML are priced correctly, all above are underpriced securities, that is, the return is higher than what is expected for a given level of risk ( $\beta_i$ ). Similarly, all stocks under the SML are overpriced. For example, above we determined that IBM has a  $\beta_{i=IBM} = 0.895774$ , but its average excess return is negative, -0.00073141. Then, IBM is under the SML line and, thus, IBM is overvalued.



The SML answers the question: Which stocks deserve higher returns? Assets with higher exposure to market risk –i.e., higher  $\beta_i$ .

A linear cross-sectional DGP consistent with the CAPM is:  $r_i - r_f = \alpha + \beta_i \ \lambda + \varepsilon_i, \qquad i = 1, ..., N$ 

Testing implication of the SML for the cross-section of stock returns:

H<sub>0</sub> (CAPM holds in the CS):  $\alpha = 0$  and  $\lambda = E[r_{m,t} - r_f]$ 

H<sub>1</sub> (CAPM rejected in the CS):  $\alpha \neq 0$  and/or  $\lambda \neq E[r_{m,t} - r_f]$ .

Again, we have a **joint** test. As metioned above, there are different ways to approach this simultaneous estimation, a common approach is the Fama-MacBeth (1973) two-step estimation.

Fama-French (1992, 1993) adapted the Fama-MacBeth (1973) procedure, to produce a wellknow approach to test the relevance of the CAPM in explaining the cross-section of stock returns. The first step, or *first pass*, involves a time-series regression to estimate the betas for each asset (IBM, GE, MSFT, etc.), the second step, or *second pass*, regresses the average excess return of eah asset (IBM, GE, MSFT, etc.) on the estimated betas (OLS **b**'s, also called "*factor loadings*"). That is:

#### (1) First Pass

Using the time series (*T* observations), run a CAPM regression to estimate  $\beta_i$  for each asset i = 1, ..., N.

$$r_{i,t} - r_{f,t} = \alpha_i + \beta_i (r_{M,t} - r_{f,t}) + \varepsilon_{i,t}, \qquad t = 1, \dots, T \quad \Rightarrow \text{ Get } N \text{ b}_i \text{'s.}$$

#### (2) Second Pass

Using the  $N b_i$ 's as regressors, estimate

$$(\bar{r}_i - \bar{r}_f) = \alpha + b_i \lambda + \varepsilon_i,$$
  $i = 1, ...,$ 

Ν

where  $(\bar{r}_i - \bar{r}_f)$  is the average excess return of asset *i* in our sample.

According to the CAMP, we expect  $\lambda > 0$ .

The usual execution of almost all 2-step procedures involves:

Since returns are estimated with a lot of noise, portfolios are used, not individual securities.
 The estimation takes into account the possible change over time of beta coefficients, by estimating the coefficients every 5 or 10 years.

**Example**: We test the CAPM, in the cross-section, using the 2-step Fama-French approach. We use returns of 25 Fama-French portfolios, sorted by Size (ME) and Book-to-Market (BM). We downloaded the returns of the 25 portfolios, along the 3-Fama-French factors, from Ken French's website:

 $https://mba.tuck.dartmouth.edu/pages/faculty/ken.french/data\_library.html \# Research.$ 

Note: the files are zip files. I unzipped them and put them on my website.

FF\_p\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/FF\_25\_portfolios.csv", head=TRUE, sep=",") FF\_f\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/FF\_3\_factors.csv", head=TRUE, sep=",")

# Extract variables from imported data

}

Mkt_RF_fm <- FF_f_da\$Mkt_RF	<pre># extract Market excess returns (in %)</pre>
HML_fm <- FF_f_da\$HML	# extract HML returns (in %)
SMB fm <- FF f da\$SMB	# extract HML returns (in %)
$RF_{fm} \leq FF_{fa} = FF_{fa}$	# extract Risk-free rate (in %)
$Y_p <- FF_p_da[,2:26] - RF_fm$	# Compute excess returns of 25 portfolios
T <- length(HML_fm)	# Number of observations (1926:July on)
$x0 \leq matrix(1,T,\overline{1})$	# Vector of ones, represents constant in X
$x \le cbind(x0, Mkt_RF_fm)$	# Matrix X (Tx2)
$k \leq -ncol(Y_p)$	
## First Pass (CAPM to get get N	$\mathbf{b}_i$ )
Allbs = NULL	# Initialize empty (a space to put betas)
for (i in seq(1,k,1)){	
$y \le Y p[,i] $ # select Y (portfolie	0)
$b \le solve(t(x)) \times t(x) \times t(x)$	*%y # OLS regression = $(X'X)^{(-1)} X'y$
Allbs =cbind(Allbs,b)	# accumulate b as rows

beta\_ret <- cbind(colMeans(Y\_p),t(Allbs)) # Mean portfolio returns along alpha & beta
estimates</pre>

```
cor(beta_ret[,1], beta_ret[,3]) # Correlation of mean portfolio return & beta
> cor(beta_ret[,1], beta_ret[,3])
[1] 0.3326008
```

plot(beta\_ret[,1], beta\_ret[,3], main="Scatterplot: Portfolio Returns & CAPM Beta", xlab="Mean Portfolio Returns ", ylab="Market Beta", pch=19)



## Second Pass (cross-sectional SML)
fit\_fm\_capm\_25 <- lm(beta\_ret[,1] ~ beta\_ret[,3])
> summary(fit\_fm\_capm\_25)

Coefficients:

Estimate Std. Error t value Pr(>|t|)(Intercept) 0.3728 0.3113 1.198 0.243 beta ret[, 3] 0.4289 0.2536 **1.691** 0.104  $\Rightarrow$  Not significant: Beta plays no role!

Residual standard error: 0.2 on 23 degrees of freedom Multiple R-squared: 0.1106, Adjusted R-squared: 0.07195 F-statistic: 2.861 on 1 and 23 DF, p-value: 0.1043

<u>Conclusion</u>: The CAPM's beta is not significant. That is, in the cross-section, Beta plays no role in determining expected returns for assets.  $\P$ 

• Fama and French (1992, 1993) estimate variations of the DGP with more factors. They find that  $\beta$  is weakly significant or not significant ("*Beta is dead*") in explaining the C-S of stock returns. They even report the wrong sign for beta in the second pass.

But, other researchers dispute the "Beta is dead" finding, criticizing the selection of estimation period, construction of portfolios, number of factors, statistical problems like measurement error and incorrect SE, etc.

The debate about  $\beta$  & what (& how many) factors to include in the DGP continues.

# **The 3-Factor Fama-French Model**

As mentioned above, the CAPM is routinely rejected. A popular alternative model is the empirically derived 3-factor Fama-French (1993) model, which adds two rsik factors, related to firm characteristics, to the market factor:

a) *Size* factor (**SMB**): It is measured as the difference between the returns on diversified portfolios of small stocks (*small size portfolio*) and big stocks (*big size portfolio*) = long **S**mall & short **B**ig.

b) *Value* factor or book-to-market factor (**HML**): It is measured as the difference between the returns on diversified portfolios of high stocks (*high B/M portfolio*) and low B/M stock (*low B/M portfolio*) = long High & short Low.

Using monthly data from July 1927- July 2024 (98 years), we have the following returns for the three Fama-French macro-factors:

	$(r_{M,t} - r_{f,t})$	SMB <sub>t</sub>	$HML_t$
Min	-29.1300	-17.2000	-13.8800
Median	1.0600	0.0700	0.1200
Mean	0.6845	0.1801	0.3449
Max	38.8500	36.5600	35.6100
SD	5.33431	3.176139	3.568389

The annualized average returns (& SD) for Market, SMB, and HML are, respectively, **8.21%** (18.48%), **2.16%** (11%) and **4.14%** (12.36%). High premiums, but with high estimation uncertainty (high SD).

Below, we plot the SMB and HML cumulative returns



The SMB or size factor performed very well up to about 1982, generating over 600% returns. Then, on average, the reverse occurred: large-cap stocks outdid small caps. The value factor performed extremely well up to financial crisis 2008, generating over 4,500% returns. Value investing was very profitable. Since, then, on average, the pattern has changed.

• The three (macro-) factors are, in theory, "*factor mimicking portfolios*," that is, portfolios with exposure only to the factor in question (market, size, or value), and no exposure to any other factor.

The correlation matrix below show that, though not high, the correlations are not zero, especially with the market.

	Mkt_RF	SMB	HML
Mkt_RF	1.00000	0.31507	0.22697
SMB	0.31507	1.00000	0.12044
HML	0.22697	0.12044	1.00000

<u>Note</u>: For a long time, before the early 1980s, any significant factor, beyond the market factor, was considered a "CAPM *anomaly*."

# The 3-Factor Fama-French Model: Construction of Factors

The portfolios are formed as follows:

Step 1. At the end of June of year *t*, sort the stock returns by attribute (size of Size, or B/M).

**Step 2**. Split the sorted assets by attribute into 3 equal/value-weighted portfolios (3 *tercile* portfolios). Split can be thinner (quintile portfolios) or based on more complicated sorts, for example, using 6 portfolios constructed by intersecting 2 size portfolios & 3 value portfolios.

**Step 3**. At the end of each month (week or day), from July of year t to June of year t + 1, based on the portfolios constructed in **Step 1**, compute the returns of each of the split portfolios.

**Step 4** Form a "hedge portfolio": long the top portfolio (say, top tercile) and short the bottom portfolio (say, bottom tercile).

<u>Note</u>: The portfolios for July of year t to June of t+1 include all NYSE, AMEX, and NASDAQ stocks for which there is data for June of t.

This approach to construct factors is very popular. Using this procedure it is possible to construct other factors, for example, a PE factor or a Profitability factor, where we sort stock by PE or Profitability.

# The 3-Factor Fama-French Model: Intrepretation of Coefficients

A linear DGP generating the 3-factor Fama-French model is:

 $r_{i,t} - r_f = \alpha_i + \beta_1 (r_{m,t} - r_f) + \beta_2 \operatorname{SMB}_t + \beta_3 \operatorname{HML}_t + \varepsilon_{i,t}.$ 

That is, according to this model, excess returns are driven by their sensitivity to the market, to size, and to value stocks, as measured by the book-to-market ratio. The estimated parameters in

the time-series regressions are called *"factor loadings,"* they measure the sensitivity of asset *i* to changes in the factor.

• Interpretation of coefficients:

-  $\beta_1$  has the same as the interpretation in the CAPM, it measures the relation between asset *i* risk and market risk.

-  $\beta_2$  measures how tilted asset *i* is towards small stock (in general,  $\beta_2 > 0$  means that returns of asset *i* behaves like small stocks.

-  $\beta_3$  measures how tilted asset *i* is towards value stock (in general,  $\beta_3 > 0$  means that returns of asset *i* behave like high book-to-market stocks).

# The 3-Factor Fama-French Model: Testing the model (in the Time Series)

Like the CAPM, the 3-factor Fama-French model produces expected excess returns:  $E[r_{i,t} - r_{f,t}] = \beta_1 E[r_{m,t} - r_f] + \beta_2 E[SMB_t] + \beta_3 E[SMB_t]$ 

A significant constant,  $\alpha_i$ , would be evidence against this model: something is missing in the model. Thus, using the time series of asset *i* to estimate the 3-factor FF model DGP with a constant, we test the FF model for asset *i*:

H<sub>0</sub> (3-factor FF model true) =  $\alpha_i$ 

H<sub>1</sub> (3-factor FF model not true) =  $\alpha_i$ 

This test can be implemented by a simple t-test on the estimated  $\alpha_i$ .

**Example (continuation)**: Using the time-series, we test the significance of the 3-factor Fama-French model for IBM returns:

```
SFX da <-
read.csv("http://www.bauer.uh.edu/rsusmel/4397/Stocks FX 1973.csv",head=TRUE,sep=",")
x ibm <- SFX da$IBM
                                   # Extract IBM price data
x Mkt RF <- SFX da$Mkt RF
                                   # Extract Market excess returns (in %)
x SMB <- SFX da$SMB
                                   # Extract SMB factor returns (in %)
x HML <- SFX da$HML
                                   # Extract HML factor returns (in %)
                            # Extract Risk-free rate factor returns (in %)
x RF <- SFX da$RF
Mkt RF <- x Mkt RF[-1]/100
                                   # Adjust size (take one observation out )
SMB \le x SMB[-1]/100
                                   # Adjust size (take one observation out )
HML \le x HML[-1]/100
                                   # Adjust size (take one observation out )
RF \le x RF[-1]/100
T \leq - length(x ibm)
                                   # Sample size
\ln ibm < \log(x ibm[-1]/x ibm[-T])
                                         # IBM log returns (lost one observation)
ibm x <- lr ibm - RF
fit ibm ff3 <- lm(ibm x \sim Mkt RF + SMB + HML)
> summary(fit ibm ff3)
                                          # print lm results
Coefficients:
             Estimate
                            Std. Error t value Pr(>|t|)
                            0.002488 -2.046 0.0412 *
             -0.005089
(Intercept)
```

Mkt_RF	0.908299	0.056722 16.013	<2e-16 **	**
SMB	-0.212460	0.084112 -2.526	0.0118 *	$\Rightarrow$ significant at 5% level.
HML	-0.171500	0.084682 -2.025	0.0433 *	$\Rightarrow$ significant at 5% level.

Residual standard error: 0.05848 on 565 degrees of freedom Multiple R-squared: 0.3389, Adjusted R-squared: 0.3354 F-statistic: 96.55 on 3 and 565 DF, p-value: < 2.2e-16

<u>Conclusion</u>: Consistent with the 3-factor Fama-French model, Mkt\_RF, SMB and HML are drivers of the expected returns for IBM. The signs of  $\beta_2 \& \beta_3$ : IBM behaves like a large & low B/M firm.

Note 1: The constant is significant, that is, there is an "extra" component of expected returns not explained by the 3 F-F factors.

<u>Note 2</u>: The CAPM is also rejected for IBM, since there are, beyond the market factor, other significant factors.

• Now, to gauge the behavior of the 3-factor model, we plot fitted IBM values and compare with actual IBM values.





We observe some periods with good fit –mainly early and late periods- and some periods with poor fit –mainly the middle period.

• In the context of the 3-factor Fama-French model, we test again if IBM's risk has a one-to-one relation with the market risk. That is, we test if the market beta ( $\beta_1$ ) is equal to 1. Formally, we test:

H<sub>0</sub>:  $\beta_1 = 1$ H<sub>1</sub>:  $\beta_1 \neq 1$ 

Using the previous estimation, we have:

 $\Rightarrow t_k = (b_k - \beta_k^0) / \text{Est. SE}(b_k)$   $\hat{t}_1 = (0.9082989 - 1) / 0.05672206 = -1.616674$  $\Rightarrow |\hat{t}_1 = -1.616674 | < 1.96 \Rightarrow \text{Cannot reject H}_0 \text{ at 5\% level.}$ 

Conclusion: IBM bears the same market risk as the market.

<u>R Note</u>: You should get the same numbers using R's *lm* and extracting information from lm:

b_ibm <- fit_ibm_ff3\$coefficients	# Extract from lm function OLS coefficients
SE_ibm <- sqrt(vcov(fit_ibm_ff3))	# SE from fit_ibm (also a kx1 vector)
$t_beta1 \le (b_ibm[2] - 1)/SE_ibm[2]$	# t-stat for H <sub>0</sub> : $\beta_1 = 1$ .
> t_beta1	
[1] -1.616674	
p_val <- 1- pnorm(abs(t_beta1))	<pre># pvalue for t_beta1 (one-sided)</pre>
> p_val * 2	# two-sided test (multiply one-side p-value by 2)
[1] <b>0.1059487.</b> ¶	

 A cross-section test of the 3-factor FF model is a joint test: H<sub>0</sub>: α<sub>1</sub> = α<sub>2</sub> = .... = α<sub>N</sub> = 0 H<sub>1</sub>: at least one α<sub>i</sub> ≠ 0.

 A Fama-MacBeth (1973) two-step procedure is usually implemented.

# The 3-Factor Fama-French Model: Testing Multi-factor Models (Cross-Section)

The 2-step framework can be extended to include more factors in both steps, for example, Fama-French (1993) introduce their well-known 3-factor model to test if beta is a significant driver of expected excess returns in the cross-section. Fama and French (1993) added to the CAPM a size factor (SMB) and a value factor (HML). Then, for asset *i* we have

$$r_{i,t} - r_{f,t} = \alpha_i + \beta_{1,i} (r_{M,t} - r_{f,t}) + \beta_{2,i} SMB_t + \beta_{3,i} HML_t + \varepsilon_{i,t}, \quad t = 1, ..., T$$

In this case, the Fama-MacBeth two-step procedure involves:

#### (1) First pass

Using the time series (*T* observations), run a regression with the 3 Fama-French factors (Market, SMB, HML) to estimate 3  $\beta_i$ 's for each asset i = 1, ..., N.

$$\begin{aligned} r_{i,t} - r_{f,t} &= \alpha_i + \beta_{1,i} \ (r_{M,t} - r_{f,t}) + \beta_{2,i} \ SMB_t + \ \beta_{3,i} \ HML_t \ + \varepsilon_{i,t}, \qquad t = 1, \ \dots, T \\ &\implies \text{Get } N \ \mathbf{b}_i = [\mathbf{b}_{1,i}, \ \mathbf{b}_{2,i}, \ \mathbf{b}_{3,i}]. \end{aligned}$$

(2) Second Pass

Using the  $N \mathbf{b}_i$ 's as regressors, estimate

 $(\bar{r}_i - \bar{r}_f) = \alpha + b_{1,i} \lambda_1 + b_{2,i} \lambda_2 + b_{3,i} \lambda_3 + \varepsilon_i,$ where  $(\bar{r}_i - \bar{r}_f)$  is the average excess return of asset *i* in our sample.

Using the estimates from the second pass, we test

H<sub>0</sub> (FF holds in the CS):  $\alpha = 0 \& \lambda_j > 0$ , j = 1, 2, k = 3H<sub>1</sub> (FF rejected in the CS):  $\alpha \neq 0$  and/or <u>Note</u>: In equilibrium,  $\lambda_1 = \mathbb{E}[r_{m,t} - r_f]$ ,  $\lambda_2 = \mathbb{E}[SMB_t] \& \lambda_3 = \mathbb{E}[HML_t]$ .

**Example**: We test the 3-factor Fama-French model, in the cross-section, using the 2-step Fama-McBeth method. We use returns of **25 Fama-French portfolios** (sorted by Size & BM), downloaded, along the 3-Fama-French factors from Ken French's website.

FF\_p\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/FF\_25\_portfolios.csv", head=TRUE, sep=",") FF\_f\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/FF\_3\_factors.csv", head=TRUE, sep=",")

```
# Extract variables from imported data
Mkt RF fm <- FF f da$Mkt RF
                                 # extract Market excess returns (in %)
HML fm <- FF f da$HML
                                 # extract HML returns (in %)
SMB fm <- FF f da$SMB
                                 # extract HML returns (in %)
RF fm \leq FF f da RF
                                 # extract Risk-free rate (in %)
Y p <- FF p da[,2:26] - RF fm
                                        # Compute excess returns of 25 portfolios
T \leq - length(HML fm)
                                 # Number of observations (1926: July on)
x_0 < -matrix(1,T,1)
                                 # Vector of ones, represents constant in X
x <- cbind(x0, Mkt RF, SMB, HML)
                                        # Regressors (vector of ones + 3 Factors)
k \leq ncol(Y p)
## First Pass
Allbs = NULL
                                 # Initialize empty (a space to put betas)
for (i in seq(1,k,1))
y < -Y p[,i]
              # select Y (portfolio)
 # OLS regression = (X'X)^{(-1)} X'y
 Allbs =cbind(Allbs,b)
                                 # accumulate b as rows
beta ret <- cbind(colMeans(Y p),t(Allbs)) # Mean portfolio returns along alpha & beta
estimates
> cor(beta ret)
                           Mkt RF
                                                      HML
                                        SMB
             1.000000
                           0.5242033
                                        -0.2502789
                                                      0.26183205
                                                                   0.69403377
                                        -0.7533964
             0.524203
                           1.0000000
                                                      -0.3289295
                                                                   -0.02888856
Mkt RF
                           -0.7533967
                                        1.0000000
                                                      0.1747401
                                                                   0.11693857
             -0.250279
SMB
             0.261832
                           -0.328929
                                        0.1747402
                                                      1.0000000
                                                                   0.03922282
HML
             0.694034
                           -0.028888
                                        0.1169386
                                                      0.0392228
                                                                    1.0000000
```

plot(beta\_ret[,1], beta\_ret[,3], main="Scatterplot: Portfolio Returns & Market Beta", xlab="Mean Portfolio Returns ", ylab="Market Beta", pch=19)



<u>Conclusion</u>: The 3-factor FF model has a significant constant, which strong evidence against the model –i.e., something is missing. The negative coefficient of the market beta goes against theory. Question: Is Beta dead? Or, maybe, something is not right in the way we structure the test (incorrect model, problems with the data, the period, etc)? ¶

#### The 3-Factor Fama-French Model: Remarks & Extension

There is a big debate about the number of factors and how they are "discovered."

Q: How where these factors determined to be drivers of stock returns? By looking at data characteristics, not theory. As seen in the previous graphs, by the mid-1990s there was evidence that small firms outperformed big firms and that high BM firms outperformed low BM firms. Thus, data mining issues are likely present. Data mining issues are likely present. Q Are these 3 factors the definitive number of factor?

No. There have been over 200 factors proposed, counting interactions among factors, the potential number can be, easily, in the thousands. Many of these factors are likely a product of data mining, which is a problem for correct statistical inference and, more important, for out-of-sample forecasting. Feng, Giglio and Xiu (2020), who propose a methodology to select factors that explain the cross-section of expected return, call their paper "Taming the **Factor Zoo**."

• In 2014, Fama and French added two additional factors to their 3-factor model: RMW & CMA. - RMW measures the return of the portfolio of most profitable firms ("robust") minus the returns of portfolio least profitable ("weak").

- CMA measures the return of a portfolio of firms that invest conservatively minus the returns portfolio of firms that invest aggressively.

Again, the 5-factor FF model produces expected excess returns:

 $E[r_{i,t} - r_f] = \beta_1 E[r_{m,t} - r_f] + \beta_2 E[SMB_t] + \beta_3 E[SMB_t] + \beta_4 E[RMW_t] + \beta_5 E[CMA_t]$ 

There is debate regarding the validity or usefulness of this extension, especially, outside the U.S. market.

#### **OLS Estimation – Linear Algebra Interpretation**

• Disturbances and Residuals

In the population:  $E[\mathbf{X}' \boldsymbol{\epsilon}] = 0.$ In the sample:  $\mathbf{X}' \mathbf{e} = \mathbf{X}'(\mathbf{y} - \mathbf{X}\mathbf{b}) = \mathbf{X}'\mathbf{y} - \mathbf{X}'\mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y}$  $= 1/T(\mathbf{X}' \mathbf{e}) = 0.$ 

• We have two ways to look at **y**:

 $\mathbf{y} = \mathbf{E}[\mathbf{y}|\mathbf{X}] + \varepsilon = \text{Conditional mean} + \text{disturbance}$  $\mathbf{y} = \mathbf{X}\mathbf{b} + \mathbf{e} = \text{Projection} + \text{residual}$ 



# **OLS Estimation – Important Matrices: M**

Important Matrices

(1) "Residual maker	۶ <sup>د</sup> م	
<b>M</b> =	$I_{T} - X(X'X)^{-1}X'$	(TxT matrix)
My =	$= \mathbf{y} - \mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \ \mathbf{y} = \mathbf{y} - \mathbf{X}\mathbf{b} = \mathbf{e}$	(residuals)
MX	$= (\mathbf{I}_{\mathrm{T}} - \mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}') \mathbf{X} = 0$	
- <b>M</b> is symmetric	$-\mathbf{M}=\mathbf{M}'$	

	1.4	111	
- M is idempotent		$-\mathbf{M}^{*}\mathbf{M} = \mathbf{M}$	
- <b>M</b> is singular		$-\mathbf{M}^{-1}$ does not exist.	$\Rightarrow$ rank( <b>M</b> ) = $T - k$

- Special case:  $\mathbf{X} = \mathbf{i}$ 

$$\mathbf{M}^{0} = \mathbf{I} - \mathbf{i}(\mathbf{i}' \mathbf{i})^{-1} \mathbf{i}' = \mathbf{I} - \mathbf{i} \mathbf{i}'/T \qquad - \text{ since } \mathbf{i}' \mathbf{i} = T$$
  
$$\mathbf{M}^{0} \mathbf{y} = \mathbf{y} - \mathbf{i}(\mathbf{i}' \mathbf{i})^{-1} \mathbf{i}' \mathbf{y} = \mathbf{y} - \mathbf{i} \overline{\mathbf{y}} \qquad - \text{ since } \mathbf{i}' \mathbf{y}/T = \overline{\mathbf{y}}$$

$$\mathbf{M}^{\mathbf{0}} \boldsymbol{y} = \begin{bmatrix} y_1 \\ y_2 \\ \vdots \\ y_T \end{bmatrix} - \begin{bmatrix} 1 \\ 1 \\ \vdots \\ 1 \end{bmatrix} \bar{y} = \begin{bmatrix} y_1 - \bar{y} \\ y_2 - \bar{y} \\ \vdots \\ y_T - \bar{y} \end{bmatrix}$$

Interpretation of M<sup>0</sup>: De-meaning matrix.

(2) "Projection matrix"

 $\mathbf{P} = \mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \qquad (T\mathbf{x}T \text{ matrix})$   $\mathbf{P}\mathbf{y} = \mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y} = \mathbf{X}\mathbf{b} = \mathbf{\hat{y}} \qquad (\text{fitted values})$   $\mathbf{P}\mathbf{y} = \text{Projection of } \mathbf{y} \text{ into the } column \ space \ (\text{dimension } k) \text{ of } \mathbf{X}.$   $\mathbf{P}\mathbf{X} = (\mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}') \ \mathbf{X} = \mathbf{X}$   $\mathbf{P}\mathbf{X} = \text{Projection of } \mathbf{X} \text{ into } \mathbf{X} = \mathbf{X}.$   $\mathbf{P}\mathbf{M} = \mathbf{M}\mathbf{P} = \mathbf{0}$   $\text{Note: } \mathbf{M} = \mathbf{I}_{T} - \mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' = \mathbf{I}_{T} - \mathbf{P}$ 

- P is symmetric	$-\mathbf{P}=\mathbf{P}'$	
- P is idempotent	$-\mathbf{P}^*\mathbf{P}=\mathbf{P}$	
- P is singular	$-\mathbf{P}^{-1}$ does not exist.	$\Rightarrow$ rank( <b>P</b> ) = k

# **Results when X Contains a Constant Term**

Let the first column of X be a column of ones. That is  $\mathbf{X} = [\textbf{i}, \textbf{x}_2, \, ..., \, \textbf{x}_K]$ 

• Recall  $\mathbf{i}' \mathbf{z} = \sum_{i=1}^{T} z_i$ , where  $\mathbf{z}$  and  $\mathbf{i}$  are  $T \ge 1$ .

Then,

(1) Residuals sum to zero.

Since **X**' e = 0 $= \begin{bmatrix} 1 & 1 & \cdots & 1 \\ x_{21} & x_{22} & \cdots & x_{2T} \\ \vdots & \vdots & \ddots & \vdots \\ x_{k1} & x_{k2} & \cdots & x_{kT} \end{bmatrix} * \begin{bmatrix} e_1 \\ e_2 \\ \vdots \\ e_T \end{bmatrix} = 0 \qquad \Rightarrow \sum_i^T e_i = 0$ 

 $\Rightarrow$  **x**<sub>1</sub>'  $\boldsymbol{e} = \sum_{i=1}^{T} \boldsymbol{e}_{i} = 0$  –the residuals sum to zero.

#### (2) Regression line passes through the means

Recall we can write y = fitted value + residual y = Xb + e

Pre-multiply by 
$$\mathbf{i}'$$
:  $\mathbf{i}' \mathbf{y} = \mathbf{i}' \mathbf{X} \mathbf{b} + \mathbf{i}' \mathbf{e}$   

$$\Rightarrow \sum_{i}^{T} y_{i} = \sum_{i}^{T} \{b_{1} \ 1 + b_{2} \ x_{2i} + \dots + b_{k} \ x_{ki} \} + \sum_{i}^{T} e_{i}$$

$$\Rightarrow \sum_{i}^{T} y_{i} = b_{1} \ \sum_{i}^{T} \ 1 + b_{2} \ \sum_{i}^{T} x_{2i} + \dots + b_{k} \ \sum_{i}^{T} x_{ki}$$

$$\Rightarrow \sum_{i}^{T} y_{i} = b_{1} \ T + b_{2} \ \sum_{i}^{T} \ x_{2i} + \dots + b_{k} \ \sum_{i}^{T} \ x_{ki}$$

Dividing both sides by *T*:

$$\sum_{i}^{T} y_{i}/T = b_{1} + b_{2} \sum_{i}^{T} x_{2i}/T + \dots + b_{k} \sum_{i}^{T} x_{ki}/T$$
$$\bar{y} = b_{1} + b_{2} \bar{x}_{2} + \dots + b_{k} \bar{x}_{k}$$
$$\Rightarrow \bar{y} = \bar{x}' \mathbf{b}$$

That is, the regression line passes through the means.

Note: These results are only true if X contains a constant term!

#### **Goodness of Fit of the Regression**

After estimating the model (A1), we would like to judge the adequacy of the model. There are two ways to do this:

- Visual: Plots of fitted values and residuals, histograms of residuals.
- Numerical measures: R<sup>2</sup>, adjusted R<sup>2</sup>, AIC, BIC, etc.

Numerical measures. In general, they are simple and easy to compute. We call them *goodness-of-fit* measures. Most popular: R<sup>2</sup>.

Definition: Variation

In the context of a model, we consider the *variation* of a variable as the movement of the variable, usually associated with movement of another variable. Total variation = Total sum of squares  $(TSS) = \sum_{i} (y_i - \bar{y})^2$ 

We want to decompose TSS in two parts: one explained by the regression and one unexplained by the regression.

TSS = 
$$\sum_{i} (y_{i} - \bar{y})^{2} = \sum_{i} (y_{i} - \hat{y}_{i} + \hat{y}_{i} - \bar{y})^{2}$$
  
=  $\sum_{i} (y_{i} - \hat{y}_{i})^{2} + \sum_{i} (\hat{y}_{i} - \bar{y})^{2} + 2 \sum_{i} (y_{i} - \hat{y}_{i}) (\hat{y}_{i} - \bar{y})^{2}$   
=  $\sum_{i} e_{i}^{2} + \sum_{i} (\hat{y}_{i} - \bar{y})^{2}$ 

Since

$$\sum_{i}(y_i - \hat{y}_i)(\hat{y}_i - \bar{y}) = \sum_{i}e_i(\hat{y}_i - \bar{y}) = 0$$

 $Or mtext{TSS} = RSS + SSR$ 

RSS: Residual Sum of Squares (also called SSE: SS of errors) SSR: Regression Sum of Squares (also called ESS: *explained* SS)

#### **Goodness of Fit of the Regression – Linear Algebra**

Recall that we can use the de-meaning matrix  $\mathbf{M}^{0}$  to write  $\mathbf{y} - \mathbf{i} \ \overline{\mathbf{y}} = \mathbf{M}^{0} \mathbf{y} \ (T \ge 1 \text{ vector})$  where  $\mathbf{M}^{0} = \mathbf{I} - \mathbf{i} (\mathbf{i}' \ \mathbf{i})^{-1} \mathbf{i}'$ 

Using linear algebra we also get the decomposition of TSS. Now,  $TSS = \sum_{i} (y_i - \bar{y})^2 = y' \mathbf{M}^0 \mathbf{y} = y' \mathbf{M}^0 \mathbf{M}^0 \mathbf{y} = y' \mathbf{M}^0 \mathbf{y}.$ 

We want to decompose the total variation of y (assume  $X_1 = i$  –a constant.)

y = Xb + e,

then,

$$\begin{split} \mathbf{M}^0 \ \mathbf{y} &= \mathbf{M}^0 \mathbf{X} \mathbf{b} + \mathbf{M}^0 \mathbf{e} = \mathbf{M}^0 \mathbf{X} \mathbf{b} + \mathbf{e} \\ \mathbf{y}' \mathbf{M}^0 \mathbf{y} &= \mathbf{b}' (\mathbf{X}' \ \mathbf{M}^0) (\mathbf{M}^0 \mathbf{X}) \mathbf{b} + \mathbf{e}' \mathbf{e} \\ &= \mathbf{b}' \mathbf{X}' \ \mathbf{M}^0 \mathbf{X} \mathbf{b} + \mathbf{e}' \mathbf{e} . \\ \mathbf{TSS} &= \mathbf{SSR} + \mathbf{RSS} \end{split}$$
 (deviations from means) ( $\mathbf{M}^0$  is idempotent &  $\mathbf{e}' \ \mathbf{M}^0 \mathbf{X} = \mathbf{0}$ )

#### A Goodness of Fit Measure: R-squared

We want to have a measure that describes the fit of a regression. Simplest measure: the standard error of the regression (SER)

SER =  $\sqrt{\frac{\text{RSS}}{\text{T}-k}}$   $\implies$  SER depends on units. Not good!

• R-squared (R<sup>2</sup>)

1 = SSR/TSS + RSS/TSS R<sup>2</sup> = SSR/TSS = Regression variation/Total variation R<sup>2</sup> = b'X'M<sup>0</sup>Xb/y'M<sup>0</sup>y = 1 - e' e/y'M<sup>0</sup>y =  $(\hat{y} - \hat{i} \, \bar{y})' (\hat{y} - \hat{i} \, \bar{y})/(y - \hat{i} \bar{y})' (y - \hat{i} \bar{y}) = [\hat{y}'\hat{y} - T \, \bar{y}^2]/[y'y - T \bar{y}^2]$ 

As introduced here,  $R^2$  lies between 0 and 1 (& it is independent of units of measurement!). It measures how much of total variation (TSS) is explained by regression (SSR): the higher  $R^2$ , the better.

<u>Note</u>:  $\mathbb{R}^2$  is bounded by zero and one only if:

- (a) There is a constant term in X –we need e'  $M^0X=0!$
- (b) The line is computed by linear least squares.

• Main problem with R<sup>2</sup>: Adding regressors

Given the above interpretation of  $\mathbb{R}^2$ , it seems an appropriate criteria to select a model: If we have several models, the model with the higher  $\mathbb{R}^2$  should be selected. However,  $\mathbb{R}^2$  favors the addition of "*irrelevant*" explanatory variables.

It can be shown that  $R^2$  never falls when regressors (say z) are added to the regression. This occurs because RSS decreases with more information.

<u>Problem</u>: Judging a model based on  $R^2$  tends to *over-fitting* –i.e., in our linear model, including too many explanatory variables.

• Comparing Regressions

When  $R^2$  is used as a criteria for model selection, make sure the denominator in  $R^2$  is the same i.e., same left hand side variable. For example,  $R^2$  will not be an appropriate criteria to select between a linear vs. loglinear specifications of the dependent variable,  $y_i$ . Loglinear will almost always appear to fit better because taking logs reduces variation.

• Linear Transformation of data does not change R<sup>2</sup>.

- Based on X,  $\mathbf{b} = (\mathbf{X'X})^{-1}\mathbf{X'y}$ . Suppose we work with  $\mathbf{X^*} = c\mathbf{X}$ , instead (c is a constant).  $\mathbf{P^* y} = \mathbf{X^* b^*} = c\mathbf{X} (c\mathbf{X' cX})^{-1}c\mathbf{X'y}$   $= c\mathbf{X} (c^2 \mathbf{X'X})^{-1}c\mathbf{X' y}$   $= \mathbf{X} (\mathbf{X'X})^{-1}\mathbf{X'y} = \mathbf{Py}$  $\Rightarrow$  same fit, same residuals, same  $\mathbf{R}^2$ !

# **Adjusted R-squared**

To avoid over-fitting,  $R^2$  is modified with a penalty for number of parameters: Adjusted- $R^2$ 

$$\overline{R}^2 = 1 - \frac{(T-1)}{(T-k)} (1 - R^2) = 1 - \frac{(T-1)}{(T-k)} \frac{\text{RSS}}{\text{TSS}} = 1 - \frac{s^2}{\text{TSS}/(T-1)}$$
  

$$\Rightarrow \text{maximizing } \overline{R}^2 \iff \text{minimizing } [\text{RSS}/(T-k)] = s^2$$

There is a trade-off in  $s^2$ : higher k decreases the numerator, RSS, but it also decreases the denominator, (T - k), the *degrees of freedom*.

 $\overline{R}^2$  includes a penalty for variables that do not add much fit. Can fall when a variable is added to the equation.

<u>Technical note</u>:  $\overline{R}^2$  will rise when a variable, say  $\mathbf{z}$ , is added to the regression if and only if the tratio on  $\mathbf{z}$  is larger than one in absolute value.

Theil (1957) shows that, under certain assumptions (an important one: the true model is being considered), if we consider two linear models:

and choose the model with smaller  $s^2$  (or, larger Adjusted R<sup>2</sup>), we will select the true model, M<sub>1</sub>, on average.

In this sense, we say that "maximizing Adjusted R<sup>2</sup>" is an *unbiased* model-selection criterion.

#### **Other Goodness of Fit Measures**

There are other goodness-of-fit measures that also incorporate penalties for number of parameters (degrees of freedom). We minimize these measures.

Information Criteria

- Amemiya:  $[\mathbf{e'e}/(T-k)] * (1 + k/T) = s^2 * (1 + k/T)$ 

- Akaike Information Criterion (AIC)  $AIC = -2/T(\ln L - k)$  L: Likelihood  $\Rightarrow$  if normality AIC =  $\ln(\mathbf{e}^{*}\mathbf{e}/T) + (2/T) k$  (+constants)
- Bayes-Schwarz Information Criterion (BIC) BIC =  $-(2/T \ln L - [\ln(T)/T] k)$  $\Rightarrow$  if normality AIC =  $\ln(\mathbf{e}^{2}\mathbf{e}/T) + [\ln(T)/T] k$  (+constants)

**Example**: 3 Factor F-F Model (continuation) for IBM returns:

$b \le solve(t(x)\%*\% x)\%*\% t(x)\%*\%y$	$\# \mathbf{b} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \mathbf{y} \text{ (OLS regression)}$
e <- y - x%*%b	# regression residuals, e
$k \leq -ncol(x)$	# Number of parameters estimated
RSS <- as.numeric( $t(e)$ %*%e)	# RSS
$R2 \le 1 - as.numeric(RSS)/as.numeric(t(y)\%*\%y)$	# R-squared
$Adj_R2 <-1 - (T-1)/(T-k)*(1-R2)$	# Adjusted R-squared
$AIC \le log(RSS/T) + 2*k/T$	# AIC under N(.,.) -i.e., under (A5)

> R2 [1] 0.338985  $\Rightarrow$  The 3 factors explain 34% of the variability of IBM returns. > Adj\_R2 [1] 0.3354752 > AIC [1] -5.671036.

<u>R Note</u>: We can extract R<sup>2</sup> and Adjusted R<sup>2</sup> from the lm function: r2\_fit\_ff3r <- summary(fit\_ibm\_ff3)\$r.squared adjr2\_fit\_ff3r <- summary(fit\_ibm\_ff3)\$adj.r.squared. ¶

#### **Maximum Likelihood Estimation**

<u>Idea</u>: Assume a particular distribution with unknown parameters. Maximum likelihood (ML) estimation chooses the set of parameters that maximize the likelihood of drawing a particular sample.

**Example:** Suppose we have a sample with N realizations of a coin flip. The coin was flipped N=100 times: 60 heads (H) & 40 tails (T).

We know the distribution of the RV X = number of H & T from N coin flips. It follows a binomial distribution, with parameter p, the probability of a head. Then the probability of having x heads in N trials is given by:

$$P[X = x, N|p] = {\binom{N}{x}}p^{x}(1-p)^{N-x}$$

MLE estimates p as the probability that maximizes what we observed in our particular sample. In our case, our intuition suggests that the MLE is p = 0.60.

To check our	intuition that $p = 0.60$ , we compute $P[X = 60, N = 100]$ for different $p$ :
p = 0.50	$P[X = 60, N = 100 p] = {100 \choose 60} \cdot 50^{60} (\cdot 50)^{40} = 0.010844$
p = 0.55	$P[X = 60, N = 100 p] = {\binom{100}{60}} \cdot 55^{60} (\cdot 45)^{40} = 0.048803$
p = 0.60	$P[X = 60, N = 100   p] = \binom{100}{60} \cdot 60^{60} (\cdot 40)^{40} = 0.081219$
p = 0.65	$P[X = 60, N = 100 p] = {\binom{100}{60}} \cdot 65^{60} (\cdot 35)^{40} = 0.047392$
p = 0.70	$P[X = 60, N = 100 p] = \binom{100}{60} \cdot 70^{60} (\cdot 30)^{40} = 0.008491$

It checks! But, in general, it will be easier to find the *p* that maximizes P[X = x, N|p] using calculus. That is, taking the 1<sup>st</sup> derivative of P[X = x, N|p] with respect to *p*, set the first order condition and solve for *p* (we get  $\hat{p}_{MLE} = x/N$ ).

Formally speaking, we form a function that describes the likelihood of observing the sample results. In the previous example, for X = x:

$$L(X = x, N|p) = {\binom{N}{k}} p^{x} (1-p)^{N-x}$$

Then, we maximize L(X = x, N|p) with respect to p.

More general, let's consider a sample  $(X_1, X_2, ..., X_N)$  which is drawn from a pdf  $f(X|\theta)$ , where  $\theta$  are k unknown parameters. If the  $X_i$ 's are independent with pdf  $f(X_i|\theta)$ , the joint probability for the whole sample is just:

$$L(\mathbf{X}|\theta) = f(X_1, X_2, \dots, X_N|\theta) = f(X_1|\theta) * f(X_2|\theta) * \dots * f(X_N|\theta)$$
$$= \prod_{i=1}^N f(X_i|\theta)$$

The function  $L(X|\theta)$  –also written as  $L(X;\theta)$ – is called the *likelihood function*. This function can be maximized with respect to  $\theta$  to produce maximum likelihood estimates:  $\hat{\theta}_{MLE}$ .

It is often convenient to work with the *Log of the likelihood* function. That is,  $\ln L(X|\theta) = \sum_{i=1}^{N} \ln f(X_i|\theta)$  Then, we maximize as usual:

1<sup>st</sup>-derivative 
$$\Rightarrow \frac{\partial \ln L(X|\theta)}{\partial \theta} = \sum_{i=1}^{N} \frac{\partial \ln f(X_i|\theta)}{\partial \theta} = \sum_{i=1}^{N} \frac{f'(X_i|\theta)}{f(X_i|\theta)}$$
  
f.o.c.  $\Rightarrow \frac{\partial \ln L(X|\hat{\theta}_{MLE})}{\partial \theta} = 0$ 

Usually, the first order conditions are solved using numerical optimization.

**Example:** Let the sample be  $X = \{5, 6, 7, 8, 9, 10\}$  drawn from a Normal( $\mu$ , 1). The probability of each of these points based on the unknown mean,  $\mu$ , can be written as:

$$f(5|\mu) = \frac{1}{\sqrt{2\pi}} \exp\left[-\frac{(5-\mu)^2}{2}\right]$$
$$f(6|\mu) = \frac{1}{\sqrt{2\pi}} \exp\left[-\frac{(6-\mu)^2}{2}\right]$$
$$\vdots$$
$$f(10|\mu) = \frac{1}{\sqrt{2\pi}} \exp\left[-\frac{(10-\mu)^2}{2}\right]$$

Assume that the sample is independent. Then, the joint pdf function can be written as:

$$L(X|\mu) = f(5|\mu) * f(6|\mu) * \dots * f(10|\mu)$$
  
=  $\frac{1}{(2\pi)^{6/2}} exp\left[-\frac{(5-\mu)^2}{2} - \frac{(6-\mu)^2}{2} - \dots - \frac{(10-\mu)^2}{2}\right]$ 

The value of m that maximizes the likelihood function of the sample can then be defined by  $max L(X|\mu)$ .

It easier, however, to maximize the Log likelihood,  $\ln L(X|\mu)$ . That is,

$$\max_{\mu} \ln(L(X|\mu)) = -\frac{6}{2} \ln(2\pi) + \left[-\frac{(5-\mu)^2}{2} - \frac{(6-\mu)^2}{2} - \dots - \frac{(10-\mu)^2}{2}\right]$$

1<sup>st</sup>-derivative 
$$\Rightarrow \frac{\partial}{\partial \mu} \left[ K - \frac{(5-\mu)^2}{2} - \frac{(6-\mu)^2}{2} - \dots - \frac{(10-\mu)^2}{2} \right]$$

f.o.c. 
$$\Rightarrow (5 - \hat{\mu}_{MLE}) + (6 - \hat{\mu}_{MLE}) + \dots + (10 - \hat{\mu}_{MLE}) = 0$$

Solving for  $\hat{\mu}_{MLE}$ :

$$\hat{\mu}_{MLE} = \frac{5+6+7+8+9+10}{6} = 7.5 = \bar{x}$$

<u>Note</u>: The MLE estimator  $\hat{\mu}_{MLE}$  is equal to the sample mean. This is good for the sample mean: MLE has very good properties! ¶

The ML estimation approach is very general. In the context of the CLM, we need a model (A1) and a pdf for the errors, for example, normality, to apply ML. Now, if the model is not correctly specified, the estimates are sensitive to misspecification.

A lot of applications in finance and economics: Time series, volatility (GARCH and stochastic volatility) models, factor models of the term structure, switching models, option pricing, logistic models (mergers and acquisitions, default, etc.), trading models, etc.

In general, we rely on numerical optimization to get MLEs.

# **Maximum Likelihood Estimation: Properties**

ML estimators (MLE) have very appealing properties:

(1) *Efficiency*. Under general conditions, they achieve lowest possible variance for an estimator.

(2) *Consistency*. As the sample size increases, the MLE converges to the population parameter it is estimating:

$$\hat{\theta}_{MLE} \xrightarrow{p} \theta$$

(3) *Asymptotic Normality:* As the sample size increases, the distribution of the MLE converges to the normal distribution.

$$\hat{\theta}_{MLE} \xrightarrow{a} N(\theta, [n\mathbf{I}(\theta)]^{-1})$$

where  $I(\theta)$  is the information matrix:

$$E\left[\left(\frac{\partial \log L}{\partial \theta}\right)\left(\frac{\partial \log L}{\partial \theta}\right)^{\mathrm{T}}\right] = I(\theta) \ (kxk \text{ matrix})$$
  
$$\Rightarrow \qquad \mathrm{SE}[\hat{\theta}_{MLE,k}|\mathbf{X}] = \mathrm{sqrt}\{\mathrm{diag}([\mathbf{I}(\theta|X)^{-1}]_{kk})\}$$

(4) *Invariance*. The ML estimate is invariant under functional transformations. That is, if  $\hat{\theta}_{MLE}$  is the MLE of  $\theta$  and if  $g(\theta)$  is a function of  $\theta$ , then  $g(\hat{\theta}_{MLE})$  is the MLE of  $g(\theta)$ .

(5) Sufficiency. If a single sufficient statistic exists for  $\theta$ , the MLE of  $\theta$  must be a function of it. That is,  $\hat{\theta}_{MLE}$  depends on the sample observations only through the value of a sufficient statistic.

# **Maximum Likelihood Estimation: Numerical Optimization**

We have a function  $f(X|\theta) = \ln L(X|\theta)$ , with *k* unknown parameters. We use numerical optimization to estimate  $\theta$ . Numerical optimization are methods that search over the parameter space of  $\theta$  looking for the values that optimize –i.e., maximize or minimize– the function  $f(X|\theta)$ .

In R, the functions *optim* & *nlm* do numerical optimization. Both **minimize** any non-linear function  $f(X|\theta)$ . Recall that max  $f(X|\theta) = \min - f(X|\theta)$ . Then, in practice, we numerically minimize the negative of the likelihood function, or  $\ln L(X|\theta) * (-1)$ .

**Example**: In Example I above, we numerically minimize  $\ln L(X|\mu) * (-1)$ .

Most common optimization algorithms are based on the **Newton-Raphson method** (N-R). It is an iterative algorithm:

- At iteration j + 1, based on information from the previous iteration j, N-R update the estimate of  $\boldsymbol{\theta}$ .

- N-R stops when the values of  $\boldsymbol{\theta}$  at *j* is similar to the value at j - 1.

In our ML case, at iteration j + 1, N-R computes  $\theta_{j+1}$  (or *updates*  $\theta_j$ ) based on  $\theta_j$  plus an update. The update is based on the first and second derivatives of  $\ln L(X|\theta)$ .

• NR's i + 1 iteration:

 $\boldsymbol{\theta}_{j+1} = \boldsymbol{\theta}_j - \mathbf{A}_j^{-1} * \frac{\partial lnL}{\partial \theta} |_j \qquad (= \boldsymbol{\theta}_j + \text{update})$ 

 $\frac{\partial \ln L}{\partial \theta}|_{j} = (k \ge 1) \text{ Vector of 1st derivatives of } \ln L(X|\theta), \text{ evaluated at iteration } j, \text{ with parameter } \theta_{j}.$  $\mathbf{A}_{j} = (k \ge k) \text{ Matrix of 2nd derivatives of } \ln L, \text{ evaluated at } \theta_{j}.$ 

<u>Note</u>: At iteration j = 1, we input initial values for  $\theta_{j=0}$ , called  $\theta_0$ , which we use to compute  $A_0$  and  $\frac{\partial lnL}{\partial \theta}|_0$ .

The vector of first derivatives of ln L is called the Score. The matrix of second derivatives is called the Hessian.

- To run *optim* or *nlm*, we need to specify:
  - Initial values for the parameters,  $\theta_0$ .
  - Function to be minimized (in Example I,  $\ln L(X|\mu) * (-1)$ ).
  - Data used.
  - Other optional inputs: Choice of method, hessian calculated, etc.
- More on this topic in Lecture 10.

Maximum Likelihood Estimation: Estimating  $\mu \& \sigma^2$  from a Normal Sample

Now, we generalize the previous example to an *i.i.d.* sample  $X = \{X_1, X_2, ..., X_T\}$  drawn from a Normal( $\mu$ ,  $\sigma^2$ ). Then, the joint pdf function is:

$$L(X|\mu) = \frac{1}{(2\pi\sigma^2)^{-T/2}} \exp\left[-\frac{(x_1-\mu)^2}{2\sigma^2} - \frac{(x_2-\mu)^2}{2\sigma^2} - \dots - \frac{(x_T-\mu)^2}{2\sigma^2}\right]$$

Then, taking logs, we have:

$$\ln L = -\frac{T}{2}\ln(2\pi\sigma^2) - \frac{1}{2\sigma^2}\sum_{i=1}^{T} (X_i - \mu)^2 = -\frac{T}{2}\ln 2\pi - \frac{T}{2}\ln \sigma^2 - \frac{1}{2\sigma^2}(X - \mu)'(X - \mu)$$

Then, taking logs, we have:

$$\ln L = -\frac{T}{2}\ln(2\pi\sigma^2) - \frac{\sum_{i=1}^{T}(x_i - \mu)^2}{2\sigma^2} = -\frac{T}{2}\ln 2\pi - \frac{T}{2}\ln\sigma^2 - \frac{(X - \mu)'(X - \mu)}{2\sigma^2}$$

Taking first derivatives:

$$\frac{\partial \ln L}{\partial \mu} = -\frac{\sum_{i=1}^{T} 2(x_i - \mu)(-1)}{2\sigma^2} = \frac{\sum_{i=1}^{T} (x_i - \mu)}{\sigma^2}$$
$$\frac{\partial \ln L}{\partial \sigma^2} = -\frac{T}{2\sigma^2} + \frac{\sum_{i=1}^{T} (x_i - \mu)^2}{2\sigma^4}$$

We can write the first derivatives as a vector, the *gradient*, whose length is the number of unknown parameters in the likelihood –i.e., size of  $\theta$ . In this case, a 2x2 vector:

$$\frac{\partial \ln L}{\partial \theta} = \begin{bmatrix} \frac{\partial \ln L}{\partial \mu} \\ \frac{\partial \ln L}{\partial \sigma^2} \end{bmatrix} = \begin{bmatrix} \frac{\sum_{i=1}^{T} (x_i - \mu)}{\sigma^2} \\ -\frac{T}{2\sigma^2} + \frac{\sum_{i=1}^{T} (x_i - \mu)^2}{2\sigma^4} \end{bmatrix}$$

In the case of a log likelihood function, the vector of first derivatives is called the Score.

When we set the Score equal to **0**, we have the set of first order conditions (f.o.c.).Then, we have the f.o.c. and jointly solve for the ML estimators:

(1) 
$$\frac{\partial \ln L}{\partial \mu} = \frac{1}{\widehat{\sigma}_{MLE}^2} \sum_{i=1}^T (X_i - \hat{\mu}_{MLE}) = 0 \quad \Rightarrow \hat{\mu}_{MLE} = \frac{1}{T} \sum_{i=1}^T X_i = \bar{X}$$

<u>Note</u>: The MLE of  $\mu$  is the sample mean. Therefore, it is unbiased.

(2) 
$$\frac{\partial \ln L}{\partial \sigma^2} = -\frac{T}{2\hat{\sigma}_{MLE}^2} + \frac{1}{2\hat{\sigma}_{MLE}^4} \sum_{i=1}^T (X_i - \hat{\mu}_{MLE})^2 = 0$$
$$\Rightarrow \hat{\sigma}_{MLE}^2 = \frac{1}{T} \sum_{i=1}^T (X_i - \bar{X})^2$$

<u>Note</u>: The MLE of  $\sigma^2$  is not  $s^2$ . Therefore, it is biased! But, it is consistent.

**Example:** Using  $X = \{5, 6, 7, 8, 9, 10\}$ , now drawn from a Normal( $\mu, \sigma^2$ ).

$$\hat{\mu}_{MLE} = X = 7.5$$

$$\hat{\sigma}_{MLE}^2 = \frac{\sum_{i=1}^6 (x_i - 7.5)^2}{6} = \frac{17.5}{6} = 2.916667$$

$$\hat{\sigma}_{MLE} = \text{sqrt}(2.916667) = 1.707825$$
Note 1:  $s^2 = \frac{17.5}{(6-1)} = 3.5$ 

<u>Note 2</u>: The computation of MLE for the mean parameter  $\hat{\mu}_{MLE}$  is independent of the computation of the MLE for the variance  $\hat{\sigma}_{MLE}^2$ .

• To obtain the variance of  $\hat{\theta}_{MLE} = [\hat{\mu}_{MLE}, \hat{\sigma}_{MLE}^2]$  we invert the information matrix for the whole sample  $\mathbf{I}(\theta|X)$ . Recall,

$$\widehat{\theta}_{MLE} \xrightarrow{a} N(\theta, \mathbf{I}(\theta|X)^{-1})$$

where  $I(\theta|X)$  is the *Information matrix* for the whole sample. It is generally calculated as:

 $E\left[-\left(\frac{\partial^2 \ln L(\theta|X)}{\partial \theta \partial \theta'}\right)\right] = I(\theta|X), \qquad (kxk \text{ matrix})$ where the matrix of second derivatives is the Hessian matrix, **H**:  $\frac{\partial^2 \ln L(\theta|X)}{\partial \theta \partial \theta'} = \mathbf{H} \qquad (kxk \text{ matrix})$ 

The inverse of the Hessian gives the variance of the MLE estimator:  $\operatorname{Var}(\hat{\theta}_{MLE}) = E[-\mathbf{H}]^{-1} = I(\theta)^{-1}$  (kxk matrix)

In practice, we use numerical optimization packages (say, *nlm* in R), which minimize a function. Thus, we *minimize* the *negative* log  $L(\theta|X)$  and, thus, to get  $Var[\hat{\theta}_{MLE}]$  we do not need to multiply **H** by (-1).

Then, MLE standard error of parameter k is given by:  $SE[\hat{\theta}_{MLE,k}|\mathbf{X}] = sqrt\{diag([\mathbf{H}^{-1}]_{kk})\}$ 

**Example:** For  $X = \{5, 6, 7, 8, 9, 10\} \sim N(\mu, \sigma^2)$ , code to get MLEs. mu <- 0 # assumed mean (initial value) sig <- 1 # assumed sd (initial value)  $x 6 \le c(5, 6, 7, 8, 9, 10)$ # Step 1 - Create Likelihood function likelihood  $lf \leq function(x)$ # Create a prob function with mu & sig as arguments mu < x[1]sig < x[2]sum(log(dnorm(x 6, mu, sd=sig))) } negative likelihood  $lf \leq function(x)$  # R uses a minimization algorithm, change sign mu < x[1]sig <- x[2]sum(log(dnorm(x 6, mu, sd=sig))) \* (-1)} negative likelihood lf(x)

```
# Step 2 - Maximize Log Likelihood function (or Minimize negative Likelihood function)
results lf <- nlm(negative likelihood lf, x, stepmax=4)
                                                              # nlm minimizes the function
> results lf
                                       # displays nlm results
$minimum
[1] 11.72496
                                       <= Minimized value of function
$estimate
                                               \leq MLEs for \mu \& \sigma^2 (= \hat{\mu}_{MLF} \& \hat{\sigma}_{MLF}^2)
[1] 7.500000 1.707825
$gradient
[1] -1.846772e-07 -7.986103e-08 \leq \approx 0 if we're at a minimum
$code
                                       <= = 1 if we program stopped at a minimum
[1] 1
```

\$iterations [1] 34	<= Number of iterations
par_max <- results_lf\$estimate > par_max	# Extract estimates # Should be equal to sample mean
[1] 7.500000 1.707825 > likelihood_lf(par_max) [1] -11.72496	# Check max value of likelihood function

# Step 3 - Standard Errors (by inverting the Hessian)
results lf <- nlm(negative likelihood lf, x, stepmax=4, hessian=TRUE)</pre>

par_hess <- results_lf\$hessian > par_hess	# Extract Hessian # Show Hessian
[,1] [,2]	
[1,] 2.0571428731 -0.0009030531	
[2,] -0.0009030531 4.1122292411	
<pre>cov_lf &lt;- solve(coeff_hess) &gt; cov_lf     [,1] [,2]</pre>	<ul><li># invert Hessian to get cov(MLEs)</li><li># Show covariance matrix</li></ul>
[1,] <b>0.4861111542</b> 0.0001067509 [2,] 0.0001067509 <b>0.2431771280</b>	
se_lf <- sqrt(diag(cov_lf)) >se_lf [1] <b>0.6972167 0.4931299</b>	# Compute standard errors of MLEs
# t-tests	
> par_max[1]/se_lf[1]	# t-ratio for mu
$\begin{array}{c} [1] 10.75706\\ \text{par_max}[2]/\text{se_lf}[2]\\ [1] 3.463236. \ \P \end{array}$	# t-ratio for sigma2

# Maximum Likelihood Estimation: Linear Model Example

We will work the previous example with matrix notation. Suppose we assume:

or  $y_i = \mathbf{x}_i' \mathbf{\beta}^{T} + \varepsilon_i, \qquad \varepsilon_i \sim N(0, \sigma^2)$  $\mathbf{y} = \mathbf{X}\mathbf{\beta} + \varepsilon, \qquad \mathbf{\varepsilon} \sim N(0, \sigma^2 I_T)$ 

where  $x_i$  is a kx1 vector of exogenous numbers and  $\beta$  is a kx1 vector of unknown parameters. Then, the joint likelihood function becomes:

$$L = \prod_{i=1}^{T} \frac{1}{\sqrt{2\pi\sigma^2}} exp\left(-\frac{\varepsilon_i^2}{2\sigma^2}\right) = (2\pi\sigma^2)^{-T/2} \prod_{i=1}^{T} exp\left(-\frac{\varepsilon_i^2}{2\sigma^2}\right)$$

Taking logs, we have the log likelihood function:

$$\ln L = -\frac{T}{2} \ln 2 \pi \sigma^2 - \frac{1}{2\sigma^2} \sum_{i=1}^{T} \varepsilon_i^2 = -\frac{T}{2} \ln 2 \pi \sigma^2 - \frac{1}{2\sigma^2} (y - X\beta)' (y - X\beta)$$

The joint likelihood function becomes:

$$\ln L = -\frac{T}{2} \ln 2\pi\sigma^{2} - \frac{1}{2\sigma^{2}} \sum_{i=1}^{T} \varepsilon_{i}^{2} =$$
  
=  $-\frac{T}{2} \ln(2\pi) - \frac{T}{2} \ln(\sigma^{2}) - \frac{1}{2\sigma^{2}} (y - X\beta)' (y - X\beta)$ 

We take first derivatives of the log likelihood w.r.t.  $\beta$  and  $\sigma^2$ :

$$\frac{\partial \ln L}{\partial \beta} = -\frac{1}{2} \sum_{i=1}^{T} 2\varepsilon_i x_i' / \sigma^2 = -\frac{1}{\sigma^2} X' \varepsilon$$
$$\frac{\partial \ln L}{\partial \sigma^2} = -\frac{T}{2\sigma^2} - (-\frac{1}{2\sigma^4}) \sum_{i=1}^{T} \varepsilon_i^2 = (\frac{1}{2\sigma^2}) [\frac{\varepsilon' \varepsilon}{\sigma^2} - T]$$

Using the f.o.c., we jointly estimate  $\beta$  and  $\sigma^2$ :

$$\frac{\partial \ln L}{\partial \beta} = -\frac{1}{\sigma^2} \mathbf{X}' \mathbf{\varepsilon} = \frac{1}{\sigma^2} \mathbf{X}' (\mathbf{y} - \mathbf{X} \widehat{\boldsymbol{\beta}}_{MLE}) = 0 \quad \Rightarrow \widehat{\boldsymbol{\beta}}_{MLE} = (\mathbf{X}' \mathbf{X})^{-1} \mathbf{X}' \mathbf{y}$$
$$\frac{\partial \ln L}{\partial \sigma^2} = (\frac{1}{2\widehat{\sigma}_{MLE}^2}) [\frac{\mathbf{e}' \mathbf{e}}{\widehat{\sigma}_{MLE}^2} - T] = 0 \quad \Rightarrow \widehat{\sigma}_{MLE}^2 = \frac{\mathbf{e}' \mathbf{e}}{T} = \sum_{i=1}^T \frac{(y_i - \mathbf{X}_i \widehat{\boldsymbol{\beta}}_{MLE})^2}{T}$$

Under (A5) –i.e., normality for the errors–, we have that  $\hat{\beta}_{MLE} = b$ . This is a good result for OLS **b**. ML estimators have very good properties: Efficiency, consistency, asymptotic normality and invariance.

 $\hat{\sigma}_{MLE}^2$  is biased, but given that it is an ML estimator, it is efficient, consistent and asymptotically normally distributed.

**Example:** We estimate the 3 F-F factor model for IBM.

SFX\_da <read.csv("https://www.bauer.uh.edu/rsusmel/4397/Stocks\_FX\_1973.csv",head=TRUE,sep=",") x\_ibm <- SFX\_da\$IBM x\_Mkt\_RF<- SFX\_da\$Mkt\_RF x\_SMB <- SFX\_da\$MB x\_HML <- SFX\_da\$SMB x\_HML <- SFX\_da\$HML x\_RF <- SFX\_da\$RF T <- length(x\_ibm) lr\_ibm <- log(x\_pfe[-1]/x\_pfe[-T]) x0 <- matrix(1,T-1,1) Mkt\_RF <- x\_Mkt\_RF[-1]/100

```
SMB <- x SMB[-1]/100
HML \le x HML[-1]/100
RF \le x RF[-1]/100
ibm x <- lr ibm - RF
X <- cbind(x0, Mkt RF, SMB, HML)
# Step 1 - Negative Likelihood function
likelihood lf \leq function(theta, y, X) 
N \leq nrow(X)
k \leq -ncol(X)
beta <- theta [1:k]
sigma2 <- theta[k+1]^2
e <- y - X%*%beta
logl <- .5*N*log(2*pi)-.5*N*log(sigma2)- ((t(e)%*%e)/(2*sigma2))
return(-logl)
}
theta \leq c(0,1,1,1,1)
                                          # initial values
likelihood lf(theta,ibm x,X)
       [,1]
[1,] -599.0825
# Step 2 - Maximize (or Minimize negative Likelihood function)
results lf <- nlm(likelihood lf, theta, hessian=TRUE, y=ibm x, X=X)
                                                                       # nlm minimizes 1 f
par max <- results lf$estimate
                                                 # Extract estimates
                                                 # Should be equal to OLS results
> par max
[1] -0.0005907974 0.8676052091 -0.6815947799 -0.2284249895 0.0557422421
> likelihood lf(par max,ibm x,X)
                                                 # Check max value of likelihood function
[,1]
[1,] -835.3316
# Compare with OLS results
fit ibm ff3 <- lm(ibm x \sim Mkt RF + SMB + HML)
> summary(fit ibm ff3)
Coefficients:
                                                            Pr(\geq |t|)
                 Estimate
                                    Std. Error
                                                  t value
(Intercept)
                 -0.0005903
                                    0.0023793
                                                  -0.248
                                                            0.80416
Mkt RF
                 0.8676042
                                    0.0542554
                                                  15.991
                                                            < 2e-16 ***
SMB
                 -0.6815950
                                    0.0804542
                                                  -8.472
                                                            < 2e-16 ***
                                                  -2.820
                                                            0.00497 **
HML
                                    0.0809992
                 -0.2284263
```

---

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

#### **Maximum Likelihood Estimation: Score and Information Matrix**

<u>Definition</u>: Score (or efficient score)

$$S(X;\theta) = \frac{\delta \log(L(X|\theta))}{\delta \theta} = \sum_{i=1}^{n} \frac{\delta \log(f(x_i|\theta))}{\delta \theta}$$

 $S(X; \theta)$  is called the *score* of the sample. It is the vector of partial derivatives (the gradient), with respect to the parameter  $\theta$ . If we have k parameters, the score will have a kx1 dimension.

<u>Definition</u>: Fisher information for a single parameter for observation *i*:  $E\left[\left(\frac{\partial \log(f(x_i|\theta))}{\partial x_i}\right)^2\right] = I(\theta)$ 

$$E\left[\left(\frac{\partial \log(f(x_{l}|\theta))}{\partial \theta}\right)\right] = I(\theta)$$

 $I(\theta)$  is sometimes just called *information*. It measures the shape of the log  $f(X|\theta)$ .

The concept of information can be generalized for the k-parameter case. In this case, for the whole sample:

$$E\left[\left(\frac{\partial \log L}{\partial \theta}\right)\left(\frac{\partial \log L}{\partial \theta}\right)^{T}\right] = \boldsymbol{I}(\theta)$$

This is *kxk* matrix.

If L is twice differentiable with respect to  $\theta$ , and under certain regularity conditions, then the information may also be written as

$$E\left[\left(\frac{\partial \log L}{\partial \boldsymbol{\theta}}\right)\left(\frac{\partial \log L}{\partial \boldsymbol{\theta}}\right)^{\mathrm{T}}\right] = E\left[-\left(\frac{\delta^{2}\log(L(X \mid \boldsymbol{\theta})))}{\partial \boldsymbol{\theta} \partial \boldsymbol{\theta}'}\right)\right] = \mathbf{I}(\boldsymbol{\theta})$$

 $I(\theta)$  is called the *information matrix* (negative Hessian). It measures the shape of the likelihood function.

• The inverse of the information matrix for the whole sample is the Variance of  $\hat{\theta}_{MLE}$ . That is,  $\operatorname{Var}(\hat{\theta}_{MLE}) = I(\theta)^{-1}$ 

Sometimes, the notation for the information matrix for the whole sample is  $I(\theta|X)$ .

<u>Remark</u>: In practice, we use the inverse of the Hessian, evaluated at  $\hat{\theta}_{MLE}$ , as the estimator of the variance. R calculates the Hessian in all optimization packages (for example, *nlm* or *optim*). In the previous example, we extracted the Hessian from the *nlm* function with

**Example:** We assume:

 $\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\varepsilon}, \qquad \boldsymbol{\varepsilon} \sim N(0, \sigma^2 I_T)$ 

Taking logs, we have the log likelihood function:

$$\ln L = -\frac{T}{2} \ln 2\pi\sigma^2 - \frac{1}{2\sigma^2} \sum_{i=1}^T \varepsilon_i^2 = -\frac{T}{2} \ln 2\pi - \frac{T}{2} \ln \sigma^2 - \frac{1}{2\sigma^2} (y - X\beta)' (y - X\beta)$$

The score function is –first derivatives of log L w.r.t.  $\theta = (\beta, \sigma^2)$ :

$$\frac{\partial \ln L}{\partial \beta} = -\frac{1}{2\sigma^2} (-2 \mathbf{X}' \mathbf{y} + 2 \mathbf{X} \mathbf{X} \boldsymbol{\beta}) = \frac{1}{\sigma^2} \mathbf{X}' (\mathbf{y} - \mathbf{X} \boldsymbol{\beta})$$
$$\frac{\partial \ln L}{\partial \sigma^2} = -\frac{T}{2\sigma^2} - (-\frac{(\mathbf{y} - \mathbf{X} \boldsymbol{\beta})'(\mathbf{y} - \mathbf{X} \boldsymbol{\beta})}{2\sigma^4}) = (\frac{1}{2\sigma^2}) [\frac{\boldsymbol{\varepsilon}' \boldsymbol{\varepsilon}}{\sigma^2} - T]$$

Then, we take second derivatives to calculate  $I(\theta)$ :

$$\frac{\partial^2 \ln L}{\partial \beta \partial \beta'} = -\frac{X'X}{\sigma^2} - kxk \text{ matrix.}$$

$$\frac{\partial^2 \ln L}{\partial \beta \partial \sigma^2'} = -\frac{1}{\sigma^4} (X'y - XX\beta) = -\frac{1}{\sigma^4} (X'\varepsilon)$$

$$\frac{\partial^2 \ln L}{\partial \sigma^2 \partial \sigma^2'} = -\frac{1}{2\sigma^4} \left[\frac{\varepsilon'\varepsilon}{\sigma^2} - T\right] + \left(\frac{1}{2\sigma^2}\right) \left(-\frac{\varepsilon'\varepsilon}{\sigma^4}\right) = -\frac{1}{2\sigma^4} \left[2\frac{\varepsilon'\varepsilon}{\sigma^2} - T\right] - \text{scalar}$$

Using linear algebra notation:

$$I(\theta) = E\left[-\frac{\partial \ln L}{\partial \theta \partial \theta}\right] = \begin{bmatrix} (\frac{1}{\sigma^2} X' X) & 0\\ 0 & \frac{T}{2\sigma^4} \end{bmatrix}$$
 a  $(k+1)x(k+1)$  matrix.

To get SE for  $\hat{\theta}_{MLE}$ , we invert the (k+1)x(k+1) information matrix.

<u>Technical Note</u>: The information matrix is block-diagonal, the inverse is the inverse of the diagonal blocks. Then,

$$I(\theta|X)^{-1} = \begin{bmatrix} \sigma^2 (X'X)^{-1} & 0\\ 0 & \frac{2\sigma^4}{T} \end{bmatrix}$$

Then,

$$\operatorname{Var}[\widehat{\boldsymbol{\beta}}_{MLE}] = \widehat{\sigma}_{MLE}^2 (\boldsymbol{X}'\boldsymbol{X})^{-1}$$
$$\operatorname{Var}[\widehat{\sigma}_{MLE}^2] = 2 \ \widehat{\sigma}_{MLE}^4 / \mathrm{T.} \P$$

**Example:** We continue the previous IBM example, computing MLE SEs for linear model

# <b>Step 3</b> - Compute S.E. by inverting Hessian	
par_hess <- results_lf\$hessian	# Extract Hessian
> par_hess	# Show Hessian matrix

[,1] [,2] [,3] [,4] [,5] [1,] 183123.21311034.3403801300.5280632452.9161743-3.243494e+02[2,] 1034.3404390.199568371.3131499-55.6126338-6.913297e-01[3,] 300.528171.3131499170.5839168-26.9486009-3.023956e-01[4,] 452.9162-55.6126338-26.9486009165.2938181-2.928687e-01[5,] -324.3494-0.6913297-0.3023956-0.29286873.629895e+05

cov\_lf <- solve(par\_hess)</td># invert Hessian to get covariancese\_lf <- sqrt(diag(cov\_lf))</td># Compute standard errors (compare with OLS SE)> se\_lf[1] 0.002370939 0.054063912 0.080170161 0.080713227 0.001659791

	OLS		MLE	
	Coefficients	S.E.	Coefficients	S.E.
Intercept	-0.00509	0.00238	-0.00509	0.00237
Mkt_RF	0.86761	0.05425	0.86761	0.05406
SMB	-0.68159	0.08045	-0.68159	0.08017
HML	-0.22842	0.08100	-0.22842	0.08071

• Summary: OLS vs MLE

Conclusion: Same coefficients as expected. Very similar results for the S.E. ¶

# **Data Problems**

"If the data were perfect, collected from well-designed randomized experiments, there would hardly be room for a separate field of econometrics." Zvi Griliches (1986, Handbook of Econometrics)

Three important data problems:

- (1) Missing Data very common, especially in cross sections and long panels.
- (2) Outliers unusually high/low observations.
- (3) Multicollinearity there is perfect or high correlation in the explanatory variables.

• In general, data problems are exogenous to the researcher. We cannot change the data or collect more data.

# **Missing Data**

General Setup

We have an indicator variable,  $s_i$ . If  $s_i = 1$ , we observe  $Y_i$ , and if  $s_i = 0$  we do not observe  $Y_i$ .

Note: We always observe the missing data indicator  $s_i$ .

Suppose we are interested in the population mean  $\theta = E[Y_i]$ .

With a lot of information -large *T*-, we can learn  $p = E[s_i]$  and  $\mu_1 = E[Y_i | s_i = 1]$ , but nothing about  $\mu_0 = E[Y_i | s_i = 0]$ .

We can write:  $\theta = p * \mu_1 + (1 - p) * \mu_0$ .

<u>Problem</u>: Even in large samples we learn nothing about  $\mu_0$ . Without additional information and/or assumptions there is no much we can say about  $\theta$ .

Now, suppose the variable of interest is binary:  $Y_i \in \{0, 1\}$ . We also have an explanatory variable of  $Y_i$ , say  $W_i$ .

Then, the natural (not data-informed) lower and upper bounds for  $\mu_0$  are 0 and 1 respectively. This implies bounds on  $\theta$ :

 $\theta \in [\theta_{\text{LB}}, \theta_{\text{UB}}] = [p * \mu_1, p * \mu_1 + (1 - p) * \mu_0].$ 

These bounds are *sharp*, in the sense that without additional information we cannot improve on them.

If from variable  $W_i$  we can infer something about the missing values, these bounds can be improved.

# **Missing Data – CLM**

Now, suppose we have the CLM:  $y_i = x_i' \beta + \varepsilon_i$ 

We use the selection indicator,  $s_i$ , where  $s_i = 1$  if we can use observation *i*. After some algebra we get,

$$\mathbf{b} = \boldsymbol{\beta} + \left(\sum_{i=1}^{T} s_i \ \boldsymbol{x}_i' \boldsymbol{x}_i/T\right)^{-1} \left(\sum_{i=1}^{T} s_i \ \boldsymbol{x}_i' \boldsymbol{\varepsilon}_i/T\right)$$

• For unbiased (and consistent) results, we need  $E[s_i \mathbf{x}_i' \varepsilon_i] = 0$ , implied by  $E[\varepsilon_i | s_i \mathbf{x}_i'] = 0$  (\*)

In general, we find that when  $s_i = h(x_i)$ , that is, the selection is a function of  $x_i$ , we have an inconsistent OLS **b**. This situation is called *selection bias*.

#### **Example of Selection Bias:** Determinants of Hedging.

A researcher only observes companies that hedge. Estimating the determinants of hedging from this population will bias the results!  $\P$ 

If missing observations are randomly (exogenously) "selected," it is likely safe to ignore problem. Rubin (1976) calls this assumption "*missing completely at random*" (or MCAR).

In general, MCAR is rare. In general, it is more common to see "*missing at random*," where missing data depends on observables (say, education, sex) but one item for individual *i* is NA (Not Available).

If in the regression we "control" for the observables that influence missing data (not easy), it is OK to delete the whole observation for i.

# **Missing Data – Usual Solutions**

Otherwise, we can:

a. Fill in the blanks –i.e., *impute* values to the missing data- with averages, interpolations, or values derived from a model.

b. Use (inverse) probability weighted estimation. Here, we inflate or "over-weight" unrepresented subjects or observations.

c. Heckman selection correction. We build a model for the selection function,  $h(x_i)$ .

# **Outliers**

Many definitions: Atypical observations, extreme values, conditional unusual values, observations outside the expected relation, etc.

In general, we call an *outlier* an observation that is numerically different from the data. But, is this observation a "mistake," say a result of measurement error, or part of the (heavy-tailed) distribution?

In the case of normally distributed data, roughly 1 in 370 data points will deviate from the mean by 3\*SD. Suppose T=1,000 and we see 9 data points deviating from the mean by more than 3\*SD indicates outliers. We expect 3 data points to deviate by more than 3\*SD. Which of the 9 observations can be classified as an outlier?

<u>Problem with outliers</u>: They can affect estimates. For example, with small data sets, one big outlier can seriously affect OLS estimates.

# **Outliers: Identification**

- Informal identification method:
- Eyeball: Look at the observations away from a scatter plot.

**Example:** Plot residuals for the 3 FF factor model for IBM returns **e\_ibm** <- residuals(**fit\_ibm\_ff3**) # Extract IBM residuals from FF regression **fit\_ibm\_ff3**. plot(**e\_ibm**, type ="l", col="blue", main ="IBM Residuals from 3 FF Factor Model", xlab="Date", ylab="IBM residuals")



• Formal identifications methods:

- *Standardized residuals*, e<sub>i</sub>/SD(e<sub>i</sub>): Check for errors that are 2\*SD (or more) away from the expected value.

- *Leverage statistics*: It measures the difference of an independent data point from its mean. High leverage observations can be potential outliers. Leverage is measured by the diagonal values of the  $\mathbf{P}$  matrix:

$$h_t = 1/T + (x_t - \boldsymbol{a}_{\mathcal{X}})/[(T-1)s_x^2]$$

• Formal identifications methods:

- *Standardized residuals*, ei/SD(ei): Check for errors that are 2\*SD (or more) away from the expected value.

Example: Plot standardized residuals for IBM residuals

x\_stand\_resid <- e\_ibm /sd(e\_ibm) # standardized residuals plot(x\_stand\_resid, type ="l", col="blue", main ="IBM Standardized Residuals from 3 FF Factor Model", xlab="Date", ylab="IBM residuals")



- *Leverage statistic*: It measures the difference of an independent data point from its mean. High leverage observations can be potential outliers. Leverage is measured by the diagonal values of the  $\mathbf{P}$  matrix:

$$h_t = 1/T + (x_t - \bar{x})/[(T-1)s_x^2].$$

But, an observation can have high leverage, but no *influence*.

- *Influence statistic: Dif beta*. It measures how much an observation influences a parameter estimate, say b<sub>j</sub>. Dif beta is calculated by removing an observation, say *i*, recalculating b<sub>j</sub>, say b<sub>j</sub>(-*i*), taking the difference in betas and standardizing it. Then,

$$Dif beta_{j(-i)} = [b_j - b_j(-i)]/SE[b_j].$$

- *Influence statistic: Distance D (as in Cook's D).* It measures the effect of deleting an observation on the fitted values, say  $\hat{y}_{j}$ .

$$D_j = \sum_j \left[ \hat{y}_j - \hat{y}_j(-i) \right] / [k * MSE],$$

where k is the number of parameters in the model and MSE is mean square error of the regression model ((MSE=RSS/T)..

The identification statistics are usually compared to some ad-hoc cut-off values. For example, for Cook's D, if  $D_i > 4/T \Rightarrow$  observation *i* is considered a (potential) highly influential point.

The analysis can also be carried out for groups of observations. In this case, we would be looking for blocks of highly influential observations.

# **Outlier Identification: Leverage & Influence**



Deleting the observation in the upper right corner has a clear effect on the regression line. This observation has *leverage* and *influence*.

#### **Outliers: Summary of Rules of Thumb**

General rules of thumb (ad-hoc thresholds) used to identify outliers:

Measure	Value
abs(stand resid)	> 2
leverage	>(2k+2)/T
abs( <i>Dif Beta</i> )	> 2/sqrt(T)
Cook's D	> 4/T

In general, if we have 5% or less observations exceeding the ad-hoc thresholds, we tend to think that the data is OK.
Example: Cook's D for IBM returns using the 3 FF Factor Model y <- ibm\_x x <- cbind(x0, Mkt\_RF, SMB, HML) dat\_xy <- data.frame(y, x) fit\_ibm\_ff3 <- lm(y ~ x - 1) cooksd <- cooks.distance(fit\_ibm\_ff3) # plot cook's distance plot(cooksd, pch="\*", cex=2, main="Influential Obs by Cooks distance") # add cutoff line abline(h = 4\*mean(cooksd, na.rm=T), col="red") # add cutoff line # add labels text(x=1:length(cooksd)+1, y=cooksd, labels=ifelse(cooksd>4\*mean(cooksd, na.rm=T), names(cooksd),""), col="red") # add labels

# influential row numbers
influential <- as.numeric(names(cooksd)[(cooksd > 4\*mean(cooksd, na.rm=T))])
# print first 10 influential observations.
head(dat\_xy[influential, ], n=10L)



> # print first 10 influential observations. >head(dat\_xy[influential, ],n=10L)

V1 Mkt\_RF SMB HML 8 -0.16095068 1 0.0475 0.0294 0.0219 94 0.01266444 1 0.0959 -0.0345 -0.0835 227 -0.04237227 1 0.1084 -0.0224 -0.0403 237 -0.19083575 1 0.0102 0.0205 -0.0210 239 -0.30648638 1 0.0153 0.0164 0.0252 282 0.07787100 1 -0.0597 -0.0383 0.0445 286 0.20734626 1 0.0625 -0.0389 0.0117 291 0.15218986 1 0.0404 -0.0565 -0.0006 306 0.13928315 1 -0.0246 -0.0512 -0.0096 315 0.16196934 1 0.0433 0.0400 0.0253 Note: There are easier ways to plot Cook's D and identify the suspect outliers. The package olsrr can be used for this purpose too.

**Example:** Different tools to check for outliers for residual in the FF model for IBM returns. We will use the package *olsrr* --install it with **install.packages()**. install.packages("olsrr")

library(olsrr) e ibm <- residuals(fit ibm ff3) x stand resid <- e ibm /sd(e\_ibm) sum(x stand resid > 2)x lev <- ols leverage(fit ibm ff3) sum(x lev > (2\*k+2)/T)sum(cooksd > 4/T)ols plot resid stand(fit ibm ff3) ols plot cooksd bar(fit ibm ff3) ols plot dfbetas(fit ibm ff3)

# need to install package olsrr # standardized residuals # Rule of thumb count (5% count is OK) # leverage residuals # Rule of thumb count (5% count is OK) # Rule of thumb count (5% count is OK) # Plot standardized residuals # Plot Cook's D measure # Plot Difference in betas

> sum(x stand resid > 2) [1] 13 #5%? = 13/569 = 0.0228> sum(x lev > (2\*k+2)/T)[1] 32 #5%? = 32/569 = 0.0562> sum(cooksd > 4/T) # 5%? = 38/569 = **0.0668** [1] 38





>ols plot cooksd bar(fit ibm ff3)

# Plot Cook's D measure



Conclusion: Using standardized residuals, we get some evidence of outliers.

## **Outliers: What to Do?**

Typical solutions:

- Use a non-linear formulation or apply a transformation (log, square root, etc.) to the data.

- Remove suspected observations. (Sometimes, there are theoretical reasons to remove suspect observations. Typical procedure in finance: remove public utilities or financial firms from the analysis.)

- Winsorization of the data (cut an  $\alpha$ % of the highest and lowest observations of the sample).

- Use dummy variables.

- Use LAD (quantile) regressions, which are less sensitive to outliers.
- Weight observations by size of residuals or variance (robust estimation).

General rule: Present results with or without outliers.

### Multicollinearity

The X matrix is *singular* (perfect collinearity) or *near singular* (multicollinearity). - *Perfect collinearity* Not much we can do. OLS will not work  $\Rightarrow$  X'X cannot be inverted. The model needs to be reformulated.

#### - Multicollinearity.

OLS will work.  $\beta$  is still unbiased. The problem is in  $(X'X)^{-1}$ ; that is, in the Var[b|X]. Let's see the effect on the variance of particular coefficient,  $b_k$ .

Recall the estimated Var[ $b_k$ |**X**] is the *k*th diagonal element of  $\sigma^2$ (**X**'**X**)<sup>-1</sup>.

Let define  $\mathbb{R}^{2_{k}}$  as the  $\mathbb{R}^{2}$  in the regression of  $\mathbf{x}_{k}$  on the other regressors,  $\mathbf{X}_{(-k)}$ . Then, we can show the estimated  $\operatorname{Var}[\mathbf{b}_{k}|\mathbf{X}]$  is

$$\operatorname{Var}[\mathbf{b}_{k}|\mathbf{X}] = \frac{s^{2}}{\left[(1-R_{k}^{2})\sum_{i=1}^{n}(x_{ik}-\overline{x}_{k})^{2}\right]}.$$

 $\Rightarrow$  the higher  $R^{2_{k}}$  –i.e., the fit between  $\mathbf{x}_{k}$  and the rest of the regressors–, the higher Var[ $\mathbf{b}_{k}|\mathbf{X}$ ].

### **Multicollinearity: Signs**

Signs of Multicollinearity:

- Small changes in **X** produce wild swings in **b**.
- High R<sup>2</sup>, but **b** has low t-stats –i.e., high standard errors
- "Wrong signs" or difficult to believe magnitudes in **b**.

There is no *cure* for collinearity. Estimating something else is not helpful (transforming regressors, principal components, etc.).

There are "measures" of multicollinearity, such as the

- *K*# = *Condition number* = max(singular value)/min(singular value)
- *Variance inflation factor* =  $VIF_k = 1/(1 R^2_k)$ .

<u>Rule of thumb for Condition number</u>: If K# > 30 such matrix cannot be inverted reliably. Thus, **X** shows severe multicollinearity.

## **Multicollinearity: VIF and Condition Index**

Belsley (1991) proposes to calculate the VIF and the condition number, using Rx, the correlation matrix of the standardized regressors:

VIF<sub>k</sub> = diag(Rx<sup>-1</sup>)<sub>k</sub> Condition Index =  $\kappa_k$  = sqrt( $\lambda_1 / \lambda_k$ ) where  $\lambda_1 > \lambda_2 > ... > \lambda_p > ...$  are the ordered eigenvalues of Rx.

Belsley's (1991) rules of thumb for  $\kappa_k$ :

- below 10	$\Rightarrow$ good
- from 10 to 30	⇒ concern
- greater than 30	$\Rightarrow$ trouble
- greater than 100	$\Rightarrow$ disaster.

Another common rule of thumb: If  $VIF_k > 5$ , concern.

Best approach: Recognize the problem and understand its implications for estimation.

<u>Note</u>: Unless we are very lucky, some degree of multicollinearity will always exist in the data. The issue is: when does it become a problem?

### **Multicollinearity: Example**

**Example:** Check for multicollinearity for IBM returns 3-factor model library(olsrr) ols\_vif\_tol(fit\_ibm\_ff3) ols\_eigen\_cindex(fit\_ibm\_ff3)

>	ols_vif_tol(	fit_ibm_ff3	)
1	Variables	Tolerance	VIF
1	xMkt_RF	0.8901229	1.123440
2	xSMB	0.9147320	1.093216
3	xHML	0.9349904	1.069530

> ols eigen cindex(fit ibm ff3)

	Eigenvalue	Condition Index	x intercept	xMkt_RF	xSMB	xHML	
1	1.4506645	1.000000	0.01557614	0.24313961	0.2120	01760	0.1518949
2	1.0692689	1.164770	0.66799183	0.01432250	0.0017	89253	0.2129328
3	0.7967889	1.349310	0.16184731	0.01239755	0.5764	32492	0.4107435
4	0.6832777	1.457085	0.15458473	0.73014033	0.2097	76495	0.2244287

Conclusion: Multicollinearity does not seem to be a problem.

## Lecture 4 – Appendix A: Rules for Vector Derivatives

#### (1) Linear function

Consider the linear function:  $y = f(z) = z' \gamma + \omega$ 

where  $\boldsymbol{x}$  and  $\boldsymbol{\gamma}$  are *k*-dimensional vectors and  $\boldsymbol{\omega}$  is a constant.

We derive the gradient in matrix notation as follows:

1. Convert to summation notation:  $f(\mathbf{z}) = \sum_{i}^{k} z_i \gamma_i = z_1 \gamma_1 + z_2 \gamma_2 + \ldots + z_k \gamma_k$ 

2. Take partial derivative w.r.t. element  $x_j$ :  $\frac{\partial}{\partial z_j} \left[ \sum_{i}^{k} z_i \gamma_i \right] = \gamma_j$ 

3. Put all the partial derivatives in a vector:

$$\nabla f(\mathbf{z}) = \begin{bmatrix} \frac{\partial f(\mathbf{z})}{\partial z_1} \\ \vdots \\ \frac{\partial f(\mathbf{z})}{\partial z_k} \end{bmatrix} = \begin{bmatrix} \gamma_1 \\ \vdots \\ \gamma_k \end{bmatrix}$$

4. Convert to matrix notation:  $\nabla f(\mathbf{z}) = \mathbf{y}$ 

### (2) Quadratic form

Consider a quadratic form:  $\mathbf{q} = f(\mathbf{x}) = \mathbf{x}^* \mathbf{A} \mathbf{x}$ 

where x is kx1 vector and A is a kxk matrix, with  $a_{ji}$  elements.

Steps:

1. Convert to summation notation:

$$f(\mathbf{x}) = \mathbf{x}^{*} \begin{bmatrix} \sum_{j=1}^{k} a_{j1} x_{j} \\ \vdots \\ \sum_{j=1}^{k} a_{jk} x_{j} \end{bmatrix} = \sum_{i=1}^{k} \sum_{j=1}^{k} x_{i} a_{ji} x_{j}$$

(we rewrite  $\sum_{j}^{k} \sum_{i}^{k} x_{i} a_{ji} x_{j} = \sum_{i}^{k} a_{ii} x_{i}^{2} + \sum_{i}^{k} \sum_{j \neq i}^{k} x_{i} a_{ji} x_{j}$ ) 2. Take partial derivative with element  $x_{i}$ :

Take partial derivative w.r.t. element 
$$x_j$$
:  

$$\frac{\partial}{\partial x_j} \left[ \sum_{i}^{k} \sum_{j}^{k} x_i a_{ji} x_j \right] = 2 a_{jj} x_j + \sum_{i \neq j}^{k} x_i a_{ji} + \sum_{i \neq j}^{k} a_{ij} x_i$$

$$= \sum_{i}^{k} x_i a_{ji} + \sum_{i}^{k} a_{ij} x_i$$

## Appendix B: Expectation of a RV and Rules of Expectations

Let *X* denote a *discrete RV* with probability function p(x), then the expected value of *X*, E[X], is defined to be:

 $E[X] = \sum_{i} x_{i} p(x_{i})$ and if *X* is *continuous* with probability density function *f*(*x*):

$$E[X] = \int_{-\infty}^{\infty} x f(x) dx$$

• For the continuous case, the expected value of g(X), E[g(X)], is:

$$[K] = \int_{-\infty}^{\infty} g(x) f(x) dx$$

E[2Note: The discrete case is a simple adaptation.

 $g(x) = x \qquad \Rightarrow E[g(x)] = E[x]$   $g(x) = (x - \mu)^2 \qquad \Rightarrow E[g(x)] = E[(x - \mu)^2]$   $g(x) = (x - \mu)^k \qquad \Rightarrow E[g(x)] = E[(x - \mu)^k]$ **Examples**:

• We derive the rules for the continuous case. That is,

$$\mathbf{E}[X] = \int_{-\infty}^{\infty} g(x) f(x) dx$$

- Rule 1. E[c] = c, where c is a constant. <u>Proof</u>: g(x) = cThen,  $E[g(x)] = E[c] = \int_{-\infty}^{\infty} c f(x) dx = c \int_{-\infty}^{\infty} f(x) dx = c$ 

- Rule 2. E[c + dX] = c + dE[X], where c & d are constants. Proof: q(x) = c + dXThen,  $E[g(x)] = E[c + dX] = \int_{-\infty}^{\infty} (c + dx) f(x) dx$  $= c \int_{-\infty}^{\infty} f(x) dx + d \int_{-\infty}^{\infty} x f(x) dx$ = c + d E[X]

- **Rule 3**. Var[X] =  $\mu_2^0 = E[(X - \mu)^2] = E[X^2] - [E(X)]^2 = \mu_2 - \mu^2$ Proof:  $g(x) = (x - \mu)^2$  $Var[X] = E[(X - \mu)^{2}] = \int_{-\infty}^{\infty} (x - \mu)^{2} f(x) dx$  $= \int_{-\infty}^{\infty} (x^2 - 2x\mu + \mu^2) f(x) dx$ =  $\int_{-\infty}^{\infty} x^2 f(x) dx - \int_{-\infty}^{\infty} 2x\mu f(x) dx + \int_{-\infty}^{\infty} \mu^2 f(x) dx$ =  $\int_{-\infty}^{\infty} x^2 f(x) dx - 2\mu \int_{-\infty}^{\infty} x f(x) dx + \mu^2 \int_{-\infty}^{\infty} f(x) dx$  $= E[X^2] - 2 \mu E(X) + \mu^2 = \mu_2 - \mu^2$ 

<u>Note</u>: If  $\mu = 0$ , then  $\operatorname{Var}[X] = E[X^2]$ 

- **Rule 4**.  $Var[a X + b] = a^2 Var[X]$ Proof: Do it yourself. Define  $g(x) = (aX + b - E[aX + b])^2$ Then, simplify *b*, apply square and use Rule 2.

• Suppose excess returns for asset *i*,  $r_{i,t} - r_f$ , are driven by the following linear model (DGP) behind the CAPM):

 $(r_{i,t}-r_f) = \alpha_i + \beta_i (r_{m,t}-r_f) + \varepsilon_{i,t},$ where

 $r_{m,t} - r_f$  = excess return on the market portfolio at time *t*.  $\beta_i$  = the sensitivity to market (systematic) risk.  $\varepsilon_{i,t}$  = idiosyncratic error term, with mean 0 & unrelated to  $r_{m,t}$ .

Then,

$$E[(r_{i,t} - r_f)] = E[\alpha_i] + \beta_i E[(r_{m,t} - r_f)] + E[\varepsilon_{i,t}]$$
 (by Rule 2)  

$$E[(r_{i,t} - r_f)] = \alpha_i + \beta_i E[(r_{m,t} - r_f)] + E[\varepsilon_{i,t}]$$
 (by Rule 1)  

$$E[(r_{i,t} - r_f)] = \alpha_i + \beta_i E[(r_{m,t} - r_f)]$$
 -by assumption about mean 0 of  $\varepsilon_{i,t}$ 

The CAPM implies that  $\alpha_i = 0$ .

Also, by **Rule 4**, 
$$\operatorname{Var}[(r_{i,t} - r_f)] = (\beta_i)^2 \operatorname{Var}[(r_{m,t} - r_f)] + \operatorname{Var}[\varepsilon_{i,t}]$$

**Example:** We estimate  $E[r_{i=IBM} - r_f] \& Var[r_{i=IBM} - r_f]$  for IBM, using OLS estimates for  $\alpha_i \& \beta_i \& Var[\varepsilon_{i,t}]$  and sample estimates for  $E[r_{m,t} - r_f] \& Var[r_{m,t} - r_f]$ .

Estimates:

 $b_1$  (Intercept) = -0.00579,

$$b_2 = 0.89577,$$

Estimated  $\operatorname{Var}[\varepsilon_{i,t}] = 0.003484$ 

Mean  $[r_{m,t} - r_f] = 0.0056489$ 

Estimated  $Var[r_{m,t} - r_f] = 0.002148$ 

Then, our estimates for  $E[r_{i=IBM} - r_f] \& Var[r_{i=IBM} - r_f]$  are:  $E[r_i - r_f] = b_1 + b_2 E[(r_{m,t} - r_f)]$  -0.00579 + 0.89577 \* 0.0056489 = -0.000729 (-0.0729%)  $Var[r_i - r_f] = b_2^2$  Est.  $Var[r_{m,t} - r_f] + Var[\varepsilon_{i,t}] = 0.89577^2 * 0.002148 + 0.003484 = .0052076$  $\Rightarrow SD[r_i - r_f] = sqrt(.0052076) = 0.07216$  (7.22%)

# Lecture 4 - OLS: Sampling, and Bootstrapping

### **OLS Estimation - Assumptions**

CLM Assumptions (A1) DGP:  $\mathbf{y} = \mathbf{X} \ \beta + \boldsymbol{\varepsilon}$  is correctly specified. (A2)  $\mathbf{E}[\boldsymbol{\varepsilon}|\mathbf{X}] = 0$ (A3)  $\operatorname{Var}[\boldsymbol{\varepsilon}|\mathbf{X}] = \sigma^2 \mathbf{I}_T$ (A4)  $\mathbf{X}$  has full column rank – rank( $\mathbf{X}$ )=*k*-, where  $T \ge k$ .

• From assumptions (A1), (A2), and (A4)  $\Rightarrow \mathbf{b} = (\mathbf{X'X})^{-1}\mathbf{X' y}$ 

We define  $\mathbf{e} = \mathbf{y} - \mathbf{X}\mathbf{b}$   $\Rightarrow \mathbf{X}'\mathbf{e} = \mathbf{X}' (\mathbf{y} - \mathbf{X}\mathbf{b}) \mathbf{X}'\mathbf{y} - \mathbf{X}'\mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y} = \mathbf{0}$ 

• Now, we will study the properties of **b**.

### **Sampling Distribution of b**

Small sample = For all sample sizes -i.e., for all values of T (or N).

 $\mathbf{b} = \mathbf{\beta} + (\mathbf{X}'\mathbf{X})^{\mathsf{T}}\mathbf{X}'\mathbf{\varepsilon} \implies \mathbf{b}$  is a vector of random variables.

• Properties

(1)  $E[\mathbf{b} | \mathbf{X}] = \mathbf{\beta}$ (2)  $Var[\mathbf{b} | \mathbf{X}] = E[(\mathbf{b} - \mathbf{\beta}) (\mathbf{b} - \mathbf{\beta})' | \mathbf{X}] = \sigma^2 (\mathbf{X}'\mathbf{X})^{-1}$ (3) Gauss-Markov Theorem: **b** is BLUE (MVLUE). (4) If (A5)  $\varepsilon | \mathbf{X} \sim N(\mathbf{0}, \sigma^2 \mathbf{I}_T) \implies \mathbf{b} | \mathbf{X} \sim N(\mathbf{\beta}, \sigma^2 (\mathbf{X}'\mathbf{X})^{-1}) \implies \mathbf{b}_k | \mathbf{X} \sim N(\mathbf{\beta}_k, \sigma^2 (\mathbf{X}'\mathbf{X})_{kk})^{-1}$ 

(<u>Note</u>: the last implication is derived from the fact that the marginal distributions of a multivariate normal are also normal.)

Note: Under (A5), b is also the MLE. Thus, it has all the nice MLE properties: efficiency, consistency, sufficiency and invariance!

### **Sampling Distribution of b**

Recall that a sample statistic like **b** is a function of RVs. Then, it has a statistical distribution.

In general, in finance, we observe only *one* sample mean (actually, our only sample). But, *many* sample means are possible from the DGP.

• A sampling distribution is a distribution of a statistic over all possible samples.

Let's generate some y<sub>i</sub>'s using a DGP and, then, some **b**'s. Using:

 $\mathbf{b} = \mathbf{\beta} + (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{\varepsilon} = \mathbf{\beta} + \Sigma_i \mathbf{v}_i'\varepsilon_i$ 

Set  $\beta$  = .4; then, the DGP is:  $\mathbf{y} = (.4) \mathbf{X} + \boldsymbol{\epsilon}$ (1) Generate  $\mathbf{X}$  (to be treated as numbers). Say  $\mathbf{X} \sim N(2,4)$   $\Rightarrow x_1 = 3.22, x_2 = 2.18, x_3 = -0.37, \dots, x_T = 1.71$ (2) Generate  $\boldsymbol{\epsilon} \sim N(0,1)$   $\Rightarrow$  draws  $\varepsilon_1 = 0.52, \varepsilon_2 = -1.23, \varepsilon_3 = 1.09, \dots, \varepsilon_T = -0.09$ (3) Generate  $\mathbf{y} = .4 \mathbf{X} + \boldsymbol{\epsilon}$   $\Rightarrow y_1 = .4 * 3.22 + 0.52 = 1.808$   $y_2 = .4 * 2.18 + (-1.23) = -0.358$   $y_3 = .4 * (-0.37) + 1.09 = 0.942$   $\dots$   $y_T = .4 * 1.71 + (-0.09) = 0.594$ (4) Generate  $\mathbf{b} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y} = \Sigma_i (x_i - \bar{x}) (y_i - \bar{y}) / \Sigma_i (x_i - \bar{x})^2$ 

• We want to generate many **b**'s. Steps

- (1) Generate **X** (to be treated as numbers). Say  $\mathbf{X} \sim N(2,4)$
- (2) Generate  $\boldsymbol{\varepsilon} \sim N(0,1)$
- (3) Generate  $\mathbf{y} = .4 \mathbf{X} + \boldsymbol{\varepsilon}$
- (4) Generate  $\mathbf{b} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y} = \Sigma_i (\mathbf{x}_i \bar{\mathbf{x}}) (\mathbf{x}_i \bar{\mathbf{y}})/\Sigma_i (\mathbf{x}_i \bar{\mathbf{x}})^2$

Conditioning on step (1), we can repeat (2)-(4) B times, say 1,000 times. Then, we are able to generate a sampling distribution for  $\mathbf{b}$ .

We can, obviously, play with *T*; say *T*=100; 1,000; 10,000.

We can check:  $E[\mathbf{b}|\mathbf{X}] = (1/B) \Sigma_i \mathbf{b}_i = \mathbf{\beta}$ ?

We can calculate the variance of Var[**b**|**X**].

### Sampling Distribution of b – Code in R

Steps (1)-(4) in R to generate **b**, with a sample of size T=100:> T <- 100</td># sample size> x <- rnorm(T,2,2)</td># generate **x** from a N(2, 2<sup>2</sup>).> ep <- rnorm(T,0,1)</td># generate errors from a N(0, 1).> y <- .4\*x + ep</td># generate **y**> b <-solve(t(x)%\*% x)%\*% t(x)%\*%y</td># OLS regression

We run these commands B (say, B=1,000) times to get the sampling distribution of **b**. Then, we can calculate means, variances, skewness and kurtosis coefficients, etc.

• Script to generate the sampling distribution for B=1,000 & T=100:

Allbs=NULL	#Initialize vector that collects the b
T <- 100	
x <- rnorm(T,2,2)	# generate x
reps=1000	# number of repetitions (B)
for (i in seq(1,reps,1)){	# "for" loop starts
ep <- rnorm(T,0,1)	# generate errors, ep
y <4*x+ep	#generate y
$b \le solve(t(x)\%*\% x)\%*\% t(x)\%*\%y$	#OLS regression
Allbs=rbind(Allbs,b)	#accumulate b as rows
}	# loop ends
mb <- mean(Allbs)	
varb <- var(Allbs)	
hist(Allbs[,1],main="Histogram for OLS C	oefficients", xlab="b Coefficients")

For T=100 B = 1,000 Mean[b] = 0.3995132 SD[b] = 0.02613134

## Histogram for OLS Coefficients



For T=1,000 B = 1,000 Mean[b] = 0.3999375 SD[b] = 0.022086

#### Histogram for OLS Coefficients



## **Bootstrapping (Again!)**

*Bootstrapping* is the practice of estimating the properties of an estimator -say, its variance- by measuring those properties when sampling from an approximating distribution (the *bootstrap DGP*).

<u>Idea</u>: We use the data at hand -the empirical distribution (ED)- to estimate the variation of statistics that are themselves computed from the same data. Recall that, for large samples drawn from F, the ED approximates the CDF of F very well.

Thus, an easy choice for an approximating distribution is the ED of the observed data. That is, the ED becomes a "*fake population*."

John Fox (2005, UCLA): "The population is to the sample as the sample is to the bootstrap samples."

## **Bootstrapping: Empirical Bootstrap (Again!)**

Suppose we have a dataset with N i.i.d. observations drawn from F:

 $\{x_1, x_2, x_3, ..., x_N\}$  -"fake population."

From the ED,  $F^*$ , we sample with replacement *N* observations:  $\{x_1^*, x_2^*, x_3^*, ..., x_N^*\}$  - a bootstrap sample

This is an *empirical bootstrap sample*, which is a resample of the same size N as the original data, drawn from  $F^*$ .

For any statistic  $\theta$  computed from the original sample data, we can define a statistic  $\theta^*$  by the same formula, but computed instead using the resampled data.



• We compute many  $\hat{\theta}^*$ , by resampling many times from  $F^*$ . Say, we resample  $\theta^* B$  times:  $\{\hat{\theta}_1^*, \hat{\theta}_2^*, \hat{\theta}_3^*, ..., \hat{\theta}_B^*\}.$ 

From this collection of  $\hat{\theta}^*$ 's, we learn about statistic  $\theta$ : we compute moments, C.I.'s, etc.

**Bootstrap Steps:** 

- 1. From the original sample, draw random sample with size N.
- 2. Compute statistic  $\theta$  from the resample in 1:  $\hat{\theta}_1^*$ .
- 3. Repeat steps 1 & 2 B times  $\Rightarrow$  Get B statistics:  $\{\hat{\theta}_1^*, \hat{\theta}_2^*, \hat{\theta}_3^*, ..., \hat{\theta}_B^*\}$
- 4. Compute moments, draw histograms, etc. for these *B* statistics.

• Results:

**1.** With a large enough *B*, the LLN allows us to use the  $\hat{\theta}^*$ 's to estimate the distribution of  $\hat{\theta}$ ,  $F(\hat{\theta})$ .

2. The variation in  $\hat{\theta}$  is well approximated by the variation in  $\hat{\theta}^*$ .

Result 2 is the one we used in Lecture 2 to estimate the size of a C.I.

### **Bootstrapping: Variations**

If the ED is used for the draws, the method is usually called the *nonparametric bootstrap*. If a distribution is assumed, say a t-distribution, and we draw from this distribution, the method is called the *parametric bootstrap*.

• If the y's and the x's are sampled together, this method is sometimes called the *paired bootstrap* –for example, in a regression or to bootstrap a correlation coefficient.

• If blocks of data are sample together, the method is called *block bootstrap* –for example, in the presence of correlated data, typical of time series or spatial data.

### **Bootstrapping: Why?**

Question: Why do we need a bootstrap?

- *N* is "small," asymptotic assumptions do not apply.
- DGP assumptions are violated.
- Distributions are complicated.

The main appeal is its simplicity and its *consistent* results.

## **Bootstrapping in Econometrics**

Bootstrapping provides a very general method to estimate a wide variety of statistics. It is most useful when:

(1) Reliance on "formulas" is problematic because the formula's assumptions are dubious.

- (2) A formula holds only as  $T \rightarrow \infty$ , but our sample is not very big.
- (3) A formula is complicated or it has not even been worked out yet.

The most common econometric applications are situations where you have a consistent estimator of a parameter of interest, but it is hard or impossible to calculate its standard error or its C.I.

<u>Technical note</u>: Bootstrapping is easiest to implement if the estimator is "smooth,"  $\sqrt{T}$ -consistent, and based on an *i.i.d.* sample. In other situations, it is more complicated.

### **Bootstrapping in Econometrics: Example**

You are interested in the relation between CEO's education (**X**) and firm's long-term performance (**y**). You have 1,500 observations on both variables. You estimate the correlation coefficient,  $\rho$ , with its sample counterpart, *r*. You find the correlation to be very low.

Q: How reliable is this result? The distribution of r is complicated. You decide to use a bootstrap to study the distribution of r. Note that to compute r, we need to bootstrap pairs, then, we use a paired bootstrap.

Randomly construct a sequence of *B* samples (all with T=1,500). Say, B<sub>1</sub> = {(x<sub>1</sub>,y<sub>1</sub>), (x<sub>3</sub>,y<sub>3</sub>), (x<sub>6</sub>,y<sub>6</sub>), (x<sub>6</sub>,y<sub>6</sub>), ..., (x<sub>1458</sub>,y<sub>1458</sub>)}  $\Rightarrow r_1$ B<sub>2</sub> = {(x<sub>5</sub>,y<sub>5</sub>), (x<sub>7</sub>,y<sub>7</sub>), (x<sub>11</sub>,y<sub>11</sub>), (x<sub>12</sub>,y<sub>12</sub>), ..., (x<sub>1486</sub>,y<sub>1486</sub>)}  $\Rightarrow r_2$ .... B<sub>B</sub> = {(x<sub>2</sub>,y<sub>2</sub>), (x<sub>2</sub>,y<sub>2</sub>), (x<sub>2</sub>,y<sub>2</sub>), (x<sub>3</sub>,y<sub>3</sub>), ..., (x<sub>1499</sub>,y<sub>1499</sub>)}  $\Rightarrow r_B$ 

We rely on the observed data. We take it as our "fake population" and we sample from it *B* times. We have a collection of *bootstrap subsamples*.

The sample size of each bootstrap subsample is the same (T). Thus, some elements are repeated.

Now, we have a collection of estimators of  $\mathbf{p}_i$ 's: { $r_1$ ,  $r_2$ ,  $r_3$ , ...,  $r_B$ }. We can do a histogram and get an approximation of the probability distribution. We can calculate its mean, variance, kurtosis, confidence intervals, etc.

# Bootstrapping in Econometrics: Estimating the mean & correlation coefficient

**Example:** We bootstrap the mean returns of IBM, using monthly data 1973-2020, with B = 1,000. (You need to install R package *boot*.)

#### sim size = 1000

# bootstrapping with sim\_size replications
boot.samps <- boot(data=ibm\_x, statistic=mean\_p, R=sim\_size)</pre>

#### ORDINARY NONPARAMETRIC BOOTSTRAP

Call: boot(data = ibm\_x, statistic = mean\_p, R = sim\_size)

Bootstrap Statistics : original bias std. error t1\* -0.0006990633 5.021474e-07 0.002964358

> boot.samps\$t[1:10] # Show first 10 bootstrapped mean
[1] -0.0066684274 0.0011648002 -0.0010053505 -0.0024989738 -0.0025442486
[6] 0.0007935133 -0.0039867127 0.0030962313 -0.0017929592 -0.0023480292

# Elegant histogram

> hist(boot.samps\$t,main="Histogram for Bootstrapped Means",

+ xlab="Means", breaks=20)

Histogram for Bootstrapped Means



**Example:** We bootstrap the correlation between the returns of IBM & the S&P 500, using monthly data 1973-2020, with B = 1,000.

sim\_size = 1000
x\_sp <- SFX\_da\$SP500
lr\_sp <- log(x\_sp[-1]/x\_sp[-T])
dat\_spibm <- data.frame(lr\_sp, lr\_ibm)</pre>

### ORDINARY NONPARAMETRIC BOOTSTRAP

Call: boot(data = dat\_spibm, statistic = **cor\_xy**, R = **sim\_size**)

Bootstrap Statistics : original bias std. error t1\* 0.5894632 -0.001523914 0.03406313

> boot.samps\$t[1:10] #show first 10 bootstrapped correlations coeff
[1] 0.5863186 0.5898572 0.6473122 0.6473249 0.5311525 0.5734280 0.6241236 0.5790740
[9] 0.5790095 0.5932918

> mean(boot.samps\$t)	#our estimate of the correlation
[1] 0.5879392	
<pre>&gt; sd(boot.samps\$t)</pre>	#SD of the correlation estimate
[1] 0.03406313	

# Elegant histogram

> hist(boot.samps\$t,main="Histogram for Bootstrapped Correlations",

+ xlab="Correlations", breaks=20)



Simple 95% percentile method C.I. new <- sort(boot.samps\$t)</li>
> new[25]
[1] 0.5151807
> new[975]
[1] 0.6495722

<u>Note</u>: You get same results using boot.ci(boot.samps, type = "**perc**")

Empirical boostrap method C.I. (our preferred method)
 > boot.ci(boot.samps, type="basic")
 BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
 Based on 1000 bootstrap replicates

CALL : boot.ci(boot.out = boot.samps, type = "**basic**")

Intervals : Level Percentile 95% (0.5293, 0.6637) Calculations and Intervals on Original Scale.

## **Bootstrapping: How many bootstraps?**

It is not clear. There are many theorems on asymptotic convergence, but there are no clear rules regarding *B*. There are some suggestions.

Efron and Tibsharani's (1994) textbook recommends *B*=200 as enough. (Good results with *B* as low as 25!)

Davidson and Mackinnon's (2001) textbook suggests steps to select B. In the D&M simulations, on average, B is between 300 and 2,400.

Wilcox's (2010) textbook recommends "599 [...] for general use."

<u>Rule of thumb</u>: Start with B=100, then, try B=1,000, and see if your answers have changed by much. Increase bootstraps until you get stability in your answers.

**Example:** We bootstrap the correlation between IBM returns and S&P 500 returns, using B = 100.

> # view bootstrap results
> boot.samps
ORDINARY NONPARAMETRIC BOOTSTRAP

Call: boot(data = dat\_spibm, statistic = cor\_xy, R = sim\_size)

Bootstrap Statistics : original bias std. error t1\* 0.5898636 -0.00115623 0.03449216

> mean(boot.samps\$t)
[1] 0.5887074
> sd(boot.samps\$t)
[1] 0.02885868. ¶



Example: We bootstrap the correlation between IBM returns and S&P 500 returns, using B = 25.
# view bootstrap results
boot.samps

ORDINARY NONPARAMETRIC BOOTSTRAP

Call: boot(data = dat spibm, statistic = cor xy, R = sim size)

Bootstrap Statistics : original bias std. error t1\* 0.5898636 -0.00115623 0.03449216

```
> mean(boot.samps$t)
[1] 0.5847676
> sd(boot.samps$t)
[1] 0.03449216
```

Conclusion: Results do not change that much. ¶

### **Bootstrapping: Bias**

You can estimate the bias of the bootstrap of a parameter, say **b**:  $Bias(\mathbf{b}) = (1/B)\Sigma_r \mathbf{b}(\mathbf{r}) - \mathbf{b}$ 

<u>Note</u>: In the OLS case, **b** is an unbiased estimator, but as an estimate, the bias can be non-zero. This estimate must be analyzed along the SE's.

**Example:** In the previous bootstrapping correlations exercise, R displays the bias: Bootstrap Statistics :

original bias std. error t1\* 0.5898636 -0.001244376 0.03455582. ¶

## **Bootstrapping: Linear Model - Var[b]**

Some assumptions in the CLM are not reasonable –for example, (A3) assuming homoscedasticity or no serial correlation, or if (A5) assuming normality. If we assuming normality (A5), we also assume the sampling distribution of **b**. But if data is not normal, the normality of the sampling distribution of **b** only apply for large N–i.e., asymptotic results.

We can use a bootstrap to estimate the sampling distribution of **b**. It can give us a better idea of the small sample distribution. Then, we can estimate the Var[**b**].

Monte Carlo (MC=repeated sampling) method:

1. Estimate model using full sample (of size T)  $\Rightarrow$  we get **b** 

- 2. Repeat B times:
  - Draw T observations from the sample, with replacement
  - Estimate  $\boldsymbol{\beta}$  with  $\mathbf{b}(\mathbf{r})$ .
- 3. Estimate variance with  $V_{boot} = (1/B) [\mathbf{b}(r) - \mathbf{b}] [\mathbf{b}(r) - \mathbf{b}]^{\prime}$
- In the case of one parameter, say  $\mathbf{b}_1$ : Estimate variance with  $Var_{boot}[\mathbf{b}_1] = (1/B)\Sigma_r [\mathbf{b}_1(r) - \mathbf{b}_1]^2$

You can also estimate Var[**b**<sub>1</sub>] as the variance of **b**<sub>1</sub> in the bootstrap Var<sub>boot</sub>[**b**<sub>1</sub>] =  $(1/B)\Sigma_r [\mathbf{b}_1(\mathbf{r}) - \text{mean}(\mathbf{b}_{1-\mathbf{r}})]^2$ ; mean(**b**<sub>1-r</sub>) =  $(1/B)\Sigma_r \mathbf{b}_1(\mathbf{r})$ 

<u>Note</u>: Obviously, this method for obtaining standard errors of parameters is most useful when no formula has been worked out for the standard error (SE), or the formula is complicated –for example, in some 2-step estimation procedures–, or the assumptions behind the formula are not realistic.

## **Bootstrapping: Linear Model - Estimating Var[b]**

**Example:** We bootstrap the SE for **b** for IBM returns using the 3 FF Factor Model. We use the R package *lmboot*, which needs to be installed with the **install.packages()** function.

```
library(lmboot)
                                                 # need to run before
install.packages("Imboot")
y \le ibm x
x \le cbind(x0, Mkt RF, SMB, HML)
dat yx <- data.frame(y, x)
                                          # Imboot needs an R data frame. We make one.
ff3 b <- paired.boot(y ~ x-1, data=dat yx, B = sim size)
ff3 b$origEstParam
                                          # print OLS results ("original estimates")
> ff3 b$origEstParam
         [,1]
              -0.005088944
Х
xMkt RF
              0.908298898
xSMB
              -0.212459588
xHML
              -0.171500223
# Mean values for b
mean(ff3 b$bootEstParam[,1])
                                   # print mean of bootstrap samples for constant
                                   # print mean of bootstrap samples for Mkt RF
mean(ff3 b$bootEstParam[,2])
                                   # print mean of bootstrap samples for SMB
mean(ff3 b$bootEstParam[,3])
                                   # print mean of bootstrap samples for HML
mean(ff3 b$bootEstParam[,4])
```

```
# Statistics for sampling distribution of b
summary(ff3_b$bootEstParam)  # distribution of b
```

# SD of parameter vector b
sd(ff3\_b\$bootEstParam[,1])
sd(ff3\_b\$bootEstParam[,2])
sd(ff3\_b\$bootEstParam[,3])
sd(ff3\_b\$bootEstParam[,4])

# bootstrap bias

ff3\_b\$origEstParam[1] - mean(ff3\_b\$bootEstParam[,1]) ff3\_b\$origEstParam[2] - mean(ff3\_b\$bootEstParam[,2]) ff3\_b\$origEstParam[3] - mean(ff3\_b\$bootEstParam[,3]) ff3\_b\$origEstParam[4] - mean(ff3\_b\$bootEstParam[,4])

> summary(ff3 b\$bootEstParam)

	Х		xMkt_RF	xSN	ЛB		xHML
Min.	:-0.012159	Min.	:0.7115	Min.	:-0.5175	Min.	:-0.4699
1st Qu.	:-0.006731	1st Qu.	:0.8669	1st Qu.	:-0.2890	1st Qu.	:-0.2362
Median	:-0.005074	Median	:0.9087	Median	:-0.2185	Median	:-0.1690
Mean	:-0.005008	Mean	:0.9068	Mean	:-0.2125	Mean	:-0.1710
3rd Qu.	:-0.003273	3rd Qu.	:0.9492	3rd Qu.	:-0.1415	3rd Qu.	:-0.1086
Max.	: 0.002293	Max.	:1.0854	Max.	: 0.1909	Max.	: 0.2477

> sd(ff3\_b\$bootEstParam[,1])
[1] 0.002493708
> sd(ff3\_b\$bootEstParam[,2])
[1] 0.06132218
> sd(ff3\_b\$bootEstParam[,3])
[1] 0.1108
> sd(ff3\_b\$bootEstParam[,4])
[1] 0.09729972

• Comparing OLS and Bootstrap

	OLS		Bootstrap		$\mathbf{Bias}$
	Coeff. (1)	S.E.	Coeff. (2)	S.E.	(2)-(1)
X	-0.00509	0.00249	-0.00501	0.00249	8.0765e-05
xMkt_RF	0.90829	0.05672	0.90684	0.06132	-0.0014571
xSMB	-0.21246	0.08411	-0.21245	0.11080	1.9914e-06
xHML	-0.17150	0.08468	-0.17099	0.09730	0.0005133

<u>Conclusion</u>: Very similar results for the coefficients, a bit different for S.E. Usually, we rely on the bigger S.E., in this case, for inferences we'd rely on the Booststrap S.E.

> ff3 b\$bootEstParam[1:10,] # print the first 10 of B=1,000 bootstrap samples xMkt RF xSMB xHML х [1,] -6.109007e-03 0.9186830 -0.1299534100 -0.163421636 [2,] -1.757503e-03 0.8333006 -0.2067565390 -0.147604991 [3,] -3.907573e-03 0.9746878 -0.2870744815 -0.169189619 [4,] 1.596103e-03 0.9185157 -0.2937731120 -0.296972497 [5,] -8.409239e-03 0.7309406 -0.0681714313 -0.149883639 [6,] -1.998929e-03 0.9133751 -0.3001713380 -0.315913280 [7,] -6.289286e-03 0.9441856 -0.2276894034 -0.058924929 [8,] -5.533354e-03 0.8210057 -0.2221866298 -0.078512341 [9,] -6.152301e-03 1.0389917 -0.2592958758 -0.237930809 [10,] -3.778058e-03 0.9544829 -0.1859554067 -0.217702583



From the B samples, we compute variances and SD as usual.

## **Bootstrapping: Some Remarks**

Question: How reliable is bootstrapping?

- There is still no consensus on how far it can be applied, but for now nobody is going to dismiss your results for using it.

- There is a general agreement that for normal and close to normal (and symmetric) distributions it works well.

- Bootstrapping is more problematic for skewed distributions.

- It can be unreliable for situations where there are not a lot of observations. Typical example in finance: estimation of quantiles in the tails of returns distributions.

<u>Note</u>: We presented two simple examples. There are many bootstraps variations. We will not cover them.

# Lecture 5 - Testing in the CLM

## **Review – OLS Assumptions**

CLM Assumptions (A1) DGP:  $\mathbf{y} = \mathbf{X} \beta + \boldsymbol{\varepsilon}$  is correctly specified. (A2)  $E[\boldsymbol{\varepsilon}|\mathbf{X}] = 0$ (A3)  $Var[\boldsymbol{\varepsilon}|\mathbf{X}] = \sigma^2 \mathbf{I}_T$ (A4) X has full column rank -rank(X)=k-, where  $T \ge k$ .

Issues for this lecture: Q: What happens when we impose restrictions to the DGP (A1)?

Q: How do we test restrictions in the context of OLS estimation?

## **OLS Subject to Linear Restrictions**

Restrictions: Theory imposes certain restrictions on parameters and provide the foundation of several tests. In this Lecture, we only consider linear restrictions, written as  $\mathbf{R}\boldsymbol{\beta} = \mathbf{q}$ . The dimension of **R** is *Jxk*, where *J* is the number of restrictions, and *k* is the number of parameters.  $\boldsymbol{\beta}$ , as usual, is a *kx*1 column vector. Then, **q** is a *Jx*1 column vector.

### **Examples**:

(1) Dropping variables from the equation. That is, certain coefficients in **b** forced to equal 0. For example, in the CAPM, we impose that variables  $\mathbf{x}_3$ =*SMB* and  $\mathbf{x}_4$ =*HML* are not part of the model. That is, we impose  $\beta_{SMB} = 0$  and  $\beta_{HML} = 0$ .Using the above notation:

$$\mathbf{R}\boldsymbol{\beta} = \mathbf{q} \qquad \Rightarrow \begin{bmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix} * \begin{bmatrix} \beta_1 \\ \beta_{Mkt} \\ \beta_{SMB} \\ \beta_{HML} \end{bmatrix} = \begin{bmatrix} \beta_{SMB} \\ \beta_{HML} \end{bmatrix} = \begin{bmatrix} 0 \\ 0 \end{bmatrix}$$

We have two restrictions (J=2):  $\beta_{SMB} = 0 \& \beta_{HML} = 0$ . We have k=4 parameters.

 $\Rightarrow$  **R** is a 2x4 matrix,  $\beta$  is a 4x1 vector, and **q** is a 2x1 vector.

Note: The restrcitions make the FF model into the traditional CAPM.

(2) Adding up conditions: Sums of certain coefficients must equal fixed values. Adding up conditions in demand systems. In a CAPM setting, the sum of all cross-sectional  $\beta_i$ 's should be equal to 1. For example, in the 3 Fama-French factor model, we force  $\beta_{SMB} + \beta_{HML} = 1$ .

$$\mathbf{R}\boldsymbol{\beta} = \mathbf{q} \qquad \Rightarrow \begin{bmatrix} 0 & 0 & 1 & 1 \end{bmatrix} * \begin{bmatrix} \boldsymbol{\beta}_1 \\ \boldsymbol{\beta}_{Mkt} \\ \boldsymbol{\beta}_{SMB} \\ \boldsymbol{\beta}_{HML} \end{bmatrix} = \boldsymbol{\beta}_{SMB} + \boldsymbol{\beta}_{HML} = 1$$

We have one restrictions (J=1):  $\beta_{SMB} + \beta_{HML} = 1$ . We have k=4 parameters.

 $\Rightarrow$  **R** is a 1x4 matrix (a row vector),  $\beta$  is a 4x1 vector, and **q** is a scalar.

(3) Equality restrictions: Certain coefficients must equal other coefficients. Using real vs. nominal variables in equations. For example, in the 3 Fama-French factor model, we force  $\beta_{SMB} = \beta_{HML}$ .

$$\mathbf{R}\boldsymbol{\beta} = \mathbf{q} \qquad \Rightarrow \begin{bmatrix} 0 & 0 & 1 & -1 \end{bmatrix} * \begin{bmatrix} \beta_1 \\ \beta_{Mkt} \\ \beta_{SMB} \\ \beta_{IIMI} \end{bmatrix} = 0.$$

We have one restrictions (J = 1):  $\beta_{SMB} + \beta_{HML} = 1$ . We have k = 4 parameters.

 $\Rightarrow$  **R** is a 1x4 matrix (a row vector),  $\beta$  is a 4x1 vector, and **q** is a scalar.

• Common formulation: We minimize the error sum of squares, subject to the linear restrictions. That is,

 $Min_{\mathbf{b}} \{S(x_{i}, \theta) = \Sigma_{i} \varepsilon_{i}^{2} = \varepsilon' \varepsilon = (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})' (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})\} \quad \text{ s.t. } \mathbf{R}\boldsymbol{\beta} = \mathbf{q}$ 

In practice, restrictions can usually be imposed by solving them out. Suppose we have a model:  $y_i = \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \varepsilon_i$ 

(1) Dropping variables –i.e., force a coefficient to equal zero, say  $\beta_3$ .

<u>Problem</u>: Min<sub>\beta</sub>  $\sum_{i=1}^{n} (y_i - \beta_1 x_{i1} - \beta_2 x_{i2} - \beta_3 x_{i3})^2$  s.t.  $\beta_3 = 0$ Min<sub>\beta</sub>  $\sum_{i=1}^{n} (y_i - \beta_1 x_{i1} - \beta_2 x_{i2})^2$ 

(2) Adding up. Suppose we impose:  $\beta_1 + \beta_2 + \beta_3 = 1$ . Then,  $\beta_3 = 1 - \beta_1 - \beta_2$ . Substituting in model:

$$(\mathbf{y} - \mathbf{x}_1) = \beta_1(\mathbf{x}_1 - \mathbf{x}_3) + \beta_2(\mathbf{x}_2 - \mathbf{x}_3) + \varepsilon.$$
  
Problem: Min<sub>\beta</sub>  $\sum_{i=1}^n ((y_i - x_{i3}) - \beta_1(x_{i1} - x_{i3}) - \beta_2(x_{i2} - x_{i3}))^2$ 

(3) Equality. Suppose we impose:  $\beta_2 = \beta_3$ . Substituting in model:

$$y = \beta_1 x_1 + \beta_2 x_2 + \beta_2 x_3 + \varepsilon = \beta_1 x_1 + \beta_2 (x_2 + x_3) + \varepsilon$$
  
Problem: Min<sub>\beta</sub>  $\sum_{i=1}^n (y_i - \beta_1 x_{i1} - \beta_2 x_{i2} - \beta_3 x_{i3})^2$  s.t.  $\beta_2 = \beta_3$   
Min<sub>\beta</sub>  $\sum_{i=1}^n (y_i - \beta_1 x_{i1} - \beta_2 (x_{i2} + x_{i3}))^2$ 

• Before setting the general restricted LS problem, we look at the simplest case: one explanatory variable (*x*) and one restriction ( $r\beta = q$ ).

Then, we set up the Lagrangean (recall values of Lagrange multiplier',  $\lambda$ , play no role):  $Min_{\beta,\lambda} L(\beta, \lambda) = \sum_{i=1}^{n} (y_i - x_i \beta)^2 + 2\lambda (r \beta - q)$ 

We take first derivatives of  $L(\beta, \lambda)$  with respect to  $\beta, \lambda$ :

$$\Rightarrow \quad \frac{\partial L(\beta,\lambda)}{\partial \beta} = -2\sum_{i}^{T} (y_{i} - x_{i} \beta))(-x_{i}) + 2\lambda r$$
$$\frac{\partial L(\beta,\lambda)}{\partial \lambda} = 2(r \beta - q)$$

Then, the f.o.c. are:

$$\Rightarrow -\sum_{i}^{T} (y_{i} - x_{i} b^{*}) (x_{i}) + \lambda r = 0 \qquad \Rightarrow \sum_{i}^{T} (y_{i} x_{i} - x_{i}^{2} b^{*}) = \lambda r$$
$$\Rightarrow \lambda (r b^{*} - q) = 0 \qquad \Rightarrow r b^{*} - q = 0$$

From the 1<sup>st</sup> equation:

$$\sum_{i}^{T} y_{i} x_{i}^{T} - b^{*} \sum_{i}^{T} x_{i}^{2} = \mathbf{x} \mathbf{y} - b^{*} (\mathbf{x} \mathbf{x})^{-1} = \lambda r \qquad \Rightarrow b^{*} = (\mathbf{x} \mathbf{x})^{-1} \mathbf{x}^{*} \mathbf{y} - (\mathbf{x} \mathbf{x})^{-1} \lambda r = b - (\mathbf{x} \mathbf{x})^{-1} \lambda r$$

 $b^* = b - r (\mathbf{x}'\mathbf{x})^{-1} \lambda \implies \text{Restricted OLS} = \text{OLS} + "correction"$ 

Premultiplying both sides by *r* and then subtract *q*:

 $r b^* - q = rb - r^2 (\mathbf{x} \mathbf{x})^{-1} \lambda - q$  $0 = -r^2 (\mathbf{x} \mathbf{x})^{-1} \lambda + (rb - q)$ 

Solving for  $\lambda \implies \lambda = [r^2 (\mathbf{x}'\mathbf{x})^{-1}]^{-1} (r\mathbf{b} - q)$ 

Substituting in b\*  $\Rightarrow$  b\* = b - (**x**'**x**)<sup>-1</sup> r [r<sup>2</sup> (**x**'**x**)<sup>-1</sup>]<sup>-1</sup> (rb - q)

This is the Restricted OLS estimator.

• Properties of Restricted OLS.

**Property 1.** Taking expectations of b\*:

 $E[b^*|X] = E[b|X] - (\mathbf{x'x})^{-1}r [r^2 (\mathbf{x'x})^{-1}]^{-1} E[(rb-q)|X]$ =  $\beta - (\mathbf{x'x})^{-1}r [r^2 (\mathbf{x'x})^{-1}]^{-1} (r\beta - q)$ 

Implications:

If the restriction is *true* –i.e.,  $(r\beta = q)$   $\Rightarrow E[b^*|X] = \beta$ If the restriction is *not true* –i.e.,  $(r\beta \neq q)$   $\Rightarrow E[b^*|X] \neq \beta$ 

Then, if theory imposes a correct restriction, then, b\* is *unbiased*:  $E[b^*|X] = \beta$ 

In practice, if restriction is true, the restricted and unrestricted estimators should be similar.

Note: If theory is correct, the expected shadow price is 0!  $E[\lambda|X] = [r^2 (\mathbf{x'x})^{-1}]^{-1} E[(rb - q)|X] = 0$ 

That is, you would pay nothing to release the restriction,  $r\beta = q$ .

**Property 2.** We compute the Var[b\*]. It can be shown that

 $Var[b^*|X] = Var[b|X] - \sigma^2 (\mathbf{x'x})^{-1} r [r^2 (\mathbf{x'x})^{-1}]^{-1} r (\mathbf{x'x})^{-1}$  $\Rightarrow Var[b|X] - Var[b^*|X] = \sigma^2 (\mathbf{x'x})^{-1} r [r^2 (\mathbf{x'x})^{-1}]^{-1} r (\mathbf{x'x})^{-1} > 0.$ 

 $\Rightarrow$  The restricted OLS estimator is more efficient!

<u>Remark from Properties 1 and 2</u>: It is common to select an estimator based on the MSE (=RSS/*T*). The one with the lowest MSE is said to be more "*precise*."

We can decompose the MSE of an estimator,  $\hat{\theta}$ , as: MSE[ $\hat{\theta}$ ] = Variance[ $\hat{\theta}$ ] + Squared bias[ $\hat{\theta}$ ]

For an unbiased estimator, like  $\mathbf{b} \Rightarrow MSE[\mathbf{b}] = Var[\mathbf{b}|\mathbf{X}]$ 

• Back to  $\mathbf{b}^*$ . Suppose the theory is incorrect  $\Rightarrow \mathbf{b}^*$  is biased.

There may be situations (small bias, but much lower variance) where  $\mathbf{b}^*$  is more "precise" (lower MSE) than **b**. It is possible that a practitioner may prefer imposing a wrong H<sub>0</sub> to get a better MSE.

• For the general case, with k explanatory variables and J restrictions, which we write as:

 $\mathbf{R}\boldsymbol{\beta} = \mathbf{q}$ , we have a programming problem:

Minimize wrt  $\boldsymbol{\beta}$   $L^* = (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})'(\mathbf{y} - \mathbf{X}\boldsymbol{\beta})$  s.t.  $\mathbf{R}\boldsymbol{\beta} = \mathbf{q}$ 

Quadratic programming problem: Minimize a quadratic criterion subject to a set of linear restrictions. We solve this minimizations problem using the Lagrange multiplier method.

We form the Lagrangean (the 2 is for convenience, since the value of  $\lambda$  is irrelevant for extrema):

Min <sub>b,λ</sub> f.o.c.:

$$\partial L^* / \partial \mathbf{b}' = -2\mathbf{X}'(\mathbf{y} - \mathbf{X}\mathbf{b}^*) + 2\mathbf{R}' \lambda = \mathbf{0} \Rightarrow -\mathbf{X}'(\mathbf{y} - \mathbf{X}\mathbf{b}^*) + \mathbf{R}' \lambda = \mathbf{0}$$
  
$$\partial L^* / \partial \lambda = 2(\mathbf{R}\mathbf{b}^* - \mathbf{q}) = \mathbf{0} \qquad \Rightarrow (\mathbf{R}\mathbf{b}^* - \mathbf{q}) = \mathbf{0}$$

 $L^* = (\mathbf{v} - \mathbf{X}\mathbf{\beta})'(\mathbf{v} - \mathbf{X}\mathbf{\beta}) + 2 \lambda (\mathbf{R}\mathbf{\beta} - \mathbf{q})$ 

where  $\mathbf{b}^*$  is the restricted OLS estimator.

f.o.c.: 
$$-X'(y - Xb^*) + R'\lambda = 0$$
 (1)  
(Rb\*-q) = 0 (2)

where **b**\* is the restricted OLS estimator.

Then, from the 1<sup>st</sup> equation (and assuming full rank for X):  

$$-\mathbf{X'y} + \mathbf{X'Xb^*} + \mathbf{R'\lambda} = \mathbf{0} \implies \mathbf{b^*} = (\mathbf{X'X})^{-1}\mathbf{X'y} - (\mathbf{X'X})^{-1}\mathbf{R'\lambda}$$

$$= \mathbf{b} - (\mathbf{X'X})^{-1}\mathbf{R'\lambda}$$

Premultiply both sides by  ${\bf R}$  and then subtract  ${\bf q}$ 

Rb* = Rb* - 0	= $\mathbf{R}\mathbf{b} - \mathbf{R}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{R}'\lambda$ - $\mathbf{q} = \mathbf{R}\mathbf{b} - \mathbf{R}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{R}'\lambda - \mathbf{q}$ = $-\mathbf{R}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{R}'\lambda + (\mathbf{R}\mathbf{b} - \mathbf{q})$
Solving for $\lambda$	$\Rightarrow \lambda = [\mathbf{R}(\mathbf{X'X})^{-1}\mathbf{R'}]^{-1} (\mathbf{Rb} - \mathbf{q})$
Substituting in <b>b</b> *	$\Rightarrow \mathbf{b}^* = \mathbf{b} - (\mathbf{X}'\mathbf{X})^{-1}\mathbf{R}'[\mathbf{R}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{R}']^{-1}(\mathbf{R}\ \mathbf{b} - \mathbf{q})$
Note: Restricted OL	LS = Unrestricted OLS + "correction"

## **Restricted Least Squares**

Question: How do linear restrictions affect the properties of the least squares estimator?

Restricted LS estimator:  $b^* = b - (X'X)^{-1}R'[R(X'X)^{-1}R']^{-1}(Rb - q)$ 

• Properties:

1. Unbiased? Yes, if Theory is correct!  $E[b^*|X] = \beta - (X'X)^{-1}R'[R(X'X)^{-1}R']^{-1}E[(Rb-q)|X] = \beta$ 

But, if Theory is incorrect:  $E[(\mathbf{Rb} - \mathbf{q})|\mathbf{X}] \neq \mathbf{0} \implies E[\mathbf{b}^*|\mathbf{X}] \neq \mathbf{\beta}$ .

2. Efficiency?

 $Var[\mathbf{b}^*|\mathbf{X}] = \sigma^2 (\mathbf{X}'\mathbf{X})^{-1} - \sigma^2 (\mathbf{X}'\mathbf{X})^{-1}\mathbf{R}'[\mathbf{R}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{R}']^{-1} \mathbf{R}(\mathbf{X}'\mathbf{X})^{-1}$ Var[**b**\*|**X**] = Var[**b**|**X**] - a nonnegative definite matrix < Var[**b**|**X**]

3. **b**\* may be more "precise," where precision is measured by the MSE (=RSS/T).

We can decompose the MSE of an estimator,  $\hat{\theta}$ , as:  $MSE[\hat{\theta}] = Variance[\hat{\theta}] + Squared bias[\hat{\theta}]$ For an unbiased estimator, say **b**, then,  $MSE[\mathbf{b}] = Var[\mathbf{b}|\mathbf{X}]$ 

Suppose the theory is incorrect. Then,  $\mathbf{b}^*$  is biased. There may be situations (small bias, but much lower variance) where  $\mathbf{b}^*$  is more "precise" (lower MSE) than  $\mathbf{b}$ . A practitioner may prefer imposing a wrong H<sub>0</sub> to get a better MSE.

#### **Restricted Least Squares - Interpretation**

1.  $\mathbf{b}^* = \mathbf{b} - \mathbf{Cm}$ ,  $\mathbf{m} =$  the "discrepancy vector"  $\mathbf{Rb} - \mathbf{q}$ . Note: If  $\mathbf{m} = \mathbf{0} \implies \mathbf{b}^* = \mathbf{b}$ . (Q: What does  $\mathbf{m} = \mathbf{0}$  mean?)

- 2.  $\lambda = [\mathbf{R}(\mathbf{X'X})^{-1}\mathbf{R'}]^{-1}(\mathbf{Rb} \mathbf{q}) = [\mathbf{R}(\mathbf{X'X})^{-1}\mathbf{R'}]^{-1}\mathbf{m}$ When does  $\lambda = \mathbf{0}$ ? We usually think of  $\lambda$  as a "shadow price."
- 3. Combining results:  $\mathbf{b}^* = \mathbf{b} (\mathbf{X}'\mathbf{X})^{-1}\mathbf{R}'\lambda$

4. We can show that RSS never decreases with restrictions:

 $e'e = (y - Xb)'(y - Xb) \le e^{*'}e^{*} = (y - Xb^{*})'(y - Xb^{*})$ ⇒ Restrictions cannot increase  $R^2 \Rightarrow R^2 \ge R^{2*}$ 

- Two cases
  - Case 1: Theory is correct:  $\mathbf{R}\boldsymbol{\beta} \mathbf{q} = \mathbf{0}$  (restrictions hold). **b**\* is unbiased & Var[**b**\*|**X**]  $\leq$  Var[**b**|**X**]
  - Case 2: Theory is incorrect:  $\mathbf{R\beta} \mathbf{q} \neq \mathbf{0}$  (restrictions do not hold). **b**\* is biased & Var[**b**\*|**X**]  $\leq$  Var[**b**|**X**].
- Interpretation
- The theory gives us information.

Bad information produces bias (away from "the truth.") Any information, good or bad, makes us more certain of our answer. In this context, *any* information reduces variance.

### **Testing: Parameter vs Diagnostic**

So far, the tests discussed in Lectures 3 & 4, involved parameters. We call these types of testing *parameter tests*.

When we tests the assumptions behind the CLM, for example, (A5), we perform a *diagnostic tests*.

• *Parameter testing:* We test economic H<sub>0</sub>'s. **Example:** Test  $\beta_k = 0$  -say, there is no size effect on the expected return equation. ¶

• *Diagnostic testing:* We test assumptions behind the model. In our case, assumptions (A1)-(A5) in the CLM.

**Example:** Test  $E[\varepsilon|X] = 0$  -i.e., the residuals are zero-mean, white noise distributed errors. ¶

## **Review – Significance Testing**

Fisher's *significance testing* procedure relies on the *p*-value: the probability of observing a result at least as extreme as the test statistic, under  $H_0$ .

• Fisher's Idea

1) Form  $H_0$  & decide on a *significance level* ( $\alpha$ %) to compare your test results.

- 2) Find T(X). Know (or derive) the distribution of T(X) under  $H_0$ .
- 3) Collect a sample of data  $X = \{x_1, x_2, ..., x_T\}$ .

Compute the test-statistics T(X) used to test  $H_0 \Rightarrow$  Report its *p*-value.

4) <u>Rule</u>: If *p*-value  $< \alpha$  (say, 5%)  $\Rightarrow$  test result is *significant*: Reject *H*<sub>0</sub>. If the results are "*not significant*," no conclusions are reached (no learning here).  $\Rightarrow$  Go back gather more data or modify model.

### **Review – Testing Only One Parameter**

We are interested in testing a hypothesis about one parameter in our linear model:  $y = X \beta + \varepsilon$ 

1. Set H<sub>0</sub> and H<sub>1</sub> (about only one parameter): H<sub>0</sub>:  $\beta_k = \beta_k^0$ H<sub>1</sub>:  $\beta_k \neq \beta_k^0$ 

2. Appropriate T(X): *t-statistic*. To derive the distribution of the test under H<sub>0</sub>, we will rely on assumption (A5)  $\epsilon$ |X ~ N(0,  $\sigma^2$ I<sub>T</sub>) (otherwise, results are only asymptotic).

Let  $\mathbf{b}_k = \text{OLS}$  estimator of  $\beta_k$  $\text{SE}[\mathbf{b}_k|\mathbf{X}] = \text{sqrt}\{[s^2(\mathbf{X}^*\mathbf{X})^{-1}]_{kk}\} = s_{b,k}$ 

From assumption (A5), we know that

$$\begin{aligned} \mathbf{b}_{k} | \mathbf{X} \sim \mathbf{N}(\boldsymbol{\beta}_{k}, \mathbf{v}_{k}^{2}) & \Rightarrow \text{Under H}_{0}: \mathbf{b}_{k} | \mathbf{X} \sim \mathbf{N}(\boldsymbol{\beta}_{k}^{0}, s_{b,k}^{2}). \\ \Rightarrow \text{Under H}_{0}: t_{k} = \frac{b_{k} - \beta_{k}^{0}}{s_{b,k}} | \mathbf{X} \sim t_{T-k}. \end{aligned}$$

3. Compute  $t_k$ ,  $\hat{t}$ , using  $b_k$ ,  $\beta_k^0$ , s, and  $(\mathbf{X}^*\mathbf{X})^{-1}$ . Get *p*-value( $\hat{t}$ ).

4. <u>Rule</u>: Set an  $\alpha$  level. If *p*-value( $\alpha$ t ) <  $\alpha$   $\Rightarrow$  Reject H<sub>0</sub>:  $\beta_k = \beta_k^0$ Alternatively, if  $|\hat{t}| > t_{T-k,(1-\alpha)/2}$   $\Rightarrow$  Reject H<sub>0</sub>:  $\beta_k = \beta_k^0$ .

## Review - Testing Only One Parameter: t-value

Special case:  $H_0: \beta_k = 0$  $H_1: \beta_k \neq 0.$ 

Then,

$$t_k = \frac{b_k}{s_{b,k}} \qquad \Rightarrow t_k \sim t_{T-k}$$

In this case, we call  $t_k$  the *t*-value or *t*-ratio.

Usually,  $\alpha = 5\%$ , then if  $|\hat{t}_k| > 1.96 \approx 2$ , we say the coefficient b<sub>k</sub> is "significant."

### **Review – Confidence Intervals**

The goal of the *confidence intervals* (C.I.) is to set the coverage probability to equal a  $(1 - \alpha)$ % pre-specified target.

When we know the distribution of point estimate, it is easy to construct a C.I. Under the usual assumptions for  $b_k$  we have:

 $C_n = [b_k - t_{T-k,1-\alpha/2} * \text{Estimated SE}(b_k), b_k + t_{T-k,(1-\alpha)/2} * \text{Estimated SE}(b_k)]$ 

This C.I. is symmetric around b<sub>k</sub>: length is proportional to SE(b<sub>k</sub>).

Usual  $\alpha$  levels and  $t_{T-k,(1-\frac{\alpha}{2})}$  -when T > 30, (usual case)  $t_{T-k,(1-\frac{\alpha}{2})} \approx z_{(1-\alpha/2)}$  $\alpha = 5\%$ , then  $z_{(1-\alpha/2)} = 1.96$ .  $\alpha = 2\%$ , then  $z_{(1-\alpha/2)} = 2.33$ .  $\alpha = 1\%$ , then  $z_{(1-\alpha/2)} = 2.58$ .

<u>**R** Note</u>: In R, we get  $\alpha = 5\%$ , then  $z_{(1-\alpha/2)} = 1.96$ , using qnorm(0.975).

### **Testing: The Expectation Hypothesis (EH)**

**Example**: EH states that forward/futures prices are good predictors of future spot rates:  $E_t[S_{t+T}] = F_{t,T}$ Implication of EH:  $S_{t+T} - F_{t,T} =$  unpredictable.

That is,  $E_t[S_{t+T} - F_{t,T}] = E_t[\varepsilon_t] = 0!$ 

Empirical tests of the EH are based on a regression:

$$(S_{t+T} - F_{t,T})/S_t = \alpha + \beta Z_t + \varepsilon_t,$$
 (where  $E_t[\varepsilon_t] = 0$ )

where  $Z_t$  represents any economic variable that might have power to explain  $S_t$ , for example, interest rate differentials,  $(i_d - i_f)$ .

Then, under EH,  $H_0: \alpha = 0 \text{ and } \beta = 0.$ vs  $H_1: \alpha \neq 0 \text{ and/or } \beta \neq 0.$ 

• We will informally test EH using exchange rates (USD/GBP), 3-mo forward rates and 3-mo interest rates.

```
SF_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/SpFor_prices.csv", head=TRUE, sep=",")
summary(SF_da)
x_date <- SF_da$Date
x_S <- SF_da$DBPSP
x_F3m <- SF_da$GBP3M
i_us3 <- SF_da$Dep_USD3M
i_uk3 <- SF_da$Dep_UKP3M
T <- length(x_S)
prem <- (x_S[-1] - x_F3m[-T])/x_S[-1]
int_dif <- (i_us3 - i_uk3)/100
y <- prem
x <- int_dif[-T]
fit_eh <- lm(y ~ x)
```

• We do two individual t-tests on  $\alpha \& \beta$ .

```
> summary(fit eh)
Call:
lm(formula = y \sim x)
Residuals:
Min
         1Q Median
                           3Q
                                   Max
-0.125672 -0.014576 -0.000439 0.017356 0.094283
Coefficients:
            Estimate
                        Std. Error t value Pr(>|t|)
(Intercept) -0.0001854 0.0016219 -0.114 0.90906 \Rightarrow constant not significant (|t|<2)
            -0.2157540 0.0731553 -2.949 0.00339 ** \Rightarrow slope is significant (|t|>2): Reject H<sub>0</sub>
Х
___
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.02661 on 361 degrees of freedom
Multiple R-squared: 0.02353, Adjusted R-squared: 0.02082
F-statistic: 8.698 on 1 and 361 DF, p-value: 0.003393
• 95% C.I. for b:
```

```
C_n = [b_k \pm t_{T-k,1-\alpha/2} * \text{Estimated SE}(b_k)]
```

hen,

 $C_n = \begin{bmatrix} -0.215754 - 1.96 * 0.0731553, -0.215754 + 1.96 * 0.0731553 \end{bmatrix}$ =  $\begin{bmatrix} -0.3591384, -0.07236961 \end{bmatrix}$ 

Since  $\beta = 0$  is not in C<sub>n</sub> with 95% confidence  $\Rightarrow$  Reject H<sub>0</sub>:  $\beta_1 = 0$  at 5% level.

Note: The EH is a joint hypothesis, it should be tested with a joint test! ¶

## The General Linear Hypothesis: Wald Statistic

Most of our test statistics, including joint tests, are Wald statistics.

Wald = normalized distance measure.

• One parameter:

$$t_k = \frac{b_k - \beta_k^0}{\mathrm{SE}[b_k]} = \mathrm{distance/unit}$$

• More than one parameter.

Let  $\mathbf{z} = (\text{random vector} - \text{hypothesized value})$  be the distance  $W = \mathbf{z}' [\text{Var}(\mathbf{z})]^{-1} \mathbf{z}$  (a quadratic form)

**Example:** Let  $\mathbf{z} = \mathbf{R}\mathbf{b} - \mathbf{q}$ , which under (A5) & H<sub>0</sub>:  $\mathbf{R}\boldsymbol{\beta} = \mathbf{q}$  $\mathbf{z} \sim N(\mathbf{0}, \operatorname{Var}[\mathbf{z}]),$  where  $\operatorname{Var}[\mathbf{z}] = \mathbf{R} [\operatorname{Var}[\mathbf{b}|\mathbf{X}]]^{-1} \mathbf{R'}$  Then, if  $H_0$  is correct, W should be a small number, ideally close to zero. A large value would be evidence against  $H_0$ .

We need the distribution of W to determine how "far" is from zero.

• Distribution of *W*? We have a quadratic form.

- If **z** is normal and  $\sigma^2$  known,  $W \sim \chi^2_{rank(Var[x])}$
- If z is normal and  $\sigma^2$  unknown, which we estimate with  $s^2 = \mathbf{e'e}/(T k)$ , then,  $W \sim F$
- If z is not normal and we use  $s^2$  to estimate the unknown  $\sigma^2$ , we rely on asymptotic theory, then,  $W \xrightarrow{d} \chi^2_{rank(Var[r])}$

## The General Linear Hypothesis: $H_0$ : $R\beta - q = 0$

• Suppose we are interested in testing J joint hypotheses.

**Example:** We want to test that in the 3 FF factor model that the SMB and HML factors have the same coefficients,  $\beta_{SMB} = \beta_{HML} = \beta^0$ .

We can write linear restrictions as H<sub>0</sub>:  $\mathbf{R}\boldsymbol{\beta} - \mathbf{q} = \mathbf{0}$ , where **R** is a Jxk matrix and **q** a Jx1 vector.

In the above example (J=2), we write:

$$\begin{bmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix} * \begin{bmatrix} \beta_1 \\ \beta_{Mkt} \\ \beta_{SMB} \\ \beta_{HML} \end{bmatrix} = \begin{bmatrix} \beta^0 \\ \beta^0 \end{bmatrix}$$

~

• Question: Is **Rb** – **q** close to **0**?

There are two different approaches to this question. Both have in common the property of unbiasedness for **b**.

Approach (1) We base the answer on the discrepancy vector:

 $\mathbf{m} = \mathbf{R}\mathbf{b} - \mathbf{q}.$ Then, we construct a Wald statistic:  $W = \mathbf{m'} (\operatorname{Var}[\mathbf{m}|\mathbf{X}])^{-1} \mathbf{m}$ to test if **m** is different from **0**.

**Approach (2)** We base the answer on a model loss of fit when restrictions are imposed: RSS must increase and  $R^2$  must go down. Then, we construct an F test to check if the unrestricted RSS ( $RSS_{II}$ ) is different from the restricted RSS ( $RSS_{R}$ ).

Approach (1). To test H<sub>0</sub>, we calculate the discrepancy vector:

 $\mathbf{m} = \mathbf{R}\mathbf{b} - \mathbf{q} \qquad (\text{under } (\mathbf{A5}) \& H_0: \mathbf{m} \sim N(\mathbf{0}, Var[\mathbf{m}])).$ Then, we compute the Wald statistic:  $W = \mathbf{m}' (Var[\mathbf{m}|\mathbf{X}])^{-1} \mathbf{m}$  It can be shown that  $\operatorname{Var}[\mathbf{m}|\mathbf{X}] = \mathbf{R}[\sigma^{2}(\mathbf{X}'\mathbf{X})^{-1}]\mathbf{R}'$ . Then,  $W = (\mathbf{Rb} - \mathbf{q})' \{\mathbf{R}[\sigma^{2}(\mathbf{X}'\mathbf{X})^{-1}]\mathbf{R}'\}^{-1}(\mathbf{Rb} - \mathbf{q})$  (under (A5) & H<sub>0</sub>:  $W \sim \chi_{I}^{2}$ ).

Under H<sub>0</sub> and assuming (A5) & estimating  $\sigma^2$  with  $s^2 = \mathbf{e'e'}(T-k)$ :

$$W^* = = \frac{W}{\left[\frac{\mathbf{e'e}}{T-k}\right]/\sigma^2} = (\mathbf{R}\mathbf{b} - \mathbf{q})' \{\mathbf{R}[s^2(\mathbf{X'X})^{-1}]\mathbf{R}\}^{-1}(\mathbf{R}\mathbf{b} - \mathbf{q})$$
$$\mathbf{F} = W^*/J \sim F_{J,T-k}$$

If (A5) is not assumed, the results are only asymptotic:  $J * F \xrightarrow{d} \chi_J^2$ 

<u>Technical Note</u>: Why  $F = W^*/J$  follows an F distribution? The F-distribution is a ratio of two independent  $\chi_I^2$  and  $\chi_T^2$  RV divided by their degrees of freedom:

$$F = \frac{\chi_J^2 / J}{\chi_T^2 / T} \sim F_{J,T}.$$

(1) Numerator:  $W = (\mathbf{R}\mathbf{b} - \mathbf{q})' \{\mathbf{R}[\sigma^2(\mathbf{X}'\mathbf{X})^{-1}]\mathbf{R}'\}^{-1}(\mathbf{R}\mathbf{b} - \mathbf{q}) \sim \chi_J^2$ 

(2) Denominator:  $(T-k) * s^2/\sigma^2 = \mathbf{e'e}/\sigma^2 \sim \chi^2_{T-k}$ 

$$F = \frac{\chi_J^2 / J}{\chi_T^2 / T} = \frac{[(\mathbf{Rb} - \mathbf{q})' \{\mathbf{R}[\sigma^2(\mathbf{X'X})^{-1}]\mathbf{R'}\}^{-1}(\mathbf{Rb} - \mathbf{q})] / J}{[(T-k)*s^2/\sigma^2]/(T-k)} \sim F_{J,T-k}.$$

**Example:** We want to test that in the 3 FF factor model (*T*=569)

1. Ho:  $\beta_{SMB} = 0.2$  and  $\beta_{HML} = 0.6$ . H1:  $\beta_{SMB} \neq 0.2$  and/or  $\beta_{HML} \neq 0.6$ .  $\Rightarrow J = 2$ 

We define **R** (2x4) below and write  $m = R\beta - q = 0$ :

$$\begin{bmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix} * \begin{bmatrix} \beta_1 \\ \beta_{Mkt} \\ \beta_{SMB} \\ \beta_{HML} \end{bmatrix} = \begin{bmatrix} 0.2 \\ 0.6 \end{bmatrix}$$

2. Test-statistic:  $\mathbf{F} = \mathbf{W}^*/J = (\mathbf{R}\mathbf{b} - \mathbf{q})' \{\mathbf{R}[s^2(\mathbf{X}'\mathbf{X})^{-1}]\mathbf{R}'\}^{-1}(\mathbf{R}\mathbf{b} - \mathbf{q})$ Distribution under H<sub>0</sub>:  $\mathbf{F} = W^*/2 \sim F_{J=2,T-4}$  (or asymptotic,  $2^*F \xrightarrow{d} \chi_2^2$ )

3. Get OLS results, compute F,  $\hat{F}$ .

4. <u>Decision Rule</u>:  $\alpha = 0.05$  level. We reject H<sub>0</sub> if p-value(#F) < .05. Or, reject H<sub>0</sub>, if  $\hat{F} > F_{J=2,T-4,.05}$ .

```
Var_b <- vcov(fit_ibm_ff3)</pre>
                                                        # Extract Var[b]
R <- matrix(c(0,0,0,0,1,0,0,1), nrow=2)
                                                        # matrix of restrictions
                                                       # hypothesized values
q \le c(.2, .6)
m <- R%*%b - q
                                                       \# m = Estimated R*Beta - q
Var m \le R \% \% Var b \% \% t(R)
                                                       # Variance of m
                                                       # check for non-singularity
det(Var m)
W <- t(m)%*%solve(Var m)%*%m
F t <- as.numeric(W/J
                                                       # F-test statistic
                               )
qf(.95, df1=J, df2=(T - k))
                                                       # exact distribution (F-dist) if errors normal
p val <-1 - pf(F t, df1=J, df2=(T - k))
                                                       # p-value(F_t) under errors normal
p_val
F t asym <-J*F
                                                        # Asymptotic F test (a Chi-square test)
qchisq(.95, df=J)
                                                       # asymptotic distribution (chi-square)
p val \leq 1 - pchisq(F t asym, df=J)
                                                       # p-value(F t) under asymptotic distribution
p val
>F t
49.217
>
> qf(.95, df1=J, df2=(T - k))
                                                        # exact distribution (F-dist) if errors normal
                                        F t > 3.011644 \Rightarrow reject H<sub>0</sub> at 5% level
[1] 3.011644
p_val <-1 - pf(F t, df1=J, df2=(T - k))
                                                       # p-value(F t) under errors normal
> p val
[1] < 2.2e-16 `
                                       \Rightarrow reject H<sub>0</sub> at 5% level.
>F t asym
98.433
>
> qchisq(.95, df=J)
                                                        # asymptotic distribution (chi-square)
                                       F t > 5.991465 \Rightarrow reject H<sub>0</sub> at 5% level
[1] 5.991465
> p val <- 1 - pchisq(F t asym, df=J)
                                                       # p-value(F t) under asymptotic distribution
> p val
[1] < 2.2e-16
                               \Rightarrow so low it is almost zero. Extremely low chance H<sub>0</sub> is true.
Conclusion: We reject the restrctions: \beta_{SMB} = 0.2 and \beta_{HML} = 0.6.
```

<u>R Note</u>: You can use the R package *car* to test linear restrictions (linear H<sub>0</sub>). install.packages("car") library(car) linearHypothesis(fit\_ibm\_ff3, c("SMB = 0.2","HML = 0.6"), test="F") # Exact F test

Linear hypothesis test

Hypothesis: SMB = 0.2HML = 0.6Model 1: restricted model Model 2: ibm  $x \sim Mkt RF + SMB + HML$ Res.Df RSS Df Sum of Sq F Pr(>F)1 567 2.2691 2 565 1.9324 2 0.33667 **49.217** < 2.2e-16 \*\*  $\Rightarrow$  reject H<sub>0</sub> at 5% level \_\_\_\_ Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1.¶ **Example:** Now, we do a joint test of the EH.  $H_0$ :  $\alpha = 0$  and  $\beta = 0$ . Using the previous program but with: J <- 2 # number of restriction  $R \le matrix(c(1,0,0,1), nrow=2)$ # matrix of restrictions q <- c(0,0)# hypothesized values  $> F_t$ 4.1024 >> qf(.95, df1=J, df2=(T - k))# exact distribution (F-dist) if errors normal [1] 3.020661 F  $t > 3.020661 \Rightarrow$  reject H<sub>0</sub> at 5% level p val <-1 - pf(F t, df1=J, df2=(T - k))# p-value(F t) under errors normal > p val [1] 0.01731  $\Rightarrow$  reject H<sub>0</sub> at 5% level.  $> F_t$  asym 8.2047 >> qchisq(.95, df=J) # asymptotic distribution (chi-square) [1] 5.991465 F t > 5.991465  $\Rightarrow$  reject H<sub>0</sub> at 5% level # p-value(F\_t) under asymptotic distribution > p val <-1 - pchisq(F t asym, df=J)> p val` [1] 0.01653  $\Rightarrow$  reject H<sub>0</sub> at 5% level.

The R package car can do the above too: > linearHypothesis(fit\_eh, c("(Intercept) = 0", "x = 0"), test="F") # "F": exact test, with F-distr

Linear hypothesis test Hypothesis: (Intercept) = 0x = 0 

 Model 1: restricted model

 Model 2:  $y \sim x$  

 Res.Df
 RSS Df Sum of Sq
 F
 Pr(>F)

 1
 363 0.27033

 2
 361 0.26432
 2 0.0060075
 4.1024
 0.01731 \*

 -- Signif. codes:
 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

 > qf(.95, df1=J, df2=(T - k))
 # exact distribution (F-dist) if errors normal

 [1] 3.020661
 F\_t > 3.020661  $\Rightarrow$  reject H<sub>0</sub> at 5% level

<u>Conclusion</u>: We reject the joint restrictions:  $H_0$ :  $\alpha = 0$  and  $\beta = 0$ .

### The F Test: $H_0$ : $R\beta - q = 0$

Approach (2). We know that imposing the restrictions leads to a loss of fit.  $R^2$  must go down. Does it go down a lot? –i.e., significantly?

Recall (i)  $e^* = y - Xb^* = e - X(b^* - b)$ (ii)  $b^* = b - (X'X)^{-1}R'[R(X'X)^{-1}R']^{-1}(Rb - q)$  $\Rightarrow e^{*'}e^* = e'e + (b^* - b)'X'X(b^* - b)$ 

Replacing  $(\mathbf{b}^* - \mathbf{b})$  from (ii) in the above formula, we get:

$$e^{*'}e^{*} - e'e = (Rb - q)'[R(X'X)^{-1}R']^{-1}(Rb - q)$$

<u>Note</u>:  $e^{*'}e^* - e^{'}e$  is a quadratic form, then we can use a lot of results to derive its asymptotic distribution

• Recall, the F-distribution is a ratio of two independent  $\chi_J^2$  and  $\chi_T^2$  RV divided by their degrees of freedom

$$F = \frac{\chi_J^2 / J}{\chi_T^2 / T} \sim F_{J,T}$$

Then, to get to the F-test, we rely on two results:

 $-W = (\mathbf{R}\mathbf{b} - \mathbf{q})' \{ \mathbf{R}[\sigma^2 (\mathbf{X}'\mathbf{X})^{-1}]\mathbf{R}' \}^{-1} (\mathbf{R}\mathbf{b} - \mathbf{q}) \sim \chi_J^2 \text{ (if } \sigma^2 \text{ is known)} \\ -\mathbf{e'}\mathbf{e}/\sigma^2 \sim \chi_{T-k}^2.$  $\implies F = \frac{(\mathbf{e}^{*'}\mathbf{e}^{*} - \mathbf{e'}\mathbf{e})/J}{[\mathbf{e'}\mathbf{e}/(T-k)]} \sim F_{J,T-k}$ 

• We can write the F-test in terms of  $R^2$ 's. Let  $R^2 =$ unrestricted model = 1 – RSS/TSS
$$R^{*2}$$
 = restricted model fit = 1 – RSS\*/TSS

Then, dividing and multiplying F by TSS we get

 $F = \frac{(1 - R^{*2}) - (1 - R^2)/J}{(1 - R^2)/(T - k)} \sim F_{J,T-k}$ 

or

$$F = \frac{(R^2 - R^{*2})/j}{(1 - R^2)/(T - k)} \sim F_{j,T-k}.$$

### The F Test: H<sub>0</sub>: F-test of Goodness of Fit

In the linear model, with a constant  $(\mathbf{X}_1 = \mathbf{i})$ :  $\mathbf{y} = \mathbf{X} \ \mathbf{\beta} + \mathbf{\varepsilon} = \mathbf{X}_1 \ \mathbf{\beta}_1 + \mathbf{X}_2 \ \mathbf{\beta}_2 + \dots + \mathbf{X}_k \ \mathbf{\beta}_k + \mathbf{\varepsilon}$ 

• We want to test if the slopes  $X_2, ..., X_k$  are equal to zero. That is,

*H*<sub>0</sub>:  $\beta_2 = \cdots = \beta_k = 0$ *H*<sub>1</sub>: at least one  $\beta \neq 0$   $\Rightarrow J = k - 1$ 

We can write H<sub>0</sub>: 
$$\mathbf{R}\boldsymbol{\beta} - \mathbf{q} = \mathbf{0}$$
  $\Rightarrow \begin{bmatrix} \mathbf{0} & \mathbf{1} & \dots & \mathbf{0} \\ \dots & \dots & \dots & \dots \\ \mathbf{0} & \mathbf{0} & \mathbf{0} & \mathbf{1} \end{bmatrix} \begin{bmatrix} \boldsymbol{\beta}_1 \\ \boldsymbol{\beta}_2 \\ \dots \\ \boldsymbol{\beta}_k \end{bmatrix} = \begin{bmatrix} \mathbf{0} \\ \dots \\ \mathbf{0} \end{bmatrix}$ 

• We have J = k - 1. Then,

$$F = \frac{(R^2 - R^{*2})/J}{(1 - R^2)/(T - k)} \sim F_{k-1,T-k}.$$

For the restricted model,  $R^{*2} = 0$ . Then,

$$F = \frac{R^2/(k-1)}{(1-R^2)/(T-k)} \sim F_{k-1,T-k}$$

Recall *ESS/TSS* is the definition of  $R^2$ . *RSS/TSS* is equal to  $(1 - R^2)$ .

$$F = \frac{R^2/(k-1)}{(1-R^2)/(T-k)} = \frac{\frac{ESS}{TSS}/(k-1)}{\frac{RSS}{TSS}/(T-k)} = \frac{ESS/(k-1)}{RSS/(T-k)}$$

This test statistic is called the *F*-test of goodness of fit. It is reported in all regression packages as part of the regression output. In R, the lm function reports it as "*F*-statistic."

**Example:** We want to test if all the FF factors (Market, SMB, HML) are jointly significant (J=3), using monthly data 1973 – 2020 (T=569). y <- ibm\_x T <- length(x) x0 <- matrix(1,T,1) x <- cbind(x0,Mkt\_RF, SMB, HML) > F\_goodfit [1] 96.58204  $\Rightarrow$  F\_goodfit > F<sub>3,565,.05</sub> = 2.62067  $\Rightarrow$  Reject H<sub>0</sub>. ¶

<u>Conclusion</u>: We strongly reject the restrctions:  $\beta_{Mtk} = \beta_{SMB} = \beta_{HML} = 0.$ 

### The F Test: General Case – Example

DCC

F

In the linear model

 $\mathbf{y} = \mathbf{X} \boldsymbol{\beta} + \boldsymbol{\varepsilon} = \boldsymbol{\beta}_1 + \mathbf{X}_2 \boldsymbol{\beta}_2 + \mathbf{X}_3 \boldsymbol{\beta}_3 + \mathbf{X}_4 \boldsymbol{\beta}_4 + \boldsymbol{\varepsilon}$ 

We want to test if the slopes  $X_3$ ,  $X_4$  are equal to zero. That is,

H<sub>0</sub>:  $\beta_3 = \beta_4 = 0$ H<sub>1</sub>:  $\beta_3 \neq 0$  or  $\beta_4 \neq 0$  or both  $\beta_3$  and  $\beta_4 \neq 0$ 

We can use,  $F = (e^{*'}e^* - e'e)/J / [e'e/(T-k)] \sim F_{J,T-K}$ .

DCC

Define  $\mathbf{y} = \mathbf{X} \ \mathbf{\beta} + \mathbf{\varepsilon} = \mathbf{\beta}_1 + \mathbf{X}_2 \ \mathbf{\beta}_2 + \mathbf{\varepsilon}$  (Restricted RSS = RSS<sub>R</sub>, with  $k_R$  parameters)  $\mathbf{y} = \mathbf{\beta}_1 + \mathbf{X}_2 \ \mathbf{\beta}_2 + \mathbf{X}_3 \ \mathbf{\beta}_3 + \mathbf{X}_4 \ \mathbf{\beta}_4 + \mathbf{\varepsilon}$  (Unrestricted RSS = RSS<sub>U</sub>, with  $k_U$ parameters)

Then,

$$=\frac{\frac{KSS_R - KSS_U}{(k_U - k_R)}}{\frac{RSS_U}{(T - k_U)}} \sim F_{J,T-k}, \text{ where } J = (k_U - k_R), \text{ and } T - k = T - k_U.$$

### The F Test: Are SMB and HML Priced Factors?

**Example:** We want to test if the additional FF factors (SMB, HML) are significant, using monthly data 1973 – 2020 (T=569). That is, we test H<sub>0</sub>:  $\beta_{SMB} = \beta_{HML} = 0$ .

Unrestricted Model (Fama-French 3-factor model): (U)  $(r_{IBM,t} - r_f) = \beta_0 + \beta_1 (r_{m,t} - r_f) + \beta_2 SMB_t + \beta_3 HML_t + \varepsilon_t$ 

Hypothesis: H<sub>0</sub>:  $\beta_2 = \beta_3 = 0$ H<sub>1</sub>:  $\beta_2 \neq 0$  and/or  $\beta_3 \neq 0$ 

Then, the Restricted Model (CAPM): (R)  $(r_{IBM,t} - r_f) = \beta_0 + \beta_1 (r_{m,t} - r_f) + \varepsilon$ 

Test: 
$$F = \frac{(RSS_R - RSS_U)/J}{RSS_U/(T - k_u)} \sim F_{J,T-k}$$
, where  $J = (k_U - k_R) = 4 - 2 = 2$ 

• The unrestricted model was already estimated in Lecture 3. For the restricted model:

```
y \le ibm x
x_0 < -matrix(1,T,1)
x r \leq cbind(x0,Mkt RF)
                                                          # Restricted X vector
k \leq -ncol(x)
T \leq nrow(x)
k2 \leq ncol(x)
                                                          # Restricted OLS regression
b2 \le solve(t(x r)\%*\% x r)\%*\% t(x r)\%*\%
e2 \le y - x r\% * b2
RSS2 \leq as.numeric(t(e2)\%*\%e2)
> RSS = 1.932442
                                                          # RSSU
> RSS2 = 1.964844
                                                          # RSS<sub>R</sub>
> J < -k - k2
                                                          # J = degrees of freedom of numerator
> F test <- ((RSS2 - RSS)/J)/(RSS/(T-k))
>F test
[1] 4.736834
> qf(.95, df1=J, df2=(T-k))
                                                          # F<sub>2,565,.05</sub> value (\approx 3)
[1] 3.011672
                                                  \Rightarrow Reject H<sub>0</sub>.
> p val <- 1 - pf(F test, df1=J, df2=(T-k))
                                                          # p-value of F test
> p val
[1] 0.009117494
                                                          \Rightarrow p-value is small \Rightarrow Reject H<sub>0</sub>.
```

<u>Conclusion</u>: We strongly reject the restrctions:  $\beta_{SMB} = \beta_{HML} = 0.$ 

<u>R Note</u>: There is package in R, *Intest*, that performs this test, *waldtest*, (and many others, used in this class). You need to install it first: install.packages("Intest"). For the *waldtest*, the default reports the *F-test* with the F distribution.

<u>Remark</u>: The models need to be nested.

**Example:** We test if the additional FF factors (SMB, HML) are significant, using monthly data 1973 – 2020 (T=569).

library(lmtest)
fit\_ibm\_ff3 <- lm (y ~ Mkt\_RF + SMB + HML)
fit\_ibm\_capm <- lm (y ~ Mkt\_RF)
waldtest(fit\_ibm\_ff3, fit\_ibm\_capm)
Wald test</pre>

# Unrestricted Model # Restricted Model Model 1:  $y \sim Mkt_RF + SMB + HML$ Model 2:  $y \sim Mkt_RF$ Res.Df Df F Pr(>F) 1 565 2 567 -2 4.7368 0.009117 \*\*  $\Rightarrow$  p-value is small: Reject H<sub>0</sub>.

# Trilogy of Asymptotic Tests: LR, Wald, and LM

In practice, so far, to test joint hypothesis H<sub>0</sub>:  $\mathbf{R}\boldsymbol{\beta} = \mathbf{q}$ , we have relied on the asymptotic distribution of the Wald test. We constructed the Wald test based on the unrestricted estimation (OLS), which in the produces **b** and  $s^2$ .

Then, the Wald test:

 $W^* = (\mathbf{R}\mathbf{b} - \mathbf{q})' \{ \mathbf{R}[s^2(\mathbf{X}'\mathbf{X})^{-1}]\mathbf{R} \}^{-1} (\mathbf{R}\mathbf{b} - \mathbf{q}) \xrightarrow{d} \chi_J^2.$ 

In more general terms, we construct the Wald test based on the unrestricted estimation, which produces  $\hat{\theta}^{U}$ .

There are two other popular tests that are asymptotically equivalent -i.e., they have the same asymptotic distribution: the Likelihood Ratio (LR) and the Lagrange Multiplier (LM) tests.

• The LR test is based on the (log) Likelihood. It needs two ML estimations: the unrestricted estimation, producing  $\hat{\theta}_{ML}$ , and the restricted estimation, producing  $\hat{\theta}^R$ . Below we define the LR test:

$$LR = 2[\log(L(\hat{\theta}_{ML})) - \log(L(\hat{\theta}^R))] \xrightarrow{d} \chi_J^2$$

Note: MLE requires assuming a distribution, usually, a normal.

<u>Technical note</u>: The LR test is a *consistent test*. An asymptotic test which rejects  $H_0$  with probability one when the  $H_1$  is true is called a *consistent test*. That is, a consistent test has asymptotic power of 1. The LR test is a consistent test.

**Example:** We use a likelihood ratio test to check if the additional FF factors (SMB, HML) are significant, using monthly data 1973 – 2020 (T=569).

library(Imtest)
fit\_ibm\_ff3 <- lm (ibm\_x ~ Mkt\_RF + SMB + HML)
fit\_ibm\_capm <- lm (ibm\_x ~ Mkt\_RF)
lrtest(fit\_ibm\_ff3, fit\_ibm\_capm)
Likelihood ratio test
Model 1: ibm\_x ~ Mkt\_RF + SMB + HML
Model 2: ibm\_x ~ Mkt\_RF
#Df LogLik Df Chisq Pr(>Chisq)

# 1 5 810.03 2 3 805.30 -2 9.4616 0.008819 \*\* $\Rightarrow$ p-value is small: Reject H<sub>0</sub>. ¶

• The LM test needs only one estimation: the restricted estimation, producing  $\hat{\theta}^R$ . If the restriction is true, then the slope of the objective function (say, the Likelihood) at  $\hat{\theta}^R$  should be zero. The slope is called the Score,  $S(\hat{\theta}^R)$ . The LM test is based on a Wald test on  $S(\hat{\theta}^R) = 0$ .

$$LM = S(\hat{\theta}^R)' [Var(S(\hat{\theta}^R)]^{-1}S(\hat{\theta}^R) \xrightarrow{d} \chi_J^2$$

It turns out that there is a much simpler formulation for the LM test, based on the residuals of the restricted model. We will present this version of the test in Lecture 6.



If the likelihood function were quadratic then LR = LM = W. In general, however, W > LR > LM.

### **Testing Remarks: Pre-testing**

A special case of omitted variables.

 $\mathbf{v} =$ 

- First, a researcher starts with an unrestricted model (U):

$$X$$
β + ε. (U)

- Then, based on ("preliminary") tests –say, an *F*-test- a researcher decides to use restricted estimator,  $\mathbf{b}^*$ . That is,

 $\mathbf{y} = \mathbf{X}\mathbf{\beta} + \mathbf{\epsilon}.$  s.t.  $\mathbf{R}\mathbf{\beta} = \mathbf{q}$  (R)

- We can think of the estimator we get from estimating R as:

 $\mathbf{b}_{\text{PT}} = \mathbf{I}_{\{0, c\}}(F) \ \mathbf{b}^* + \mathbf{I}_{\{c, \infty\}}(F) \ \mathbf{b},$ 

where  $I_{\{0,c\}}$  is an indicator function:

 $I_{\{c,\infty\}}(F) = 0$ , if *F*-stat in *R* (rejection region) –say, F > c,

 $I_{\{0, c\}}(F) = 1$  if *F*-stat in  $\mathbb{R}^C$ -say, F < c.

*c* : critical value chosen for testing H<sub>0</sub>:  $\mathbf{R}\boldsymbol{\beta} = \mathbf{q}$ , using the *F*-stat.

• The *pre-test estimator* is a rule, which chooses between the restricted estimator, **b**\*, or the OLS estimator, **b**:

 $\mathbf{b}_{PT} = \mathbf{I}_{\{0, c\}}(F) \ \mathbf{b}^* + \mathbf{I}_{\{c, \infty\}}(F) \ \mathbf{b},$ where  $\mathbf{b}^* = \mathbf{b} - (\mathbf{X}'\mathbf{X})^{-1}\mathbf{R}'[\mathbf{R}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{R}']^{-1}(\mathbf{R}\mathbf{b} - \mathbf{q})$ 

• Two "negative" situations:

(1) H<sub>0</sub>:  $\mathbf{R}\boldsymbol{\beta} = \mathbf{q}$  is true. The *F*-test will incorrectly reject H<sub>0</sub>  $\alpha$ % of the time. That is, in  $\alpha$ % of the repeated samples, we have "irrelevant variables"

 $\Rightarrow$  OLS **b**: No bias, but inefficient estimator.

(2) H<sub>0</sub>:  $\mathbf{R}\boldsymbol{\beta} = \mathbf{q}$  is false. The *F*-test will correctly reject H<sub>0</sub> a % of times equal to the power  $\pi$  of the test. That is,  $(100 - \pi)$ % of the time,  $\mathbf{R}\boldsymbol{\beta}=\mathbf{q}$  will be incorrectly imposed, we have "omitted variables:"

 $\Rightarrow$  OLS **b**\*: bias, but small variance!

The failure of the OLS estimator to have the properties under correct specification is called *pretest bias*.

Pre-testing (also called *sequential estimation*, *data mining*) is common in practice. In general, it is ignored –and not even acknowledged.

Main argument to ignore pre-testing: We need some assumptions to decide which variables are included in a model. Is the probability that pre-testing yields an incorrect set of X greater than the probability of selecting the "correct" assumption?

David Hendry, a well known thinker of these methodological issues, does not see pre-testing in the discovery stage as a problem. For him, pre-testing at that stage is part of the *process of discovery*.

<u>Practical advise</u>: Be aware of the problem. Do not rely solely on stats to select a model –use economic theory as well.

• Do not use same sample evidence to generate an H<sub>0</sub> and to test it!

**Example**: The Fama-French factors have been "discovered" using the CRSP/Compustast database for a long, long time. Thus, testing the Fama-French factors using the CRSP/Compustat is not advisable!

(You can test them with another dataset, for example, get international data.) ¶

### Testing Remarks: Significance level, a

So far, we have assumed that the distribution of the test statistic –say the *F*-statistic– under  $H_0$  is known exactly, so that we have what is called an *exact test*.

Technically, the *size of a test* is the supremum of the rejection probability over all DGPs that satisfy H<sub>0</sub>. For an exact test, the size equals the *nominal level*,  $\alpha$  –i.e., the Prob[Type I error] =  $\alpha$ .

Usually, the distribution of a test is known only approximately *(asymptotically)*. In this case, we need to draw a distinction between the nominal level,  $\alpha$  (*nominal size*), of the test & the actual *rejection probability (empirical size)*, which may differ greatly from the nominal level.

Simulations are needed to gauge the empirical size of tests.

# Testing Remarks: A word about a

Ronald Fisher, before computers, tabulated distributions. He used a .10, .05, and .01 percentiles. These tables were easy to use and, thus, those percentile became the de-facto standard  $\alpha$  for testing H<sub>0</sub>.

"It is usual and convenient for experimenters to take 5% as a standard level of significance." – Fisher (1934).

Given that computers are powerful and common, why is p = 0.051 unacceptable, but p = 0.049 is great? There is no published work that provides a theoretical basis for the standard thresholds.

Rosnow and Rosenthal (1989): " ... surely God loves .06 nearly as much as .05."

<u>Practical advise</u>: In the usual Fisher's null hypothesis (significance) testing, significance levels,  $\alpha$ , are arbitrary. Make sure you pick one, say 5%, and stick to it throughout your analysis or paper.

• Report *p*-values, along with CI's. Search for economic significance.

Questions: .10, .05, or .01 significance?

Many tables will show \*, \*\*, and \*\*\* to show .10, .05, and .01 significance levels –for example, lm() in R. Throughout the paper, the authors will point out the different significance levels. In these papers, it is not clear what  $\alpha$  is the paper using for inference.

We can think of these stars (or *p*-values) as ways of giving weights to H<sub>0</sub> relative to H<sub>1</sub>.

# Testing Remarks: A word about H<sub>0</sub>

In applied work, we only learn when we reject H<sub>0</sub>; say, when the *p*-value  $< \alpha$ . But, rejections are of two types:

- Correct ones, driven by the power of the test

- Incorrect ones, driven by Type I Error ("statistical accident," luck).

It is important to realize that, however small the *p*-value, there is always a finite chance that the result is a pure accident. At the 5% level, there is 1 in 20 chances that the rejection of  $H_0$  is just luck.

Since negative results are difficult to publish (*publication bias*), there is an unknown but possibly large number of false claims taken as truths.

**Example:** If  $\alpha = 0.05$ , proportion of false H<sub>0</sub>=10%, and  $\pi = .50$ , **47.4%** of rejections are true H<sub>0</sub> -i.e., "false positives." ¶

### **Testing Remarks: Mass significance**

We have a model. We perform k different tests, say k t-tests, each with a nominal significance level of  $\alpha$ :

 $\alpha$  = Prob (Rejecting for a given test |H<sub>0</sub> for this test is true)

The *overall significance* of the test procedure is, however, given by  $\alpha^* = \text{Prob}$  (Rejecting at least one test | all H<sub>0</sub> are true).

The probability of rejecting at least one  $H_0$  is obviously greater than of rejecting a specific test. This is the problem of *mass significance*.

• Two cases (1) Independent tests  $\alpha^* = 1 - (1 - \alpha)^k$  &  $\alpha = 1 - (1 - \alpha^*)^{1/k}$ 

(2) Dependent tests:  $\alpha^* \le k\alpha$  &  $\alpha \ge \alpha^*/k$  $\Rightarrow$  close to the "independent" values for small  $\alpha$ , but can differ for large  $\alpha$ .

Example:	$\alpha = 0.05$ and $k = 5$	$\Rightarrow \alpha^*(\text{Indep}) = .23 \& \alpha^*(\text{Dep}) = .25$
	$\alpha = 0.05 \text{ and } k = 20$	$\Rightarrow \alpha^*(\text{Indep}) = .64 \& \alpha^*(\text{Dep}) = 1$
	$\alpha^* = 0.05 \text{ and } k = 5$	$\Rightarrow \alpha(\text{Indep}) = .0102 \& \alpha(\text{Dep}) = .01$
	$\alpha^* = 0.20 \text{ and } k = 5$	$\Rightarrow \alpha(\text{Indep}) = .044 \& \alpha(\text{Dep}) = .04$
	$\alpha^* = 0.20$ and $k = 20$	$\Rightarrow \alpha(\text{Indep}) = .011 \& \alpha(\text{Dep}) = .01. \P$

• David Hendry's suggestions:

In repeated *parametric testing* (overall level 5%):

- Only accept variables as important when their *p*-values are less than 0.001, preferably smaller

- Maybe look for other ways of choosing variables, say AIC.

In repeated *diagnostic testing* (overall level 20%), we should only accept there is no misspecification if

- All *p*-values are greater than 0.05, or

- Most *p*-values are greater than 0.10 with a few in the range 0.02 to 0.10

### **Non-nested Models and Tests**

So far, all our tests (t-, F- & Wald tests) have been based on *nested models*, where the R model is a restricted version of the U model.

### Example:

Model U	$y = X\beta + W\delta + \varepsilon$	(Unrestricted)
Model R	$y = X\beta + \xi$	(Restricted)

Model U becomes Model R under H<sub>0</sub>:  $\delta = 0$ . We know how to select a model, based on a statistical test, for example, using a Wald test. ¶

• Sometimes, we have two rival models to choose between, where neither can be nested within the other -i.e., neither is a restricted version of the other. In this situation, we call the models *non-nested*.

### **Example:**

Model 1	$y = X\beta + W\delta + \varepsilon$	
Model 2	$y = X\beta + Z\gamma + \xi.$	ſ

If the dependent variable is the same in both models (as is the case here), we can simply use Adjusted- $R^2$  to rank the models and select the one with the largest Adjusted- $R^2$ .

We can also use AIC and/or BIC.

But, we can also use more sophisticated testing-based methods: Encompassing test and J-test.

# **Non-nested Models: Encompassing Test**

Alternative approach: Encompassing

(1) Form a composite or *encompassing* model that nests both rival models –say, Model 1 and Model 2. This is the unrestricted Model (ME).

(2) Test the relevant restrictions of each rival model against ME. We do two F-tests, where the restricted models are Model 1 and Model 2.

If we reject the restrictions against one Model, say Model 1, and we cannot reject the restrictions against the other, Model 2, we are done: We select the Model that the F test do not reject restrictions (Model 2).

Assuming the restrictions cannot be rejected, we prefer the model with the lower F statistic for the test of restrictions.

Note: We test a hybrid model. Also, multicollinearity may appear.

Example: We have:Model 1 $Y = X\beta + W\delta + \varepsilon$ Model 2 $Y = X\beta + Z\gamma + \xi$ Then, the Encompassing Model (ME) is:ME: $Y = X\beta + W\delta + Z\gamma + \varepsilon$ 

Now test, separately, the hypotheses (1)  $\delta = 0$  and (2)  $\gamma = 0$ . That is,

F-test for H<sub>0</sub>:  $\gamma = 0$ : **ME** (U Model) vs Model 1 (R Model). F-test for H<sub>0</sub>:  $\delta = 0$ : **ME** (U Model) vs Model 2 (R Model).

If we reject H<sub>0</sub>:  $\gamma = \mathbf{0} \Rightarrow$  Evidence against Model 1 (statistically different from ME). If we reject H<sub>0</sub>:  $\delta = \mathbf{0} \Rightarrow$  Evidence against Model 2 (statistically different from ME).

### Non-nested Models and Tests: IFE or PPP?

Two of the main theories to explain the behaviour of exchange rates,  $S_t$ , are the International Fisher Effect (IFE) and the Purchasing Power Parity (PPP). We use the direct notation for  $S_t$ , that is, units of domestic currency (DC) per 1 unit of foreign currency (FC).

**IFE** states that, in equilibrium, changes in exchange rates (e) are driven by the interest rates differential between the domestic currency, id, and the foreign currency, if:. A DGP consistent with IFE is:

$$\mathbf{e} = \alpha^1 + \beta^1 \left( \mathbf{i}_d - \mathbf{i}_f \right) + \varepsilon^1$$

**PPP**, in its Relative version, states that that, in equilibrium, e are driven by the inflation rates differential between the domestic Inflation rate, I<sub>d</sub>, and the foreign Inflation rate, I<sub>f</sub>. A GDP consistent with IFE is:

$$e = \alpha^2 + \beta^2 (I_d - I_f) + \epsilon^2$$

Both theories are non-nested, thus, we need a non-nested method to select a model.

**Example**: What drives log changes in exchange rates for the USD/GBP (e):  $(i_d - i_f)$  or  $(I_d - I_f)$ ? The USD is the DC; the GBP is the FC. Both non-nested models are:

 $\begin{array}{ll} \textbf{IFE Model:} & e = \alpha^1 + \beta^1 \ (\textbf{i}_d - \textbf{i}_f) + \epsilon^1 \\ \textbf{PPP Model:} & e = \alpha^2 + \beta^2 \ (\textbf{I}_d - \textbf{I}_f) + \epsilon^2 \end{array}$ 

SF\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/SpFor\_prices.csv", head=TRUE, sep=",")
x\_date <- SF\_da\$Date
x\_S <- SF\_da\$GBPSP
x\_F3m <- SF\_da\$GBP3M
i\_us3 <- SF\_da\$Dep\_USD3M
i\_uk3 <- SF\_da\$Dep\_UKP3M
cpi\_uk <- SF\_da\$UE\_CPI
cpi\_us <- SF\_da\$US\_CPI
T <- length(x\_S)
int\_dif <- (i\_us3[-1] - i\_uk3[-1])/100
lr\_usdgbp <- log(x\_S[-1]/x\_S[-T])
I\_us <- log(cpi\_us[-1]/cpi\_us[-T])
I\_uk <- log(cpi\_uk[-1]/cpi\_uk[-T])
inf\_dif <- (I\_us - I\_uk)</pre> Encompassing Model (ME or "Unrestricted Model")  $\mathbf{e} = \alpha + \beta_1 \left( \mathbf{i_d} - \mathbf{i_f} \right) + \beta_2 \left( \mathbf{I_d} - \mathbf{I_f} \right) + \varepsilon^1$ # Encompassing Model and Test **fit me** <- lm(lr usdgbp ~ int dif + inf dif) # ME estimation > summary(**fit me**) Coefficients: Std. Error t value Pr(>|t|)Estimate (Intercept) -0.0009633 0.0016210 -0.594 0.5527 int dif -0.0278510 0.0741189 -0.376 0.7073  $\Rightarrow$  cannot reject H<sub>0</sub>:  $\beta_1 = 0$ . inf dif 0.7444711 0.3429106 2.171 0.0306 \*  $\Rightarrow$  reject H<sub>0</sub>:  $\beta_2 = 0$ . ---Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 0.02662 on 360 degrees of freedom Multiple R-squared: 0.01316, Adjusted R-squared: 0.007673 F-statistic: 2.399 on 2 and 360 DF, p-value: 0.09221

Conclusion: The encompasing test favors the PPP Model.

Note: Two F-tests are needed, but for the one variable case, the t-tests are equivalent.

<u>R Note</u>: The package in R, *Intest*, performs this test, *encomptest*. Recall you need to install it first: install.packages("Intest"). The test reported is an *F*-test ~  $F_{1,T-k}$ , which, in this case with only one variable in each Model, is equal to  $(t_{T-k})^2$ .

library(lmtest) fit_m1 <- lm(lr_usdgbp ~ int_dif) fit_m2 <- lm(lr_usdgbp ~ inf_dif) > encomptest(fit_m1, fit_m2)		# Restricted Model 1 # Restricted Model 2
1: lr_usdgbp ~ int_dif Model 2: lr_usdgbp ~ inf_dif Model E: lr_usdgbp ~ int_dif + inf_dif		
Res.Df Df F Pr(>F) M1 vs. ME 360 -1 <b>4.7134 0.03058</b> * M2 vs. ME 360 -1 0.1412 0.70732.	¶	⇒ reject H <sub>0</sub> : $β_2 = 0$ . Check: $(2.171)^2 = 4.713$

### Non-nested Models: J-test

We present the most popular test for non-nested models, the Davidson-MacKinnon (1981)'s *J*-test.

We start with two non-nested models. Say,

Model 1: $Y = X\beta + \varepsilon$ Model 2: $Y = Z\gamma + \xi$ 

<u>Idea</u>: If Model 2 is true, then the fitted values from the Model 1, when added to the 2nd equation, should be insignificant.

• Steps:

- (1) Estimate Model 1  $\Rightarrow$  obtain fitted values: **Xb**.
- (2) Add **Xb** to the list of regressors in Model 2
  - $\Rightarrow \mathbf{Y} = \mathbf{Z}\boldsymbol{\gamma} + \boldsymbol{\lambda}\mathbf{X}\mathbf{b} + \boldsymbol{\xi}$
- (3) Do a *t-test* on  $\lambda$ . A significant *t*-value would be evidence against Model 2, favoring Model 1. (4) Repeat the procedure for the models the other way round.
  - (4.1) Estimate Model 2  $\Rightarrow$  obtain fitted values: Zc.
    - (4.2) Add **Zc** to the list of regressors in Model 1:

$$\Rightarrow \mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\lambda} \mathbf{Z}\mathbf{c} + \boldsymbol{\varepsilon}$$

(4.3) Do a *t-test* on  $\lambda$ . A significant *t*-value would be evidence against Model 1 and in favor of Model 2.

(5) Rank the models on the basis of this test.

• The "best situation" is when we reject only one  $H_0$ :  $\lambda = 0$ . In this case, it is very clear which model to select. But, tt is possible that we cannot reject both models. This is possible in small samples, even if one model, say Model 2, is true.

It is also possible that both *t-tests* reject H<sub>0</sub> ( $\lambda \neq 0 \& \lambda \neq 0$ ). This is not unusual. McAleer's (1995), in a survey, reports that out of 120 applications all models were rejected 43 times.

<u>Technical Note</u>: As some of the regressors in step (3) are stochastic, Davidson and MacKinnon (1981) show that the *t-test* is *asymptotically* valid.

• One would also want to examine the diagnostic test results when choosing between two models.

### Non-nested Models: J-test – IFE or PPP?

**Example**: Now, we test IFE Model vs PPP Model 2, for changes in the USD/GBP exchange rate using the *J*-test.

Model 1 (IFE):	$\mathbf{e} = \boldsymbol{\alpha}^1 + \boldsymbol{\beta}^1 \left( \mathbf{i}_d - \mathbf{i}_f \right) + \boldsymbol{\varepsilon}^1$
Model 2 (PPP):	$\mathbf{e} = \alpha^2 + \beta^2 \left( \mathbf{I}_d - \mathbf{I}_f \right) + \epsilon^2$

```
y <- lr_usdgbp

fit_m1 <- lm( y ~ int_dif)

summary(fit_m1)

y_hat1 <- fitted(fit_m1)

fit_J1 <- lm( y ~ inf_dif + y_hat1)

summary(fit_J1)
```

```
fit_m2 <- lm( y ~ inf_dif)
summary(fit_m2)
y_hat2 <- fitted(fit_m2)
fit_J2 <- lm( y ~ int_dif + y_hat2)</pre>
```

summary(fit J2) fit m1 <- lm(  $y \sim int dif$ ) y hat1 <- fitted(fit m1) fit J1 <-  $lm(formula = y \sim inf dif + y hat1)$ > summary(fit J1) **Residuals:** 10 Median 30 Min Max -0.136310 -0.014168 0.000351 0.017227 0.092421 Coefficients: Estimate Std. Error t value Pr(>|t|)(Intercept) 0.0001497 0.0025556 0.059 0.9533 0.7444711 0.3429106 2.171 0.0306 \* inf dif y hat1 1.2853298 3.4206106 **0.376** 0.7073  $\Rightarrow$  cannot reject H<sub>0</sub>:  $\lambda$ =0. (Good for Model 2) Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' '1 Residual standard error: 0.02662 on 360 degrees of freedom Multiple R-squared: 0.01316, Adjusted R-squared: 0.007673 F-statistic: 2.399 on 2 and 360 DF, p-value: 0.09221 it m2 <- lm(  $y \sim inf dif$ ) y hat2 <- fitted(fit m2) fit  $J2 \le lm(formula = y \sim int dif + y hat2)$ > summary(fit J2) **Residuals**: Min 1Q Median 3Q Max -0.136310 -0.014168 0.000351 0.017227 0.092421 Coefficients: Estimate Std. Error t value Pr(>|t|)0.8529 (Intercept) -0.000304 0.0016409 -0.186 int dif -0.027851 0.0741189 -0.376 0.7073 y hat2 1.0066945 0.4636932 2.171 0.0306 \*  $\Rightarrow$  Reject H<sub>0</sub>:  $\lambda$ =0. (Again, good for Model 2) Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 0.02662 on 360 degrees of freedom Multiple R-squared: 0.01316, Adjusted R-squared: 0.007673

Conclusion: We only rejected one H<sub>0:  $\lambda = 0$ . Then, the J-test selects the PPP Model (Model 2).</sub>

F-statistic: 2.399 on 2 and 360 DF, p-value: 0.09221

<u>R Note</u>: The *lmtest* package also performs this test, with the function *jtest*. Recall that you need to install it first: install.packages("lmtest").

library(Imtest) fit\_m1 <- lm(lr\_usdgbp ~ int\_dif) fit\_m2 <- lm(lr\_usdgbp ~ inf\_dif) > jtest(fit\_m1, fit\_m2) J test Model 1: lr\_usdgbp ~ int\_dif Model 2: lr\_usdgbp ~ inf\_dif Estimate Std. Error t value Pr(>|t|)M1 + fitted(M2) 1.0067 0.4637 2.1710 0.03058 \*  $\Rightarrow$  Reject H<sub>0</sub>:  $\lambda=0$ . (Model 2 selected) M2 + fitted(M1) 1.2853 3.4206 0.3758 0.70732. ¶

### Non-nested Models: J-test – Application

We want to test				
<b>H</b> <sub>0</sub> : $y = X\beta + \varepsilon_0$ <b>H</b> <sub>1</sub> : $\ln y = (\ln X) \gamma + \varepsilon_1$		(additive) vs		
		(multiplicative)		
We look at the <i>J</i> -test				
- Step 1: OLS on H	1: get $\hat{\gamma}$			
OLS y = X	$(\beta + \lambda_1 \exp{\ln(X) \hat{\gamma})}$	$\mathbf{x} + \{ \hat{\mathbf{y}} \} + \mathbf{\epsilon}$		$\Rightarrow$ <i>t</i> - <i>test</i> on $\lambda_1$
- Step 2: OLS on H	<b>o</b> : get <b>b</b>			
OLS ln y =	$= (\ln X) \gamma + \lambda_0 X \mathbf{b} -$	3 ⊣		$\Rightarrow$ <i>t</i> - <i>test</i> on $\lambda_0$
Situations:				
(1) Both OK: $\lambda_1$	$= 0$ and $\lambda_0 = 0$	$\Rightarrow$ get	t more data	
(2) Only 1 is OK: $\lambda_1 \neq 0$ and $\lambda_0 = 0$		(multi	plicative is	OK);
$\lambda_0$	$\neq 0$ and $\lambda_1 = 0$	(addit	ive is OK)	
(3) Both rejected: $\lambda_1 \neq 0$ and $\lambda_0 \neq 0$		$\Rightarrow$ nev	$\Rightarrow$ new model is needed.	

### Non-nested Models: J-test - Remarks

The *J*-test was designed to test non-nested models (one model is the true model, the other is the false model), not for choosing competing models –the usual use of the test.

The *J*-test is likely to over reject the true (model) hypothesis when one or more of the following features is present:

i) A poor fit of the true model

ii) A low/moderate correlation between the regressors of the 2 models

iii) The false model includes more regressors than the correct model.

Davidson and MacKinnon (2004) state that the *J*-test will over-reject, *often quite severely* in finite samples when the sample size is small or where conditions (i) or (iii) above are obtained.

# Lecture 6 – Specification, Forecasting & Model Selection

# **OLS Estimation - Assumptions**

Brief Review of CLM Assumptions (A1) DGP:  $\mathbf{y} = \mathbf{X} \ \beta + \boldsymbol{\varepsilon}$  is correctly specified. (A2)  $\mathrm{E}[\boldsymbol{\varepsilon}|\mathbf{X}] = 0$ (A3)  $\mathrm{Var}[\boldsymbol{\varepsilon}|\mathbf{X}] = \sigma^2 \mathbf{I}_{\mathrm{T}}$ (A4) X has full column rank -rank(X)=k-, where T  $\geq k$ .

Question: What happens when (A1) is not correctly specified?

First, we look at (A1), in the context of linearity. Are we omitting a relevant regressor? Are we including an irrelevant variable? What happens when we impose restrictions in the DGP?

Second, in (A1), we allow some non-linearities in its functional form.

# **Specification Errors: Omitted Variables**

Omitting relevant variables: Suppose the correct model (DGP) is  $\mathbf{y} = \mathbf{X}_1 \mathbf{\beta}_1 + \mathbf{X}_2 \mathbf{\beta}_2 + \mathbf{\epsilon}$  -the "long regression," with  $\mathbf{X}_1 \& \mathbf{X}_2$ .

But, we compute OLS omitting  $X_2$ . That is,

 $\mathbf{y} = \mathbf{X}_1 \boldsymbol{\beta}_1 + \boldsymbol{\epsilon}$  —the "short regression."

We have two *nested* models: one model becomes the other, once a restriction is imposed. In the above case, the true model becomes "the short regression" by imposing the restriction  $\beta_2 = 0$ .

Question: What are the implications of using the wrong model, with omitted variables? We already know the answer, we are imposing a wrong restriction: the restricted estimator,  $\mathbf{b}^*$ , is biased, but it is more efficient.

# **Specification Errors: Omitted Variables**

Some easily proved results:  $E[\mathbf{b}_1|\mathbf{X}] = E[(\mathbf{X}_1'\mathbf{X}_1)^{-1}\mathbf{X}_1' \mathbf{y}] = E[(\mathbf{X}_1'\mathbf{X}_1)^{-1}\mathbf{X}_1' (\mathbf{X}_1\beta_1 + \mathbf{X}_2\beta_2 + \boldsymbol{\epsilon})]$   $= \beta_1 + (\mathbf{X}_1'\mathbf{X}_1)^{-1}\mathbf{X}_1'\mathbf{X}_2\beta_2 \neq \beta_1.$ 

Thus, unless  $X_1'X_2 = 0$ ,  $b_1$  is *biased*. The bias can be **huge**. It can reverse the sign of a price coefficient in a "demand equation."

(2)  $\operatorname{Var}[\mathbf{b}_1|\mathbf{X}] \leq \operatorname{Var}[\mathbf{b}_{1,2}|\mathbf{X}]$ , where  $\mathbf{b}_{1,2}$  is the OLS estimator of  $\boldsymbol{\beta}_1$  in the long regression (the true model).

Thus, we get a smaller variance when we omit  $X_2$ .

<u>Interpretation</u>: Omitting  $X_2$  amounts to using extra information –i.e.,  $\beta_2 = 0$ . Even if the information is wrong, it reduces the variance.

(3) Mean Squared Error (MSE =  $\frac{RSS}{T}$ )

If we use MSE as precision criteria for selecting an estimator, **b**<sub>1</sub> may be more "precise." Precision = Mean squared error (MSE) = Variance + Squared bias.

Smaller variance but positive bias. If bias is small, a practitioner may still favor the short regression.

<u>Note</u>: Suppose  $X_1'X_2 = 0$ . Then the bias goes away. Interpretation, the information is not "right," it is irrelevant: **b**<sub>1</sub> is the same as **b**<sub>1.2</sub>.

**Example:** We fit an ad-hoc model for U.S. short-term interest rates  $(i_{US,t})$  that includes inflation rate  $(I_{US,t})$ , changes in the USD/EUR  $(e_t)$ , money growth rate  $(m_{US,t})$ , and unemployment  $(u_{US,t})$ , using monthly data from 1975:Jan-2020: Jul. That is,

$$i_{US,t} = \beta_0 + \beta_1 I_{US,t} + \beta_2 e_t + \beta_3 m_{US,t} + \beta_4 u_{US,t} + \varepsilon_i$$

Fger da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/FX USA GER.csv", head=TRUE, sep=",") us CPI <- Fger da\$US CPI us M1 <- Fger da\$US M1 us i <- Fger da\$US I3M us GDP <- Fger da\$US GDP ger CPI <- Fger da\$GER CPI us u <- Fger da\$US UN S ger <- Fger da\$USD EUR  $T \leq - length(us CPI)$ us I <- log(us CPI[-1]/us CPI[-T]) # US Inflation: (Log) Changes in CPI us mg  $<-\log(us M1[-1]/us M1[-T])$ # US Money Growth: (Log) Changes in M1 # (Log) Changes in USD/EUR e ger  $<-\log(S \text{ ger}[-1]/S \text{ ger}[-T])$ us i  $1 \le us$  i[-1] # Adjust sample size of untransformed data us u  $1 \le us u[-1]$ # Adjust sample size of untransformed data us i  $0 \le us$  i[-T] # lagged interest rates, by removing T observation xx i <- cbind(us I, e ger, us mg, us u 1) # X matrix fit  $i \leq lm(us i 1 \sim xx i)$ > summary(fit i) Coefficients: Estimate Std. Error t value Pr(>|t|)0.52177 4.073 5.34e-05 \*\*\* (Intercept) 2.12516 xx i us I 410.03733 37.17344 **11.030** < 2e-16 \*\*\*

xx i us mg -50.07811 15.04907 -3.3280.000935 \*\*\*  $\Rightarrow$  significant. xx i us u 10.22673 0.08346 2.717 0.006805 \*\*  $\Rightarrow$  significant. Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 3.113 on 542 degrees of freedom Multiple R-squared: 0.2276, Adjusted R-squared: 0.2219 F-statistic: 39.93 on 4 and 542 DF, p-value: < 2.2e-16• Now, we include lagged interest rates xx\_i <- cbind(us\_I,e\_ger, us\_mg, us\_u\_1, us\_i\_0) # X matrix with lagged interest rates fit i 1<- lm(us i  $1 \sim xx$  i) summary(fit i 1) Coefficients: Estimate Std. Error t value Pr(>|t|)(Intercept) 0.101007 0.079458 1.271 0.20420 16.367138 6.144709 **2.664** 0.00796 \*\* xx ius I xx ie ger 3.112901 0.691673 4.501 8.3e-06 \*\*\* xx ius mg 1.231633 2.284528 0.539 0.59003  $\Rightarrow$  now, not significant. xx ius u 1 -0.015444 0.012632 -1.2230.22199  $\Rightarrow$  now, not significant. xx i us i 0 0.22673 0.08346 2.717 0.00681 \*\*  $\Rightarrow$  significant effect on other coeff.

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' '1

Residual standard error: 3.113 on 542 degrees of freedom Multiple R-squared: 0.2276, Adjusted R-squared: 0.2219

<u>Note</u>: Lagged ius (ius, t-1) is very significant & changes significance of other variables. It may point out to a general misspecification in (A1).  $\P$ 

# **Omitted Variables Example: Gasoline Demand**

We have a linear model for the demand for gasoline (G) as function of price (PG) and income (Y):

 $\mathbf{G} = \mathbf{PG} \ \beta_1 + \mathbf{Y} \ \beta_2 + \boldsymbol{\varepsilon},$ 

Q: What happens when you wrongly exclude Income (Y)?

 $\mathbf{E}[\mathbf{b}_1|\mathbf{X}] = \beta_1 + \beta_2$ 

In time series data,  $\beta_1 < 0, \beta_2 > 0$  (usually) Cov[Price, Income] > 0 in time series data.  $\Rightarrow$  The short regression will overestimate the price coefficient. In a simple regression of G (demand) on a constant and PG, the Price Coefficient ( $\beta_1$ ) should be negative.

**Example:** Estimation of a 'Demand' Equation: Shouldn't the Price Coefficient be Negative? Taken from Green's graduate Econometrics textbook



• If a multiple regression is done, incorporating income, Y, theory works!

Ordinary	least squares regress	sion		
LHS=G	Mean	= 22	5.09444	
	Standard deviation	= 50	).59182	
	Number of observs.	=	36	
Model size	Parameters	=	3	
	Degrees of freedom	=	33	
Residuals	Sum of squares	= 1472	2.79834	
	Standard error of e	=	6.68059	
Fit	R-squared	=	.98356	
	Adjusted R-squared	=	.98256	
Model test	F[ 2, 33] (prob)	= 987.1	(.0000)	
Variable  Coe	efficient Standard	Error t-:	ratio P[	T >t]
Constant   -	-79.7535*** 8.672	255 -	-9.196	.0000
Y	.03692*** .001	.32 2	28.022	.0000
PG   -	- <b>15.1224***</b> 1.880	)34 -	-8.042	.0000
+				

Note: Income is helping us to identify a demand equation –i.e., with a negative slope for the price variable.  $\P$ 

# **Specification Errors: Irrelevant Variables**

Irrelevant variables. Suppose the correct model is  $\mathbf{y} = \mathbf{X}_1 \mathbf{\beta}_1 + \mathbf{\epsilon}$  -the "short regression," with  $\mathbf{X}_1$  But, we estimate

 $y = X_1\beta_1 + X_2\beta_2 + \varepsilon$  —the "long regression."

Some easily proved results: Including irrelevant variables just reverse the omitted variables results: It increases variance -the cost of not using information-; but does not create biases.

⇒ Since the variables in  $X_2$  are truly irrelevant, then  $β_2 = 0$ , so E[**b**<sub>1.2</sub>|**X**] =  $β_1$ .

• A simple example Suppose the correct model is:  $y = \beta_1 + \beta_2 X_2 + \varepsilon$ But, we estimate:  $y = \beta_1 + \beta_2 X_2 + \beta_3 X_3 + \varepsilon$ 

• Results:

- Unbiased: Given that  $\beta_3 = 0 \implies E[b_2|X] = \beta_2$
- Efficiency:

$$Var[b_2|X] = \frac{\sigma^2}{\sum (X_{2i} - \bar{X}_2)^2} \times \frac{1}{1 - r_{X_2,X_3}^2} > \frac{\sigma^2}{\sum (X_{2i} - \bar{X}_2)^2}$$

where  $r_{X_2 X_3}$  is the correlation coefficient between X<sub>2</sub> and X<sub>3</sub>.

<u>Note</u>: These are the results in general. Note that if  $X_2$  and  $X_3$  are uncorrelated, there will be no loss of efficiency after all.

### **Testing Model Specification: Nested Models**

In both previous cases, we have two nested models, one is the restricted version of the other. For example, in the case of omitted variables:

(U)	$\mathbf{y} = \mathbf{X} \ \mathbf{\beta}_1 + \mathbf{Z} \ \mathbf{\beta}_2 + \mathbf{\epsilon}$	-the "long regression,"
(R)	$y = X \beta_1 + \epsilon$	-the "short regression."

To test H<sub>0</sub> (No omitted variables):  $\beta_2 = 0$ , we can use the F-test:

$$F = \frac{(RSS_R - RSS_U)/J}{RSS_U/(T-k)} \sim F_{J,T-K}.$$

**Example**: In the previous Lecture, we performed this F-test to test if in the 3-factor FF model for IBM returns, SMB and HML were significant, which they were. That is, we showed that the usual CAPM formulation for IBM returns had omitted variables: SMB and HML.

#### **Testing Model Specification with an LM Test**

Note that we can tests for omitted variables with Wald tests and LR tests. The F-test and the LR test of H<sub>0</sub>:  $\beta_2 = 0$  require two estimations: the Unrestricted model and the Restricted model.

There is another test of H<sub>0</sub>:  $\beta_2 = 0$ , that only uses the restricted model as the basis for testing: The Lagrange Multiplier (LM) test, which we introduced in Lecture 5.

In this section, we present the simpler formulation of the LM test, which is based on the residuals of the restricted model,  $e_R$ .

<u>Simple intuition</u>. Everything that is omitted from (& belongs to!) a model should appear in the residuals ( $e_R$ ). Suppose we consider a model driven by  $X_1$  and Z:

$$\mathbf{y} = \mathbf{X}_1 \,\mathbf{\beta}_1 + \mathbf{Z} \mathbf{\beta}_2 + \mathbf{\beta}_2$$

But, we use a simpler model, which omits the J variables,  $\mathbb{Z}$ :

$$y = X_1 \beta_1 + \varepsilon$$

The LM test checks if the restricted residuals,  $e_R$ , can be explained by the *J* omitted variables Z. We use a simple regression of  $e_R$  against Z to check for misspecification in the estimated model.

• LM test steps:

(1) Run restricted model ( $\mathbf{y} = \mathbf{X} \boldsymbol{\beta}_1 + \boldsymbol{\epsilon}$ ). Get restricted residuals,  $\mathbf{e}_{R}$ .

(2) (Auxiliary Regression). Run the regression of  $e_R$  on all the *J* omitted *m* variables, **Z**, and the *k* included variables, **X**. In our case:

 $e_{R,i} = \alpha_0 + \alpha_1 x_{i,1} + \dots + \alpha_k x_{i,k} + \gamma_1 z_{i,1} + \dots + \gamma_J z_{i,J} + v_i$  $\Rightarrow \text{Keep the } \mathbb{R}^2 \text{ from this regression, } \mathbb{R}^2_{eR}.$ 

(3) Compute LM-statistic:

 $\mathrm{LM} = T \ast R^2_{eR} \xrightarrow{d} \chi^2_J.$ 

<u>Technical Note</u>: We include the original variables in (2),  $\mathbf{X}$ , in the auxiliary regression to get the convenient form for the LM-test, as shown by Engle (1982).

The LM Test is very general. It can be used in many settings, for example, to test for nonlinearities, interactions among variables, autocorrelation or heteroscedasticity (discussed later).

Asymptotically speaking, the LM Test, the LR Test and the Wald Test are equivalent –i.e, they have the same limiting distribution,  $\chi_J^2$ . In small *T*, they can have different conclusions. In general, however, we find: W > LR > LM. That is, the LM test is more conservative (cannot reject more often) and the Wald test is more aggressive.

Coefficients:

	Estimate	Std. Error t value $Pr(> t )$		
(Intercept)	0.0007021	0.0024875 0.282	0.7779	
Mkt_RF	0.0125253	0.0567221 0.221	0.8253	
SMB	-0.2124596	0.0841119 -2.526	0.0118 *	
HML	-0.1715002	0.0846817 -2.025	0.0433 *	

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 0.05848 on 565 degrees of freedom Multiple R-squared: 0.01649, Adjusted R-squared: 0.01127 F-statistic: 3.158 on 3 and 565 DF, p-value: 0.02438

**R2\_r** <- summary(fit\_lm)\$r.squared # extracting R<sup>2</sup> from fit\_lm > R2\_r [1] 0.01649104

 $LM\_test <- R2\_r * T$ > LM\\_test [1] 9.383402  $\Rightarrow$  LM\_test > qchisq (.95,df=2)  $\Rightarrow$  Reject H<sub>0</sub>.

```
qchisq(.95, df = 2)# chi-squared (df=2) value at 5% levelp_val <-1 - pchisq(LM_test, df = 2)# p-value of LM_test> p_val= p-val[1] 0.009171071\Rightarrow p-value is small \Rightarrow Reject H<sub>0</sub>: SMB & HML not in model.
```

<u>Conclusion</u>: We strongly reject the CAPM (one factor model), since the LM tests strongly suggests that SMB and HML should be in the model.  $\P$ 

<u>Note</u>: In Lecture 5 we performed the same test with the Wald test (using the F distribution), the p-value was **0.0091175**. (This almost exact coincidence is <u>not</u> always the case.)

### **Functional Form: Linearity in Parameters**

Linear in variables and parameters:

 $y = \beta_1 + \beta_2 X_2 + \beta_3 X_3 + \beta_4 X_4 + \varepsilon$ . So far, this is the linear model we have used. OLS estimates all parameters:  $\beta_1$ ,  $\beta_2$ ,  $\beta_3$ , &  $\beta_4$ .

Non-linear in variables, but linear in parameters -i.e., *intrinsic linear*:

 $\mathbf{y} = \beta_1 + \beta_2 \mathbf{X}_2^2 + \beta_3 \sqrt{\mathbf{X}_3} + \beta_4 \log \mathbf{X}_4 + \boldsymbol{\varepsilon}$ 

Define:  $Z_2 = X_2^2$ ,  $Z_3 = \sqrt{X_3}$ , &  $Z_4 = \log X_4$ 

Then, the non-linear model becomes a linear model:

 $\mathbf{y} = \beta_1 + \beta_2 \mathbf{Z}_2 + \beta_3 \mathbf{Z}_3 + \beta_4 \mathbf{Z}_4 + \boldsymbol{\varepsilon}$ 

Again, OLS can be used to estimate all  $\beta_1$ ,  $\beta_2$ ,  $\beta_3$ , &  $\beta_4$ .

Suppose we have:

$$\mathbf{y} = \beta_1 + \beta_2 \mathbf{X}_2 + \beta_3 \mathbf{X}_2^2 + \boldsymbol{\varepsilon}$$

The model allows for a quadratic relation between y and X<sub>2</sub>:



Let  $X_3 = X_2^2$ , then, the model is intrinsic linear:  $y = \beta_1 + \beta_2 X_2 + \beta_3 X_3 + \varepsilon$ 

**Example:** We want to test if a measure of market risk  $(Mkt_{Ret} - r_f)^2$  enters as an additional explanatory variable in the 3- factor FF model for IBM returns.

The model is non-linear in (Mkt<sub>Ret</sub> – r<sub>f</sub>), but still intrinsic linear:  $IBM_{Ret} - r_f = \beta_0 + \beta_1 (Mkt_{Ret} - r_f) + \beta_2 SMB + \beta_3 HML + \beta_4 (Mkt_{Ret} - r_f)^2 + \epsilon$ 

We can do OLS, by redefining the variables: Let  $X_1 = (Mkt_{Ret} - r_f)$ ;  $X_2 = SMB$ ;  $X_3 = HML$ ;  $X_4 = X_1^2$ . Then,

$$\mathbf{y} = \beta_0 + \beta_1 \mathbf{X}_1 + \beta_2 \mathbf{X}_2 + \beta_3 \mathbf{X}_3 + \beta_4 \mathbf{X}_4 + \boldsymbol{\varepsilon}$$

Mkt\_RF2 <- Mkt\_RF^2 fit\_ibm\_ff\_2 <- lm (ibm\_x ~ Mkt\_RF + SMB + HML + Mkt\_RF2) summary(fit\_ibm\_ff\_2)

Coefficients:

	Estimate	Std. Error	t value Pr(> t )	
(Intercept)	-0.004765	0.002854	-1.670 0.0955.	
Mkt_RF	0.906527	0.057281	15.826 <2e-16 ***	
SMB	-0.215128	0.084965	-2.532 0.0116 *	
HML	-0.173160	0.085054	-2.036 0.0422 *	
Mkt_RF2	-0.143191	0.617314	-0.232 0.8167	$\Rightarrow$ Not significant!

<u>Conclusion</u>: A t-test cannot reject H<sub>0</sub>:  $\beta_4$ =0. That is, there is no evidence that  $(Mkt_{Ret} - r_f)^2$  is an explanatory variable for IBM excess returns.

• Now, we can also check with an LM test if all variables squares  $((Mkt_{Ret} - r_f)^2, SMB^2, and HML^2)$  are omitted from the 3-factor FF model for IBM returns.

Mkt\_RF2 <- Mkt\_RF^2 SMB2 <- SMB^2  $\begin{array}{ll} \textbf{HML2} <- \ \textbf{HML^{2}} \\ fit\_r <- \ \textbf{Im} \ (ibm\_x \sim Mkt\_RF + SMB + \textbf{HML}) \\ resid\_r <- \ fit\_r\$residuals \\ fit\_lm <- \ \textbf{Im} \ (resid\_r \sim Mkt\_RF + SMB + \textbf{HML} + Mkt\_RF2 + SMB2 + \textbf{HML2}) \\ \textbf{R2\_r} <- \ summary(fit\_lm)\$r.squared \\ LM\_test <- \ \textbf{R2\_r} * T \\ > \ LM\_test \\ \textbf{[1] 2.453822} \\ p\_val <- 1 - pchisq(LM\_test, df = 3) \ \# p-value of \ LM\_test \\ > \ p\_val \\ \textbf{[1] 0.4836944} \qquad \Rightarrow p-value is higher than standard levels \Rightarrow Cannot Reject H_0. \\ \end{array}$ 

<u>Conclusion</u>: The LM test cannot reject the 3-factor F-F model, since all squared terms are not jointly significant.  $\P$ 

• Nonlinear in parameters:

$$\mathbf{y} = \beta_1 + \beta_2 \mathbf{X}_2 + \beta_3 \mathbf{X}_3 + \beta_2 \beta_3 \mathbf{X}_4 + \boldsymbol{\varepsilon}$$

This model is nonlinear in parameters since the coefficient of  $X_4$  is the product of the coefficients of  $X_2$  and  $X_3$ . OLS cannot be used to estimate all parameters.

Some nonlinearities in parameters can be linearized by appropriate transformations, but not this one. This is not an intrinsic linear model. Different estimation techniques should be used in these cases.

Intrinsic linear models can be estimated using OLS. Sometimes, transformations are needed. Suppose we start with a power function:

$$\boldsymbol{y} = \beta_1 \boldsymbol{X}^{\beta_2} \boldsymbol{\varepsilon}$$

• The errors enter in multiplicative form. Then, using logs:

 $\log \mathbf{y} = \log \beta_1 \mathbf{X}^{\beta_2} \boldsymbol{\varepsilon} = \log \beta_1 + \beta_2 \log \mathbf{X} + \log \boldsymbol{\varepsilon},$ 

or

$$\mathbf{y}' = \beta_1' + \beta_2 \mathbf{X}' + \boldsymbol{\varepsilon}',$$

where  $y' = \log y$ ,  $X' = \log X$ ,  $\beta'_1 = \log \beta_1$ ,  $\varepsilon' = \log \varepsilon$ 

Now, we have an intrinsic linear model: OLS can be used to estimate all the parameters.

Similar intrinsic model can be obtained if  $y = e^{\beta_1 + \beta_2 X + \varepsilon}$ 

<u>Note</u>: Recall that we can only use logs when y has positive values. In general, we use logs when we believe the independent variable has an exponential or power formulation, typical behavior for nominal variables, like sales, revenue or prices.

• Not all models are intrinsic linear. For example:

$$y = \beta_1 X^{\beta_2} + \varepsilon$$

$$\log \mathbf{y} = \log(\beta_1 \mathbf{X}^{\beta_2} + \boldsymbol{\varepsilon})$$

We cannot linearize the model by taking logarithms. There is no way of simplifying  $\log(\beta_1 X^{\beta_2} + \varepsilon)$ .

We will have to use some nonlinear estimation technique for these situations. (ML can estimate this model.)

### **Functional Form: Linear vs Log specifications**

Two popular models, especially in Corporate Finance: linear or log?

Model 1 - Linear model:  $\mathbf{y} = \beta_1 + \beta_2 \mathbf{X} + \varepsilon$ Model 2 - (Semi-) Log model:  $\log \mathbf{y} = \beta_1 + \beta_2 \mathbf{X} + \varepsilon$ 

Box–Cox transformation:  $\frac{Y^{\lambda}-1}{\lambda} = \beta_1 + \beta_2 \mathbf{X} + \varepsilon$ 

$$\frac{\frac{\lambda}{Y^{\lambda}-1}}{\frac{\lambda}{\lambda}} = Y - 1 \qquad \text{when } \lambda = 1$$
$$\frac{\frac{Y^{\lambda}-1}{\lambda}}{\frac{\lambda}{\lambda}} = \log(Y) \qquad \text{when } \lambda \to 0$$

Putting  $\lambda = 0$  gives the (semi-)log model (think about the limit of  $\lambda$  tends to zero.). The Box-Cox transformation is flexible. We can estimate  $\lambda$  to test if  $\lambda$  is equal to 0 or 1. It is possible that it is neither!

### **Functional Form: Ramsey's RESET Test**

To test the specification of the functional form, Ramsey designed a simple test. We start with the fitted values from our (A1) model:

$$\hat{\mathbf{y}} = \mathbf{X}\mathbf{b}.$$

Then, we add  $\mathbf{\hat{y}}^2$  to the regression specification:  $y = X \beta + \mathbf{\hat{y}}^2 \gamma + \varepsilon$ 

If  $\hat{\mathbf{y}}^2$  is added to the regression specification, it should pick up quadratic and interactive nonlinearity, if present, without necessarily being highly correlated with any of the *X* variables.

We test	H <sub>0</sub> (linear functional form): $\gamma = 0$
	H <sub>1</sub> (non linear functional form): $\gamma \neq 0$
	$\Rightarrow$ <i>t-test</i> on the OLS estimator of $\gamma$ .

If the *t*-statistic for  $\mathbf{\hat{y}}^2$  is significant  $\Rightarrow$  evidence of nonlinearity.

The RESET test is intended to detect nonlinearity, but not be specific about the most appropriate nonlinear model (no specific functional form is specified in  $H_1$ ).

Example: We want to test the functional form of the 3 FF Factor Model for IBM returns, using
monthly data 1973-2020.
fit\_ibm\_ff3 <- lm(ibm\_x ~ Mkt\_RF + SMB + HML)
y\_hat <- fitted(fit\_ibm\_ff3)
y\_hat2 <- y\_hat^2
fit\_ramsey <- lm(ibm\_x ~ Mkt\_RF + SMB + HML + y\_hat2)
> summary(fit\_ramsey)

y_hat2	-0.289197	0.763526	-0.379	0.7050	$\Rightarrow$ Not significant!
HML	-0.173276	0.084875	-2.042	0.0417 *	
SMB	-0.217268	0.085128	-2.552	0.0110 *	
Mkt_RF	0.903783	0.058003	15.582	<2e-16 ***	
(Intercept)	-0.004547	0.002871	-1.584	0.1137	
	Estimate	Std. Error	t value	Pr(> t )	
Coefficients:					

<u>R Note</u>: The *lmtest* package performs this test, *resettest*, (and many others, used in this class, encompassing, jtest, waldtest, etc). You need to install it first: install.packages("lmtest"), then call the library(lmtest).

<u>Note</u>: The test reported is an *F*-test ~  $F_{1,T-k}$ , which is equal to  $(t_{T-k})^2$ . The *p*-values should be the same.

library(Imtest) > resettest(fit\_ibm\_ff3, power=2, type="fitted") RESET test data:  $y \sim Mkt_RF + SMB + HML$ RESET = 0.14346, df1 = 1, df2 = 564, p-value = 0.705  $\Rightarrow$  cannot reject H<sub>0</sub>. Check: (-0.379)<sup>2</sup> = 0.1434.

<u>Conclusion</u>: The RESET test does not find evidence of non-linearities (or, in general, of misspecification) in the 3-factor F-F model, since the squared fitted values are not significant at the 5% level.  $\P$ 

# **Qualitative Variables and Functional Form**

Suppose that you want to model CEO compensation as a function of education. You have data on annual total CEO compensation (*Comp*), annual returns, annual sales, CEO's age, CEO's previous experience, and the CEO's last degree (education). We have qualitative data.

One approach to see the impact of education on the CEO compensation model is to run individual regressions for each last degree –i.e., BA/BS; MS/MA/MBA; Doctoral:

Undergrad degree	$Comp_i = \beta_{0-u} + \beta_{1-u'} \mathbf{z}_i + \varepsilon_{u,i}$
Masters degree	$Comp_i = \beta_{0-m} + \beta_{1-m'} \mathbf{z}_i + \varepsilon_{m,i}$
Doctoral degree	$Comp_i = \beta_{0-d} + \beta_{1-d}' \mathbf{z}_i + \varepsilon_{d,i}$

where the  $z_i$  is a vector of the CEO *i*'s age and previous experience and his/her firm's *annual* returns and annual sales. We observe the impact of education through the different coefficients in each regression. A potential problem with this approach is that we may end up with three small samples (and imprecise estimations).

An alternative approach that uses the whole sample in the estimation is to combine the three regressions in one. To do this, we use a "*dummy variable*" –also called, *indicator variable*–, which is a variable that points whether an observation belongs to a category or class or not. For example:

$D_{C,i} = 1$	if observation <i>i</i> belongs to category C (say, male.)
= 0	otherwise.

Simple process: First, we define dummy/indicator variables for Masters & doctoral degrees:

degree

$D_{m,i} = 1$	if at least Masters
= 0	otherwise.
$D_{d,i} = 1$	if doctoral degree
= 0	otherwise.

Then, we introduce the dummy/indicator variables in the compensation model:

 $Comp_{i} = \beta_{0} + \beta_{1}'\mathbf{z}_{i} + \beta_{2} D_{m,i} + \beta_{3} D_{d,i} + \gamma_{1}'\mathbf{z}_{i} D_{m,i} + \gamma_{2}'\mathbf{z}_{i} D_{d,i} + \varepsilon_{i}$ 

Not, this model uses all the sample to estimate the parameters. It is flexible:

- Model for undergrads only  $(D_{m,i} = 0 \& D_{d,i} = 0)$ :  $Comp_i = \beta_0 + \beta_1 '\mathbf{z}_i + \varepsilon_i$ - Model for Masters degree only  $(D_{m,i} = 1 \& D_{d,i} = 0)$ :  $Comp_i = (\beta_0 + \beta_2) + (\beta_1 + \gamma_1)'\mathbf{z}_i + \varepsilon_i$ - Model for Doctoral degree only  $(D_{m,i} = 1 \& D_{d,i} = 2)$ :

$$Comp_{i} = (\beta_{0} + \beta_{2} + \beta_{3}) + (\beta_{1} + \gamma_{1} + \gamma_{2})'\mathbf{z}_{i} + \varepsilon_{i}$$

The parameters for the different categories are:

- Constant:

Constant for undergrad degree:  $\beta_0$ Constant for Masters degree:  $\beta_0 + \beta_2$ 

Constant for Doctoral degree:  $\beta_0 + \beta_2 + \beta_3$ 

- Slopes:

Slopes for undergrad degree:  $\beta_1$ Slopes for Masters degree:  $\beta_1 + \gamma_1$ Slopes for Doctoral degree:  $\beta_1 + \gamma_1 + \gamma_2$ 

We can test the effect of education on CEO compensation:

(1) H <sub>0</sub> : No effect of grad degree: $\beta_3 = \beta_2 = 0$ & $\gamma_1 = \gamma_2 = 0$	$\Rightarrow$ <i>F</i> -test.
(2) H <sub>0</sub> : No effect of Masters degree on constant: $\beta_2 = 0$	$\Rightarrow$ <i>t</i> - <i>test</i> .
(3) H <sub>0</sub> : No effect of doctoral degree: $\beta_3 = 0 \& \gamma_2 = 0$	$\Rightarrow$ <i>F</i> -test.
(4) H <sub>0</sub> : No effect of Dr degree on marginal effect: $\gamma_2 = 0$	$\Rightarrow$ <i>F</i> -test.

• We may have more than one qualitative category (last degree above) in our data that we may want to introduce in our model.

**Example**: Suppose we also have data for CEO graduate school. Now, we can create another qualitative category, "quality of school", defined as Top 20 school, to test if a Top 20 school provides "more value." To do this, we use  $D_{TOP}$  to define if any schooling is in the Top 20.

$$D_{TOP,i} = 1$$
 if CEO *i*'s school is a Top 20 school  
= 0 otherwise.

The model becomes:

 $Comp_{i} = \beta_{0} + \beta_{1}'z_{i} + \beta_{2}D_{m,i} + \beta_{3}D_{d,i} + \beta_{4}D_{TOP,i} + \gamma_{1}'z_{i}D_{m,i} + \gamma_{2}'z_{i}D_{d,i} + \gamma_{3}'z_{i}D_{TOP,i} + \varepsilon_{i}$ 

In this setting, we can test the effect of a Top20 education on CEO compensation: (1) H<sub>0</sub>: No effect of Top20 degree:  $\beta_4 = 0$  and  $\gamma_3 = 0 \Rightarrow F$ -test.

• The omitted category is the reference or control category.

- In our first example, with only educational degrees, the reference category is undergraduate degree. - In the second example, with educational degrees and quality of school (Top20 dummy), the reference category is undergraduate degree with no Top 20 education.

• *Dummy trap*. If there is a constant, the numbers of dummy variables per qualitative variable should be equal to the number of categories minus 1. If you put the number of dummies variables equals the number of categories, you will create perfect multicollinearity.

### **Dummy Variables as Seasonal Factors**

A popular use of dummy variables is in estimating seasonal effects. We may be interested in studying the January effect in stock returns or if the returns of oil companies (say, Exxon or BP) are affected by the seasons, since during the winter people drive less and during the summer.

In this case, we define dummy/indicator variables for Summer, Fall and Winter (the base case is, thus, Spring):

$D_{Sum,i} = 1$	if observation <i>i</i> occurs in Summer
= 0	otherwise.
$D_{Fall,i} = 1$	if observation <i>i</i> occurs in Fall
= 0	otherwise.
$D_{Win,i} = 1$	if observation <i>i</i> occurs in Winter
= 0	otherwise.

Then, letting **Z** be the vector of the three FF factors  $((r_{m,t} - r_f), SMB_t, HML_t)$ , and assuming the seasons only affect the constant, we have:

 $(r_i - r_f) = \beta_0 + \beta_1' \mathbf{z}_i + \beta_2 D_{Sum,i} + \beta_3 D_{Fall,i} + \beta_4 D_{Win,i} + \varepsilon_i$ 

**Example:** In the context of the 3-factor FF model, we test if Exxon's excess returns (XOM) are affected by seasonal (quarters) factors:

```
(r_{XOM,i} - r_f) = \beta_0 + \beta_1 z_i + \beta_2 D_{Sum,i} + \beta_3 D_{Fall,i} + \beta_4 D_{Win,i} + \varepsilon_i
x_xom <- SFX da$XOM
                                                              # Extract XOM prices
T \leq - length(x xom)
\ln xom < \log(x xom[-1]/x xom[-T])
xom x \leq - lr xom - RF
T \leq - length(xom x)
Summ \leq \text{rep}(c(0,0,0,0,0,0,1,1,1,0,0,0), \text{round}(T/12)+1)
                                                              # Create Summer dummy
Fall \leq- rep(c(0,0,0,0,0,0,0,0,0,1,1,1), round(T/12)+1)
                                                              # Create Fall dummy
                                                              # Create Winter dummy
Wint \leq \operatorname{rep}(c(1,1,1,0,0,0,0,0,0,0,0,0), \operatorname{round}(T/12)+1)
T1 <- T+1
Fall 1 <- Fall[2:T1]
                                                              # Adjusting sample (starts in Feb)
Wint 1 <- Wint[2:T1]
Summ 1 <- Summ[2:T1]
fit xom s <- lm(xom x \sim Mkt RF + SMB + HML + Fall 1 + Wint 1 + Summ 1)
summary(fit xom s)
> summary(fit xom s)
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.002445 0.003485 0.702 0.4832 \Rightarrow constant for reference category (Spring)\approx0.
           0.761816 0.040602 18.763 < 2e-16 ***
Mkt RF
SMB
           -0.261925 0.060575 -4.324 1.81e-05 ***
HML
           0.370623 0.060049 6.172 1.29e-09 ***
Fall 1
           -0.006609 0.004947 -1.336 0.1822
Wint 1
           -0.011283 0.004928 -2.290 0.0224 * \Rightarrow significant. Reject H<sub>0</sub>: No Winter effect.
Summ 1 -0.007100 0.004944 -1.436 0.1515
```

<u>Interpretation</u>: In the Winter quarter, Exxon excess returns decrease, relative to the Spring, by **1.13%**. But since Spring's (& Fall's & Winter's) effect is non-significant, the decrease is in absolute terms.

<u>Conclusion</u>: The t-value for the Winter dummy (Wint\_1) is significant at the 5% level. That is, we reject  $H_0$ : No seasonal effect on XOM excess returns.

• We can test if all quarters jointly matter. That is,  $H_0$ :  $\beta_2 = \beta_3 = \beta_4 = 0$ . We do an F-test:

 $\begin{array}{l} fit\_u <- lm \ (xom\_x \sim Mkt\_RF + SMB + HML + Fall\_1 + Wint\_1 + Summ\_1) \\ fit\_r <- lm \ (xom\_x \sim Mkt\_RF + SMB + HML) \\ resid\_u <- lit\_u\$residuals \\ RSS\_u <- sum((resid\_u)^2) \\ resid\_r <- lit\_r\$residuals \\ RSS\_r <- sum((resid\_r)^2) \\ f\_test <- ((RSS\_r - RSS\_u)/2)/(RSS\_u/(T-4)) \\ > f\_test \end{array}$ 

[1] 2.706574
> p\_val <- 1 - pf(f\_test,df1=3, df2=T-3) # p-value of F-test</li>
> p\_val
[1] 0.05504357 ⇒ p-value is "marginal." Cannot reject H₀: No joint seasonal effect.

Conclusion: We cannot reject, at the 5% level, H<sub>0</sub>: No joint seasonal effect.

• Suppose we are also interested in checking if the slopes –i.e., the marginal effects- are affected by the Winter quarter. Then, we fit:

 $(r_{XOM,i} - r_f) = \beta_0 + \beta_1' z_i + \beta_2 D_{Sum,i} + \beta_3 D_{Fall,i} + \beta_4 D_{Win,i} + \gamma_1' z_i D_{Win,i} + \varepsilon_i$   $Mkt_W <- Mkt_RF*Wint_1$   $SMB_W <- SMB*Wint_1$   $HML_W <- HML*Wint_1$   $fit_xom_s2 <- lm(xom_x \sim Mkt_RF + SMB + HML + Mkt_W + SMB_W + HML_W + Fall_1$   $+ Wint_1 + Summ_1)$   $> summary(fit_xom_s2)$ 

Coefficients:

	Estimate	Std. Error t value	$Pr(\geq  t )$	
(Intercept)	0.003127	0.003478 0.899	0.368962	
Mkt_RF	0.695762	0.048202 14.434	< 2e-16 ***	
SMB	-0.291199	0.075197 -3.872	0.000120 ***	
HML	0.270262	0.077416 3.491	0.000519 ***	
Mkt_W	0.208912	0.091972 2.271	0.023497 *	$\Rightarrow$ significant effect on Mkt's slope
SMB_W	0.064753	0.126138 0.513	0.607911	
HML_W	0.198753	0.124261  1.599	0.110278	
Fall_1	-0.006795	0.004934 -1.377	0.169038	
Wint_1	-0.013747	0.005000 -2.750	0.006159 **	$\Rightarrow$ significant effect on constant.
Summ 1	-0.007492	0.004928 -1.520	0.129012	-

<u>Interpretation</u>: The only factor interacting significantly with Winter is the Market factor. Then, we have two significantly different slopes:

- In the Winter, the Market slope is: 0.695762 + 0.208912 = 0.903674

- In all other quarters, the Market is: 0.695762

It looks like in the Winter, XOM behaves closer to the Market, while in all other quarters, it is significantly less risky than the market.

• Now, we perform a joint test for interacting Winter effects in the model:: > f\_test [1] 6.505231 p\_val <- 1 - pf(f\_test, df 1= 3, df2=T-7) # p-value of F-test > p\_val [1] 0.0007923967  $\Rightarrow$  p-value < .05, then, we reject H<sub>0</sub> (joint Winter interactive effect):  $\gamma_1 = 0$ . Conclusion: We strongly reject, at the 5% level, H<sub>0</sub>: No joint Winter interactive effect. ¶

### **Dummy Variables: Is There a January Effect?**

The January Effect is a hypothesis that states that the stock market has an unusually high return during the first month of the year. This result can be traced to an observation made in 1942 U.S. investment banker Sidney Wachtel. Wachtel noticed higher returns for small stocks than for large stock in January, a result later examined by Kiem (1983), who found that the January return premium was evident for small stocks. In one of the earlier studies, Rozeff and Kinney (1976) found seasonal patterns in an equal-weighted index of NYSE prices over the period 1904-74. Specifically, the average monthly return in January was 3.5%, while other months averaged 0.5% percent. A very strong result that shows a clearly predictable pattern, which goes against the Efficient Markets Hypothesis. Since then, a lot of work has been done: The evidence suggests that, in recent years, the January effect has dissipated.

**Example:** We want to test the January effect on IBM stock returns, where because of tax reasons/window dressing, stocks go down in December and recover in January. The test can be done by adding a dummy variable to the 3-factor FF model:

$D_{J,t} = 1$	if observation t occurs in January
= 0	otherwise.

Then, we estimate the expanded 3-factor FF model:

 $(r_t - r_f) = \beta_0 + \beta_1 (r_{m,t} - r_f) + \beta_2 SMB_t + \beta_3 HML_t + \beta_4 D_{J,t} + \varepsilon_t$ 

We test H<sub>0</sub>(No January effect):  $\beta_4 = 0 \implies t$ -test.

Alternatively, we can estimate do an LM test on the residuals of the 3-factor FF model and check if  $D_{I,t}$  is significant.

 $T \le \text{length}(\text{ibm}_x)$ Jan <- rep(c(1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0), (round(T)/12+1)) # Create January dummy  $T2 \le T+1$ Jan\_1 <- Jan[2:T2] # Adjust sample fit\_ibm\_ff3 <- lm (ibm\_x ~ Mkt\_RF + SMB + HML) # Restricted Regression resid\_r <- fit\_ibm\_ff3\$residuals # Keep residuals (e<sub>R</sub>) fit\_Jan <- lm (resid\_r ~ Mkt\_RF + SMB + HML + Jan\_1) # Auxiliary Regression > summary(fit\_Jan) Coefficients: Estimate Std. Error t value Pr(>|t|) (Intercept) -0.002111 0.002561 -0.824 0.41027 Mt + DF = 0.005100 0.056405 = 0.0020 0.026(1)

Mkt_RF	-0.005198	8 0.056405	-0.092	0.92661
SMB	-0.026306	0.084063	-0.313	0.75445
HML	-0.014914	0.083606	-0.178	0.85848
Jan 1	0.026966	0.008906	3.028	0.00258 **

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 0.058 on 565 degrees of freedom Multiple R-squared: 0.01597, Adjusted R-squared: 0.009 F-statistic: 2.292 on 4 and 565 DF, p-value: 0.05841

Given this result, we modify the 3-factor FF and add the January Dummy to the FF model:

fit\_ibm\_ff3\_Jan <- lm (ibm\_x ~ Mkt\_RF + SMB + HML + Jan\_1)
> summary(fit ibm ff3 Jan)

Coefficients:

Estimate Std. Error t value Pr(>|t|)(Intercept) -0.007302 0.002561 -2.851 0.00452 \*\* Mkt\_RF 0.905182 0.056405 16.048 < 2e-16 \*\*\* SMB -0.247691 0.084063 -2.946 0.00335 \*\* HML -0.154093 0.083606 -1.843 0.06584 . Jan\_1 0.026966 0.008906 3.028 0.00258 \*\* ---Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 0.058 on 565 degrees of freedom Multiple R-squared: 0.3499, Adjusted R-squared: 0.3453 F-statistic: 76.01 on 4 and 565 DF, p-value: < 2.2e-16

<u>Interpretation</u>: We have two constants (excess return, Jensen's alpha): Feb - Dec: -0.7302% (significant). January: -0.7302% + 2.6966% = 1.9664% (significant).

When the January dummy was not in the model, we had: -0.005191, which is close to an average of the constants (=  $-0.007302 \times 11 + 0.019664$ )/12 = -0.00505).

<u>Interpretation</u>: During January IBM has an additional **2.6966%** excess returns. This is a big number. Today, the evidence for the January effect is much weaker than in this case.

---

<u>Note</u>: In the FF model we expect the constant to be very small ( $\approx 0$ ). In this case, it is not zero. Like in the case of the CAPM, a significant constant is evidence against the 3-factor model of Fama-French. Maybe we have a misspecified (A1).

## **Dummy Variable for One Observation**

 $D_i$ 

We can use a dummy variable to isolate a single observation.

= 1 for observation j.= 0 otherwise.

Define **d** to be the dummy variable in question.

 $\mathbf{Z} =$ all other regressors.  $\mathbf{X} = [\mathbf{Z}, \mathbf{D}_I]$ 

Multiple regression of **y** on **X**. We know that

**X'**e = 0 where e = the column vector of residuals.  $\Rightarrow D_i'e = 0 \Rightarrow e_i = 0$  (perfect fit for observation *j*).

This approach can be used to deal with (eliminate) outliers.

**Example:** In Dec 1992, IBM reported record losses and gave a very bleak picture of its future. The stock tumbled -30.64% that month. We check the effect of that extreme observation, a potential outlier, on the 3-factor FF model + January dummy:

Coefficients:

	Estimate	Std. Error	t value	$\Pr(> t )$
(Intercept)	-0.006772	0.002502	-2.707	0.00699 **
Mkt_RF	0.908775	0.055054	16.507	< 2e-16 ***
SMB	-0.239213	0.082059	-2.915	0.00370 **
HML	-0.138629	0.081647	-1.698	0.09008 .
Jan_1	0.026163	0.008694	3.009	0.00273 **
dec_1992	-0.306202	0.056710	-5.399	9.86e-08 *** (same value of observation)

Conclusion: Potential "Outlier" has no major effect on coefficients.

### Chow Test: Testing the effect of Categories on a Model

It is common to have a qualitative variable with two categories, say education (MS/MBA or not). Before modelling the data, we can check if only one regression ("pooling") model applies to both categories.

We use the Chow Test (an F-test) –Chow (1960, *Econometrica*).

Steps:

(1) Run OLS with all the data, with no distinction between schools (Pooled regression or Restricted regression). Keep RSS<sub>R</sub>.

(2) Run two separate OLS, one for each school (Unrestricted regression). Keep RSS<sub>1</sub> and RSS<sub>2</sub>  $\Rightarrow$  RSS<sub>U</sub> = RSS<sub>1</sub> + RSS<sub>2</sub>.

(Alternative, we can run just one regression with the dummy variable).

(3) Run a standard F-test (testing Restricted vs. Unrestricted models):

$$F = \frac{(RSS_R - RSS_U)/(k_U - k_R)}{(RSS_U)/(T - k_U)} = \frac{(RSS_R - [RSS_1 + RSS_2])/k}{(RSS_1 + RSS_2)/(T - 2k)}$$

Recall that under normality –i.e., under (A5) – the distribution of the F test is:  $F \sim F_{J,T-k}$ , where  $J = (k_U - k_R)$ .

<u>Remark</u>: If (A5) is not assumed and it we have a large number of observations in both categories, it is common to approximate the asymptotic distribution of the Chow test with  $J * F \xrightarrow{d} \chi_J^2 / J$ . Note, that there are many papers arguing that the approximation is not very accurate.

**Example:** Who visits doctors more: Men or Women? Data: German Health Care Usage Data, with 7,293 Individuals. Time Periods: Varying Number. Variables in the file are:

Data downloaded from Journal of Applied Econometrics Archive. This is an unbalanced panel with 7,293 individuals. There are altogether **27,326** observations. The number of observations ranges from 1 to 7 per family. (Frequencies are: 1=1525, 2=2158, 3=825, 4=926, 5=1051, 6=1000, 7=987).

The dependent variable of interest is:

DOCVIS = number of visits to the doctor in the observation period

The explanatory variables are: HHNINC = household nominal monthly net income in German marks / 10000. (4 observations with income=0 were dropped) GENDER\_F = gender (1 = female) HHKIDS = children under age 16 in the household = 1; otherwise = 0 EDUC = years of schooling AGE = age in years MARRIED= marital status (1 = if married) WHITEC = 1 if has "white collar" job Health\_Da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/german\_health.csv", head=TRUE, sep=",")

x\_fem <- Health\_Da\$Gender\_F
x\_age <- Health\_Da\$age
x\_edu <- Health\_Da\$educ
x\_hhinc <- Health\_Da\$hhninc/100
x\_hhkids <- Health\_Da\$hhkids
x\_married <- Health\_Da\$married
x\_white\_col <- Health\_Da\$whitecollar
x\_docvis <- Health\_Da\$docvis</pre>

fit\_doc\_vis <- lm (x\_docvis ~ x\_age + x\_edu + x\_married + x\_white\_col + x\_hhkids +
x\_hhinc)
summary(fit\_doc\_vis)</pre>

• OLS Estimation for ALL. Keep RSS<sub>ALL</sub> = **858,435** (=5.606<sup>2</sup> \* 27,315)

Coefficients:

	Estimate	Std. Error	t value	Pr(> t )		
(Intercept)	2.683700	0.249282	10.766	< 2e-16	* * *	
x age	0.061810	0.003444	17.947	< 2e-16	* * *	
x edu	-0.118858	0.015573	-7.632	2.38e-14	* * *	
x married	-0.090716	0.089056	-1.019	0.308		
x white col	-0.115412	0.076540	-1.508	0.132		
x hhkids	-0.492028	0.080014	-6.149	7.89e-10	* * *	
x_hhinc	-0.015429	0.002046	-7.539	4.87e-14	* * *	
Signif. code	es: 0 `***	· 0.001 · * ·	*′ 0.01 `	`*′ 0.05 `	`.' 0.1 `'	1
Residual sta	andard erro	or: <b>5.606</b> or	n <b>27,315</b> d	degrees of	f freedom	
Multiple R-s	squared: (	.02949, Ad-	justed R-	-squared:	0.02928	
F-statistic:	: 138.3 on	6 and 27315	DF, p-	-value: <	2.2e-16	
			-			

• OLS Estimation for Women only. Keep RSSw = 478,894.2 (= 6.052^2 \* 13,075)

## Run a regression with only Women data. Use Allgen to collect relevant data for women only. We will do a for loop and keep data if  $x_{fem}$  is greater than 0.

xx <- cbind(x\_fem, x\_docvis, x\_age, x\_edu, x\_married, x\_white\_col, x\_hhkids, x\_hhinc)

```
Allgen = NULL # Initialize empty (to collect variables by one sex (f/m) only)
i <- 1
T <- length(x_fem)
k <- ncol(xx)
```

```
for (i in 1:T) {
if (xx[i,1] > 0) {
 Allgen = rbind(Allgen, xx[i,2:k])
 }
}
y g \leq Allgen[,1]
                     # Dependent variable: doctor's visits by women only
x g \leq Allgen[,2:(k-1)]
T f <-length(y g)
fit doc vis f \le lm (y g \sim x g)
summary(fit doc vis f)
Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
                 2.999559 0.453506 6.614 3.88e-11 ***
(Intercept)
                             0.005719 8.632 < 2e-16 ***
x gx age
                 0.049366
                -0.048141 0.027011 -1.782
                                                  0.0747 .
x gx edu
x gx married
                -0.119853 0.133846 -0.895
                                                  0.3706
x gx white col -0.006734
                             0.124768 -0.054
                                                  0.9570
x gx hhkids
                -0.636619
                                        -4.941 7.87e-07 ***
                             0.128844
x gx hhinc
                -0.015651
                             0.003174 -4.932 8.25e-07 ***
Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 `' 1
Residual standard error: 6.052 on 13075 degrees of freedom
Multiple R-squared: 0.01984, Adjusted R-squared:
                                                       0.01939
F-statistic: 44.11 on 6 and 13075 DF, p-value: < 2.2e-16
• OLS Estimation for Men only. Keep RSS_M = 372,818.1
# Use above code, but change for loop (now, keep data if x fem less than 1)
for (i in 1:T) \{
if (xx[i,1] < 1) {
 Allgen = rbind(Allgen, xx[i,2:k])
 }
}
Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
                             0.290792
                                         6.195 5.98e-10 ***
                 1.801539
(Intercept)
x gx age
                 0.067656
                             0.004421
                                        15.302 < 2e-16 ***
x gx edu
                -0.105462 0.018814 -5.605 2.12e-08 ***
x gx married
                0.022278 0.121467 0.183 0.854480
x gx white col -0.367075 0.096300 -3.812 0.000139 ***
x gx hhkids
                -0.428916 0.102070 -4.202 2.66e-05 ***
                -0.015438 0.002629 -5.872 4.40e-09 ***
x gx hhinc
```

Signif. codes: 0 `\*\*\*' 0.001 `\*\*' 0.01 `\*' 0.05 `.' 0.1 `' 1 Residual standard error: 5.118 on 14233 degrees of freedom Multiple R-squared: 0.03602, Adjusted R-squared: 0.03561 F-statistic: 88.63 on 6 and 14233 DF, p-value: < 2.2e-16

• Chow Test:  

$$F = \frac{(RSS_R - [RSS_1 + RSS_2])/k}{(RSS_1 + RSS_2)/(T - 2k)} = \frac{(858, 435 - [372, 818.1 + 478, 894.2)]/7}{(372, 818.1 + 478, 894.2)/(27, 323 - 14)}$$

$$= 31.1178$$

F(7, 27309) = 2.009925  $\Rightarrow$  reject H<sub>0</sub> at 5% level.

Conclusion: There is strong evidence that men and women do not have the same behavior.

### **Functional Form: Structural Change**

Suppose there is an event that we think had a big effect on the behaviour of our model. Suppose the event occurred at time  $T_{SB}$ . We think that the before and after behaviour of the model is significantly different. For example, the parameters are different before and after  $T_{SB}$ . That is,

$$y_{i} = \beta_{0}^{1} + \beta_{1}^{1} X_{1,i} + \beta_{2}^{1} X_{2,i} + \beta_{3}^{1} X_{3,i} + \varepsilon_{i} \qquad \text{for } i \leq T_{SB}$$
  
$$y_{i} = \beta_{0}^{2} + \beta_{1}^{2} X_{1,i} + \beta_{2}^{2} X_{2,i} + \beta_{3}^{2} X_{3,i} + \varepsilon_{i} \qquad \text{for } i > T_{SB}$$

The event caused *structural change* in the model. *T*<sub>SB</sub> separates the behaviour of the model in two regimes/categories ("*before*" & "*after*".)

A Chow test can be used to check if one model applies to both regimes:  $y_i = \beta_0 + \beta_1 X_{1,i} + \beta_2 X_{2,i} + \beta_3 X_{3,i} + \varepsilon_i$  for all *i* 

Under H<sub>0</sub> (No *structural change*), the parameters are the same for all *i*.

• We test H<sub>0</sub> (No *structural change*):  $\beta_0^1 = \beta_0^2 = \beta_0$   $\beta_1^1 = \beta_1^2 = \beta_1$   $\beta_2^1 = \beta_2^2 = \beta_2$   $\beta_3^1 = \beta_3^2 = \beta_3$ H<sub>1</sub> (*structural change*): For at least k (= 0, 1, 2, 3):  $\beta_k^1 \neq \beta_k^2$ 

What events may have this effect on a model? A financial crisis, a big recession, an oil shock, Covid-19, new taxes and regulations, etc.

Testing for structural change is the more popular use of the Chow test.
Chow tests have many interpretations: tests for structural breaks, pooling groups, parameter stability, predictive power, etc.

One important consideration: T may not be large enough. For example, we may think that Covid-19 had a structural effect on the behaviour of tech companies. We may not have enough data to run an F-test.

We structure the Chow test to test H<sub>0</sub> (No *structural change*) as usual.

• Steps for Chow (Structural Change) Test:

(1) Run OLS with all the data, with no distinction between regimes (Restricted or pooled model): Keep RSS<sub>R</sub>.

(2) Run two separate OLS, one for each regime (Unrestricted model): Before Date *T*<sub>SB</sub>.. Keep RSS<sub>1</sub>. After Date T<sub>SB</sub>... Keep RSS<sub>2</sub>.  $\Rightarrow$  RSS<sub>U</sub> = RSS<sub>1</sub> + RSS<sub>2</sub>.

(3) Run a standard F-test (testing Restricted vs. Unrestricted models):  $F = \frac{(RSS_R - RSS_U)/(k_U - k_R)}{(RSS_U)/(T - k_U)} = \frac{(RSS_R - [RSS_1 + RSS_2])/k}{(RSS_1 + RSS_2)/(T - 2k)}$ 

**Example:** We test if the Oct 1973 oil shock in quarterly GDP growth rates had an structural change on the GDP growth rate model. We use Federal Reserve data (FRED) from 1947:I to 2023:II (T = 304).

We model the GDP growth rate,  $y_t$ , with an AR(1) model, that is, GDP growth rate depends only on its own lagged growth rate:

 $y_t = \beta_0 + \beta_1 y_{t-1} + \varepsilon_t$ GDP da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/GDP q.csv", head=TRUE, sep=",") x date <- GDP da\$DATE x  $gdp \leq GDP da GDP$ x dummy <- GDP da\$D73  $T \leq - \text{length}(x \text{ gdp})$ t s <- 108  $\# T_{SB} = Oct \ 1973$  $\ln gdp \le \log(x gdp[-1]/x gdp[-T])$  $T \leq - length(lr gdp)$  $\ln gdp0 \leq \ln gdp[-1]$  $\ln gdp1 \leq \ln gdp[-T]$ t s <- t s -1 # Adjust t s (we lost the first observation)  $y \le lr gdp0$  $x1 \leq lr gdp1$  $T \leq - length(y)$ 

x0 <- matrix(1,T,1) x <- cbind(x0,x1) k <- ncol(x)				
<pre># Restricted Model (Pooling fit_ar1 &lt;- lm(lr_gdp0 ~ lr_g summary(fit_ar1) e_R &lt;- fit_ar1 \$residuals RSS_R &lt;- sum(e_R^2)</pre>	all data) dp1)	<ul> <li># AR(1) Model</li> <li># Restricted Regression</li> <li># regression residuals, e</li> <li># RSS Restricted</li> </ul>		
# Unrestricted Model (Two n	regimes)			
y_1 <- y[1:t_s] x_u1 <- x[1:t_s,] fit_ar1_1 <- lm(y_1 ~ x_u1 - e1 <- fit_ar1_1\$residuals RSS1 <- sum(e1^2)	- 1)	# AR(1) Regime 1 # Regime 1 regression residuals, e # RSS Regime 1		
$kk = t_s+1$ y_2 <- y[kk:T] x_u2 <- x[kk:T]		# Starting date for Regime 2		
$\begin{array}{l} x\_u2 < x[kk,1,j] \\ fit\_ar1\_2 <- lm(y\_2 ~ x\_u2 - e2 <- fit\_ar1\_2\$residuals \\ RSS2 <- sum(e2^2) \end{array}$	- 1)	<ul><li># AR(1) Regime 2</li><li># Regime 2 regression residuals, e</li><li># RSS Regime 2</li></ul>		
F <- ((RSS_R - (RSS1+RSS2))/k)/((RSS1+RSS2)/(T - 2*k)) > F				
[1] 4.391997 $p_{val} <-1 - pf(F, df1 = 2, df)$	$2 = T - 2^{*}k$	# p-value of F_test		
[1] <b>0.0131817</b>	$\Rightarrow$ small p-va	lues: Reject H <sub>0</sub> (No structural change). ¶		

**Example**: 3 Factor Fama-French Model for IBM (continuation) Q: Did the dot.com bubble (end of 2001) affect the structure of the FF Model? Sample: Jan 1973 – June 2020 (T = 569). Pooled RSS = 1.9324Jan 1973 – Dec 2001 RSS = RSS<sub>1</sub> = 1.3307 (T = 342) Jan 2002 – June 2020 RSS = RSS<sub>2</sub> = 0.5791 (T = 227)

	Constant	Mkt-rf	SMB	HML	RSS	Т
1973-2020	-0.0051	0.9083	-0.2125	-0.1715	1.9324	569
1973-2001	-0.0038	0.8092	-0.2230	-0.1970	1.3307	342
2002 - 2020	-0.0073	1.0874	-0.1955	-0.3329	0.5791	227

$$F = \frac{[RSS_R - (RSS_1 + RSS_2)]/k}{(RSS_1 + RSS_2)/(T - k)} = \frac{[1.9324 - (1.3307 + 0.5792)]/4}{(1.3307 + .05791)/(569 - 2*4)} = 1.6627$$
  

$$\Rightarrow \text{ Since } F_{4,565,.95} = 2.39, \text{ we cannot reject } H_0.$$

<u>Conclusion</u>: We do not find evidence that the 3-factor F-F model IBM excess returns suffered a structural break in January 2002.

<u>R Note</u>: The R package *sctrucchange* estimates the Chow test. (As usual, you need to install package first.)

>library(sctrucchange)
> t\_s <- 342
> sctest(ibm\_x ~ Mkt\_RF + SMB + HML, type = "Chow", point = t\_s)

Chow test

data:  $ibm_x \sim Mkt_RF + SMB + HML$ F = 1.6627, p-value = 0.2787. ¶

#### **Functional Form: Structural Change – Modeling with Dummy Variables**

Under the H<sub>0</sub> (No *structural change*), we can pool the data into one model. That is, the parameters are the same under both regimes. We fit the same model for all *i*, for example, with vector  $x_i$  of explanatory variables:

$$y_i = \beta_0 + \beta_1' \boldsymbol{x_i} + \boldsymbol{\varepsilon_i}$$

If the Chow test rejects  $H_0$ , we need to reformulate the model. A typical reformulation includes a dummy variable ( $D_{SB,i}$ ). For example:

$$y_i = \beta_0 + \beta_1' \boldsymbol{x}_i + \beta_2 D_{SB,i} + \gamma_1' \boldsymbol{x}_i D_{SB,i} + \varepsilon_i$$

where

 $D_{SB,i} = 1$  if observation *i* occurred after  $T_{SB} = 0$  otherwise.

**Example:** Given the Chow Test, we want to incorporate the effect of the October 1973 oil shock in GDP growth rates. We include a dummy variable in the model, say  $D_{73}$ :

 $D_{73,i} = 1$  if observation *i* occurred after October 1973 = 0 otherwise.

Then,

$$y_i = \beta_0 + \boldsymbol{\beta}_1' \boldsymbol{x}_i + \beta_2 D_{73,i} + \boldsymbol{\gamma}_1' \boldsymbol{x}_i D_{73,i} + \varepsilon_i$$

In the model, the oil shock affects the constant and the slopes. - Constant:

> Before oil shock  $(D_{73}:=0)$ :  $\beta_0$ After oil shock  $(D_{73}:=1)$ :  $\beta_0 + \beta_2$

- Slopes:

Before oil shock  $(D_{73}:=0)$ :  $\beta_1$ After oil shock  $(D_{73}:=1)$ :  $\beta_1 + \gamma_1$ 

We can estimate the above model and do an F-test to test if H<sub>0</sub> (No *structural change*):  $\beta_2 = 0 \& \gamma_1 = 0$ .

**Example:** We introduce an Oct 1973 dummy in the AR(1) GDP growth rate model.

Coefficients:

EstimateStd. Errort valuePr(>|t|)(Intercept)0.0091390.0019394.7123.75e-06 \*\*\* $lr_gdp1$ 0.4570110.0907165.0388.15e-07 \*\*\*D730.0034990.0023621.4820.13947 $\Rightarrow$  no significant effect on constant $lr_gdp1_D73$ -0.3160050.114197-2.7670.00601 \*\* $\Rightarrow$  significant effect on slope---Signif. codes:0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 0.01234 on 300 degrees of freedom Multiple R-squared: 0.09523, Adjusted R-squared: 0.08618 F-statistic: 10.53 on 3 and 300 DF, p-value: 1.333e-06

<u>Conclusion</u>: After the oil shock, the slope significantly changed from 0.457011 to 0.141006 (= 0.457011 + (-0.316005)).

### **Chow Test: Structural Change – Asymptotics and Wald Test**

Before, when we presented the Chow test, we used the F-distribution, which will be appropriate under (A5). In general, we rely on the asymptotic distribution -i.e., we do not rely on (A5).

It is common to approximated the distribution of the Chow test, under H<sub>0</sub>, (& if the number of observations pre- and post-break are large), with

$$J * F \xrightarrow{d} \chi_J^2$$
 (sometimes written as  $F \xrightarrow{d} \chi_J^2 / J$ ).

• It is also possible to do a Wald test to test H<sub>0</sub>, using only the unrestricted estimators. Steps:

1) Run two separate OLS, one for each regime (Unrestricted model):

Before Date $I_{SB}$ . Keep <b>b</b> <sub>1</sub> & Var  <b>b</b>	1]
---	----

After Date  $T_{SB}$ . Keep  $\mathbf{b}_2 \& \operatorname{Var}[\mathbf{b}_2]$ 

2) Compute the Wald test:

$$W = (\mathbf{b}_1 - \mathbf{b}_2)' \left\{ \frac{T}{T_{SB}} * \operatorname{Var}[\mathbf{b}_1] + \frac{T}{(T - T_{SB})} * \operatorname{Var}[\mathbf{b}_2] \right\}^{-1} (\mathbf{b}_1 - \mathbf{b}_2)$$

where T is the sample size and  $T_{SB}$  the observation of tested date.

Under H<sub>0</sub> (& if the number of observations pre- and post-break are large), the Wald test follows:  $W \xrightarrow{d} \chi_I^2$ 

### **Chow Test: Structural Change – Unknown Break**

The previous example, computes the Chow test assuming that we know exactly when the break occurred –say, October 73, Dec 2001 or January 2009.

That is, the results are *conditional* on the assumed breaking point.

In general, breaking points are unknown, we need to estimate them.

One quick approach is to do a rolling Chow test –that is we run the Chow test for all dates in the sample– and pick the date that maximizes the F-test. However, technically speaking, we cannot run the Chow test for all observations in our sample. We do not have enough observations to estimate reliable parameters on both sides of the potential breaking points.

A solution to this problem is to "trim" the data, we start to check for a breaking point at date  $\tau_{min}$ , and we finish to check at date  $\tau_{max}$ . Usually, we set  $\tau_{min}$  and  $\tau_{max}$  by leaving a percentage,  $\pi$ , of the initial of observations and final observations. We call  $\pi$  the "trimming parameter," usually set to equal 10% or 15% of the observations.

This rolling Chow test was proposed by Quandt (1958):  $QLR_T = \max_{\tau \in \{\tau_{min}, \dots, \tau_{max}\}} F_T(\tau)$ 

The max (supremum) is taken over all potential breaks in ( $\tau_{min}$ ,  $\tau_{max}$ ). For example,  $\tau_{min} = T^*.15$ ;  $\tau_{max} = T^*.85$ ; then we trim 30% of the observations ( $\pi_0 = 15\%$  in each side) to run the test. That is, we are looking for the structural break on the middle 70% of the sample.

It is also possible to run the Wald test version of the Chow test for all possible dates, again, selecting the date that maximizes

$$QLR_T = \max_{\tau \in \{\tau_{min}, \dots, \tau_{max}\}} W_T(\tau)$$

• The first  $QLR_T$  is called the **SupF** test, the second the **SupW**.

The problem with testing over all potential breaks is that the technical conditions under which the asymptotic distribution is derived are not met in this setting (the F-test are correlated, they are not independent).

Andrews (1993) showed that under appropriate conditions, the QLR statistic, also known as Suptest (W, F, LR) statistic, has a *non-standard limiting distribution* ("*non-standard*" = no existing table; needs a new one).

The distribution depends on the number of parameters of the model, k, which we are tested for stability, trimming values,  $\pi_0$ , which only affect the distribution through  $\lambda = (1-\pi_0)^2/\pi_0^2$ .

• And rews (1993) tabulated the non-standard distribution of the SupW for different k,  $\alpha$ , and trimming values ( $\pi_0$ ).

For example, for k=4,  $\pi_0 = \tau_{\min}/T = (1-\tau_{\max}/T) = .15$ , &  $\alpha=.05$ , the critical value is = 16.45. For the k=2 and same trimming and  $\alpha$  values, the critical value is = 11.79.

Critical values of the QLR test Distribution, taken from Andrews (1993). <u>Note</u>: p = # of parameters (*k*),  $\pi_0 =$  trimming value. (Ignore  $\lambda$ .) Andrews (2003) issued a slightly corrected Table.

<u>Note</u>: It is usual to test the **SupF**, using the critical values of **SupW**, by dividing the **SupW** critical values by k. Then, for the critical value for the SupF test for k=2,  $\pi_0 = .15$  and  $\alpha = .05$ , the critical value is **5.89** (= **11.79**/2).

#### TABLE I

ASYMPTOTIC CRITICAL VALUES

			p = 1			p = 2			p = 3			p = 4			p = 5	
$\pi_0$	λ	10%	5%	1%	10%	5%	1%	10%	5%	1%	10%	5%	1%	10%	5%	1%
.50	1.00	2.71	3.84	6.63	4.61	5.99	9.21	6.25	7.81	11.34	7.78	9.49	13.28	9.24	11.07	15.09
.49	1.08	3.47	4.73	7.82	5.42	6.86	10.30	7.19	8,83	12.58	8.93	10.63	14.64	10.39	12.28	16.34
.48	1.17	3.79	5.10	8.26	5.80	7.31	10.71	7.64	9.29	13.05	9.42	11.17	15.17	10.96	12.88	16.83
.47	1.27	4.02	5.38	8.65	6.12	7.67	11.01	7.98	9.62	13.39	9.82	11.63	15.91	11.40	13.27	17.32
.45	1.49	4.38	5.91	9.00	6.60	8.11	11.77	8.50	10.15	14.23	10.35	12.27	16.64	12.05	14.00	18.06
.40	2.25	5.10	6.57	9.82	7.45	9.02	12.91	9.46	11.17	14.88	11.39	13.32	17.66	13.09	15.16	19.23
.35	3.45	5.59	7.05	10.53	8.06	9.67	13.53	10.16	12.05	15.71	12.10	14.12	18.54	13.86	15.93	19.99
.30	5.44	6.05	7.51	10.91	8.57	10.19	14.16	10.76	12.58	16,24	12,80	14.79	19.10	14.58	16.48	20.67
.25	9.00	6.46	7.93	11.48	9.10	10.75	14.47	11.29	13.16	16.60	13.36	15.34	19.78	15.17	17.25	21.39
.20	16.00	6.80	8.45	11.69	9.59	11.26	15.09	11.80	13.69	17.28	13.82	15.84	20.24	15.63	17.88	21.90
.15	32.11	7.17	8.85	12.35	10.01	11.79	15.51	12.27	14.15	17,68	14.31	16.45	20.71	16.20	18.35	22.49
.10	81.00	7.63	9.31	12.69	10.50	12.27	16.04	12.81	14.62	18.28	14.94	16.98	21.04	16.87	18.93	23.34
.05	361.00	8.19	9.84	13.01	11,20	12.93	16,44	13.47	15.15	19.06	15.62	17.56	21.54	17.69	19.61	24.18
			p = 6			p = 7			p = 8			p = 9			p = 10	
$\pi_0$	λ	10%	5%	1%	10%	5%	1%	10%	5%	1%	10%	5%	1%	10%	5%	1%
.50	1.00	10.64	12.59	16.81	12.02	14.07	18.48	13.36	15.51	20.09	14.68	16.92	21.67	15.99	18.31	23.21
.49	1.08	11.81	13.74	18.32	13.27	15.52	19.93	13.29	15.63	20.53	16.17	18.56	23.05	17.35	19.79	24.62
.48	1.17	12.42	14.45	19.12	13.92	16.14	20.64	13.89	16.31	21.14	16.82	19.25	23.83	18.08	20.35	25.75
.47	1.27	12.90	14.86	19.64	14.32	16.63	21.14	14.43	16.74	21.72	17.26	19.74	24.80	18.67	20.92	26.43
.45	1.49	13.53	15.59	20.45	14.97	17.38	22.32	15.05	17.53	22.28	18.10	20.59	25.52	19.39	21.78	27.30
.40	2.25	14.71	16.91	21.60	16,23	18.41	23.35	16.26	18.73	23.63	19.56	22.12	26.86	20.74	23.15	28.86
.35	3.45	15.56	17.75	22.33	17.09	19.34	24.10	17.06	19.46	24.64	20.49	22.93	27.77	21.87	24.17	29.76
.30	5.44	16.32	18.46	23.06	17.74	20.01	24.86	17.90	20.36	25,64	21.27	23.65	28.50	22.73	25.05	30.74
.25	9.00	17.00	19.07	23.65	18.38	20.63	25.11	18.61	20.95	26.10	21.93	24.31	29.23	23.32	25.80	31.32
.20	16.00	17.56	19.64	24.27	19.04	21.07	25.72	19.17	21.47	26.76	22.54	24.91	29.92	24.00	26,42	31.98
.15	32.11	18.12	20.26	24.79	19,69	21.84	26.23	19.82	22.13	27.25	23.15	25.47	30.52	24.62	27.03	32.33
.10	81.00	18.78	20.82	25.21	20.32	22.51	26.91	20.45	22.87	27.69	23.77	26.16	31.15	25.39	27.87	32.95
.05	361.00	19.49	21.56	25.96	21.02	23.22	27.53	21.23	23.60	28.77	24.64	26.94	31.61	26.24	28.63	33.86

**Example (continuation)**: We search for breaking points for GDP growth rate in AR(1) model. We use package *desk*. (You can also use library *strucchange*, but it runs the SupW (F= SupW/2), you need to use Andrews (1993) Table.)

```
library(desk)
pie <- .15
T0 <- round(T * pie)
T1 <- round(T *(1-pie))
my.qlr <- qlr.test(lr_gdp0 ~ lr_gdp1, from = T0, to = T1, sig.level = 0.05, details = TRUE)
> my.qlr  # Print test results
```

```
QLR-Test for structural breaks at unknown date
```

Hypotheses:

HO: No break in t = 46...262 Some break in t = 46...262

Test resul <sup>.</sup> f.value <mark>21.1441</mark>	ts: lower.cv 5.86	upper.cv 6.085	p.value < <mark>1e-04</mark>	sig.level 0.05	HO rej.
my.qlr\$breakpoint			# Extract b	reakpoint obs	ervation
x_date[my.qlr\$breakpoint]			# Print date	e	

Below, we plot all F-tests starting at T\*15 (observation 46):



QLR Test: GDP growth rate AR(1) Model - 1947-2024

Maximum F is 21.1441 occurs in Jan 2009 (observation #250). Then, using Andrews' tabulated SupWcritical value of 11.79 and dividing by k=2, to obtain 5.89, we have  $\widehat{QLR} = 21.1441 > 5.89 (=11.79/2) \implies \text{Reject H}_0 \text{ at } 5\% \text{ level.}$ 

<u>**R** Note</u>: The function qlr.test (package *desk*) also computes the p-value, using Hansen's (1997) approximation. You can use the p-value, as usual, to evaluate (in this case, reject)  $H_0$ .

<u>Conclusion</u>: We find strong evidence that the AR(1) GDP growth model suffered a structural break during the sample. The QLR tests points toward January 2009 as the date of the structural break, not October 1973!.  $\P$ 

**Example:** We search for breaking points for IBM returns in the 3-factor FF model.

No break	in t = 92	H0: 519 Sc	ome break	in t = 92	H1: .519
Test resul f.value	ts: lower.cv	upper.cv	p.value	sig.level	HO rej

Below, we plot all starting at T\*15:

QLR Test: IBM 3-factor FF Model - 1973 - 2023



Maximum F is 4.5302 occurs in September 2012 (observation #477), the p-value is given as 0.0243. That means, we reject H. Using Andrews' tabulated critical value for SupW to get 4.11, we have

 $\widehat{QLR} = 4.5302 > 4.11 \ (=16.45/4) \implies \text{reject H}_0 \text{ at 5\% level.}$ 

<u>Conclusion</u>: We do find evidence that the 3-factor F-F model IBM excess returns suffered a structural break during the sample.  $\P$ 

### Chow Test: Structural Change – Script in R

Chow Test for different breaking points, starting at T1. y <- ibm\_x; x1 <- Mkt\_RF x2 <- SMB x3 <- HML T <- length(x1) x0 <- matrix(1,T,1) x <- cbind(x0,x1,x2,x3) k <- ncol(x) b <- solve(t(x)%\*% x)%\*% t(x)%\*%y #b

# b = (X'X)-1 X' y (OLS regression)

```
# regression residuals, e
e <- y - x%*%b
RSS R <- as.numeric(t(e)%*%e)
                                                     # RSS for Restricted (no structural change)
T1 \le round(T * 1/5)
                                                     # Trim .20 of data
t <- T1
                                                     # t will be the counter for loop. Starts at T1.
T2 \le round(T * 4/5)
                                                     # Trim .20 of data
T sam <- T2 - T1
All F \leq matrix(0,T \text{ sam},1)
                                                     # Matrix to accumulate the (T2-T1) F-tests
while (t \le T2) {
                                                     # Start while loop with counter t
y 1 < -y[1:t]
x u1 <- x[1:t,]
b 1 \le \text{solve}(t(x \ u1)) \% \% x \ u1) \% \% t(x \ u1) \% \% y \ 1
                                                             \# b = (X'X)-1 X' y (OLS regression)
e1 <- y 1 - x u1%*%b 1
                                                             # regression residuals, e
RSS1 <- as.numeric(t(e1)%*%e1)
                                                             # RSS for regime 1
kk = t+1
y_2 <- y[kk:T]
x u_2 <- x[kk:T,]
b_2 \le solve(t(x_u2)\%*\% x_u2)\%*\% t(x_u2)\%*\%y_2
                                                             \# b = (X'X)-1 X' y (OLS regression)
e2 <- y 2 - x u2%*%b 2
                                                             # regression residuals, e
RSS2 <- as.numeric(t(e2)\%*\%e2)
                                                             # RSS for regime 2
F \le ((RSS_R - (RSS1 + RSS2))/k)/((RSS1 + RSS2)/(T - 2*k))
kt <- t - T1 +1
                                                             # kt is an index that start at 1
All F[kt] \leq F
                                                             # add F-test to All F according to kt
t = t+1
}
```

plot(All\_F, col="red",ylab ="F-test", xlab ="Break Point") title("F-test at different Break Points") F\_max <- max(All\_F)

# Find the maximum F-test (QLR)

### **Chow Test: Structural Change – Remarks**

The results are *conditional* on the breaking point -say, October 73 or Dec 2001.

The breaking point is usually unknown. It needs to be estimated.

It can deal only with one structural break -i.e., two categories!

The number of breaks is also unknown.

Characteristics of the data (heteroscedasticity –for example, regimes in the variance- and unit roots (high persistence) complicate the test.

In general, only asymptotic (consistent) results are available.

Related to the next section, missing structural breaks in deterministic parameters (intercepts, trends, etc.) can be a cause of forecast failure –see simulations by Clements and Hendry (1999).

There are many modern tests that take care of these issues, but usually also with *non-standard* distributions.

### **Forecasting and Prediction**

Objective: Forecast Distinction: Ex post vs. Ex ante forecasting - Ex post: RHS data are observed - Ex ante (true forecasting): RHS data must be forecasted

Prediction and Forecast

- Prediction: Explaining an outcome, which could be a future outcome.

- Forecast: A particular prediction, focusing in a future outcome.

Example:	Prediction:	Given <b>x</b> <sup>0</sup>	$\Rightarrow$ predict $y^0$ .
	Forecast:	Given $\mathbf{x}^{0}_{t+1}$	$\Rightarrow$ predict $y_{t+1}$ .

• Two types of predictions:

- In sample (prediction): The expected value of y (in-sample), given the estimates of the parameters. In sample prediction produces fitted values,  $\hat{y}$ .

- Out of sample (forecasting): The value of a future y that is not observed by the sample.

Notation: Let T be the forecast origin and l is the forecast horizon.

- Prediction for T made at T:  $\hat{Y}_T$ .

- Forecast for T+l made at  $T: \hat{Y}_{T+l}, \hat{Y}_{T+l|T}, \hat{Y}_{T}(l)$ .

-  $\hat{Y}_T(l)$ : *l-step ahead* forecast = Forecasted value  $Y_{T+l}$  at time T.

• Any prediction or forecast needs an information set,  $I_T$ . This includes data, models and/or assumptions available at time *T*. The predictions and forecasts will be conditional on  $I_T$ .

For example, in-sample,  $I_T = \{x^0\}$  to predict  $y^0$ .

Or in a time series context,  $I_T = {x^0_{T-1}, x^0_{T-2}, ..., x^0_{T-q}}$  to predict  $y_{t+l}$ .

Then, the forecast is just the conditional expectation of  $Y_{T+l}$ , given the observed sample:  $\hat{Y}_{T+l} = E[Y_{T+l}|X_T, X_{T-1}, \dots, X_1]$ 

**Example**: If  $X_T = Y_T$ , then, the one-step ahead forecast is:  $\hat{Y}_{T+1} = E[Y_{T+1}|Y_T, Y_{T-1}, \dots, Y_1]$ .

• The conditional expectation of  $Y_{T+l}$  is, in general, based on a model, the experience of the forecaster or a combination of both.

**Example**: We base the conditional expectation for excess returns on the 3 FF factor model:  $\hat{Y}_{T+l} = E[(\beta_0 + \beta_1 (r_{m,t+l} - r_f) + \beta_2 SMB_{t+l} + \beta_3 HML_{t+l}|I_T$ 

<u>Note</u>: The forecast of  $Y_{T+l}$  also needs a forecast for the driving variables in the model. We need a forecast for  $E[(r_{m,t+l} - r_f)| I_T]$ ,  $E[SMB_{t+l}| I_T]$ , and  $E[HML_{t+l}|I_T]$ .

In general, we will need a model for  $\hat{X}_{T+l}$ . Things can get complicated very quickly.

Keep in mind that the forecasts are a random variable. Technically speaking, they can be fully characterized by a pdf.

In general, it is difficult to get the pdf for the forecast. In practice, we get a point estimate (the forecast) and a C.I.

• Q: What is a good forecast? We need metrics to evaluate the forecasting performance of different models. In general, the evaluation of forecasts relies on MSE.

Later in this class, when we cover time series (Brooks Chapter 6), we go deeper into forecasting.

# Forecasting and Prediction: Variance-bias Trade-off

We start with general model (DGP): (A1) DGP:  $y = f(\mathbf{X}, \theta) + \boldsymbol{\varepsilon}$ .

Given  $\mathbf{x}^0$ , we predict  $y^0$ , using the expectation:  $E[y|\mathbf{X}, \mathbf{x}^0] = f(\mathbf{x}^0, \theta)$ ; which we estimate with  $\hat{y}^0 = f(\mathbf{x}^0, \hat{\theta})$ .

The realization  $y^0$  is just:

$$y^0 = f(\mathbf{x}^0, \theta) + \varepsilon^0$$

With  $y^0$  observed, we compute the prediction error:  $\hat{y}^0 - y^0$  and its associated expected squared error, which can be written as:

$$E[(\hat{y}^0 - y^0)^2] = E[(\hat{y}^0 - (f(\mathbf{x}^0, \theta) + \varepsilon^0))^2]$$
  
= 
$$E[(f(\mathbf{x}^0, \hat{\theta}) - f(\mathbf{x}^0, \theta)) - \varepsilon^0)^2]$$
  
= 
$$Var[\hat{y}^0] + [Bias(\hat{y}^0)]^2 + Var[\varepsilon]$$

We want to minimize this squared error. Note that there is nothing a forecaster can do regarding the last term, called the *irreducible error*. All efforts are devoted to minimize the sum of a variance and a squared bias (the MSE). This creates the *variance-bias trade-off* in forecasting.

It is possible that biased forecast can produce a lower MSE than an unbiased one. In this lecture, we based our forecasts on OLS estimates, which under the CLM assumptions, produce unbiased forecast.

<u>Note</u>: The variance-bias trade-off is always present in forecasting. In general, more flexible models have less bias and more variance. The key is to pick an "optimal" mix of both.

### **Forecasting and Prediction: Point Estimate**

Prediction: Given  $\mathbf{x}^0 \Rightarrow$  predict  $\mathbf{y}^0$ .

Given the CLM, we h	nave:
Expectation:	$\mathrm{E}[\mathbf{y} \mathbf{X},\mathbf{x^0}] = \mathbf{\beta'x^0};$
Predictor:	$\widehat{y}^0 = \mathbf{b'} \mathbf{x}^0$
Realization:	$y^0 = \boldsymbol{\beta' x^0} + \boldsymbol{\epsilon}^0$

<u>Note</u>: The predictor includes an estimate of  $\varepsilon^0$ :  $\hat{y}^0 = \mathbf{b}^2 \mathbf{x}^0 + \text{estimate of } \varepsilon^0$ . (Estimate of  $\varepsilon^0=0$ , but with variance.)

• Associated with the prediction (a point estimate), there is a forecast error:

 $\hat{y}^0 - y^0 = \mathbf{b'x^0} - \boldsymbol{\beta'x^0} - \boldsymbol{\varepsilon}^0 = (\mathbf{b} - \boldsymbol{\beta})'\mathbf{x^0} - \boldsymbol{\varepsilon}^0$ 

and a variance:

$$\begin{aligned} \operatorname{Var}[(\hat{y}^0 - y^0) | \mathbf{x}^0] &= \operatorname{E}[(\hat{y}^0 - y^0)' (\hat{y}^0 - y^0) | \mathbf{x}^0] \\ &= \mathbf{x}^{0'} \operatorname{Var}[(\mathbf{b} - \mathbf{\beta}) | \mathbf{x}^0] \mathbf{x}^0 + \sigma^2 \end{aligned}$$

**Example**: We have already estimated the 3 Factor Fama-French Model for IBM returns: > summary(fit\_ibm\_ff3)

	Estimate	Std. Error	t value	$Pr(\geq  t )$
(Intercept)	-0.005089	0.002488	-2.046	0.0412 *
Mkt_RF	0.908299	0.056722	16.013	<2e-16 ***
SMB	-0.212460	0.084112	-2.526	0.0118 *
HML	-0.171500	0.084682	-2.025	0.0433 *

Suppose we are given  $\mathbf{x}^0 = [1.0000 - 0.0189 - 0.0142 - 0.0027]$ Then,  $\hat{\alpha}^0 = -0.005080 + 0.008200 * (-0.0180) - 0.212460 * -0.0142 - 0.171500 * (-0.0180) - 0.0142 - 0.0142 - 0.0142 - 0.0171500 * (-0.0180) - 0.0142 - 0.0042 - 0.00$ 

 $\hat{y}^0 = -0.005089 + 0.908299 * (-0.0189) - 0.212460 * -0.0142 - 0.171500 * (-0.0027) = -0.01877582$ 

Suppose we observe  $y^0 = 0.1555214$ . Then, the forecast error is  $\hat{y}^0 - y^0 = -0.01877582 - 0.1555214 = -0.1742973$ 

> ef\_0 [,1] [1,] -0.1742973. ¶

# **Forecasting and Prediction: Confidence Intervals**

How do we estimate the uncertainty behind the forecast? Form a confidence interval.

Two cases: (1) If  $\mathbf{x}^0$  is given –i.e., constants. Then,  $\operatorname{Var}[\hat{y}^0 - y^0 | \mathbf{x}^0] = \mathbf{x}^{0'} \operatorname{Var}[\mathbf{b} | \mathbf{x}^0] \mathbf{x}^0 + \sigma^2$  $\Rightarrow$  Form confidence interval as usual.

<u>Note</u>: In out-of-sample forecasting,  $\mathbf{x}^{0}$  is unknown, it has to be estimated.

(2) If  $\mathbf{x}^0$  has to be estimated, then we use a random variable. What is the variance of the product? One possibility: Use bootstrapping.

• Assuming  $\mathbf{x}^0$  is known, the variance of the forecast error is  $\sigma^2 + \mathbf{x}^0$ ,  $\operatorname{Var}[\mathbf{b}|\mathbf{x}^0] = \sigma^2 + \sigma^2[\mathbf{x}^0, (\mathbf{X}^*\mathbf{X})^{-1}\mathbf{x}^0]$ 

If the model contains a constant term, this is

$$\operatorname{Var}[e^{0}] = \sigma^{2} \left[ 1 + \frac{1}{N} + \sum_{j=1}^{K-1} \sum_{k=1}^{K-1} (x_{j}^{0} - \bar{x}_{j})(x_{k}^{0} - \bar{x}_{k})(Z'M^{0}Z)^{jk} \right]$$

(where Z is X without  $x_1=i$ ). In terms squares and cross products of deviations from means.

<u>Note</u>: Large  $\sigma^2$ , small *N*, and large deviations from the means, decrease the precision of the forecasting error.

<u>Interpretation</u>: Forecast variance is smallest in the middle of our "experience" and increases as we move outside it.

Then, the  $(1 - \alpha)$ % C.I. is given by:  $[\hat{y}^0 \pm t_{T-k,\alpha/2} * \operatorname{sqrt}(\operatorname{Var}[e^0])]$ 

As  $\mathbf{x}^0$  moves away from its mean, the C.I increases, this is known as the "butterfly effect."



**Example (continuation):** We want to calculate the variance of the forecast error: for the given  $\mathbf{x}^0 = [1.0000 - 0.0189 - 0.0142 - 0.0027]$ Recall we got  $\hat{\mathbf{y}}^0 = \mathbf{b}^* \mathbf{x}^0 = -0.01877587$ 

Then,

Estimated Var $[\hat{y}^0 - y^0 | \mathbf{x}^0] = \mathbf{x}^{0'}$  Var $[\mathbf{b} | \mathbf{x}^0] = \mathbf{x}^{0'}$  Var $[\mathbf{b} | \mathbf{x}^0]$ 

```
var_ef_0 <- t(x_0)%*% Var_b%*% x_0 + Sigma2
> var_ef_0
       [,1]
[1,] 0.003429632
> sqrt(var_ef_0)
       [,1]
[1,] 0.05856306
```

<u>Check</u>: What is the forecast error if  $\mathbf{x}^0 = \text{colMeans}(\mathbf{x})$ ?

```
# (1-alpha)% C.I. for prediction (alpha = .05)
CI_lb <- y_f0 - 1.96 * sqrt(var_ef_0)
> CI_lb
>[1] -0.1335594
CI_ub <- y_f0 + 1.96 * sqrt(var_ef_0)
> CI_ub
>[1] 0.09600778
```

That is, CI for prediction: [-0.13356; 0.09601] with 95% confidence. A wide interval, which makes clear the uncertainty surrounding the point forecast:  $\hat{y}^0 = -0.01877587$ . ¶

# **Forecasting and Prediction – Model Validation**

*Model validation* refers to establishing the statistical adequacy of the assumptions behind the model -i.e., (A1)-(A5) in this lecture. Predictive power can be used to do model validation.

In the context of prediction and forecasting, model validation is done by fitting a model insample, but keeping a small part of the sample, the *hold-out-sample*, to check the accuracy of OOS forecasts.

Hold out sample: We estimate the model using only a part of the sample (say, up to time T<sub>1</sub>). The rest of the observations, the hold out sample, (T - T<sub>1</sub> observations) are used to check the predictive power of the model –i.e., the accuracy of predictions, by comparing  $\hat{y}^0$  with actual  $y^0$ .



Steps to measure forecast accuracy:

**Step 1**. Select a (long) part of the sample (say, first  $T_1$  observations) to estimate the parameters of the model. (Get in-sample forecasts,  $\hat{y}$ .) We call this sample, the *estimation period*.

**Step 2**. Keep a (short) part of the sample (say,  $(T - T_1)$  observations) to check the model's forecasting skills, This is the *validation step*. (Get OSS  $\hat{y}^0$ , but  $y^0$  is known.) Since  $y^0$  is known calculate true MSE or MAE. For example:

$$MSE = \frac{1}{(T - T_1)} \sum_{i=(T_1 + 1)}^{(T - T_1)} (\hat{y}_i^0 - y_i^0)^2$$

Step 3. If happy with Step 2, proceed to do out-of-sample forecasts.

<u>Note</u>: In the Machine Learning literature, the terminology used for model validation is slightly different.

**Step 1** is called "*training*," the data used (first  $T_1$  observations) are called *training data/set*. In this step, we estimate the parameters of the model, subject to the assumptions, for example, (A1)-(A4).

**Step 2** has the same name, *validation (or "single-split" validation)*. This step can be used to "*tune (hyper-)parameters.*" In our CLM, we can "tune" the model for departures of (A1)-(A4), for example, by including more or different variables (A1) and re-estimating the model accordingly using "training data" alone. We choose the model with lower MSE or MAE.

<u>Remark</u>: The idea of this step is to *simulate* out-of-sample accuracy. But, the "tuned" parameters selected in Step 2 are fed back to Step 1.

**Step 3** *tests* the true out-of-sample forecast accuracy of model selected by **Step 1 & Step 2**. This last part of the sample is called "*testing sample*."

# **Forecasting and Prediction – Cross Validation**

**Step 2** is used as a testing ground of the model before performing OOS forecasting. There are many ways to approach the validation step.

Instead of a single split, split the data in *K* parts. This is called *K-fold cross-validation*. For j = 1, 2, ..., *K*, use all folds but fold *j* to estimate model; use fold *j* to check model's forecasting skills by computing MSE,  $MSE_i$ . The *K*-fold CV estimate is an average of each fold MSE's:

$$CV_K = \frac{1}{K} \sum_{j=1}^K MSE_j$$

Usual choices for K are 5 & 10. This is an arbitrary choice.

Random and non-random splits of data can be used. The non-random splits are used for some special cases, such as qualitative data, to make sure the splits are "representative."

• Use a single observation for validation. This is called *leave-one-out cross-validation* (LOOCV), which is a special case of *K-fold cross-validation* with K = T. That is, use (T - 1) observations for estimation, and, then, use the observation left out, i = 1, ..., T, to compute  $MSE_{(-i)}$ , which is just  $(\hat{y}_{(-i)} - y_i)^2$ , where  $\hat{y}_{(-i)}$  is the prediction for observation *i* based on the full sample but observation *i*. Then, compute:

$$CV_n = \frac{1}{n} \sum_{i=1}^n MSE_{(-i)}$$

Instead of just one, it is possible to leave p observations for validation. This is called *leave-p-out cross-validation* (LpOCV).

<u>Remark</u>: In time series, since the order of the data matters, cross validation is more complicated. In general, rolling windows are used.

```
Example: We do cross-validation on the 3-factor Fama-French Model for IBM returns with
K=5:
y \le ibm x
ff cv data <- data.frame(Mkt RF, SMB, HML)
####### CV: Cross-Validation K-fold Code Function #######
CV \leq - function(dats, n.folds){
 folds <- list() # flexible object for storing folds
 fold.size <- nrow(dats)/n.folds
 remain <- 1:nrow(dats)
                              # all obs are in
 for (i in 1:n.folds){
  select <- sample(remain, fold.size, replace = FALSE)
                                                             #randomly sample fold size from
remaining obs)
  folds[[i]] <- select
                              # store indices ( write a special statement for last fold if 'leftover
points')
    if (i == n.folds)
   folds[[i]] <- remain
  }
remain <- setdiff(remain, select)
                                      # update remaining indices to reflect what was taken out
remain
 }
 results \leq- matrix(0,1,n.folds)
                                             # Vector to accumulate accuracy measures (MSE)
 for (i in 1:n.folds){
  # fold i
  indis <- folds[[i]]
                                             # unpack into a vector
  estim <- dats[-indis, ]
                                             # split into estimation (train) & validation (test) sets
  test <- dats[indis, ]</pre>
  lm.model \le lm(y[-indis] \sim ., data = estim)
                                                     # OLS with estimation data
  pred <- predict(lm.model, newdata = test)</pre>
                                                     # predicted values for fold not used
  MSE \le mean((y[indis] - pred)^2)
                                             # MSE (any other evaluation measure can be used)
                                             # Accumulate MSE in vector
  results[[i]]<- MSE
 }
 return(results)
}
CV ff 5 \leq CV(\text{ff cv data}, 5)
> CV ff 5
       [,1]
               [,2]
                        [,3]
                                 [,4]
                                         [,5]
[1,] 0.003578998 0.00329871 0.002058409 0.004202831 0.004524011
> mean(CV ff 5)
```

#### **[1] 0.00346262** K T <- length(y)

LOOCV\_ff\_5 <- CV(ff\_cv\_data, K\_T) > mean(LOOCV\_ff\_5) [1] 0.003516136. ¶

# **Evaluation of Forecasts: Measures of Accuracy**

Summary measures of out-of-sample forecast accuracy, after *m* forecasts:

Mean Error  $=\frac{1}{m}\sum_{i=T+1}^{T+m}(\hat{y}_i - y_i) = \frac{1}{m}\sum_{i=T+1}^{T+m}e_i$ Mean Absolute Error (MAE)  $=\frac{1}{m}\sum_{i=T+1}^{T+m}|\hat{y}_i - y_i| = \frac{1}{m}\sum_{i=T+1}^{T+m}|e_i|$ Mean Squared Error (MSE)  $=\frac{1}{m}\sum_{i=T+1}^{T+m}(\hat{y}_i - y_i)^2 = \frac{1}{m}\sum_{i=T+1}^{T+m}e_i^2$ Root Mean Square Error (RMSE)  $=\sqrt{\frac{1}{m}\sum_{i=T+1}^{T+m}e_i^2}$ Theil's U-stat  $=\frac{\sqrt{\frac{1}{m}\sum_{i=T+1}^{T+m}e_i^2}}{\sqrt{\frac{1}{T}\sum_{i=1}^{T+m}y_i^2}}$ 

Theil's U statistics has the interpretation of an R<sup>2</sup>. But, it is not restricted to be smaller than 1.

The lower the above criteria, say MSE, the better the forecasting ability of our model.

• An OOS R<sup>2</sup> can be computed as:  

$$R_{OOS}^{2} = 1 - \frac{MSE_{A}}{MSE_{N}}$$
with  $MSE_{A} = \sum_{t=1}^{m} (y_{t+\tau} - \hat{y}_{t+\tau})^{2}$   
 $MSE_{N} = \sum_{t=1}^{m} (y_{t+\tau} - \bar{y}_{t})^{2}$ 

where  $\tau$  is the forecasting horizon. (See Goyal and Welch (2008) for a well-known finance application.)

• Again, cross-validation measures can be used to evaluate forecasting performance.

**Example**: We want to check the forecast accuracy of the 3 FF Factor Model for IBM returns. We estimate the model using only 1973 to 2017 data (T=539), leaving 2018-2020 (m=30 observations) for validation of predictions.

T0 <-1 T1 <-539 T2 <-T1+1 y1 <-y[T0:T1] x1 <-x[T0:T1, ] fit2 <-lm(y1~x1-1) # End of Estimation Period # End of Estimation Period # Start of Validation Period# Estimation Period Regression from T0 to T1 b1 <- fit2\$coefficients
> summary(fit2)

# Extract OLS coefficients from regression

	Estimate	Std. Error	t value Pr(> t )
x1	-0.003848	0.002571	-1.497 0.13510
x1Mkt_RF	0.865579	0.059386	14.575 < 2e-16 ***
x1SMB	-0.224914	0.085505	-2.630 0.00877 **
x1HML	-0.230838	0.090251	-2.558 0.01081 *

We condition on the observed data (no model to predict FF factors used) from 2018: Jan to 2020: Jun.

x 0 < x[T2:T,]# Validation data y 0 < -y[T2:T]# Validation data y f0 <- x 0%\*% b1 # Forecast ef 0 < -y = 0# Forecasat error mse ef  $0 \le \operatorname{sum}(ef 0^2)/\operatorname{nrow}(x 0)$ # MSE > mse ef 0 [1] 0.003703207 mae ef  $0 \le sum(abs(ef 0))/nrow(x 0)$ # MAE > mae ef 0 [1] 0.04518326 That is,MSE = **0.003703207** MAE = **0.04518326** • Plot of actual IBM returns and forecasts. plot(y\_f0, type="l", col="red", main = "IBM: Actual vs. Forecast (2018-2020)", xlab = "Obs", ylab = "Forecast") lines(y 0, type = "l", col = "blue")legend("topleft", legend = c("Actual", "Forecast"), col = c("blue", "red"), lty = 1) vs. Forecast (2018-2020) IBM: Actual Actual Forecast 2 50 Forecast 8 99 0 5 10 15 20 25 30 Obs

From the plot, some forecasts are very good, some are very bad.

# **Evaluation of forecasts: Testing Accuracy**

Above, we have competing forecasting models and we computed measures of accuracy for each model. So far we have implicitly judged the model with the best (usually, the lower) measure of accuracy as the best forecasting model. But, measures of accuracy are RV, thus, in order to say one model forecasts better than other, we need a test.

Suppose two competing forecasting procedures produce a vector of errors:  $e^{(1)} \& e^{(2)}$ . We decide to use the expected MSE as the criterion to judge the forecasting accuracy of a model.

• We want to test	$H_0: MSE(1) = MSE(2)$
	H <sub>1</sub> : MSE(1) $\neq$ MSE(2).

<u>Assumptions</u>: forecast errors are unbiased, normal, and uncorrelated. If forecasts are unbiased, then MSE = Variance.

Consider, the pair of RVs:  $(e^{(1)} + e^{(2)}) \& (e^{(1)} - e^{(2)})$ . Now,  $E[(e^{(1)} + e^{(2)})(e^{(1)} - e^{(2)})] = \sigma_1^2 - \sigma_2^2$ 

That is, we test H<sub>0</sub> by testing that the two RVs are not correlated! Under H<sub>0</sub>,  $E[(e^{(1)} + e^{(2)})(e^{(1)} - e^{(2)})] = 0.$ This idea is due to Morgan, Granger and Newbold (MGN, 1977).

• There is a simpler way to do the MGN test. Let,

 $z_t = e^{(1)} + e^{(2)}$   $x_t = e^{(1)} - e^{(2)}$ (1) Do a regression:  $z_t = \beta x_t + \varepsilon_t$ (2) Test H<sub>0</sub>:  $\beta = 0 \implies$  a simple *t*-test.

The MGN test statistic is exactly the same as that for testing the null hypothesis that  $\beta = 0$  in this regression (recall:  $\mathbf{b} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y}$ ). This is the approach taken by Harvey, Leybourne and Newbold (1997).

If the assumptions are violated, these tests have problems.

A non-parametric HLN variation: Spearman's rank test for zero correlation between  $x_t$  and  $z_t$ .

**Example**: We produce IBM returns one-step-ahead forecasts for 2018-2020 using the 3 FF Factor Model for IBM returns:

$$(r_i - r_f)_t = \beta_0 + \beta_1 (r_m - r_f)_t + \beta_2 SMB_t + \beta_3 HML_t + \varepsilon_t$$

Taking expectations at time t+1, conditioning on time t information set,  $I_t = \{(Mkt_{Ret} - r_f)_t, SMB_t, HML_t\}$ 

 $E[(r_i - r_f)_{t+1}|I_t] = \beta_0 + \beta_1 E[(r_m - r_f)_{t+1}|I_t] + \beta_2 E[SMB_{t+1}|I_t] + \beta_3 E[HML_{t+1}|I_t]$ 

In order to produce forecast, we will make a naive assumption: The best forecast for the FF factors is the previous observation. Then,

$$\mathbb{E}[(r_i - r_f)_{t+1}|I_t] = \beta_0 + \beta_1 (r_m - r_f)_t + \beta_2 SMB_t + \beta_3 HML_t.$$

Now, replacing the  $\beta$  by the estimated **b**, we have our one-step-ahead forecasts. We produce one forecast at a time.

We compare the forecast accuracy relative to a random walk model for IBM returns. That is,  $E[(r_i - r_f)_{t+1}|I_t] = (r_i - r_f)_t$ 

Using R, we create the forecasting errors for both models and MSE: # By assumption on the X, it starts at T1. x  $01 \le x[T1:(T-1),]$  $y \ 0 < -y[T2:T]$ y f0 <- x 01%\*% b1 # b1 coefficients from previous regresssion  $\# e^{(2)}$ ef 0 < -y f0 - y 0 mse ef  $0 \le sum(ef 0^2)/nrow(x_0)$ > mse ef 0 # MSE(2) [1] 0.01106811  $# e^{(1)}$ ef rw  $0 \le y[T1:(T-1)] - y 0$ mse ef rw  $0 \le sum(ef rw 0^2)/nrow(x 0)$ > mse ef rw 0 # MSE(1)  $\leq =$  (1) is the higher MSE. [1] 0.02031009 • Now, we create  $z_t = e^{(1)} + e^{(2)}$ , &  $x_t = e^{(1)} - e^{(2)}$ . Then, regress:  $z_t = \beta x_t + \varepsilon_t$  and test H<sub>0</sub>:  $\beta = 0$ .  $z \text{ mgn} \leq ef rw 0 + ef 0$ x mgn <- ef rw 0 - ef 0 fit mgn <- lm(z mgn ~ x mgn) > summary(fit mgn) Coefficients: Estimate Std. Error t value Pr(>|t|)(Intercept) 0.05688 0.03512 1.619 0.117 2.77770 0.58332 4.762 5.32e-05 \*\*\* x mgn

<u>Conclusion</u>: We reject that both MSE are equal  $\Rightarrow$  MSE of RW is higher. ¶

### **Evaluation of forecasts: MSE/MAE?**

MSE and MAE are very popular criteria to judge the forecasting power of a model. However, it may not be the best measure for everybody.

Richard Levich's textbook compares forecasting services to the freely available forward rate. He finds that forecasting services may have some ability to predict direction (appreciation or depreciation).

For some investors, the direction is what really matters, since direction determines potential profits, not the error.

**Example:** Two forecasts: Forward Rate (Ft,T) and Forecasting Service (FS)

$$\begin{split} S_t &= .7330 \text{ USD/CAD} & (\text{Today's market spot rate.}) \\ F_{t,T=1-month} &= .7335 \text{ USD/CAD} & (\text{Today's market forward rate.}) \\ E_{FS,t} \left[S_{t+1-month}\right] &= .7342 \text{ USD/CAD.} &\Rightarrow \text{Today, FS forecasts an appreciation of CAD.} \\ \underline{\text{Investor's strategy}} \text{: Buy CAD forward if FS forecasts CAD appreciation, greater than the implied by the forward rate.} \end{split}$$

Based on the FS forecast, Ms. Sternin decides to buy CAD forward at Ft,1-month.

(A) Suppose that the CAD appreciates to  $S_{t+1} = .7390$  USD/CAD.

 $MAE_{FS} = |.7390 - .7342| = .0052 USD/CAD.$ 

Investor makes a profit of .7390 - .7335 = USD .055 USD.

(B) Suppose that the CAD depreciates to  $S_{t+1} = .7315$  USD/CAD.

 $MAE_{FS} = |.7315 - .7342| = .0027 USD/CAD. \Rightarrow smaller MAE!$ 

Investor takes a loss of .7315 - .7335 = USD -.0020.

<u>Conclusion</u>: A small forecast error is not that relevant for investor, the direction of the error matter much more.  $\P$ 

# **Forecasting Application: Fundamental Approach**

There are two pure approaches to forecasting. Based on how we select the "driving" variables  $X_t$ , we have:

- Fundamental (based on data considered fundamental)
- Technical analysis (based on data that incorporates only past prices)

• Fundamental Approach to Forecast Exchange Rates,  $S_t$  (USD/JPY)

Suppose we have an built an economic model,  $S_t = f(X_t)$ , where  $X_t$  is a dataset regarded as *fundamental* economic variables:

- GNP growth rate,
- Current Account,
- Interest rates,
- Inflation rates, etc.

The economic model usually incorporates:

- Statistical characteristics of data (seasonality, autocorrelation, etc.)
- Experience of the forecaster (what information to use, lags, etc.)

 $\Rightarrow$  Mixture of art and science.

Then, based on the economic model, we generate at time t a forecast for the exchange rate next period,  $S_{t+1}$ :

 $E_t[S_{t+1}] = E_t[f(X_{t+1})] = g(X_t).$ 

• Steps for building a forecast:

(1) The economic model (also called *structural model*) provides the structure for the forecasts. The economic model is the starting point of the fundamental approach.

(2) Once we selected the economic model, we proceed to estimate the parameter on the model. We need to collect data and decide on how to estimate the model (OLS, MLE, etc.).

(3) Then, we test the model. We have to make sure that we have a good model. If the model survives the tests, then we use the model to forecast.

(4) We evaluate the forecasts by comparing the economic model's performance with the performance of other models, for example, in our case, a simpler model, the Random Walk model (RWM). The RMW is found to be very good model for forecasting  $S_t$  in the short-run. The forecasts for the RWM are given by:

$$\operatorname{Et}[S_{t+1}] = S_t$$



FIGURE 6.2 - Steps for building a forecast

• Fundamental Forecasting: Steps (example:  $S_t = USD/JPY$ )

(1) Select a Model: Based on Theory (IFE, & Asset Approach) we model percentage changes in FX rates,  $e_{f,t} = \log(S_t) - \log(S_{t-1})$ :

 $\begin{aligned} e_{f,t} &= \beta_0 + \beta_1 (i_{US,t} - i_{JAP,t}) + \beta_2 (y_{US,t} - y_{JAP,t}) + \beta_3 (m_{US,t} - m_{JAP,t}) + \varepsilon_t \\ & E_t[e_{f,t+1}] = \beta_0 + \beta_1 E_t[i_{US,t+1} - i_{JAP,t+1}] + \beta_2 E[y_{US,t+1} - y_{JAP,t+1}] + \beta_3 E[m_{US,t+1} - m_{JAP,t+1}] \\ &\implies E_t[S_{t+1}] = S_{t+1}^F = S_t * (1 + E_t[e_{f,t+1}]) \end{aligned}$ 

(2) Collect data:  $S_t$ ,  $X_t$  (Interest rates (i), GDP growth rates (y) and money growth (m) data needed.)

(3) Estimation of Model (using *estimation period*): OLS  $\Rightarrow$  get **b**.

(4) Generate forecasts. Assumptions about  $X_t$  are needed.

$$E_{t}[\boldsymbol{X}_{t+1}] = \delta_{1} + \delta_{2} (\boldsymbol{X}_{t}) \quad \text{-an AR(1) model.}$$
  

$$\Rightarrow \quad E_{t}[\boldsymbol{e}_{f,t+1}] = E_{t}[\boldsymbol{X}_{t+1}]' \mathbf{b}$$
  

$$\Rightarrow \quad E_{t}[\boldsymbol{S}_{t+1}] = \boldsymbol{S}_{t}^{*} (1 + E_{t}[\boldsymbol{e}_{f,t+1}])$$

(5) Evaluation of Forecasts: MSE (& compare with RW's MSE).

Model's Forecast Error<sub>t+1</sub> =  $E_t[S_{t+1}] - S_{t+1}$ RW's Forecast Error<sub>t+1</sub> =  $S_t - S_{t+1}$ 

**Example:** (1) & (2) Based on above model, I collect quarterly data (FX\_USA\_JAP.csv) from 1978:II – 2020:II. I read the data and transform it to estimate model:

# Step (2) – Read Data	
FX_da <- read.csv("https://www.ba	auer.uh.edu/rsusmel/4397/FX_USA_JAP.csv", head=TRUE,
sep=",")	
us_I <- FX_da\$US_INF	<pre># Extract US Money growth (mus) data from FX_da</pre>
us_i <- FX_da\$US_I3M	# Extract US 3-mo Interest rate (ius) data
us_y <- FX_da\$US_GDP_g	# Extract US GDP growth (yus) data
us_tb <- FX_da\$US_CA_c	# Extract US Current account change (tbus) data
jp_I <- FX_da\$JAP_INF	# Extract Japan Inflation (I <sub>US</sub> ) data
jp_mg <- FX_da\$JAP_MI_c	# Extract Japan Money growth (mJP) data
jp_i <- FX_da\$JAP_I3M	# Read Japan 3-mo Interest rate (ijp) data
jp_y <- FX_da\$JAP_GDP_g	# Extract Japan GDP growth y <sub>JP</sub> ) data
jp_tb <- FX_da\$JAP_CA_c	# Extract Japan Current account change (tb <sub>JP</sub> ) data
e_f <- FX_da\$JPY.USD_c	# Extract changes in JPY/USD ( $e_{f,t}$ )
# Step (2) – Transform variables (cro	eate differentials)
inf_dif <- us_I - jp_I	<pre># Define inflation rate differential (inf_dif)</pre>
int_dif <- us_i - jp_i	<pre># Define interest rate differential (int_dif)</pre>
mg_dif <- us_mg - jp_mg	<pre># Define money growth rate differential (mg_dif)</pre>
y dif <- us y - jp y	# Define income growth rate differential (y dif)

y_dif <- us_y - jp_y	# Define income growth rate differential (mg_and
tb_dif <- us_tb - jp_tb	# Define Trade balance differential (tb_dif)
xx <- cbind(int_dif, mg_dif, y_dif) T <- length(e_f) T_est <- 161 e_f1 <- e_f[1:T_est] xx_1 <- xx[1:T_est,]	<ul> <li># Define final observation for <i>estimation period</i>.</li> <li># Adjust sample size to T_est</li> <li># Adjust sample size to T_est</li> </ul>

# Step (3) - Estimation of model(using only estimation period (T=161): Get b.
fit\_ef <- lm(e\_fl ~ xx\_1)
> summary(fit\_ef)

Call: lm(formula =  $e fl \sim xx l$ )

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) 1.7246 0.6971 2.474 0.0144 *
xx_lint_dif -0.5281 0.2478 -2.131 0.0346 *
xx_lmg_dif 0.1104 0.1912 0.577 0.5647
xx_ly_dif -0.2034 0.4538 -0.448 0.6546
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Residual standard error: 6.293 on 157 degrees of freedom Multiple R-squared: 0.04673, Adjusted R-squared: 0.02851 F-statistic: 2.565 on 3 and 157 DF, p-value: 0.05661

# Step (4) – Generate Forecasts. Need first to estimate model for X variables. (using *estimation period* data only)

```
• AR(1) for (i_{US,t} - i_{JAP,t})
int dif lag1 <- int dif[1:T est-1]
                                                        \# Lag (ius,t – iJAP,t)
int dif lag0 <- int dif[2:T est]
                                                        # Adjust sample size (lost one observation
above)
fit int \leq - \ln(int \operatorname{dif} \operatorname{lag0} \sim int \operatorname{dif} \operatorname{lag1})
                                                        # Fit AR(1) model
> summary(fit int)
Coefficients:
        Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.22774 0.11074 2.057 0.0414 *
int dif lag1 0.87537 0.03772 23.210 <2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 1.045 on 158 degrees of freedom
Multiple R-squared: 0.7732, Adjusted R-squared: 0.7718
F-statistic: 538.7 on 1 and 158 DF, p-value: < 2.2e-16
• AR(1) for (m_{US,t} - m_{JAP,t})
mg dif lag1 <- mg dif[1:T est-1]
                                                         \# Lag (m<sub>US,t</sub> – m<sub>JAP,t</sub>)
mg dif lag0 <- mg dif[2:T est]
                                                         # Adjust sample size (lost one observation)
fit mg \leq - lm(mg dif lag0 \sim mg dif lag1)
                                                         # Fit AR(1) model
> summary(fit mg)
```

Coefficients:

Estimate Std. Error t value Pr(>|t|)(Intercept) -0.008708 0.216621 -0.040 0.967986 mg dif lag1 0.296597 0.076124 3.896 0.000144 \*\*\* Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 2.74 on 158 degrees of freedom Multiple R-squared: 0.08766, Adjusted R-squared: 0.08188 F-statistic: 15.18 on 1 and 158 DF, p-value: 0.000144 • AR(1) for  $(y_{US,t} - y_{JAP,t})$  $y_dif_lag1 <- y_dif[1:T_est-1]$ # Lag (y<sub>US,t</sub> – y<sub>JAP,t</sub>) # Adjust sample size (lost one observation above) y dif lag0 <- y dif[2:T est] fit  $y \le lm(y \text{ dif } lag0 \sim y \text{ dif } lag1)$ # Fit AR(1) model > summary(fit y) Coefficients: Estimate Std. Error t value Pr(>|t|)(Intercept) 0.166258 0.086575 1.920 0.0566. y\_dif\_lag1 -0.008828 0.077255 -0.114 0.9092 Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 1.08 on 158 degrees of freedom Multiple R-squared: 8.263e-05, Adjusted R-squared: -0.006246 F-statistic: 0.01306 on 1 and 158 DF, p-value: 0.9092 • Now, we can do *one-step-ahead* forecast for the X variables: T val  $\leq T$  est+1 # start of Validation period xx cons  $\leq$  rep(1,T-T val+1) # create the constant vector int dif 0 <- cbind(xx cons,xx[T val:T,1]) %\*% fit int\$coeff # 8 forecasts for  $(i_{US,t} - i_{JAP,t})$ mg\_dif\_0 <- cbind(xx\_cons,xx[T\_val:T,2]) %\*% fit\_mg\$coeff #8 forecasts for  $(m_{US,t} - m_{JAP,t})$ y dif\_0 <- cbind(xx\_cons,xx[T\_val:T,3]) %\*% fit\_y\$coeff # 8 forecasts for  $(y_{US,t} - y_{JAP,t})$ • Finally, we compute the *one-step-ahead* forecast for **e** and MSE: e\_Mod\_0 <- cbind(xx\_cons,int\_dif\_0,mg\_dif\_0,y\_dif\_0)%\*%fit\_ef\$coeff # Model's forecast f e Mod  $\leq$  e f[T val:T] - e Mod 0 # Model's forecast error mse e f <- sum(f e  $Mod^2$ )/(T-T val+1) # Model's MSE >mse e f [1] 3.974203

• Compute the *one-step-ahead* forecast for RW Model and MSE e:

• Compare MSEs: The RW model has a better MSE (usual finding).

• A MGN test is usually done. But, we have only m=8 observations, we can do the test, but the results are very likely not to be taken seriously.

# Step (5) - Evaluation of Forecasts
• MGN/HLN test:
z\_mgn <- e\_Mod + e\_RW
x\_mgn <- e\_Mod - e\_RW
fit\_mgn <- lm(z\_mgn ~ x\_mgn)
> summary(fit\_mgn)

Coefficients:

Estimate Std. Error t value Pr(>|t|)(Intercept)1.3552.6800.5060.631x\_mgn1.7982.759**0.651**0.539 $\Rightarrow$  not significant, but unreliable (*small sample*).

Residual standard error: 3.026 on **6 degrees of freedom**  $\Rightarrow$  *very small* df to make inferences. Multiple R-squared: 0.05322, Adjusted R-squared: -0.1046 F-statistic: 0.3373 on 1 and 6 DF, p-value: 0.5826

• Suppose you are happy with the Model, you believe the difference in MSEs is not significant), now you generate out-of-sample forecasts.

# Step (6) – Out-of-sample one-step-ahead forward forecast for  $S_t$ : Et=2020:II[ $S_{t+1=2020:III}$ ] =  $S_{t=2020:II}$ \* (1 + Et=2020:II[ $e_{f,t+1=2020:III}$ ])

We observe  $S_t$  today (2020:II):  $S_{t=2020:II} = 100.77$  JPY/USD, which we invert since we work with direct quotes:  $S_{t=2020:II} = 0.009279$  USD/JPY.

We need to forecast the independent variables, based on AR(1) results,  $\mathbf{X}_{t} = \{(i_{US,t} - i_{JAP,t}), (y_{US,t} - y_{JAP,t}), (m_{US,t} - m_{JAP,t})\}$ 

• Forecasting  $(i_{US,t+1} - i_{JAP,t+1})$ : Et=2020:II[ $(i_{US,t} - i_{JAP})_{t+1=2020:III}$ ] int\_dif\_p1 <- cbind(1,int\_dif[T]) %\*% fit\_int\$coeff# int\_dif\_p1 = Et=2020:II[ $(i_{US,t} - i_{JAP})_{t+1=2020:III}$ ] > int\_dif\_p1 [,1] [1,] 0.4684645

• Forecasting  $(m_{US,t} - m_{JAP,t})$ :  $E_{t=2020:III}(m_{US,t} - m_{JAP})_{t+1=2020:III}$ ]

```
mg dif p1 <- cbind(1,m dif[T]) %*% fit m$coeff #mg dif p1=E_{t=2020:III}(m_{US} - m_{JAP})_{t+1=2020:III}
> mg_dif_p1
      [,1]
[1,] 4.921977
• Forecasting (y_{US,t} - y_{JAP,t}): E_{t=2020:III}(y_{US,t} - y_{JAP})_{t+1=2020:III}
y_dif_p1 <- cbind(1,y_dif[T]) %*% fit_y$coeff
                                                             # y dif p_1 = E_{t=2020:III}(y_{US,t} - y_{JAP})_{t+1=2020:III}
> y dif p1
      [,1]
[1,] 0.176617
• Forecasting E<sub>t=2020:II</sub>[S<sub>t+1=2020:III</sub>]
S <- 0.009279
                                                            # Today's value of S_{t=2020:II}
e f p1 <- cbind(1,int dif p1,mg dif_p1,y_dif_p1)%*%fit_ef$coeff # Today's forecast: et=2020:III
                                                            # Print forecast for e_{f,t+1=2020:III}
> e_f_p1
       [,1]
                                           \Rightarrow 2% depreciation of USD against JPY in 3<sup>rd</sup> Quarter.
[1,] 1.984401
                                                            # Today's forecast for S_{t+1=2020:III}
S pl \leq S^{(1+e)} f pl/100)
> S_p1 <- S^*(1+e f p1/100)
                                                    # e is in %, we divide by 100 to put it decimal from
                                                    # Print forecast for S_{t+1=2020:III}
> S p1
      [,1]
[1,] 0.009463133
                                                    \Rightarrow Model's forecast for S_{t+1=2020:III}
```

 $\Rightarrow$  Model's forecast for S<sub>t+1=2020:III</sub> = E<sub>t=2020:II</sub>[ $S_{t+1=2020:III}$ ] = 0.009463133 USD/JPY. (using the indirect quote, E<sub>t=2020:II</sub>[ $S_{t+1=2020:III}$ ] = 105.6732 JPY/USD).

• We can use the one-step-ahead forecasts to generate two-step-ahead forecasts. That is, we forecast  $E_{t=2020:II}[S_{t+1=2020:IV}]$ (=S p2 below)S1 <- S p1 # Today's forecast for St+1=2020:III int dif p2 <- cbind(1,int dif p1)%\*%fit int\$coeff # Today's forecast for  $(i_{US} - i_{JP})_{t+2}$ mg dif p2 <- cbind(1,mg dif p1)%\*%fit mg\$coeff # Today's forecast for  $(m_{US} - m_{JP})_{t+2}$ y\_dif\_p2 <- cbind(1,y\_dif\_p1)%\*%fit\_y\$coeff</pre> # Today's forecast for  $(y_{US} - y_{JP})_{t+2}$ e f p2 <- cbind(1,int dif p2,mg dif p2,y dif p2)%\*%fit ef\$coeff # forecast for et=2020:IV  $> e_f p2$ [,1]  $\Rightarrow$  1.11% depreciation of USD against JPY in 4<sup>th</sup> Quarter. [1,] 1.514363 S p2 <- S1\*(1+e f p2/100) > S p2[,1] [1,] 0.009606439

 $\Rightarrow$  Et=2020:II[ $S_{t+1=2020:III}$ ] = 0.009606439 USD/JPY.

<u>Note</u>: We can use the two-step-ahead forecast to generate *three-step-ahead* forecasts. Obviously, we can continue this process to generate *l-step-ahead* forecasts for  $S_t$  (a simple do loop will do it).

Eventually, we will collect *m* of out-of-sample forecasts (*m* one-step-ahead forecasts, *m* two-step-ahead forecasts, *m* three-step-ahead forecasts, etc.) to get an MSE and run a MGN/HLN test on them.  $\P$ 

It is possible that one model is the best in the short-term (say, up to 3 steps ahead); other is better in the medium-term (say, from 4 to 6 steps ahead); and another is best for longer-term. For example, the RW model is very good ("*unbeatable*") up to 3 months ahead. Then, other models start to produce better forecasts, especially after 6 months.

# **Forecasting Application: Fundamental Approach**

Practical Issues in Fundamental Forecasting

- Are we using the "right model?"
- Estimation of the model (OLS, MLE, other methods).
- Some explanatory variables  $(X_{t+T})$  are contemporaneous.
  - $\Rightarrow$  We also need a model to forecast the X<sub>t+T</sub> variables.

• Does Forecasting Work?

For many financial assets (stock prices, exchange rates), we expect forecasting to be difficult. The Efficient Markets Hypothesis posits that financial asset returns closely follows a "*Random Walk*" process, therefore forecasting asset returns is fruitless. Burton Malkiel in his book "*A Random Walk Down Wall Street*," first published in 1973, popularized this point.

**Example**: For exchange rates, in the short-run, the Random Walk consistently models beat structural (and other) models, like PPP, IFE, Monetary Approach: Lower MSE, MAE. That is bad news for the beaten models, since the RW forecast uses today's price to forecast any future price. No model or estimation is needed.

<u>Note</u>: Many argue that the structural models used to forecast exchange rates are not the "right model."  $\P$ 

# **Model Selection Strategies**

Specifying the DGP in (A1) is the most important step in applied work. We have assumed "correct specification," which, in practice, is an unrealistic assumption, since we do not really observed the true DGP.

A bad model can create a lot of problems: biases, wrong inferences, bad forecasts, etc.

So far, we have implicitly used a simple strategy:

(1) We started with a DGP, which we assumed to be true.

- (2) We tested some  $H_0$  (from economic theory).
- (3) We used the model (restricted, if needed) for prediction & forecasting.

Question: How do we propose and select a model (a DGP)?

Potentially, we have a huge number of possible models. We can have models with different functional form: f(.), g(.), , or h(.), and/or different explanatory variables: **X**, **Z**, **W** and dummy variables, **D**. For example, we may have four different formulations to choose from:

Model 1 $\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\epsilon}$ Model 2 $\mathbf{Y} = \mathbf{Z}\boldsymbol{\gamma} + \boldsymbol{\xi}$ Model 3 $\mathbf{Y} = (\mathbf{W}\boldsymbol{\gamma})^{\lambda} + \boldsymbol{\eta}$ Model 4 $\mathbf{Y} = \exp(\mathbf{Z} \ \mathbf{D} \ \boldsymbol{\delta}) + \boldsymbol{\epsilon}$ 

We want to select the best model, the one that is closest to the true and unobserved DGP. In practice, we aim for a "good" model, a model that passes a barrage of specification tests and has good forecasting power.

### **Model Selection Strategies: Views**

A model is a simplification. There are many approaches to specify a model:

- "Pre-eminence of theory." Economic theory should drive a model. Data is only used to quantify theory. Econometric methods offer sophisticated ways 'to bring data into line' with a particular theory.

- Purely data driven models. Success of ARIMA models (late 60s – early 70s), discussed in Lecture 6: No theory, only exploiting the time-series characteristics of the data to build models.

- Modern (LSE) view. A compromise: theory and the characteristics of the data are used to build a model.

• Theory and practice play a role in deriving a good model. David Hendry (2009) emphasizes:

"This implication is not a tract for mindless modeling of data in the absence of economic analysis, but instead suggests formulating more general initial models that embed the available economic theory as a special case, consistent with our knowledge of the institutional framework, historical record, and the data properties."

"Applied econometrics cannot be conducted without an economic theoretical framework to guide its endeavours and help interpret its findings. Nevertheless, since economic theory is not complete, correct, and immutable, and never will be, one also cannot justify an insistence on deriving empirical models from theory alone."

### **Model Selection Strategies: A Good Model**

According to David Hendry, a good model should be:

- Data admissible -i.e., modeled and observed **y** should have the same properties.

- Theory consistent	-our model should "make sense"
- Predictive valid	-we should expect out-of-sample validation
- Data coherent	-all information should be in the model. Nothing left in the errors (white noise
	errors).
- Encompassing	-our model should explain earlier models.

That is, we are searching for a statistical model that can generate the observed data (y, X), this is usually referred as *statistical adequacy*, makes theoretical sense and can explain other findings.

# **Model Selection Strategies: FAQ**

FAQ in practice:

- Should I include all the variables in the database in my model?
- How many explanatory variables do I need in my model?
- How many models do I need to estimate?
- What functional form should I be using?
- Should the model allow for structural breaks?
- Should I include dummies & interactive dummies?
- Which regression model will work best and how do I arrive at it?

# **Model Selection Strategies: Important Concepts**

*Diagnostic testing:* We test assumptions behind the model. In our case, assumptions (A1)-(A5) in the CLM.

**Example**: Test  $E[\mathbf{\epsilon}|\mathbf{X}] = 0$  -i.e., the residuals are zero-mean, uncorrelated with anything (that is, white noise distributed errors).

In selecting a model, this is a very important step. We run a lot of test to check the residuals are acceptable or the model is not misspecified: Ramsey's reset test, tests for autocorrelation, etc.

Parameter testing: We test economic H<sub>0</sub>'s.

**Example**: Test  $\beta_k = 0$  -say, there is no size effect on the expected return equation.

# **Model Selection Strategies: Two Methods**

There are several *model-selection methods*. We will consider two:

- Specific to General
- General to Specific

- Specific to General. Start with a small "restricted model," do some testing and make model bigger model in the direction indicated by the tests (for example, add variable  $x_k$  when test reject H<sub>0</sub>:  $\beta_k=0$ ).

- General to Specific. Start with a big "general unrestricted model," do some testing and reduce model in the direction indicated by the tests (for example, eliminate variable  $x_k$  when test cannot reject H<sub>0</sub>:  $\beta_k=0$ ).

# **Model Selection Strategies: Specific to General**

Steps:				
(1) Begin with a small theoretical model	– for example, the CAPM			
$\mathbf{y} = \mathbf{X} \boldsymbol{\beta} + \boldsymbol{\varepsilon}.$				
(2) Estimate the model	– say, using OLS.			
(3) Do some diagnostic testing	- are residuals white noise (uncorrelated)?			
If the assumptions do not hold, then	use:			
- More advanced econometrics	- GLS instead of OLS?			
- A more general model	– More regressors? Lags?			
(4) Test economic $H_0$ on the parameters	- Is SMB and HML significant?			
(5) Modify model in (1) in the direction of rejections of $H_0$ .				

• This strategy is known as *specific to general*. In the machine learning literature, this strategy is also called *forwards selection*.

**Example:** Specific-to-general strategy to model IBM returns: (1) We start with the 3-factor FF model for IBM:  $(r_{i=IBM} - r_f)_t = \beta_0 + \beta_1 (r_m - r_f)_t + \beta_2 SMB_t + \beta_3 HML_t + \varepsilon_t$ 

(2) Estimate the 3-factor FF model for IBM: fit\_ibm\_ff3 <- lm (ibm\_x ~ Mkt\_RF + SMB + HML) > summary(fit ibm ff3)

Coefficients:

	Estimate	Std. Error t value $Pr(> t )$					
(Intercept)	-0.005191	0.002482 -2.091	0.0369 *				
Mkt_RF	0.910379	0.056784 16.032	<2e-16 ***				
SMB	-0.221386	0.084214 -2.629	0.0088 **				
HML	-0.139179	0.084060 -1.656	0.0983.				

Residual standard error: 0.05842 on 566 degrees of freedom Multiple R-squared: 0.3393, Adjusted R-squared: 0.3358 F-statistic: 96.9 on 3 and 566 DF, p-value: < 2.2e-16

(3) Diagnostic tests: Check t-values &  $R^2$ , F-test goodness of fit, etc.

(4) LM Test to test if there is a January Effect (H<sub>0</sub>: No January effect):
> LM\_test
[1] 9.084247 ⇒ LM\_test > 3.84 ⇒ Reject H<sub>0</sub>.

(5) Given this result, we modify the 3-factor FF and add the January Dummy to the FF model: fit\_ibm\_new <- lm (ibm\_x ~ Mkt\_RF + SMB + HML + Jan\_1) > summary(fit\_ibm\_new) Coefficients: Estimate Std. Error t value Pr(>|t|)

Estimate	Sta. Error	t value $Pr(- t )$
-0.007302	0.002561	-2.851 0.00452 **
0.905182	0.056405	16.048 < 2e-16 ***
-0.247691	0.084063	-2.946 0.00335 **
-0.154093	0.083606	-1.843 0.06584.
0.026966	0.008906	<b>3.028</b> 0.00258 **
	-0.007302 0.905182 -0.247691 -0.154093 0.026966	Estimate         Std. Error           -0.007302         0.002561           0.905182         0.056405           -0.247691         0.084063           -0.154093         0.083606           0.026966         0.008906

<u>Note</u>: We can continue our search to see if an expansion of the specific model is needed. For example, we could have tested for a 2008 Financial crisis dummy or Dot.com dummy.  $\P$ 

• Some remarks based on the previous example:

• The specific-to-general method makes assumptions along the way.

(1) Very likely the starting model is based on theory and experience (HML is not significant at the usual 5% level). Not clear how to proceed from there to a more general model.

(2) We tested for a January effect and then added to the model. However, we could have tested for a Dot.com effect or for an interactive Dot.com/January effect with the 3 FF factors. Not clear when to stop the search.

(3) Selection step uses a p-value to add variables to the model. In this case, we use the standard 5% for the tests.

# Model Selection Strategies: Specific to General – Stepwise Regression

Note that in the previous example, we started with a model. What happens if we are skeptical regarding models?

• A popular implementation of the specific-to-general model selection is the *stepwise regression*, where we start with only a set of potential explanatory variables and let the data determine, starting from all potential one-variable models, which variables to add.

Overall structure of Stepwise Regression:

- The method begins with a *k* potential regressors.

- Do k one-variable regressions. Pick the one that shows the biggest t-stat or maximizes a goodness of fit measure, say, Adjusted-R<sup>2</sup>,  $\overline{R}^2$ . Suppose  $x_i$  is selected.

- Then, do (k - 1)-variable regressions all with  $x_j$ . Select the regressor (in addition to  $x_j$ ) that has the highest t-stat or that maximizes  $\overline{R}^2$ .

- Continue. But, when we start adding regressors, we usually check if the added regressor(s) change the significance of previous steps. (Note: at each step, we remove or add a regressor(s) based on t- or F-tests.)

- Stop: Additional regressors do not have *significant* t-stats/increase  $\overline{R}^2$ .

• Decisions: We need to select the *k* initial variables, the  $\alpha$  level for tests ( $\alpha = 5\%$ , 10%, 30%?) and/or the goodness of fit statistic.

<u>Remark</u>: Always keep in mind that the selected (final) model is not necessarily better than others. Type I and Type II errors are likely to occur, thus the final model may have irrelevant and/or omitted variables.

**Example:** Stepwise regression strategy to model IBM returns. We start with the 5 FF factors as candidates for IBM. We use the function *ols\_step\_forward\_p* in the *olsrr* package, which uses *p-values* to select variables. (You can also use other criteria to select the model, for example, *ols\_step\_forward\_aic* uses *aic* to select variables.) The final output is long (details = TRUE), below we present the last two results:

library(olsrr)

ff\_step\_data <- data.frame(Mkt\_RF, SMB, HML, RMW, CMA) ibm\_ff\_model <- lm(ibm\_x ~ ., data = ff\_step\_data) ols step forward p(ibm ff model, details = TRUE) # default p-value (penter) is 0.3

				2							
n	nodel	Beta	Std. E	rror S	td. Beta	t	S	Sig	lower	up	per
(Inte N	rcept) Ikt_RF SMB RMW	-0.00 0.887 -0.26 -0.12 Sel	5 0.00 7 0.05 1 0.08 8 0.11 lection	2 5 0 8 -0 4 -0 Summa	.574 .111 .042 	-1. 16 -2. -1.	999 .227 960 122	0.04 0.00 0.00 0.26	6 -0.0 00 0.' 3 -0.4 2 -0.3	)10 780 435 351	0.000 0.995 -0.088 0.096
Step	Variable Entered	R-S	quare	Adj. R-Squ	are C(	(p)	Al	[C	RN	ЛSE	
1 2 3	Mkt_RF SMB RMW	0.308 0.317 0.318	87 0 74 0 88 (	).3075 ).3151 ).3154	7.7 2.2 2.9	108 117 552	-166 -167 -167	65.555 71.054 70.320	51 0.0 48 0.0 07 0.0	)594 )590 )590	

Parameter Estimates

<u>Conclusion</u>: The Stepwise Regression method selects Market excess returns, SMB & RMW as the drivers of IBM excess returns. If we change the p-value to 0.1, RMW will be drop from final model.  $\P$ 

<u>Technical Note</u>: In general, the selection of variables based on p-values is not advised, since the distribution of the OLS coefficients is affected. We mentioned this above, when we discussed pre-testing.
# **Model Selection Strategies: General to Specific**

Begin with a *general unrestricted model* (GUM), which nests restricted models and, thus, allows any restrictions to be tested. Say:

$$\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{Z}\boldsymbol{\gamma} + \mathbf{W}^{\lambda}\boldsymbol{\delta} + (\mathbf{X}\ ^{\ast}\mathbf{W})\boldsymbol{\zeta} + (\mathbf{Z}\ ^{\ast}\mathbf{D})\boldsymbol{\psi} + \boldsymbol{\epsilon}.$$

Then, reduction of the GUM starts. Mainly using *t-tests*, and *F-tests*, we move from the GUM to a smaller, more parsimonious, specific model. If competing models are selected, encompassing tests or information criteria (AIC, BIC) can be used to select a final model. This is the *discovery stage*. After this reduction, we keep a final (restricted GUM) model:

$$\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\varepsilon}.$$

Creativity is needed for the specification of a GUM. Theory and empirical evidence play a role in designing a GUM.

## • Steps:

**Step 1** - First ensure that the GUM does not suffer from any diagnostic problems. Check residuals in the GUM to ensure that they possess acceptable properties. (For example, test for white noise in residuals, incorrect functional form, autocorrelation, etc.).

**Step 2** - Test the restrictions implied by the specific model against the general model – either by exclusion tests or other tests of linear restrictions.

**Step 3** - If the restricted model is accepted, test its residuals to ensure that this more specific model is still acceptable on diagnostic grounds.

• This strategy is called *general to specifics* ("*gets*"), *LSE*, *TTT* (Test, test, test). It was pioneered by Sargan (1964). The properties of gets are discussed in Hendy and Krolzig (2005, Economic Journal).

• The role of diagnostic testing is two-fold.

- In the *discovery steps* (Steps 1 & 2), the tests are being used as design criteria. Testing plays the role of checking that the original GUM was a good starting point after the GUM has been simplified.

- In the context of model evaluation (Step 3), the role of testing is clear cut. Suppose you use the model to produce forecasts. These forecasts can be evaluated with a test. This is the critical evaluation of the model.

**Example:** General-to-specific strategy to model IBM returns: **Step 1 -** Start with a GUM: the 3-factor FF model for IBM + January Dummy + Dot.com Dummy + non-linear & interactive effects:

$$(r_{i=IBM} - r_{f})_{t} = \beta_{0} + \beta_{1} (r_{m} - r_{f})_{t} + \beta_{2} SMB_{t} + \beta_{3} HML_{t} + \beta_{4} January_{t} + \beta_{5} (r_{m} - r_{f})_{t}^{2} + \beta_{6} SMB_{t}^{2} + \beta_{7} HML_{t}^{2} + \beta_{8} (r_{m} - r_{f})_{t} * SMB_{t} + \beta_{9} (r_{m} - r_{f})_{t} * HML_{t} + + \beta_{10} Dot.com_{t} + \beta_{11} (r_{m} - r_{f})_{t} * January_{t} + \beta_{12} HML_{t} * January_{t}$$

```
+ \beta_{13} (r_m - r_f)_t * \text{Dot.com}_t + \beta_{14} HML_t * \text{Dot.com} + \beta_{15} SMB_t * \text{Dot.com} + \varepsilon_t
```

```
Estimate GUM:

t_sb <- 342

T_s_1 <- T - t_sb

d_0 <- matrix(0, t_sb, 1)

d_1 <- matrix(1, T_s_1, 1)

Dot_com <- rbind(d_0,d_1)

Mkt_Jan <- Mkt_RF * Jan_1

HML_Jan <- HML * Jan_1

Mkt_Dot <- Mkt_RF * Dot_com

HML_Dot <- HML * Dot_com

SMB_Dot <- SMB * Dot_com
```

# Structural break date (End of 1st-regime)
# Dot.com dummy = 0 before t\_sb
# Dot.com dummy = 1 after t\_sb
# Dot.com dummy (join rows d\_0 & d\_1)

fit\_ibm\_gum <- lm (ibm\_x ~ Mkt\_RF + SMB + HML + Jan\_1 + Mkt\_RF\_2 + SMB\_2 + HML\_2 + Mkt\_HML + Mkt\_SMB + SMB\_HML + Mkt\_Jan + HML\_Jan + Mkt\_Dot + HML\_Dot + SMB\_Dot) > summary(fit ibm\_gum)

Coefficients:

	Estimate	Std. Error	t value	$Pr(\geq  t )$			
(Intercept)	-0.007836	0.003063	-2.559	0.010772	*		
Mkt_RF	0.791866	0.090474	8.752	< 2e-16 *	<b>*</b> *		
SMB	-0.295790	0.110655	-2.673	0.007738	**		
HML	-0.233942	0.135146	-1.731	0.084004	$\Rightarrow$	practice says "keep it." Judge	ement call.
Jan_1	0.031769	0.009349	3.398	0.000727	***		
Mkt_RF_2	-0.433762	0.850899	-0.510	0.610417			
SMB_2	-0.927271	1.470645	-0.631	0.528615			
HML_2	2.707992	1.670366	1.621	0.105545	$\Rightarrow$	almost 10%, I keep it. Judgen	nent call.
Mkt_HML	0.628721	1.557090	0.404	0.686531			
Mkt_SMB	0.791625	1.746939	0.453	0.650618			
SMB_HML	-1.044806	2.029091	-0.515	0.606819			
Mkt_Jan	-0.069413	0.189309	-0.367	0.714008			
HML_Jan	-0.259697	0.255484	-1.016	0.309841			
Mkt_Dot	0.323382	0.130645	2.475	0.013612	*		
HML_Dot	0.059742	0.208277	0.287	0.774342			
SMB_Dot	0.076998	0.198964	0.387	0.698910			
Signif. codes:	0 '***' 0.0	01 '**' 0.0	01 '*' 0	0.05 '.' 0.1	• 1		

Residual standard error: 0.05788 on 553 degrees of freedom Multiple R-squared: 0.3663, Adjusted R-squared: 0.3491 F-statistic: 21.31 on 15 and 553 DF, p-value: < 2.2e-16

Step 1 – Check GUM residuals for departures of (A2)-(A3). A Ramsey's reset test can be done (using the *resettest* in the *lmtest* library).

> resettest(fit\_ibm\_gum, type="fitted")
 RESET test
data: fit\_gumHomework 1 Review <br>
RESET = 1.2645, df1 = 2, df2 = 552, p-value = 0.2832

**Step 2** – Reduce Model with t-test and F-tests. Say, we keep all the variables with a p-value close to 10% (we still keep HML, using previous experience). We estimate a restricted GUM:

fit\_ibm\_gum\_r <- lm (ibm\_x ~ Mkt\_RF + SMB + HML + Jan\_1 + HML\_2 + Mkt\_Dot)
> summary(fit\_ibm\_gum\_r)

Coefficients:

	Estimate	Std. Error	t value	$\Pr(\geq  t )$
(Intercept)	-0.008696	0.002788	-3.119	0.00191 **
Mkt_RF	0.779336	0.072453	10.756	<2e-16 ***
SMB	-0.280018	0.083891	-3.338	0.00090 ***
HML	-0.250480	0.088504	-2.830	0.00482 **
Jan_1	0.028499	0.008937	3.189	0.00151 **
HML_2	1.676011	1.331161	1.259	0.20853
Mkt_Dot	0.344030	0.116685	2.948	0.00333 **
Signif. codes:	0 '***' 0.0	01 '**' 0.0	1 '*' 0	.05 `.' 0.1 ` ' 1

Residual standard error: 0.05761 on 562 degrees of freedom Multiple R-squared: 0.3618, Adjusted R-squared: 0.355 F-statistic: 53.11 on 6 and 562 DF, p-value: < 2.2e-16

Step 2 – Test the restrictions implied by the specific model against the general model. Using an F-test, we test J=9 restrictions:

H<sub>0</sub>: 
$$\beta_5 = \beta_6 = \beta_8 = \beta_9 = \beta_{10} = \beta_{11} = \beta_{12} = \beta_{14} = \beta_{15}$$
.

e u <- fit ibm gum\$residuals # GUM residuals RSS  $u \le t(e \ u)\%\%$  u e r <- fit ibm gum r\$residuals # Restricted GUM residuals RSS  $r \le t(e r)\%*\%e r$ f test gum <- ((RSS r - RSS u)/9)/(RSS u/(T-16)) #F-test > f test gum [,1] [1,] **0.4299497**  $\Rightarrow$  we cannot reject H<sub>0</sub> (f test gum < qchisq(.95,9, 553) = **1.896801**) > qf(.95, df1=9, df2=T-16)[1] **1.896801**  $p_val <-1 - pf(f_test gum, df = 9, df2=T-16)$ *# p-value* of F-test > p val[1,] **0.919105**  $\Rightarrow$  p-value is very high. No evidence for H<sub>0</sub>.

Step 2 - Further specification checks of Restricted GUM, for example, perform a Ramsey's reset test (using the *resettest* in the lmtest library). > resettest(fit gum r, type="fitted")

**RESET** test

data: fit\_ibm\_gum\_r RESET = 1.1361, df1 = 2, df2 = 561, p-value = 0.3218

**Step 3** - Test if Restricted GUM residuals are acceptable –i.e., do diagnostic tests (mainly, make sure they are white noise). If Restricted GUM passes all the diagnostic tests, it becomes the "final model."

<u>Note</u>: With the final model, we use it to justify/explain financial theory and features, and do forecasting.  $\P$ 

• Some remarks based on the previous example:

The general-to-specific method makes assumptions along the way.

(1) Select a p-value for the tests of significance in the discovery stage (we use 10%). Given that we performed 15 *t-tests*, we should not be surprised we rejected the GUM, since we had an overall significance,  $\alpha^* = .79 [= 1 - (1 - .10)^{15}]$ . Mass significance is an issue.

(2) Judgement calls are also made.

(3) The reduction of the GUM involves "pre-testing" –i.e., data mining. We are likely rejecting a true  $H_0$  (false positives) and not rejecting a true  $H_1$ , (false negatives) along the way. This increases the probability that the final model is not a good approximation. It is common to ignore (or not even acknowledge) pre-testing issues.

<u>Note</u>: Similar to stepwise regression, we can remove use p-values to remove one step at a time variables from the GUM. R can do this using the function *ols\_step\_backward\_p* in the olsrr package.

## **Model Selection Strategies: Best Subset**

Begin with a big model, with *k* regressors:

 $y = X\beta + \epsilon$ .

The idea is to select the "best" subset of the k regressors in X, where "best" is defined by the researcher, say MSE, Adjusted- $R^2$ , etc.

In theory, it requires  $2^k$  regressions. It can take a while if k is big (k < 40 is no problem). There are many tricks are used to reduce the number of regressions.

In practice, we use best subset to reduce the number of models to consider. For example, from the regressions with one-variable, keep the best one-variable model, from the regression with two-variables, keep the best two-variable model, etc.

Example: We want to select a model for IBM excess returns, using the k=3 Fama-French factors: Market excess returns (Mkt\_RF), SMB, & HML. We have 8 (=2<sup>3</sup>) models and, thus, regressions:
1) Constant;
2) Mkt\_RF (CAPM)
3) SMB
4) HML
5) Mkt\_RF & SMB
6) Mkt\_Rf & HML
7) SMB & HML
8) Mkt\_RF, SMB, & HML (the 3-factor F-F Model).

We select the model with the lower MSE. Or, we can carry two or three models of the best models to do *cross-validation* and, then, pick the best model.

Suppose we selected three model: CAPM (M1); Mkt\_RF & SMB (M2); and the 3-factor F-F Model (M3). Now, we use *K*-fold cross-validation, with K = 5. CV<sub>5</sub> M1: 0.003542756 CV<sub>5</sub> M2: **0.003505873** CV<sub>5</sub> M3: 0.003556918

<u>Note</u>: Models look very similar. Practitioners compute a SE for  $CV_K$  and use a one SE rule. If within one SE, keep simplest model (M1).¶

## **Model Selection Strategies: Properties**

A modeling strategy is *consistent* if its probability of finding the true model tends to 1 as *T* -the sample size- increases.

- Properties for strategies
- (1) Specific to General
- It is not consistent if the original model is incorrect.
- It need not be predictive valid, data coherent, & encompassing.
- No clear stopping point for an unordered search.
- (2) General to Specific
- It is consistent under some circumstances. But, it needs a large T.
- It uses data mining, which can lead to incorrect models for small T.
- The significance levels are incorrect. This is the problem of *mass significance*.

#### **Model Selection Strategies: Using Machine Learning**

So far, we have emphasized finding a DGP, that gives us a (linear) model for the conditional expectation of y. Then, using this model, we estimate its parameters to get  $\hat{y}$ . For example, a *k*-factor model:

$$\mathbf{E}[y_i|x_i] = \alpha + \beta_1 x_{1,i} + \beta_2 x_{2,i} + \ldots + \beta_k x_{k,i} \Longrightarrow \hat{y}_i$$

Machine Learning (ML) methods can be used to select a model and covariates, especially when the goal is to generate predictions,  $\hat{y}_i$ . ML models are very efficient in settings with many (hundreds or thousands) explanatory variables or covariates –i.e., large k.

<u>Note</u>: We have relied on linear models, but ML methods can allow for almost any functional form for  $E[y_i | x_i]$ . Moreover, in general, ML does not care about the interpretation of its parameters, though work is being done to derive the properties of parameters and predictions.

We start with an ML method that preserve linearity for the conditional expectation,  $E[y_i | x_i]$ , with *k* covariates:

$$E[y_i | x_i] = \beta_1 x_{1,i} + \beta_2 x_{2,i} + \beta_3 x_{3,i} + \dots + \beta_k x_{k,i} = \beta' x_i$$

OLS estimates this model by

$$\min_{\boldsymbol{\beta}} \sum_{i=1}^{N} (y_i - \boldsymbol{\beta}' \boldsymbol{x}_i)^2$$

Question: We can do OLS, which has nice properties, why do we need ML?

When k is very large, possible exceeding N, the OLS estimator may have inferior predictive properties, in terms of MSE, to those of other estimators that impose some restrictions or "penalties" on the size of the parameters in the minimization problem. These restrictions are called "*regularizations*."

In general, the bigger the size of the vector of parameter (the "*complexity*") the bigger the penalty. For example,

$$\min_{\boldsymbol{\beta}} \sum_{i=1}^{N} (y_i - \boldsymbol{\beta}' \boldsymbol{x}_i)^2 + \boldsymbol{\lambda} \operatorname{Penalty}(\operatorname{size}(\boldsymbol{\beta})) \qquad (\boldsymbol{\lambda} > 0)$$

where  $\lambda > 0$ . Different penalties for complexity give different models.

The parameter controls the strength of the penalty.

- when  $\lambda = 0$ , we have OLS
- when  $\lambda = \infty$ , we have  $\beta = 0$ .

- when  $\lambda \in (0, \infty)$ , we have a combination (or trade-of) between OLS and reducing complexity (setting coefficients to zero) and/or reducing the weights of covariates ("shrinking the coefficients" in the model.

#### Model Selection Strategies: Machine Learning – LASSO & Ridge Regression

**LASSO** or *Least Absolute Shrinkage and Selection Operator*, proposed by Tibshirani (1996), sets Penalty(size( $\beta$ )) =  $\sum_{j=1}^{k} |\beta_j|$ . That is:

$$\min_{\boldsymbol{\beta}} \sum_{i=1}^{N} (y_i - \boldsymbol{\beta}' \boldsymbol{x}_i)^2 + \boldsymbol{\lambda} \sum_{j=1}^{k} |\boldsymbol{\beta}_j|$$

Lasso, given its penalty structure, "shrinks" the  $\beta$ 's toward zero, some  $\beta$ 's will be set to exactly zero.

Unlike OLS, there is no closed form solution to Lasso minimization. but we can numerically compute the solution,  $\hat{\beta}_{Lasso}$ , to the above problem. (It is a quadratic programming from convex optimization.)

**Ridge regression** of Hoerl and Kennard (1970) sets Penalty(size( $\beta$ )) =  $\sum_{j=1}^{k} \beta_j^2$ . That is:  $\min_{\beta} \{ \sum_{i=1}^{N} (y_i - \beta' x_i)^2 + \lambda \sum_{j=1}^{k} \beta_j^2 = (y - X\beta)'(y - X\beta) + \lambda \beta' \beta \}$ 

Using linear algebra, we get a closed form solution for this problem:

$$\widehat{\boldsymbol{\beta}}_{Ridge} = (\mathbf{X}'\mathbf{X} + \lambda \mathbf{I})^{-1} \mathbf{X}' \mathbf{y}$$

Ridge regression, given its penalty structure, tends to reduce all the  $\beta$ 's.

<u>Remark</u>: Ridge regression shrinks –i.e., reduce- all coefficients *towards* zero, but lasso can remove predictors from model by shrinking (setting) the coefficients *completely* to zero. Thus, we can think of Lasso as a mechanism to select covariates –i.e., model selection.

<u>Technical note</u>: We can generalize the above estimation problem by defining the penalty using the  $L_p$ -norm notation:

$$\min_{\boldsymbol{\beta}} \sum_{i}^{N} (y_{i} - \boldsymbol{\beta}' \boldsymbol{x}_{i})^{2} + \boldsymbol{\lambda} (\| \boldsymbol{\beta} \|_{q})^{1/q}$$
$$\| \boldsymbol{\beta} \|_{q} = \sum_{i=1}^{k} |\boldsymbol{\beta}_{i}|^{q}.$$

where

For q = 1, we have Lasso. For q = 2, we have *Ridge regression*. As  $q \rightarrow 0$ , we get closer to best subset regression.

• It is also possible to combine (weight) the restrictions (LASSO & Ridge), this combination is called *Elastic net*:

 $\min_{\boldsymbol{\beta}} \sum_{i=1}^{N} (y_i - \boldsymbol{\beta}' \boldsymbol{x}_i)^2 + \boldsymbol{\lambda} \{ \alpha \sum_{j=1}^{k} |\beta_j| + (1 - \alpha) \sum_{j=1}^{k} {\beta_j}^2 \}$ where  $\alpha \in [0,1]$ .

• The parameter controls the strength of the penalty. How do we compute it?

The parameter  $\lambda$  is estimated ("tuned") through out-of-sample *K-fold cross-validation*. That is, for each  $\lambda$ , we split the data in *K* parts. For j = 1, 2, ..., K, use all folds but fold *j* to estimate model; use fold *j* to check model's forecasting skills by computing MSE,  $MSE_j$ . The *K*-fold CV estimate is an average of each fold MSE's:

 $CV_K = \frac{1}{K} \sum_{j=1}^{K} MSE_j$ 

Pick  $\lambda$  that has the smallest  $CV_K$ .

It is desirable to select the *K*-folds randomly, easier to do in cross section than in time series, where dependence creates problems.

• Both Lasso and Ridge regression estimates are not scale invariant, unlike OLS. Suppose we move  $x_k$  from percentage points to decimal. That is,  $x_k^* = x_k/100$ .

The  $x_k^*$  coefficient will be scaled as  $\beta_k^* = 100 * \beta_k$ . Then, the impact of  $x_k$  on y does not change  $(\beta_k ' x_k = \beta_k^* ' x_k^*)$ . Given the nature of the penalty –i.e., large coefficients are penalized –, we have that

 $\hat{\boldsymbol{\beta}}_{lasso,k}^{*} \neq 100 * \hat{\boldsymbol{\beta}}_{lasso,k} \qquad \& \qquad \hat{\boldsymbol{\beta}}_{Ridge,k}^{*} \neq 100 * \hat{\boldsymbol{\beta}}_{Ridge,k}$ 

To avoid these issue, it is common to standardize all covariates,  $x_k$ :

$$\mathbf{z}_k = \frac{\mathbf{x}_k - \overline{\mathbf{x}}_k}{\mathbf{s}_k}$$
 ( $\mathbf{s}_k$ : sample SD of  $\mathbf{x}_k$ )

Note: Now, all predictors have zero mean and unit variance.

We know OLS **b** is unbiased. Thus, the regularized (restricted) estimators  $\hat{\beta}_{lasso} \& \hat{\beta}_{Ridge}$  are biased; their appeal is lower variance.

In particular, for  $\hat{\beta}_{Ridge}$ , the variance is much smaller than OLS **b** when the data shows multicollinearity, something common in large cross-section models. ( $\hat{\beta}_{lasso}$  does not do as well.)

As pointed out above, the big appeal of  $\hat{\beta}_{lasso}$  is its sparcity (a smaller dimension than **b** &  $\hat{\beta}_{Ridge}$ ). We can use lasso as a model selection tool.

**Example:** In the general-to-specific example, we estimated with OLS a model with 16 parameters. Now, we estimate the model with LASSO, using the R package *glmmet* (for LASSO set alpha=1, for Ridge set alpha=0). This package uses the Matrix package and the vector of covariates need to be formatted as a matrix, using *data.matrix*. It selects lambda,  $\lambda$ , based on *k*-fold (sets *k*=10) cross-validation.

coef(opt model) # print coefficients > coef(opt model) # print coefficients 16 x 1 sparse Matrix of class "dgCMatrix" s0 (Intercept) -0.005835974 Mkt RF 0.791803084 SMB -0.079653804 HML Jan 1 0.008697714 Mkt RF 2 SMB 2 HML 2 . Mkt HML Mkt SMB SMB HML . Mkt Jan HML Jan -0.004928686 Mkt Dot 0.004006239 HML Dot . SMB Dot .

opt model  $\leq$  glmnet(x la, ibm x, alpha = 1, lambda = tuned lambda)

<u>Note</u>: As expected many coefficients are completely "shrunk" to 0. The model with the non-zero coefficients is the one that we use to predict out-of-sample –we need new data for the covariates to do this.

It is possible to compute R<sup>2</sup> for the estimated LASSO model: > y\_predicted <- predict(best\_model, s = tuned\_lambda, newx = x\_la) > sst <- sum((y - mean(y))^2) > sse <- sum((y\_predicted - y)^2) > R2 <- 1 - sse/sst > R2 [1] 0.3197774 (unrestricted OLS R2 = 0.3484) If we have new data, say x new, we set i newx=x new, above in predict function.

# Lecture 7 - Departures from CLM Assumptions & the Generalized Regression Model

# **Review of CLM Results**

Recall the CLM Assumptions (A1) DGP:  $\mathbf{y} = \mathbf{X} \ \beta + \boldsymbol{\varepsilon}$  is correctly specified. (A2)  $\mathrm{E}[\boldsymbol{\varepsilon}|\mathbf{X}] = 0$ (A3)  $\mathrm{Var}[\boldsymbol{\varepsilon}|\mathbf{X}] = \sigma^2 \mathbf{I}_{\mathrm{T}}$ (A4) X has full column rank -rank(X) = k-, where T  $\geq k$ .

• OLS estimation:

 $\begin{aligned} \mathbf{b} &= (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \mathbf{y} \\ \mathrm{Var}[\mathbf{b}|\mathbf{X}] &= \sigma^2 (\mathbf{X}'\mathbf{X})^{-1} \\ &\implies \mathbf{b} \text{ unbiased and efficient (MVUE)} \end{aligned}$ 

• If (A5)  $\boldsymbol{\varepsilon}|\mathbf{X} \sim N(\mathbf{0}, \sigma^2 \mathbf{I}_T) \implies \mathbf{b}|\mathbf{X} \sim N(\boldsymbol{\beta}, \sigma^2(\mathbf{X}, \mathbf{X})^{-1})$ Under (A5), **b** is also the MLE (consistency, efficiency, invariance, etc). (A5) gives us *finite* sample results for **b** (and for tests: *t-test*, *F-test*, Wald tests).

## **CLM: Departures from the Assumptions**

So far, we have discussed some violations of CLM Assumptions:

(1) (A1) – OLS can easily deal with some non-linearities in the DGP.

 $\Rightarrow$  as long as we have intrinsic linearity, **b** keeps its nice properties.

- Wald, F, & LM tests to check for misspecification

(2) (A4) – Perfect multicollinearity means the model needs to be changed. Multicollinearity is a potential problem. In general, exogenous to the researcher. We need to be aware of this problem.

• In this lecture, we examine assumptions (A2), (A3) and (A5). That is, we check

(i) **X** is stochastic. That is, it has a distribution.

- (ii) Var[ $\boldsymbol{\varepsilon} | \mathbf{X} ] \neq \sigma^2 \mathbf{I}_{T.}$
- (iii)  $\boldsymbol{\varepsilon} | \mathbf{X}$  is not N( $\mathbf{0}, \sigma^2 \mathbf{I}_T$ ).

# **CLM: Departures from (A2)**

The traditional derivation of the CLM assumes X as non-stochastic. In our derivation, however, we allowed X to be stochastic, but we conditioned on observing its realizations (an elegant trick, but not very realistic).

With stochastic X we need additional assumptions to get unbiasedness and consistency for the OLS **b**.

– We need independence between **X** &  $\varepsilon$ : {*x<sub>i</sub>*,  $\varepsilon_{ij}$ } *i*=1, 2, ..., *T* is a sequence of independent observations.

– We require that **X** have finite means and variances. Similar requirement for  $\boldsymbol{\varepsilon}$ , but we also require  $E[\boldsymbol{\varepsilon}] = \boldsymbol{0}$ .

Then,

$$E[\mathbf{b}] = \mathbf{\beta} + E[(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \mathbf{\epsilon}] = \mathbf{\beta} + E[(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'] E[\mathbf{\epsilon}] = \mathbf{\beta}$$

<u>Technical Note</u>: To get consistency (& asymptotic normality) for **b**, we need an additional (asymptotic) assumption regarding X:

or  $X'X/T \xrightarrow{p} Q$  (Q a pd (kxk) matrix of finite elements) P(X'X/T) = Q

Question: Why do we need this assumption in terms of a ratio divided by T? Each element of **X'X** matrix is a sum of T numbers.

$$\boldsymbol{X}'\boldsymbol{X} = \begin{bmatrix} \Sigma_{i=1}^{T} x_{i1}^{2} & \Sigma_{i=1}^{T} x_{i1} x_{i2} & \dots & \Sigma_{i=1}^{T} x_{i1} x_{iK} \\ \Sigma_{i=1}^{T} x_{i2} x_{i1} & \Sigma_{i=1}^{T} x_{i2}^{2} & \dots & \Sigma_{i=1}^{T} x_{i2} x_{iK} \\ \dots & \dots & \dots & \dots \\ \Sigma_{i=1}^{T} x_{iK} x_{i1} & \Sigma_{i=1}^{T} x_{iK} x_{i2} & \dots & \Sigma_{i=1}^{T} x_{iK}^{2} \end{bmatrix}$$

As  $T \rightarrow \infty$ , these sums will become large. We divide by T so that the sums will not be too large.

<u>Note</u>: This assumption is not a difficult one to make since the LLN suggests that the each component of  $\mathbf{X'X}/T$  goes to the mean values of  $\mathbf{X'X}$ . We require that these values are finite. – Implicitly, we assume that there is not too much dependence in  $\mathbf{X}$ .

# CLM: Departures from (A2) – Endogeneity

If there is dependence between  $\mathbf{X} \& \boldsymbol{\epsilon}$ , OLS **b** is no longer unbiased or consistent. Easy to see the biased result: we cannot longer separate  $E[(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \boldsymbol{\epsilon}]$  into a product of two expectations:

 $E[(\mathbf{X'X})^{\text{-1}}\mathbf{X'} \ \boldsymbol{\epsilon}] \neq E[(\mathbf{X'X})^{\text{-1}}\mathbf{X'}] \ E[\boldsymbol{\epsilon}]$  Then,

 $E[\mathbf{b}] = \mathbf{\beta} + E[(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}' \mathbf{\varepsilon}] \neq \mathbf{\beta}$ 

Dependence between  $X \& \varepsilon$  occurs when X is also an *endogenous variable*, like y. This is common, especially in Corporate Finance. For example, we study CEO compensation as function of size of firm, and Board composition. Board Composition and size of firm are endogenous –i.e., determined by the firm, dependent on CEO's decisions.

Inconsistency is a fatal flaw in an estimator. In these situations, we use different estimation methods. The most popular is *Instrumental Variable* (IV) *estimation*.

#### **CLM: Departures from (A2) – Asymptotics**

Now, we have a new set of assumptions in the CLM:

(A1) DGP:  $\mathbf{y} = \mathbf{X} \boldsymbol{\beta} + \boldsymbol{\epsilon}$ . (A2') X stochastic, but E[X' ] = 0 and E[ $\boldsymbol{\epsilon}$ ] = 0. (A3) Var[ $\boldsymbol{\epsilon}$ |X] =  $\sigma^2 \mathbf{I}_T$ (A4') plim (X'X/T) = Q (p.d. matrix with finite elements, rank= k)

With these new assumptions and using properties of plims and the CLT, we can show the following asymptotic results:

1. **b** and  $s^2$  are consistent. 2.  $\sqrt{T}$  (**b** -  $\beta$ )  $\xrightarrow{d}$  N(**0**,  $\sigma^2 \mathbf{Q}^{-1}$ )  $\Rightarrow$  **b**  $\xrightarrow{a}$  N( $\beta$ , ( $\sigma^2/T$ ) $\mathbf{Q}^{-1}$ ) 3. *test-t*  $\xrightarrow{d}$  N(0,1) *F-tests* & Wald tests  $\xrightarrow{d} \chi^2_J$ 

## **CLM: Departures from (A5)**

Notice that asymptotic results 2 and 3 state the asymptotic distribution of **b** and the t-, F- and Wald test. All derived from the new set of assumptions and the CLT. (A5) was not used.

That is, we relax (A5), but, now, we require *large samples*  $(T \rightarrow \infty)$ .

Note: In practice, we use the asymptotic distribution as an approximation to the finite sample -

i.e., for any *T*- distribution. This is why we used the  $\xrightarrow{a}$  notation in:

 $\mathbf{b} \xrightarrow{a} \mathrm{N}(\mathbf{\beta}, (\sigma^2/T)\mathbf{Q}^{-1})$ 

We should be aware that this approximation may not be accurate in many situations.

• Two observations regarding relaxing (A5)  $\varepsilon | \mathbf{X} \sim i.i.d. \text{ N}(\mathbf{0}, \sigma^2 \mathbf{I}_T)$ :

– Throwing away the normality for  $\mathbf{\varepsilon}|\mathbf{X}$  is not bad. In many econometric situations, normality is not a realistic assumption (daily, weekly, or monthly stock returns do not follow a normal).

– Removing the *i.i.d.* assumption for  $\boldsymbol{\varepsilon} | \mathbf{X}$  is also not bad. In many econometric situations, identical distributions are not realistic, since different means and variances are common.

Questions:

- Do we need to throw away normality for  $\boldsymbol{\varepsilon}|\mathbf{X}$ ?

Not necessarily. We can test for normality on the residuals using a Jarque-Bera test, though, for financial assets we usually reject normality is rejected, especially at the monthly, weekly, daily, and intra-daily frequencies.

- Why are we interested in large sample properties, like consistency, when in practice we have finite samples?

As a first approximation, the answer is that if we can show that an estimator has good large sample properties, then we may be optimistic about its finite sample properties. For example, if an estimator is inconsistent, we know that for finite samples it will definitely be biased.

# **CLM: Departures from (A3)**

Now, we relax (A3). The CLM assumes that errors are uncorrelated and all are drawn from a distribution with the same variance,  $\sigma^2$ . (A3) Var[ $\epsilon$ |X] =  $\sigma^2$ I<sub>T</sub>

Instead, we will assume:

(A3') Var[ $\boldsymbol{\varepsilon}|\mathbf{X}$ ] =  $\boldsymbol{\Sigma}$  (sometimes written=  $\sigma^2 \boldsymbol{\Omega}$ , where  $\boldsymbol{\Omega} \neq \mathbf{I}_T$ )

$$\boldsymbol{\Sigma} = \begin{bmatrix} \sigma_1^2 & \sigma_{12} & \cdots & \sigma_{1T} \\ \sigma_{21} & \sigma_2^2 & \cdots & \sigma_{2T} \\ \vdots & \vdots & \vdots & \vdots \\ \sigma_{T1} & \sigma_{T2} & \cdots & \sigma_T^2 \end{bmatrix}$$

• Two Leading Cases:

- Pure heteroscedasticity: We model only the diagonal elements.
- Pure cross/auto-correlation: We model only the off-diagonal elements.

#### **CLM: Departures from (A3) – Heteroscedasticity**

Pure heteroscedasticity: 
$$E[\varepsilon_i \ \varepsilon_j | \mathbf{X}] = \sigma_{ij} = \sigma_i^2 \qquad \text{if } i = j$$
$$= 0 \qquad \text{if } i \neq j$$
$$\Rightarrow \operatorname{Var}[\varepsilon_i | \mathbf{X}] = \sigma_i^2$$
$$\Sigma = \begin{bmatrix} \sigma_1^2 & 0 & \cdots & 0 \\ 0 & \sigma_2^2 & \cdots & 0 \\ \vdots & \vdots & \vdots & \vdots \\ 0 & 0 & \cdots & \sigma_T^2 \end{bmatrix}$$

This type of variance-covariance structure is common in time series, where we observe the variance of the errors changing over time or subject to different regimes (say, bear and bull regimes).

Relative to pure heteroscedasticity, LS gives each observation a weight of 1/T. But, if the variances are not equal, then some observations (low variance ones) are more informative than others.



CLM: Departures from (A3) – Cross-correlationPure cross/auto-correlation: $E[\epsilon_i \ \epsilon_j | \mathbf{X}] = \sigma_{ij}$ if  $i \neq j$  $= \sigma^2$ if i = j

$$\boldsymbol{\Sigma} = \begin{bmatrix} \boldsymbol{\sigma}^2 & \sigma_{12} & \cdots & \sigma_{1T} \\ \sigma_{21} & \boldsymbol{\sigma}^2 & \cdots & \sigma_{2T} \\ \vdots & \vdots & \vdots & \vdots \\ \sigma_{T1} & \sigma_{T2} & \cdots & \boldsymbol{\sigma}^2 \end{bmatrix}$$

This type of variance-covariance structure is common in cross sections, where errors can show strong correlations, for example, when we model returns, the errors of two firms in the same industry can be subject to common (industry) shocks. Also common in time series, where we observe clustering of shocks over time.

Relative to pure cross/auto-correlation, LS is based on simple sums, so the information that one observation (today's) might provide about another (tomorrow's) is never used.

<u>Note</u>: Heteroscedasticity and autocorrelation are different problems and generally occur with different types of data. But, the implications for OLS are the same.

#### **CLM: Departures from (A3) – Implications**

OLS **b** is still *unbiased* and *consistent*. (Proofs do not rely on (A3).

OLS **b** still follows an *asymptotic normal distribution*. It is

- Easy to show this result for the pure heteroscedasticity case using a version of the CLT that assumes only independence ;

- More complicated derivation -i.e., with new assumptions- for the cross/auto-correlation case.

But, OLS **b** is no longer BLUE. There are more efficient estimators; estimators that take into account the heteroscedasticity in the data.

Note: We used (A3) to derive our test statistics. A revision is needed!

# **Finding Heteroscedasticity**

There are several theoretical reasons why the  $\sigma_i^2$  may be related to some variables  $z_1, \ldots, z_j$  and/or  $z_1^2, \ldots, z_j^2$ .

1. Following the *error-learning models*, as people learn, their errors of behavior become smaller over time. Then,  $\sigma_i^2$  is expected to decrease.

2. As data collecting techniques improve,  $\sigma_i^2$  is likely to decrease. Companies with sophisticated data processing techniques are likely to commit *fewer errors* in forecasting customer's orders. 3. As incomes grow, people have more *discretionary income* and, thus, more choice about how to spend their income. Hence,  $\sigma_i^2$  is likely to increase with income.

4. Similarly, companies with larger profits are expected to show greater variability in their dividend/buyback policies than companies with lower profits.

Heteroscedasticity can also be the result of *outliers* (either very small or very large). The inclusion/exclusion of an outlier, especially if T is small, can affect the results of regressions.

Violations of (A1) *-model is correctly specified*-, can produce heteroscedasticity, due to omitted variables from the model or incorrect functional form (e.g., linear vs log-linear models).

*Skewness* in the distribution of one or more regressors included in the model can induce heteroscedasticity. Examples are economic variables such as income, wealth, and education.

Heteroscedasticity is usually modeled using one the following specifications:

 $-H1: \sigma_t^2$  is a function of past  $\varepsilon_t^2$  and past  $\sigma_t^2$  (ARCH models).

- H2 :  $\sigma_t^2$  increases monotonically with one (or several) exogenous variable(s) ( $z_1$ , ...,  $z_j$ ).

- H3 :  $\sigma_t^2$  increases monotonically with  $E(y_t)$ .

-H4 :  $\sigma_t^2$  is the same within *p* subsets of the data but differs across the subsets (*grouped heteroscedasticity*). This specification allows for structural breaks.

These are the usual alternatives hypothesis (H1) in the heteroscedasticity tests.

## • Visual test

In a plot of residuals against dependent variable or other variable will often produce a fan shape.



# **Testing for Heteroscedasticity**

Question: Why do we want to test for heteroscedasticity if **b** is unbiased? OLS is no longer efficient. There is an estimator with lower asymptotic variance (the GLS/FGLS estimator).

We want to test: H<sub>0</sub>: E( $\varepsilon^2 | x_1, x_2, x_3, ..., x_k$ ) = E( $\varepsilon^2$ ) =  $\sigma^2$ 

H<sub>1</sub> and the structure of the test depend on what we consider the drivers of  $\sigma_i^2$  – i.e., in the previous examples: H1, H2, H3, H4, etc.

The key is whether  $E[\varepsilon^2] = \sigma_i^2$  is related to X and/or X<sup>2</sup>. Suppose we suspect a particular independent variable, say  $x_i$ , is driving  $\sigma_i^2$ .

Then, a simple test: Check the RSS for large values of  $x_j$ , and the RSS for small values of  $x_j$ . This is the Goldfeld-Quandt (GQ) test.

# **Testing for Heteroscedasticity: GQ Test**

GQ tests H<sub>0</sub>:  $\sigma_i^2 = s^2$ 

H<sub>1</sub>: 
$$\sigma_i^2 = f(\mathbf{X}_j)$$

• Easy to compute:

- Step 1. Arrange the data from small to large values of the independent variable suspected of causing heteroscedasticity,  $x_i$ .

- Step 2. Run two separate regressions, one for small values of  $x_j$  and one for large values of  $x_j$ , omitting *d* middle observations ( $\approx 20\%$ ). Get the RSS for each regression: RSS<sub>1</sub> for small values of  $x_j$  and RSS<sub>2</sub> for large  $x_j$ 's.

- Step 3. Calculate the F ratio

$$GQ = \frac{RSS_2}{RSS_1}, \sim F_{df,df}, \text{ with } df = [(T-d) - 2(k+1)]/2$$
 (A5 holds).

If (A5) does not hold, the F distribution becomes an approximation. Other tests may be preferred.

<u>Note</u>: When we suspect more than one variable is driving  $\sigma_i^2$ , the GQ test is not very useful.

• But, the GQ test is a popular test for structural breaks (two regimes) in variance. For these tests, we rewrite **step 3** to allow for a different sample size in the sub-samples 1 and 2, since the breaking point does not have to be in the middle of the sample.

- Step 3. Calculate the F-test ratio  $GQ = [RSS_2/(T_2 - k)]/[RSS_1/(T_1 - k)]$ 

<u>Note</u>: The package *lmtest* computes this test using function *gqtest*. It splits the sample in the middle. You need to specify the *d* of middle observations not included in test. Recall, you need to install the package before using it: install.packages("lmtest").

**Example:** We test if the 3-factor FF model for IBM and GE returns shows heteroscedasticity with a GQ test, using *gqtest* in package *lmtest*.

• IBM returns library(lmtest) > gqtest(ibm\_x ~ Mkt\_RF + SMB + HML, fraction = .20) Goldfeld-Quandt test

data:  $ibm_x \sim Mkt_RF + SMB + HML$  GQ = 1.1006, df1 = 224, df2 = 223, p-value = 0.2371  $\Rightarrow$  cannot reject H<sub>0</sub> at 5% level. alternative hypothesis: variance increases from segment 1 to 2

• GE returns gqtest(ge\_x ~ Mkt\_RF + SMB + HML, fraction = .20) Goldfeld-Quandt test

data:  $ge_x \sim Mkt_RF + SMB + HML$  GQ = 2.744, df1 = 281, df2 = 281, p-value < 2.2e-16  $\Rightarrow$  reject H<sub>0</sub> at 5% level. alternative hypothesis: variance increases from segment 1 to 2. ¶

## **Testing for Heteroscedasticity: LM Tests**

Popular heteroscedasticity LM tests:

- Breusch and Pagan (1979)'s LM test (BP).

- White (1980)'s general test.

Both tests are based on OLS residuals, **e**, and calculated under H<sub>0</sub> (No heteroscedasticity):  $\sigma^2$ . The squared residuals are used to estimate  $\sigma_i^2$ . • The BP test is an LM test, derived under normality -i.e., (A5). It is a general tests designed to detect any linear forms of heteroscedasticity, driven by some variables, z. That is, the BP tests:

H<sub>0</sub>: 
$$\sigma_i^2 = \sigma^2$$
  
H<sub>1</sub>:  $\sigma_i^2 = f(\mathbf{z}_i)$ 

• The White test is an asymptotic Wald-type test, where normality is not needed. It allows for nonlinearities by using squares and cross-products of all the *x*'s in the auxiliary regression –i.e., as the drivers of  $\sigma_i^2$ . That is, the White tests:

H<sub>0</sub>: 
$$\sigma_i^2 = \sigma^2$$
  
H<sub>1</sub>:  $\sigma_i^2 = f(x_1^2, x_2^2, ..., x_J^2, x_1x_2, x_1x_3, x_2x_3, ...)$ 

## **Testing for Heteroscedasticity: BP Test**

The derivation of the BP test is complicated, it relies on the likelihood function, which is constructed under normality, and its first derivative, the score. However, the implementation of the BP test is simple, based on the squared OLS residuals,  $e_i^2$ .

• Calculation of the Breusch-Pagan test

- Step 1. Run OLS on DGP:

 $\mathbf{y} = \mathbf{X} \boldsymbol{\beta} + \boldsymbol{\epsilon}.$  -Keep  $e_i$  and compute  $\sigma_R^2 = \text{RSS}/T$ 

- Step 2. (Auxiliary Regression). Run the regression of  $e_i^2/\sigma_R^2$  on the *m* explanatory variables, z. In our example,

 $e_i^2/\sigma_R^2 = \alpha_0 + \alpha_1 z_{1,i} + \dots + \alpha_m z_{m,i} + v_i$  -Keep RSS (*RSS<sub>e</sub>*)

- Step 3. With the RSS  $(RSS_e)$  from Step 2 regression and the Total Sum of Squares (TSS) also from Step 2, compute:

 $LM = (TSS - RSS_e)/2 \xrightarrow{d} \chi_m^2.$ 

• There is version of the BP, which is robust to departures from normality. It is the "*studentized*" version of Koenker (1981). The BP test is asymptotically equivalent to a  $T^*R^2$  test, where  $R^2$  is calculated from a regression of  $e_i^2/\sigma_R^2$  on the variables Z. (Omitting  $\sigma_R^2$  from the denominator is OK.)

• We have different Steps 2 & 3:

- Step 2. (Auxiliary Regression). Run the regression of  $e_i^2$  on the *m* explanatory variables, z. In our example,

$$e_i^2 = \alpha_0 + \alpha_1 z_{1,i} + \dots + \alpha_m z_{m,i} + v_i$$
 -Keep R<sup>2</sup> ( $R_{e2}^2$ )

- Step 3. Using the  $\mathbb{R}^2$  from Step 2. Let's call it  $\mathbb{R}^2_{e2}$ . Compute

$$LM = T R_{e2}^2 \xrightarrow{d} \chi_m^2$$

**Example:** We suspect that squared Mkt  $RF(x_1)$  –a measure of the overall market's variancedrives heteroscedasticity. We do a studentized LM-BP test for **IBM** in the 3-factor FF model: fit ibm ff3 <- lm (ibm  $x \sim Mkt RF + SMB + HML$ ) # Step 1 – OLS in DGP e <- fit ibm ff3\$residuals # Step 1 – keep residuals  $e2 \le e^2$ # Step 1 – squared residuals Mkt RF  $2 \le Mkt$  RF<sup>2</sup> fit **BP** <-  $lm(e2 \sim Mkt RF 2)$ # Step 2 – Auxiliary regression Re 2 <- summary(fit BP)\$r.squared # Step  $2 - \text{keep } R^2$ LM\_BP\_test <- Re2 \* T # Step 3 – Compute LM-BP test:  $\mathbb{R}^2 * \mathbb{T}$ >LM BP test [1] 0.25038  $p_val <- 1 - pchisq(LM_BP_test, df = 1)$ # p-value of LM test > p val[1] 0.6168019

LM-BP Test: 0.25028  $\Rightarrow$  cannot reject H<sub>0</sub> at 5% level ( $\chi^2_{[1],.05} \approx 3.84$ ); with a *p*-value= .6168.

<u>R Note</u>: The *bptest* in the *lmtest* package performs a studentized LM-BP test for the same variables used in the model (Mkt, SMB and HML). For IBM in the 3-factor FF model:

> bptest(ibm\_x ~ Mkt\_RF + SMB + HML) #bptest only allows to test H<sub>1</sub>: $\sigma_i^2 = f(\mathbf{x}_i = \text{model} \text{ variables})$ 

studentized Breusch-Pagan test

data:  $ibm_x \sim Mkt_RF + SMB + HML$ BP = 4.1385, df = 3, p-value = 0.2469

LM-BP Test: 4.1385  $\Rightarrow$  cannot reject H<sub>0</sub> at 5% level ( $\chi^2_{[3],05} \approx 7.815$ ); with a *p*-value = 0.2469.

<u>Conclusion</u>: Using the Breusch-Pagan test with Mkt\_RF^2 as the driver of heteroscedasticity, we cannot reject H<sub>0</sub>. That is, we cannot reject homocedasticity for the residuals of the 3-factor FF model for **IBM excess returns**. ¶

<u>Note</u>: Heteroscedasticity in financial time series is very common. In general, it is driven by squared market returns or squared past errors, thus the default setup of R's *bptest* is not very useful.

**Example:** We suspect that squared Market returns drive heteroscedasticity. We do an LM-BP (studentized) test for **Disney**:

 $\ln \operatorname{dis} \leq \log(x \operatorname{dis}[-1]/x \operatorname{dis}[-T])$ # Log returns for DIS # Disney excess returns dis  $x \leq lr dis - RF$ fit dis ff3 <- lm (dis  $x \sim Mkt RF + SMB + HML$ ) # Step 1 – OLS in DGP (3-factor FF model) e dis <- fit dis ff3\$residuals # Step 1 – keep residuals e2 <- e dis^2 # Step 2 – squared residuals fit dis BP <-  $lm (e2 \sim Mkt RF 2)$ # Step 2 – Auxiliary regression # Step  $2 - \text{Keep } R^2$  from Auxiliary reg **Re 2** <- summary(**fit dis BP**)\$r.squared LM\_BP\_test <- Re 2 \* T # Step 3 – Compute LM Test: R^2 \* T >LM BP test [1] 14.15224 > p val <- 1 - pchisq(LM BP test, df = 1) # p-value of LM test  $> p_val$ [1] 0.0001685967

LM-BP Test: 14.15  $\Rightarrow$  reject H<sub>0</sub> at 5% level ( $\chi^{2}_{[1],.05} \approx 3.84$ ); with a *p*-value = .0001.

• We do the same test but with SMB squared for Disney:

SMB\_2 <- SMB^2 fit\_dis\_BP\_2 <- lm (e2 ~ SMB\_2) Re\_2 <- summary(fit\_dis\_BP\_2)\$r.squared LM\_BP\_test\_2 <- Re\_2 \* T > LM\_BP\_test [1] 7.564692 p\_val <- 1 - pchisq(LM\_BP\_test\_2, df = 1) # p-value of LM\_test > p\_val [1] 0.005952284

LM-BP Test: 7.56  $\Rightarrow$  reject H<sub>0</sub> at 5% level ( $\chi^2_{[1],.05} \approx$  3.84); with a *p*-value= .006.

<u>Conclusion</u>: Using the Breusch-Pagan test, we reject homocedasticity for the residuals of the 3-factor FF model for **Disney excess returns**.

• If we do use the lmtest package, we get:

> bptest(dis\_x ~ Mkt\_RF + SMB + HML)

studentized Breusch-Pagan test

data: dis\_x ~ Mkt\_RF + SMB + HML BP = **6.9935**, df = 3, p-value = **0.07211** 

LM-BP Test: 6.99  $\Rightarrow$  cannot reject H<sub>0</sub> at 5% level ( $\chi^2_{[3],05} \approx 7.815$ ); with a p-value = .07211.

Note: Again, in general, you need squared values when model heteroscedasticity in financial assets.  $\P$ 

**Example:** We suspect that squared interest rate differentials drive heteroscedasticity for residuals in the encompassing (IFE + PPP) model for changes in the USD/GBP. We do an LM-BP (studentized) test, considering the squares of interest differentials as drivers of heteroscedasticity:

```
y <- lr_usdgbp
fit_gbp <- lm (y ~ inf_dif + int_dif)

e_gbp <- fit_gbp$residuals

e_gbp2 <- e_gbp^2

int_dif_2 <- int_dif^2

fit_BP <- lm (e_gbp2 ~ int_dif_2)

Re_2 <- summary(fit_BP)$r.squared

LM_BP_test <- Re_2 * T

> LM_BP_test

[1] 21.11134

p_val <- 1 - pchisq(LM_BP_test, df = 1)  # p-value of LM_test

> p_val

[1] 4.333567e-06
```

LM-BP Test: 21.11134  $\Rightarrow$  reject H<sub>0</sub> at 5% level (*p*-value < .00001).

<u>Conclusion</u>: Using the BP LM test, we have a strong rejection of homocedasticity for the residuals of the encompassing (PPP + IFE) model for changes in the USD/GBP.  $\P$ 

# **Testing for Heteroscedasticity: White Test**

The White test derivation is also complicated, but, the usual calculation of the White test is a known one for us:

- Step 1. (Same as BP's Step 1). Run OLS on DGP:

 $\mathbf{y} = \mathbf{X} \boldsymbol{\beta} + \boldsymbol{\epsilon}$ . Keep residuals, e<sub>i</sub>.

- Step 2. (Auxiliary Regression). Regress  $e^2$  on all the explanatory variables  $(x_j)$ , their squares  $(x_i^2)$ , & all their cross products  $(x_i * x_i)$ .

For example, when the model contains k = 2 explanatory variables, the test is based on:

 $e_i^2 = \beta_0 + \beta_1 x_{1,i} + \beta_2 x_{2,i} + \beta_3 x_{1,i}^2 + \beta_4 x_{2,i}^2 + \beta_5 x_{1,i} x_{2,i} + v_i$ 

Let *m* be the number of regressors in auxiliary regression (in the above example, m = 5). Keep  $R^2$ , say  $R_{e2}^2$ .

- Step 3. Compute the statistic:  $LM = T R_{e2}^2 \xrightarrow{d} \chi_m^2$ .

**Example:** White Test for the 3-factor F-F model for **IBM excess returns** (T=569). We also run the White Test for **DIS** and **GE excess returns**. In the case of IBM excess returns we have:

 $IBM_{Ret} - r_f = \beta_0 + \beta_1 (Mkt_{Ret} - r_f) + \beta_2 SMB + \beta_3 HML + \varepsilon$ 

e <- fit ibm ff3\$residuals e ibm $2 \le e$  ibm2Mkt HML <- Mkt RF\*HML Mkt SMB <- Mkt RF\*SMB SMB HML <- SMB\*HML xx2 <- cbind(Mkt\_RF\_2, SMB\_2, HML\_2, Mkt\_HML, Mkt\_SMB, SMB\_HML) fit ibm 2 <- lm(e ibm2 ~ Mkt RF + SMB + HML + xx2) # Not including original variables in Aux Reg (Mkt RF,SMB & HML) is OK r2 e2 <- summary(fit ibm 2)\$r.squared # Keep R^2 from Auxiliary regression >r2 e2 [1] 0.0166492 lm t <- T\*r2 e2 # Compute LM test:  $R^2$  \* sample size (T) > lm t[1] 10.93483 df  $lm \leq ncol(xx2)$ >df lm [1] 6 > qchisq(.95, df = df lm) [1] 12.59159

<u>Conclusion</u>: The White Test (LM-White Test: **10.93**) cannot reject H<sub>0</sub> at 5% level for the residuals of the 3-factor F-F model. (LM-White Test  $< \chi^2_{[6],.05} \approx$  **12.59**).

• Now, we do a White Test for the 3 factor F-F model for **DIS** and **GE excess returns** (*T*=569).

- For **DIS**, we get: e\_dis <- fit\_dis\_ff3\$residuals

```
e_dis2 <- e_dis^2

fit_dis_W <- lm (e_dis2 ~ Mkt_RF + SMB + HML + xx2)

Re_2W <- summary(fit_dis_W)$r.squared

LM_W_test <- Re_2W * T

> LM_W_test

[1] 25.00148 \Rightarrow reject H<sub>0</sub> at 5% level (\chi^2_{[6],05}\approx12.59).

p_val <- 1 - pchisq(LM_W_test, df = 6) # p-value of LM_test

> p_val

[1] 0.0003412389
```

- For GE, we get: LM-White Test: 20.15 (*p*-value = 0.0026)  $\Rightarrow$  reject H<sub>0</sub> at 5% level.

<u>Conclusion</u>: Using the White test, we strongly reject homoscedasticity for the errors of the 3-factor FF model for **DIS returns** and **GE returns**.

**Example:** We do a White Test for the residuals in the encompassing (IFE + PPP) model for changes in the **USD/GBP** (T=363):

e\_gbp <- fit\_gbp\$residuals e\_gbp2 <- e\_gbp^2 int\_dif2 <- int\_dif^2; inf\_dif2 <- inf\_dif^2; int\_inf\_dif <- int\_dif\*inf\_dif fit\_gbp\_W <- lm (e\_gbp2 ~ int\_dif + inf\_dif + int\_dif2 + inf\_dif2+ int\_inf\_dif) Re\_2W <- summary(fit\_gbp\_W)\$r.squared LM\_W\_test <- Re\_2W \* T > LM\_W\_test [1] 15.46692 p\_val <- 1 - pchisq(LM\_W\_test, df = 3) # p-value of LM\_test > p\_val [1] 0.001458139  $\Rightarrow$  reject H<sub>0</sub> at 5% level

<u>Conclusion</u>: Using the White test, we strongly reject homoscedasticity for the residuals of the encompassing (PPP + IFE) for changes in the **USD/GBP.**  $\P$ 

# **Testing for Heteroscedasticity: LR Test**

We define the likelihood function, assuming normality –i.e. (A5)–, for a general case, where we have *g* different variances:

$$\ln L = -\frac{T}{2} \ln 2\pi - \sum_{i=1}^{g} \frac{T_i}{2} \ln \sigma_i^2 - \frac{1}{2} \sum_{i=1}^{g} \frac{1}{\sigma_i^2} (y_i - X_i \beta)' (y_i - X_i \beta)$$

We have two models:

(R) Restricted under H<sub>0</sub>:  $\sigma_i^2 = \sigma^2$ . From this model, we calculate ln *L* 

$$\ln L_R = -\frac{T}{2} [\ln(2\pi) + 1] - \frac{T}{2} \ln(\hat{\sigma}^2)$$

(U) Unrestricted. From this model, we calculate the log likelihood.

$$\ln L_U = -\frac{T}{2} [\ln(2\pi) + 1] - \sum_{i=1}^{g} \frac{T_i}{2} \ln \hat{\sigma}_i^2; \quad \hat{\sigma}_i^2 = \frac{1}{T_i} (y_i - X_i b)' (y_i - X_i b)$$

• Now, we can estimate the Likelihood Ratio (LR) test:

$$LR = 2(\ln L_U - \ln L_R) = T \ln \hat{\sigma}^2 - \sum_{i=1}^{g} T_i \ln \hat{\sigma}_i^2 \xrightarrow{a} \chi_{g-1}^2$$

Under the usual regularity conditions, LR is approximated by a  $\chi^2_{q-1}$ .

# **Testing for Heteroscedasticity: Remarks**

Drawbacks of the Breusch-Pagan test:

- It is sensitive to violations of the normality assumption. The studentized version of Koenker is more robust and, then, more used.

Drawbacks of the White test

- If a model has several regressors, the test can consume a lot of df's.

- In cases where the White test statistic is statistically significant, heteroscedasticity may not necessarily be the cause, but model specification errors.

- It is general. It does not give us a clue about how to model heteroscedasticity to do FGLS. The BP test points us in a direction.

- In simulations, it does not perform well relative to others, especially, for time-varying heteroscedasticity, typical of financial time series.

#### **Finding Auto-correlation**

There are several reasons why the  $\varepsilon_i$  may be related to  $\varepsilon_j$ . In general, we find autocorrelation (or serial correlation) in time series, where  $\varepsilon_{i=t}$  is correlated to  $\varepsilon_{j=t-l}$ . Typical situation: it takes time to absorb a shock, then, shocks show persistence over time.

The shocks can also be correlated over the cross-section, causing cross-correlation. For example, if an unexpected new tax is imposed on the technology sector, all the companies in the sector are going to share this shock.

Usually, we model autocorrelation using two models: autoregressive (AR) and moving averages (MA):

- In an AR model, the errors,  $\varepsilon_t$ , show a correlation over time.

- In an MA model, the errors,  $\varepsilon$  t, are a function (similar to a weighted average) of previous errors, now denoted ut's.

#### **Examples:**

- First-order autoregressive autocorrelation: AR(1)

 $\varepsilon_{t} = \rho_{1} \varepsilon_{t-1} + u_{t}$ - p<sup>th</sup>-order autoregressive autocorrelation: AR(p)  $\varepsilon_{t} = \rho_{1} \varepsilon_{t-1} + \rho_{2} \varepsilon_{t-2} + \dots + \rho_{p} \varepsilon_{t-p} + u_{t}$ 

- Third-order moving average autocorrelation: MA(3)

 $\varepsilon_t = u_t + \lambda_1 \ u_{t-1} + \lambda_2 u_{t-2} + \lambda_3 u_{t-3}$ 

<u>Note</u>: The last example is described as third-order moving average autocorrelation, denoted MA(3), because it depends on the three previous innovations as well as the current one.

#### **Finding Auto-correlation – Visual Check**

Plot data, usually residuals from a regression, to see if there is a pattern:

- **Positive autocorrelation**: A positive (negative) observation tends to be followed by a positive (negative) observation. We tend to see continuation in the series.

- **Negative autocorrelation**: A positive (negative) observation tends to be followed by a negative (positive) observation. We tend to see reversals.

- **No autocorrelation**: A positive (negative) observation has the same probability of being followed by a negative or positive (positive or negative) observation. We tend to see no pattern.

**Simulated Example:** I simulate a  $y_t$  series, with N(0,1)  $u_t$  errors:

 $y_t = \rho_1 y_{t-1} + u_t$ Three cases: (1) Positive autocorrelation:  $\rho_1 = .70$ (2) Negative autocorrelation:  $\rho_1 = -.70$ (3) No correlation:  $\rho_1 = -0$ • R code for simulation: T sim <- 200  $u \leq rnorm(200)$ # Draw T sim normally distributed errors  $y_sim \le matrix(0, T_sim, 1)$ rho <- .7 # Change to create different correlation patterns a <- 2 # Time index for observations while (a  $\leq T \sin$ ) { y sim[a] = rho \* y sim[a-1] + u[a]# y sim simulated autocorrelated values

a <- a + 1
}
plot(y\_sim, type="l", col="blue", ylab ="Simulated Series", xlab ="Time")
title("Visual Test: Autocorrelation?")</pre>







**Example:** Residual plot for the 3 factor F-F model for **IBM returns** and **GE returns**:



<u>Conclusion</u>: It looks like a small  $\rho_1$ , but not very clear pattern from the graphs.

#### **Testing for Autocorrelation: LM Test**

There are several autocorrelation tests. All autocorrelations tests described in this lecture have an AR(p) structure, as an alternative hypothesis:

$$\varepsilon_t = \rho_1 \, \varepsilon_{t-1} + \rho_2 \, \varepsilon_{t-2} + \dots + \rho_p \, \varepsilon_{t-p} + u_t$$

In this section, we describe a popular LM test, the Breusch–Godfrey (BG, 1978) LM test. BG use the AR(p) model as the basis of H<sub>1</sub> and, thus, the test:

H<sub>0</sub>:  $\rho_1 = ... = \rho_p = 0$ . H<sub>1</sub>: at least one :  $\rho_i \neq 0$ , for i = 1, 2, ..., p

Under H<sub>0</sub>, BG use OLS residuals,  $e_t$ , to construct an LM test (BG test), similar to the BP test.

• Steps for the Breusch–Godfrey (1978): – Step 1. (Same as BP's Step 1). Run OLS on DGP:  $y = X \beta + \varepsilon$ . - Keep residuals,  $e_t$ .

- Step 2. (Auxiliary Regression). Run the regression of  $e_t$  on all the explanatory variables, X, and p lags of residuals,  $e_t$ :

 $e_t = \mathbf{x}_t, \mathbf{\gamma} + \alpha_1 e_{t-1} + \dots + \alpha_p e_{t-p} + v_t$  - Keep R<sup>2</sup> ( $R_e^2$ )

- Step 3. Keep the  $\mathbb{R}^2$  from this regression. Let's call it  $\mathbb{R}^2_e$ . Then, calculate:

LM =  $(T - p) R_e^2 \xrightarrow{d} \chi_p^2$ . - (T - p) = we lost p observation by taking lags of e.

<u>Note</u>: In general, in Step 2, if we do not include  $x_t$ , the LM test is not that different.

**Example:** LM-AR Test for the 3 factor F-F model for **IBM excess returns** (p=12 lags):

```
e ibm <- fit ibm ff3$residuals
                                            # OLS residuals
p lag <- 12
                                            # Select # of lags for test (set p)
e lag <- matrix(0,T-p lag,p lag)
                                            # Matrix to collect lagged residuals
a <- 1
while (a \le p \text{ lag})
                                            # Do loop creates matrix (e lag) with lagged e
za \le e ibm[a:(T-p lag+a-1)]
 e lag[,a] \leq za
a <- a+1
}
                                                    # Adjust for new sample size: T - p lag
Mkt RF p <- Mkt RF[(p lag+1):T]
SMB p \le SMB[(p lag+1):T]
HML p \le HML[(p lag+1):T]
fit lm1 \le lm(e ibm[(p lag+1):T] \sim e lag + Mkt RF p + SMB p + HML p) # Auxiliary Reg.
r2 e1 <- summary(fit lm1)$r.squared
                                                    # get R^2 from Auxiliary Regression
> r2 e1
[1] 0.0303721
> (T-p lag)
[1] 557
lm_t <- (T-p_lag )* r2 e1
                                                    # LM-test wih p lags
> lm t
[1] 16.91726
df \leq ncol(e lag)
                                                    # degrees of freedom for the LM Test
> 1 - pchisq(lm t,df)
[1] 0.1560063
LM-AR(12) Test: 16.91726
                                            \Rightarrow cannot reject H<sub>0</sub> at 5% level (p-value > .05).
```

• If I run the test with p = 4 lags, I get LM-AR(4) Test: 2.9747 (*p*-value = 0.56)  $\Rightarrow$  cannot re

 $\Rightarrow$  cannot reject H<sub>0</sub> at 5% level (*p*-value > .05).

<u>Conclusion</u>: No evidence of autocorrelation for the residuals of the 3-factor FF model for IBM excess returns at the 5% level.  $\P$ 

<u>R Note</u>: The package *lmtest*, performs this test, *bgtest*, (and many others, used in this class, encompassing, jtest, waldtest, etc). Recall that you need to install it first: install.packages("lmtest"), then call the library(lmtest).

library(lmtest)
> bgtest(ibm\_x ~ Mkt\_RF + SMB + HML, order=12)

Breusch-Godfrey test for serial correlation of order up to 12

data:  $lr_ibm \sim Mkt_RF + SMB + HML$ LM test = 16.259, df = 12, p-value = 0.1797 (minor difference with the previous test, likely due to multiplication by *T*. Results do not change much)

<u>Note</u>: If you do not include in the Auxiliary Regression the original regressors (Mkt\_RF, SMB, HML) the test do not change much. You get LM-AR(12) Test: 16.83253  $\Rightarrow$  very similar. Not entirely correct, but it works well. ¶

**Example:** Autocorrelation is very common. If I run the test for **Disney**, **CNP**, or **GE**, instead, we get significant test results.

- For **DIS**: lr\_dis <- log(x\_dis[-1]/x\_dis[-T]) dis\_x <- lr\_dis - RF

> bgtest(dis\_x ~ Mkt\_RF + SMB + HML, order=4)
Breusch-Godfrey test for serial correlation of order up to 4

data: dis\_x ~ Mkt\_RF + SMB + HML LM test = 8.6382, df = 4, p-value =  $0.07081 \Rightarrow$  cannot reject H<sub>0</sub> at 5% level (*p-value* >.05)

> bgtest(dis\_x ~ Mkt\_RF + SMB + HML, order=12)
Breusch-Godfrey test for serial correlation of order up to 12

data: dis\_x ~ Mkt\_RF + SMB + HML LM test = 30.068, df = 12, p-value = 0.002728  $\Rightarrow$  reject H<sub>0</sub> at 5% level (*p-value* < .05)

- For GE (with 12 lags): lr\_ge <- log(x\_ge[-1]/x\_ge[-T]); ge\_x <- lr\_ge - RF > bgtest(ge\_x ~ Mkt\_RF + SMB + HML, order=4) Breusch-Godfrey test for serial correlation of order up to 4

data:  $ge_x \sim Mkt_RF + SMB + HML$ LM test = 28.257, df = 4, p-value = 0.005073  $\Rightarrow$  cannot reject H<sub>0</sub> at 5% level (*p-value* >.05)

- For CNP (with 12 lags): lr\_cnp <- log(x\_cnp[-1]/x\_cnp[-T]); cnp\_x <- lr\_cnp - RF > bgtest(cnp\_x ~ Mkt\_RF + SMB + HML, order=12) Breusch-Godfrey test for serial correlation of order up to 12

data:  $cnp_x \sim Mkt_RF + SMB + HML$ LM test = **31.718**, df = **12**, p-value = **0.00153** 

 $\Rightarrow$  reject H<sub>0</sub> at 5% level (*p*-value < .05)

<u>Conclusion</u>: Significant evidence of autocorrelation for the residuals of the 3-factor F-F model for **DIS**, **GE & CNP excess returns**.

• Question: How many lags are needed in the test?

Enough to make sure there is no auto-correlation left in the residuals. There are some popular rule of thumbs: for daily data, 5 or 20 lags; for weekly, 4 or 12 lags; for monthly data, 12 lags; for quarterly data, 4 lags.

# **Testing for Autocorrelation: Durbin-Watson**

The Durbin-Watson (1950) (DW) test for AR(1) autocorrelation:  $H_0$ :  $\rho_1 = 0$  against  $H_1$ :  $\rho_1 \neq 0$ . Based on simple correlations of  $\boldsymbol{e}$ .

$$d = \frac{\sum_{t=2}^{T} (e_t - e_{t-1})^2}{\sum_{t=1}^{T} e_t^2}$$

It is easy to show that when  $T \to \infty$ ,  $d \approx 2(1 - \rho_1)$ .

It is estimated using the sample correlation *r*.

Under H<sub>0</sub>,  $\rho_1 = 0$ . Then, *d* should be distributed randomly around 2.

Small values (close to 0) or Big values (close to 4) of d lead to rejection of  $H_0$ . The distribution depends on **X**. Durbin-Watson derived bounds for the test. Since there are better tests, in practice, the DW is used "visually:" Is d close to 2?.

<u>**R** Note</u>: The function *dwtest* from the *lmtest* package computes d (DW test) and produces also a *p*-value.

**Example:** DW Test for the 3 factor F-F model for **IBM returns** 

<b>fit_ibm_ff3</b> <- $lm(ibm_x \sim Mkt_RF + SMB + HML)$	# OLS regression
e_ibm <- fit_ibm_ff3\$residuals	# OLS residuals
$RSS \le sum(e_ibm^2)$	# RSS
$DW \le sum((e[1:(T-1)]-e[2:T])^2)/RSS$	# DW stat
> DW	
[1] 2.042728 $\Rightarrow$ DW statistic $\approx 2 \Rightarrow$ No ev	vidence for autocorrelation of order 1.
> 2*(1-cor(e[1:(T-1)],e[2:T]))	# approximate DW stat
[1] 2.048281	

• Similar finding for Disney returns:

> DW [,1] [1,] 2.1609  $\Rightarrow$  DW statistic  $\approx 2 \Rightarrow$  But, DIS suffers from autocorrelation!

 $\Rightarrow$  This is why DW are not that informative. They only test for AR(1) in residuals.

<u>R Note</u>: The package *lmtest* performs this test too, using *dwtest*:

> dwtest(y ~ Mkt\_RF + SMB + HML) DW = 2.0427, p-value = 0.7087

<u>Conclusion</u>: No evidence of first order autocorrelation in the residuals of the 3-factor F-F model for **IBM excess returns**.

**Example:** DW Test for the residuals of the encompassing model (IFE + PPP) for changes in USD/GBP: e gbp <- fit gbp\$residuals

> dwtest(fit\_gbp)

Durbin-Watson test

data: fit\_gbp DW = 1.8588, p-value = 0.08037  $\Rightarrow$  not significant at 5% level. alternative hypothesis: true autocorrelation is greater than 0

<u>Conclusion</u>: No evidence of first order autocorrelation in the residuals of the encompassing model (IFE + PPP) for changes in **USD/GBP**.  $\P$ 

## **Testing for Autocorrelation: Portmanteu tests**

Portmanteu tests are tests with a well-defined  $H_0$ , but not specific, or loosely defined,  $H_1$ . We present two Portmanteu test for autocorrelation: the Box-Pierce (1970) test and its modification, the Ljung-Box (1978) test.

- Box-Pierce (1970) test (Q test). For a series  $y_t$ , it tests H<sub>0</sub>:  $\rho_1 = ... = \rho_p = 0$  using the sample correlation:  $r_j = \frac{\hat{\gamma}_j}{\hat{\gamma}_0}$ where, using time series notation, we have:

 $\hat{\gamma}_j$  = sample covariance between  $y_t$  and  $y_{t-j} = \frac{\sum_{t=j+1}^{T} (y_t - \bar{y})(y_{t-j} - \bar{y})}{T-j}$  $\hat{\gamma}_0$  = sample variance.

In the case of analyzing residuals of a regression,  $e_t$ , we compute  $r_j$  as:

$$r_{j} = \frac{\sum_{t=j+1}^{T-j} e_{t} e_{t-j}}{\sum_{t=1}^{T} e_{t}^{2}}$$

$$H_{0:} \qquad \mathbf{Q} = T \sum_{j=1}^{p} r_{j}^{2} \xrightarrow{d} \chi_{p}^{2}.$$

Then, under Ho

- Ljung-Box (1978) test (LB test).

A variation of the Box-Pierce test. It has a small sample correction, which improves the performance of the test:

$$LB = T * (T+2) * \sum_{j=1}^{p} \frac{r_j^2}{T-j} \stackrel{d}{\longrightarrow} \chi_p^2.$$

Both statistics test whether a group of autocorrelations are different from zero. Both are general tests, not testing zero-autocorrelation at each lag.

<u>Technical note</u>: The asymptotic distribution is based on the fact that, under the null hypothesis of independent data,  $\sqrt{T} \mathbf{r} \xrightarrow{d} N(0, \mathbf{I})$ .

Both tests are widely used, especially the LB test. But, the Breusch–Godfrey (1978) LM tests conditions on **X**. Thus, LM tests are more powerful.

**Example:** Q and LB tests with p = 12 lags for the residuals in the 3-factor FF model for IBM excess returns:

```
RSS <- sum(e ibm^2)
r sum < 0
lb sum \leq -0
p lag <- 12
a <- 1
while (a \le p \text{ lag})
        za \le as.numeric(t(e ibm[(p lag+1):T]) \%*\% e ibm[a:(T-p lag+a-1)])
 r sum \leq r sum + (za/RSS)^2
                                                  \# \operatorname{sum} \operatorname{cor}(e[(p \ lag+1):T], e[a:(T-p \ lag+a-1)])^2
 lb\_sum \le lb\_sum + (za/RSS)^2/(T-a)
                                                 # sum with LB correction
a <- a + 1
}
Q \leq T^*r \text{ sum}
> Q
[1] 16.39559 (p-value = 0.1737815)
                                                          \Rightarrow cannot reject H<sub>0</sub> at 5% level.
LB \leq T^{*}(T-2)^{*}lb sum
>LB
[1] 16.46854 (p-value = 0.1707059)
                                                           \Rightarrow cannot reject H<sub>0</sub> at 5% level.
```

<u>R Note</u>: The *Box.test* function computes Q & LB: - Q test e\_ibm <- fit\_ibm\_ff3\$residuals > Box.test(e\_ibm, lag = 12, type="Box-Pierce")

**Box-Pierce** test

data:  $e_{ibm}$ X-squared = 16.304, df = 12, p-value = 0.1777

- LB test > Box.test(e\_ibm, lag = 12, type="Ljung-Box")

Box-Ljung test

data:  $e_{ibm}$ X-squared = **16.61**, df = **12**, p-value = **0.1649** 

<u>Note</u>: There is a minor difference between the previous code and the code in Box.test. They are based on how the correlations of e are computed (centered around the mean, or assumed zero mean).

<u>Conclusion</u>: Using the Q and LB tests, with different lags, we find no evidence of autocorrelation in the residuals of the 3-factor F-F model for **IBM excess returns**.

• Same tests (p = 12 lags) & same model: for Disney & GE. - For DIS (e\_dis), we get: fit\_dis\_ff3 <- lm(ge\_x ~ Mkt\_RF + SMB + HML) # OLS regression e\_ge <- fit\_dis\_ff3\$residuals # OLS residuals > Q [1] 25.563 (p-value = 0.01237)  $\Rightarrow$  reject H<sub>0</sub> at 5% level. > LB [1] 25.879 (p-value = 0.01117)  $\Rightarrow$  reject H<sub>0</sub> at 5% level. - For GE (e ge), we get

fit\_ge\_ff3 <-  $lm(ge_x \sim Mkt_RF + SMB + HML)$  # OLS regression e\_ge <- fit\_ge\_ff3\$residuals # OLS residuals > Q [1] 27.087 (*p*-value = 0.007507)  $\Rightarrow$  reject H<sub>0</sub> at 5% level. > LB [1] 27.523 (*p*-value = 0.006493)  $\Rightarrow$  reject H<sub>0</sub> at 5% level. Conclusion: Using the Q and LB test, we find evidence of autocorrelation in the residuals of the 3-factor F-F model for DIS & GE excess returns.

• Autocorrelation in financial asset returns is a usual finding in monthly, weekly and daily data.

**Example:** Same Q and LB tests (p=12 lags) for the USD/GBP residuals in the encompassing (PPP + IFE) model:

e gbp <- fit gbp\$residuals > Box.test(e gbp, lag = 12, type="Box-Pierce")

**Box-Pierce** test

data: e gbp X-squared = 19.587, df = 12, p-value = 0.0753  $\Rightarrow$  cannot reject H<sub>0</sub> at 5% level, but close.

> Box.test(e gbp, lag = 12, type="Ljung-Box")

Box-Ljung test

data: e gbp X-squared = 20.032, df = 12, p-value = 0.06649  $\Rightarrow$  cannot reject H<sub>0</sub> at 5% level, but close.

Conclusion: Using the Q and LB tests, with different lags, we find no evidence of autocorrelation in the residuals of the encompassing (PPP + IFE) model for changes in the USD/GBP. ¶

Above, we mentioned that the Q & LB tests are widely used. But, they present two main limitations:

(1) The test was developed under the independence assumption.

If there is dependence in the data, such as heteroscedasticity, the asymptotic variance of  $\sqrt{T} r$  is no longer I, but a non-diagonal matrix.

There are several proposals to "robustify" both Q & LB tests, see Diebold (1986), Robinson (1991), Lobato et al. (2001). The "robustified" Portmanteau statistic uses  $\tilde{r}_i$  instead of  $r_i$ :

$$\widetilde{\gamma}_{j} = \frac{\widehat{\gamma}_{j}^{2}}{\tau_{j}} = \frac{\sum_{t=j+1}^{T} (y_{t} - \bar{y})(y_{t-j} - \bar{y})}{\sum_{t=j+1}^{T} (y_{t} - \bar{y})^{2} (y_{t-j} - \bar{y})^{2}}$$

Thus, for Q we have:

 $\mathbf{Q}^* = T \ \sum_{i=1}^p \tilde{r}_i^2 \xrightarrow{d} \chi_n^2.$ 

(2) The selection of the number of autocorrelations p is arbitrary.

The traditional approach is to try different p values, say 3, 6 & 12. Another popular approach is to let the data "select" p, for example, using AIC or BIC, an approach sometimes referred as "automatic selection."

Escanciano and Lobato (2009) propose combining BIC's and AIC's penalties to select p in Q\* (BIC for small r and AIC for bigger r). The Auto.Q function in the R package *vrtest* computes Q\* with this automatic selection of p.

It is possible to reach very different conclusion from Q and Q\*.

**Example:** Q\* tests with automatic selection of **p** for the residuals in the 3-factor FF model for **IBM, DIS and GE excess returns**:

For IBM (e\_ibm), we get:
> library(vrtest)
> Auto.Q(e\_ibm, 12)
\$Stat
[1] 0.2781782

\$Pvalue [1] **0.5978978** 

```
- For DIS (e_dis), we get:
> Auto.Q(e_dis, 12)
$Stat
[1] 2.649553
```

\$Pvalue [1] **0.103579** 

- For GE (e\_ge), we get: > Auto.Q(e\_ge, 12) \$Stat [1] 0.6437697

\$Pvalue [1] **0.422349** 

<u>Conclusion</u>: Using the Q\* test, with automatic lag selection, we find no evidence of autocorrelation in the residuals of the 3-factor F-F model for **IBM excess returns**. Same conclusion we reached with the Q & LB tests above. But, for **DIS** residuals and **GE** residuals we get a different conclusion. Now, once we take into consideration heteroscedasticity, we cannot reject at the 5% level the null hypothesis of no autocorrelation, ¶

Time-varying volatility is very common in financial time series. We can use the Q and LB tests for autocorrelation to check for autocorrelation in squared errors,  $e_i^2$ , which based on White's idea, we use to estimate  $\sigma_i^2$ .
#### **Testing for Autocorrelation: Heteroscedasticity**

We use the Q and LB Portmanteu tests on the squared residuals to check for a particular kind of heteroscedasticity: the variance,  $\sigma_i^2$ , is driven by lagged squared errors.

H<sub>0</sub>: 
$$\sigma_i^2 = s^2$$
  
H<sub>1</sub>:  $\sigma_i^2 = f(\varepsilon_{i-1}^2, \varepsilon_{i-2}^2, ..., \varepsilon_{i-p}^2)$ 

• Of course, an LM-BP test can also be used, using lagged squared residuals as the drivers of heteroscedasticity (more on this topic in Lecture 10).

**Example:** Q and LB tests with p=12 lags for the squared residuals in the 3-factor FF model for IBM returns:

e\_ibm <- fit\_ibm\_ff3\$residuals e\_ibm2 <- e\_ibm^2

• Q test > Box.test(e ibm2, lag = 12, type="Box-Pierce")

Box-Pierce test

data: e\_ibm2 X-squared = **37.741**, df = 12, p-value = **0.0001693** 

• LB test > Box.test(e\_ibm2, lag = 12, type="Ljung-Box")

Box-Ljung test

data: e\_ibm2 X-squared = **38.435**, df = 12, p-value = **0.0001304** 

• Q and LB tests with p=12 lags for the squared residuals in the 3-factor FF model for **DIS** & **GE** returns:

- For **DIS** (dis\_x), we get > Box.test(e\_dis2, lag = 12, type="Ljung-Box")

Box-Ljung test

data: e\_dis2 X-squared = **73.798**, df = 12, p-value = **6.195e-11** 

- For GE (ge\_x), we get > Box.test(e\_ge2, lag = 12, type="Ljung-Box") Box-Ljung test

data: e\_ge2 X-squared = **115.9**, df = 12, p-value < **2.2e-16** 

<u>Conclusion</u>: Using Q and LB tests for squared residuals, we find strong evidence for timevarying heteroscedasticity in the residuals of the 3-factor F-F model for **IBM & DIS excess returns**.

## **Generalized Regression Model (GRM)**

Now, we go back to the CLM Assumptions: (A1) DGP:  $\mathbf{y} = \mathbf{X} \boldsymbol{\beta} + \boldsymbol{\varepsilon}$  is correctly specified. (A2) or (A2') (A3') Var[ $\boldsymbol{\varepsilon} | \mathbf{X} ] = \boldsymbol{\Sigma}$  (sometimes written Var[ $\boldsymbol{\varepsilon} | \mathbf{X} ] = \sigma^2 \boldsymbol{\Omega}$ )  $\boldsymbol{\Sigma} = \begin{bmatrix} \sigma_1^2 & \sigma_{12} & \cdots & \sigma_{1T} \\ \sigma_{21} & \sigma_2^2 & \cdots & \sigma_{2T} \\ \vdots & \vdots & \vdots & \vdots \\ \sigma_{T1} & \sigma_{T2} & \cdots & \sigma_T^2 \end{bmatrix}$ -a (*T*x*T*) symmetric matrix (A4) or (A4')

This is the generalized regression model (GRM), which allows the variances to differ across observations and allows correlation across observations.

OLS is still unbiased. Can we still use OLS?

## **GR Model: True Variance for b**

From (A3)  $\operatorname{Var}[\boldsymbol{\varepsilon}|\mathbf{X}] = \sigma^2 \mathbf{I}_T \implies \operatorname{Var}[\mathbf{b}|\mathbf{X}] = \sigma^2 (\mathbf{X}'\mathbf{X})^{-1}.$ 

Now, we have (A3')  $Var[\varepsilon|X] = \Sigma$ 

The true variance of **b** under (**A3'**) should be:

 $Var_{T}[\mathbf{b}|\mathbf{X}] = E[(\mathbf{b} - \boldsymbol{\beta})(\mathbf{b} - \boldsymbol{\beta})'|\mathbf{X}]$ = (**X**'**X**)<sup>-1</sup> E[**X**' $\boldsymbol{\epsilon}\boldsymbol{\epsilon}$ '**X**|**X**] (**X**'**X**)<sup>-1</sup> = (**X**'**X**)<sup>-1</sup> **X**' $\boldsymbol{\Sigma}$ **X** (**X**'**X**)<sup>-1</sup>

**Example:** We compute the true variance for the simplest case, a regression with only one explanatory variable and uncorrelated error term:

$$\mathbf{y} = \mathbf{X} \boldsymbol{\beta} + \boldsymbol{\varepsilon}$$

Then,

$$\operatorname{Var}_{\mathrm{T}}[\mathbf{b}|\mathbf{X}] = \left(\frac{1}{\sum_{i}^{T} (x_{i} - \bar{x})^{2}}\right)^{2} \sum_{i}^{T} \sigma_{i}^{2} (x_{i} - \bar{x})^{2}.$$

If we compute the OLS variance, we see how both estimators differ:

$$\operatorname{Var}[\mathbf{b}|\mathbf{X}] = \frac{\sigma^2}{\sum_{i}^{T} (x_i - \bar{x})^2} \neq \operatorname{Var}[\mathbf{b}|\mathbf{X}]$$

<u>Note</u>: In the special case that  $\sigma_i^2$  is independent of (uncorrelated with) of  $(x_i - \bar{x})^2$ , then both variances are (asymptotically) the same since

$$\sum_{i}^{T} \sigma_{i}^{2} (x_{i} - \bar{x})^{2} \xrightarrow{p} \sigma^{2} \sum_{i}^{T} (x_{i} - \bar{x})^{2}. \P$$

Under (A3'), the usual OLS estimator of Var[**b**|**X**] –i.e.,  $s^2 (\mathbf{X'X})^{-1}$ – is *biased*. If we want to use OLS for inferences (say, with *t-test* or *F-test*),, we need to estimate Var<sub>T</sub>[**b**|**X**]. That is, we need to estimate the unknown  $\Sigma$ . But, it has  $T^*(T+1)/2$  parameters. Too many parameters to estimate with only *T* observations!

#### **GR Model: Robust Covariance Matrix**

We will not be estimating  $\Sigma$ . Impossible with *T* data points.

We will estimate  $\mathbf{X'}\Sigma\mathbf{X} = \sum_{i=1}^{T} \sum_{j=1}^{T} \sigma_{ij} \mathbf{x}_i \mathbf{x}_j'$ , a (*kxk*) matrix. That is, we are estimating  $[k^*(k+1)]/2$  elements.

This distinction is very important in modern applied econometrics:

– The White estimator

- The Newey-West estimator

Both estimators produce a *consistent* estimator of Var<sub>T</sub>[**b**|**X**].

Since **b** consistently estimates  $\beta$ , the OLS residuals, **e**, are also consistent estimators of  $\varepsilon$ . We use **e** to consistently estimate  $X'\Sigma X$ .

#### **Covariance Matrix: The White Estimator**

The White estimator simplifies the estimation since it only assumes heteroscedasticity. Then,  $\Sigma$  is a diagonal matrix, with elements  $\sigma_i^2$ .

$$\boldsymbol{\Sigma} = \begin{bmatrix} \sigma_1^2 & 0 & \cdots & 0 \\ 0 & \sigma_2^2 & \cdots & 0 \\ \vdots & \vdots & \vdots & \vdots \\ 0 & 0 & \cdots & \sigma_T^2 \end{bmatrix}$$

Thus, we need to estimate:

$$\mathbf{Q}^* = (1/T) \mathbf{X'} \mathbf{\Sigma} \mathbf{X}$$

where

$$X' \Sigma X = \begin{bmatrix} \sum_{i=1}^{T} x_{1i}^2 \sigma_i^2 & \cdots & \sum_{i=1}^{T} x_{1i} x_{ki} \sigma_i^2 \\ \vdots & \ddots & \vdots \\ \sum_{i=1}^{T} x_{ki} x_{1i} \sigma_i^2 & \cdots & \sum_{i=1}^{T} x_{ki}^2 \sigma_i^2 \end{bmatrix} = \sum_{i=1}^{T} \sigma_i^2 x_i x_i'$$

Question: How do we estimate  $\sigma_i^2$ ?

We need to estimate:  $\mathbf{Q}^* = (1/T) \mathbf{X}' \mathbf{\Sigma} \mathbf{X} = (1/T) \sum_{i=1}^T \sigma_i^2 \mathbf{x}_i \mathbf{x}_i'$ 

The OLS residuals, e, are consistent estimators of  $\varepsilon$ . This suggests using  $e_i^2$  to estimate  $\sigma_i^2$ . That is,

we estimate  $\mathbf{Q}^* = (1/T) \sum_{i=1}^T \sigma_i^2 \mathbf{x}_i \mathbf{x}_i'$ with  $\mathbf{S}_0 = (1/T) \sum_{i=1}^T e_i^2 \mathbf{x}_i \mathbf{x}_i'$ 

**Example:** Back to the simplest case, a regression with only one explanatory variable, but now with a heteroscedastic error term, we have that the variance of **b** is given by:

$$\operatorname{Var}_{\mathrm{T}}[\mathbf{b}|\mathbf{X}] = \left(\frac{1}{\sum_{i}^{T} (x_{i} - \bar{x})^{2}}\right)^{2} \sum_{i=1}^{T} \sigma_{i}^{2} (x_{i} - \bar{x})^{2}$$

which we estimate using OLS residuals,  $e_i$ :

Est Var<sub>T</sub>[**b**|**X**] = 
$$\left(\frac{1}{\sum_{i=1}^{T} (x_i - \bar{x})^2}\right)^2 \sum_{i=1}^{T} e_i^2 (x_i - \bar{x})^2$$
.

White (1980) shows that a consistent estimator of  $\operatorname{Var}_{T}[\mathbf{b}|\mathbf{X}]$  is obtained if  $e_{i}^{2}$  is used as an estimator of  $\sigma_{i}^{2}$ . Taking the square root, we get a *heteroscedasticity-consistent* (HC) standard errors.

(A3') was not specified. That is, the White estimator is *robust* to a potential misspecifications of heteroscedasticity in (A3').

The White estimator allows us to make inferences using the OLS estimator  $\mathbf{b}$  in situations where heteroscedasticity is suspected, but we do not know enough to identify its nature.

<u>Note</u>: The estimator is also called the *sandwich estimator* or the *White estimator* (also known as *Eiker-Huber-White estimator*).

• Remarks:

(1) Since there are many refinements of the White estimator, the White estimator is usually referred as HC0 (or just "HC"):

HC0 =  $(X'X)^{-1} X' \text{Diag}[e_i^2] X (X'X)^{-1}$ 

(2) In large samples, SEs, t-tests and F-tests are asymptotically valid.

(3) The OLS estimator remains inefficient. But inferences are asymptotically correct.

(4) The HC standard errors can be larger or smaller than the OLS ones. It can make a difference to the tests.

(5) It is used, along the Newey-West estimator, in almost all finance papers. Included in all the packaged software programs

(6) White SEs are easy to program:

# White SE in R

```
White f \leq function(y,X,b) {
T \le length(y); k \le length(b);
yhat \leq X\%*%b
                                               # fitted values
e <- y-yhat
                                               # residuals
hhat \leq t(X)*as.vector(t(e))
                                               # xi ei
                                               # Create kxk matrix to fill with X' Diag[e_i^2] X
G \leq matrix(0,k,k)
za <- hhat[,1:k]%*%t(hhat[,1:k])
                                               # X' diag[e_i] X
G \leq -G + za
                                               # X' diag[e_i] X
                                               # X'X
F \le t(X)\%*\%X
V \leq solve(F)\%\%\%G\%\%solve(F)
                                               # S_0
white se \leq- sqrt(diag(V))
                                               # White SE
ols se <- sqrt(diag(solve(F)*drop((t(e)%*%e))/(T-k)))
l_se = list(white_se,olse se)
return(1 se)
}
```

<u>R Note</u>: The library "*sandwich*" calculates White SEs. Remember to install it first and, then, call the library before you use it.

**Example 1:** We estimate t-values using OLS and White SE, for the 3 factor F-F model for **IBM** returns:

 $(r_{i=IBM,t} - r_f) = \beta_0 + \beta_1 (r_{m,t} - r_f) + \beta_2 SMB_t + \beta_3 HML_t + \varepsilon_t$ 

```
fit ibm ff3 <- lm(ibm x \sim Mkt RF + SMB + HML)
                                                      # OLS Regression with lm
b ibm <-fit ibm ff3$coefficients
                                                      # Extract OLS coefficients
SE OLS <- sqrt(diag(vcov(fit ibm ff3)))
                                                      # Extract OLS SE from fit ibm ff3
t OLS <- b ibm/SE OLS
                                                      # Calculate OLS t-values
>b ibm
             Mkt RF
                           SMB
(Intercept)
                                     HML
-0.005191356 0.910379487 -0.221385575 -0.139179020
> SE OLS
(Intercept)
            Mkt RF
                         SMB
                                   HML
0.002482305 0.056784474 0.084213761 0.084060299
> t OLS
(Intercept)
            Mkt RF
                         SMB
                                   HML
 -2.091345 16.032190 -2.628853 -1.655705
                                               \Rightarrow HML significant at 10% level
> library(sandwich)
White <- vcovHC(fit ibm ff3, type = "HC0")
                                                      # White Variance-covariance Matrix
SE White <- sqrt(diag(White))
                                                      # White SE HC0
t White <- b ibm/SE White
> SE White
(Intercept)
                                   HML
            Mkt RF
                         SMB
```

0.002505978 0.062481080 0.105645459 0.096087035 >t White (Intercept) Mkt RF SMB HML -2.071589 14.570482 -2.095552 -1.448468 White <- vcovHC(fit ibm ff3, type = "HC3") # White SE HC3 (refinement) SE White <- sqrt(diag(White))# White SE HC0 t White <- b ibm /SE White > SE White (Intercept) Mkt RF SMB HML 0.002533461 0.063818378 0.108316056 0.098800721 >t White (Intercept) Mkt RF SMB HML -2.049116 14.265162 -2.043885 **-1.408684**  $\Rightarrow$  similar results with HC3 refinement

<u>Conclusion</u>: White SEs make a difference in the test results. HML is not longer significant at the 10% level, once we adjust the SEs for heteroscedasticity.  $\P$ 

**Example 2:** We estimate Mexican interest rates  $(i_{MX})$  with a linear model including US interest rates, changes in exchange rates (MXN/USD), Mexican inflation and Mexican GDP growth, using quarterly data 1978:II – 2020:II (T=166):

 $i_{MX,t} = \beta_0 + \beta_1 i_{US,t} + \beta_2 e_t + \beta_3 mx_I t + \beta_4 mx_y t + \epsilon_t$ 

FMX\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/FX\_USA\_MX.csv", head=TRUE, sep=",")

 $\label{eq:second} us_i <- FMX_da US_int \\ mx_CPI <- FMX_da MX_CPI \\ mx_M1 <- FMX_da MX_M1 \\ mx_i <- FMX_da MX_int \\ mx_GDP <- FMX_da MX_GDP \\ S_mx <- FMX_da MXN_USD \\ T <- length(mx_CPI) \\ mx_I <- log(mx_CPI[-1]/mx_CPI[-T]) \\ mx_mg <- log(mx_GDP[-1]/mx_GDP[-T]) \\ mx_mg <- log(mx_M1[-1]/mx_M1[-T]) \\ e_mx <- log(S_mx[-1]/S_mx[-T]) \\ us_i_1 <- us_i[-1]/100 \\ mx_i_0 <- mx_i[-T]/100 \\ \end{tabular}$ 

 $\label{eq:summary_fit_i} \begin{array}{l} \textit{fit\_i} <- lm(mx\_i\_l \sim us\_i\_l + e\_mx + mx\_I + mx\_y) \\ > \textit{summary(fit\_i)} \end{array}$ 

Coefficients:

Estimate Std. Error t value Pr(>|t|)

# US short-term interest rates (i<sub>US</sub>)
# Mexican CPI
# Mexican Money Supply (M1)
# Mexican short-term int rates (i<sub>MX</sub>)
# Mexican GDP
# S<sub>t</sub> = exchange rates (MXN/USD)

# Mexican Inflation: Log changes in CPI
# Mexican growth: Log changes in GDP
# Money growth: Log changes in M1
# Log changes in St.
# Adjust sample size.

```
(Intercept) 0.04022 0.01506 2.671 0.00834 **
us i 1
            0.85886 0.31211 2.752 0.00661 **
            -0.01064 0.02130 -0.499 0.61812
e mx
mx I
            3.34581 0.19439 17.212 < 2e-16 ***
            -0.49851 0.73717 -0.676 0.49985
mx y
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
                                                # Extract OLS coefficients
b i <- fit i$coefficients
White <- vcovHC(fit i, type = "HC0")
                                                # White Variance-covariance Matrix HC0
SE White <- sqrt(diag(White))
                                                # White SE HC0
t White <- b i/SE White
> SE White
(Intercept)
             us i 1
                       e mx
                                 mx I
                                           mx y
0.009665759 0.480130221 0.026362820 0.523925226 1.217901733
> t White
(Intercept)
             us i 1
                       e mx
                                 mx I
                                           mx y
 4.1613603 1.7888018 -0.4035554 6.3860367 -0.4093221
                                                              \Rightarrow ius,t not significant at 5%.
White3 <- vcovHC(fit i, type = "HC3")
                                                   # Using popular refinement HC3
SE White3 <- sqrt(diag(White3))
                                                   # White SE HC3
t White <- b i/SE White3
>t White3
(Intercept)
             us i 1
                       e mx
                                 mx I
                                           mx y
 3.6338983 1.5589936 -0.2117600 5.4554986 -0.3519886
                                                              \Rightarrow ius,t not longer significant
                                                                  at 10% level.
```

<u>Conclusion</u>: Again, White SEs make a difference in the test results. U.S. interest rates are not a significant driver (& big drop in t-value!), once we adjust the SE for heteroscedasticity.

#### **Newey-West Estimator**

Now, we also have autocorrelation. We need to estimate

$$\mathbf{Q}^{*} = \frac{1}{T} \mathbf{X}' \mathbf{\Sigma} \mathbf{X} = (1/T) \sum_{i=1}^{T} \sum_{j=1}^{T} \sigma_{ij} \mathbf{x}_{i} \mathbf{x}_{j}'$$
  
=  $\frac{1}{T} \sum_{i=1}^{T} \{\sigma_{i1} \mathbf{x}_{i} \mathbf{x}_{1}' + \sigma_{i2} \mathbf{x}_{i} \mathbf{x}_{2}' + \sigma_{i3} \mathbf{x}_{i} \mathbf{x}_{3}' + \dots + \sigma_{iT} \mathbf{x}_{i} \mathbf{x}_{T}'\}$ 

Newey and West (1987) follow White (1980) to produce a HAC (*Heteroscedasticity and Autocorrelation Consistent*) estimator of  $\mathbf{Q}^*$ , also referred as *long-run variance* (LRV): Use  $e_i e_j$  to estimate  $\sigma_{ij}$ .

The natural estimator of  $\mathbf{Q}^*$  becomes:

$$\mathbf{S}_{\mathrm{T}} = \frac{1}{T} \sum_{i=1}^{T} \sum_{j=1}^{T} e_i e_j x_i x_j'$$

Or using time series notation, estimator of  $\mathbf{Q}^*$ :

$$\mathbf{S}_{\mathrm{T}} = \frac{1}{T} \sum_{t=1}^{T} \sum_{s=1}^{T} \mathbf{x}_{t} \mathbf{e}_{t} \mathbf{e}_{s} \mathbf{x}_{s}'$$

**Example:** Back to the simplest case, a regression with only one explanatory variable, but now with a heteroscedastic and autocorrelated error term. We estimate the variance of **b** with:

$$\operatorname{Var}_{\mathrm{T}}[\mathbf{b}|\mathbf{X}] = \left(\frac{1}{\sum_{i=1}^{T} (x_{i} - \bar{x})^{2}}\right)^{2} \left\{ \sum_{i=1}^{T} e_{i}^{2} (x_{i} - \bar{x})^{2} + \sum_{i=1}^{T} \sum_{j=i+1}^{T} e_{i} e_{j} (x_{i} - \bar{x}) (x_{j} - \bar{x}) \right\}.$$

We add the sum of the autocovariances of  $w_i (= x_i e_i)$  to the White estimator of **X'** $\Sigma$ **X**. If (auto-) covar( $w_i, w_j$ ) are mainly positive, the NW estimator will be bigger than the White estimator. A common situation in finance. ¶

There are some restrictions that need to be imposed:

- $\mathbf{Q}^*$  needs to be a pd matrix  $\Rightarrow$  use a quadratic form.
- The double sum cannot explode  $\Rightarrow$  use decaying weights to cut the sum short, after lag L, weights are zero.

• Two components for the NW HAC estimator:

(1) Start with Heteroscedasticity Component:

 $\mathbf{S}_0 = \frac{1}{T} \sum_{t=1}^T \boldsymbol{e}_t^2 \boldsymbol{x}_t \boldsymbol{x}_t' \qquad -\text{the White estimator.}$ 

(2) Add the Autocorrelation Component

 $\mathbf{S}_{\mathrm{T}} = \mathbf{S}_{0} + \frac{1}{T} \sum_{l=1}^{L} k(l) \sum_{t=l+1}^{T} (\mathbf{x}_{t-l} e_{t-l} e_{t} \mathbf{x}_{t}' + \mathbf{x}_{t} e_{t} e_{t-l} \mathbf{x}_{t-l}')$ 

where

$$k(\frac{j}{L(T)}) = \frac{L+1-|j|}{L+1}$$
 -decaying weights (*Bartlett kernel*)

*L* is the cut-off lag, which is a function of *T*. (More data, longer *L*).

The weights are linearly decaying, suppose L=30. Then, k(1) = 30/31 = 0.9677419 k(2) = 29/31 = 0.9354839k(3) = 28/31 = 0.9032258

**Example:** Back to the simplest case, a regression with only one explanatory variable, but with a heteroscedastic and autocorrelated error term. Suppose we set L = 12, then:

$$\operatorname{Var}_{\mathrm{T}}[\mathbf{b} \,|\, \mathbf{X}] = \left(\frac{1}{\sum_{i}^{T} (x_{i} - \bar{x})^{2}}\right)^{2} \left\{ \sum_{t=1}^{T} e_{t}^{2} (x_{t} - \bar{x})^{2} + \sum_{l=1}^{L=12} \left\{ \frac{13 - |j|}{13} \right\} \sum_{t=i+1}^{T} (x_{t} - \bar{x}) e_{t} e_{t-l} (x_{t-l} - \bar{x}) \right\}$$

To compute  $S_T$ , we only add 12 autocovariances of  $w_t (= x_t e_t)$  to the White estimator,  $S_0$ .

Technical detail: Under suitable conditions, as  $L, T \to \infty$ , and  $L/T \to 0, \mathbf{S}_T \to \mathbf{Q}^*$ . The condition  $L/T \to 0$  implies that when we have more data, we need to use a longer *L*.

Once we compute  $S_T$ , we compute the true Var[b|X]:

Est.  $\operatorname{Var}_{T}[\mathbf{b}|\mathbf{X}] = (1/T) (\mathbf{X}^{*}\mathbf{X}/T)^{-1} \mathbf{S}_{T} (\mathbf{X}^{*}\mathbf{X}/T)^{-1}$  –NW's HAC Var.

Asymptotic inferences can be based on OLS **b**, with *t*-tests and Wald tests using N(0,1) and  $\chi^2$  critical values, respectively.

There are many refinements of the NW estimators. Today, all HAC estimators are usually referred as NW estimators, regardless of the weights (*kernel*) used if they produce a positive (semi-) definite covariance matrix.

• All econometric packages (SAS, SPSS, Eviews, etc.) calculate NW SE.

<u>R Note</u>: You can use the library "*sandwich*," to calculate NW SEs: library(sandwich) > NeweyWest(x, lag = NULL, order.by = NULL, prewhite = TRUE, adjust = FALSE, diagnostics = FALSE, sandwich = TRUE, ar.method = "ols", data = list(), verbose = FALSE)

You need to install the package *sandwich* and then call the library(sandwich).

#### **Example**:

# fit the 3 factor Fama French Model for IBM returns: fit\_ibm\_ff3 <- lm(ibm\_x ~ Mkt\_RF + SMB + HML)</pre>

# NeweyWest computes the NW SEs. It requires lags=L & suppression of prewhitening NeweyWest(fit\_ibm\_ff3, lag = 4, prewhite = FALSE)

Note: It is usually found that the NW SEs are downward biased.

```
You can also program the NW SEs yourself. In R:

NW_f <- function(y, X, b, lag)

{

T <- length(y);

k <- length(b);

yhat <- X%*%b

e <- y - yhat

hhat <- t(X)*as.vector(t(e))

G <- matrix(0,k,k)

a <- 0

w <- numeric(T)

while (a <= lag) {

Ta <- T - a

ga <- matrix(0,k,k)

w[lag+1+a] <- (lag+1-a)/(lag+1)</li>
```

```
za \le hhat[,(a+1):T] \% \% t(hhat[,1:Ta])
 ga \le ga + za
 G \leq G + w[lag+1+a]*ga
a < -a+1
F \le t(X)\%*\%X
V \leq solve(F)\%\%\%G\%\%solve(F)
nw se <- sqrt(diag(V))
ols se \leq sqrt(diag(solve(F)*drop((t(e)%*%e))/(T-k)))
1 \text{ se} = \text{list}(\text{nw se,ols se})
return(1 se)
}
NW f(y,X,b,lag=4)
Example 1: We estimate the 3 factor F-F model for IBM returns with NW SE with 4 lags:
> t OLS
(Intercept)
             Mkt RF
                           SMB
                                     HML
 -2.091345 16.032190 -2.628853 -1.655705
                                                  \Rightarrow SMB significant at 1% level
NW <- NeweyWest(fit ibm ff3, lag = 4, prewhite = FALSE)
                                                                # NW Var Matrix with 4 lags
SE NW <- diag(sqrt(abs(NW)))
                                                                # NW SE with 4 lags
t NW <- b ibm/SE NW
> SE NW
(Intercept)
             Mkt RF
                           SMB
                                     HML
0.002527425 0.069918706 0.114355320 0.104112705
> t NW
(Intercept)
             Mkt RF
                           SMB
                                     HML
 -2.054010 13.020543 -1.935945 -1.336811
                                                  \Rightarrow SMB close to significant at 5% level
• If we add more lags in the NW function (lag = 8)
NW <- NeweyWest(fit ibm ff3, lag = 8, prewhite = FALSE)
SE NW <- diag(sqrt(abs(NW)))
t NW <- b ibm/SE NW
> t NW
(Intercept)
             Mkt RF
                           SMB
                                     HML
 -2.033648 12.779060 -1.895993 -1.312649
                                                 \Rightarrow not very different results.
```

<u>Conclusion</u>: Newey-West SEs make a difference in the test results. Now, SMB is not longer significant at the 5% level, though borderline, once we adjust the SEs not only for heteroscedasticity and autocorrelation.  $\P$ 

**Example 2: Mexican short-term interest rates** with NW SE with **4 lags & 8 lags**.. For comparison we reproduced the regression (with OLS t-varlues and the White t-values): fit\_i <- lm(mx\_i\_l ~ us\_i\_l + e\_mx + mx\_I + mx\_y) > summary(fit\_i)

#### Coefficients:

Estimate Std. Error t value Pr(>|t|)(Intercept) 0.04022 0.01506 2.671 0.00834 \*\* us i 1 0.85886 0.31211 2.752 0.00661 \*\* -0.01064 0.02130 -0.499 0.61812 e mx 3.34581 0.19439 17.212 < 2e-16 \*\*\* mx I -0.49851 0.73717 -0.676 0.49985 mx y Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 > t White3 (Intercept) us i 1 e mx mx I mx y 3.6338983 **1.5589936** -0.2117600 5.4554986 -0.3519886  $\Rightarrow$  i<sub>US,t</sub> not significant at 10%. NW <- NeweyWest(**fit i**, **lag** = **4**, prewhite = FALSE) SE NW <- diag(sqrt(abs(NW))) t NW <- **b i**/SE NW > SE NW (Intercept) us i 1 e mx mx I mx y 0.01107069 0.55810758 0.01472961 0.51675771 0.93960295 > t NW(Intercept) us i 1 e mx mx I mx y 3.6332593 **1.5388750** -0.7222770 6.4746121 -0.5305582  $\Rightarrow$  i<sub>US,t</sub> not significant at 10%. • If we add more lags in the text (lag = 8)NW <- NeweyWest(fit i, lag = 8, prewhite = FALSE) SE NW <- diag(sqrt(abs(NW))) t NW <- **b i**/SE NW > t NW(Intercept) us i 1 mx I mx y e mx 3.0174983 **1.4318654** -0.8279016 6.5897816 -0.5825521  $\Rightarrow$  similar results.

<u>Conclusion</u>: Newey-West SEs make a difference in the test results, but in this case, the results are not that different from the White SEs.  $\P$ 

• There are many estimators of  $Q^*$  based on a specific parametric model for  $\Sigma$ . Thus, they are not *robust* to misspecification of (A3'). This is the appeal of White & NW.

NW SEs are used almost universally in academia. However:

- NW SEs perform poorly in Monte Carlo simulations:

- NW SEs tend to be downward biased.

- The finite-sample performance of tests using NW SE is not well approximated by the asymptotic theory.

- Tests have size distortions.

Question: What happens if we know the specific form of (A3')? We can do much better than using OLS with NW SEs. In this case, we can do Generalized LS (GLS), a method that delivers the most efficient estimators.

## **Generalized Least Squares (GLS)**

GRM: Assumptions (A1), (A2), (A3') & (A4) hold. That is, (A1) DGP:  $\mathbf{y} = \mathbf{X} \ \beta + \boldsymbol{\varepsilon}$  is correctly specified. (A2)  $\mathrm{E}[\boldsymbol{\varepsilon}|\mathbf{X}] = 0$ (A3')  $\mathrm{Var}[\boldsymbol{\varepsilon}|\mathbf{X}] = \boldsymbol{\Sigma} = \sigma^2 \Omega$  ( $\Omega$  is symmetric  $\Rightarrow$  T'T =  $\Omega$ ) (A4) X has full column rank –i.e., rank(X) = k–, where T  $\geq k$ .

Question: What happens if we know the specific form of (A3')? We can use this information to gain efficiency.

When we know (A3'), we transform the y and X in such a way, that we can do again OLS with the transformed data.

To do this transformation, we exploit a property of symmetric matrices, like the variancecovariance matrix,  $\Omega$ :

 $\Omega \text{ is symmetric} \qquad \Rightarrow \text{ exists } T \ni T'T = \Omega \qquad \Rightarrow T'^{-1} \Omega T^{-1} = I$ 

<u>Note</u>: Think of **T** as  $\mathbf{\Omega}^{1/2}$ 

• We transform the linear model in (A1) using  $P = \Omega^{-1/2}$  (= T<sup>-1</sup>).

$$P = \Omega^{-1/2} \implies P'P = \Omega^{-1}$$

$$Py = PX\beta + P\varepsilon \text{ or}$$

$$y^* = X^*\beta + \varepsilon^*.$$

$$E[\varepsilon^*\varepsilon^{*'}|X^*] = E[P\varepsilon \varepsilon'P'|X^*] = P E[\varepsilon\varepsilon'|X] P' = \sigma^2 P \Omega P'$$

$$= \sigma^2 \Omega^{-1/2} \Omega \Omega^{-1/2} = \sigma^2 I_T \implies \text{back to (A3)}$$

The transformed model is homoscedastic: We have the CLM framework back. Now, we can use OLS!

$$\mathbf{b}_{\text{GLS}} = \mathbf{b}^* = (\mathbf{X}^* \mathbf{X}^*)^{-1} \mathbf{X}^* \mathbf{y}^*$$
  
=  $(\mathbf{X}' \mathbf{P}' \mathbf{P} \mathbf{X})^{-1} \mathbf{X}' \mathbf{P}' \mathbf{P} \mathbf{y}$   
=  $(\mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{X})^{-1} \mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{y}$  ( $\mathbf{P}' \mathbf{P} = \mathbf{\Omega}^{-1}$ )

Remarks:

- The transformed model is homoscedastic:

$$\operatorname{Var}[\boldsymbol{\varepsilon}^* | \boldsymbol{X}^*] = \operatorname{E}[\boldsymbol{\varepsilon}^* \boldsymbol{\varepsilon}^* | \boldsymbol{X}^*] = \sigma^2 \mathbf{I}_{\mathrm{T}}$$

– We have the CLM framework back: We do OLS with the transformed model, we call this OLS estimator, the GLS estimator:

 $b_{GLS} = b^* = (X^* X^*)^{-1} X^* Y^*$ =  $(X' \Omega^{-1} X)^{-1} X' \Omega^{-1} Y$ 

– Key assumption:  $\Omega$  is known, and, thus, P is also known; otherwise we cannot transformed the model.

Big Question: Is  $\Omega$  known?

GLS: Properties (1) Unbiased.  $\mathbf{b}_{GLS} = (\mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{X})^{-1} \mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{y} = (\mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{X})^{-1} \mathbf{X}' \mathbf{\Omega}^{-1} (\mathbf{X} \boldsymbol{\beta} + \boldsymbol{\epsilon})$   $= \boldsymbol{\beta} + = (\mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{X})^{-1} \mathbf{X}' \mathbf{\Omega}^{-1} \boldsymbol{\epsilon}$  $\Rightarrow E[\mathbf{b}_{GLS} | \mathbf{X} ] = \boldsymbol{\beta}$  (Yes, GLS is unbiased!)

(2) Efficient.

**b**<sub>GLS</sub> is BLUE. The "best" variance can be derived from  $Var[\mathbf{b}_{GLS}|\mathbf{X}] = \sigma^2 (\mathbf{X}^* \mathbf{X}^*)^{-1} = \sigma^2 (\mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{X})^{-1}$ 

Then, the usual OLS variance for **b** is biased and inefficient!

<u>Note I</u>:  $\mathbf{b}_{GLS} \neq \mathbf{b}$ .  $\mathbf{b}_{GLS}$  is BLUE by construction,  $\mathbf{b}$  is not.

<u>Note II</u>: Both unbiased and consistent. In practice, both estimators will be different, but not that different. If they are very different, something is not kosher.

• Steps for GLS:

Step 1. Find transformation matrix  $\mathbf{P} = \mathbf{\Omega}^{-1/2}$  (in the case of heteroscedasticity, **P** is a diagonal matrix).

Step 2. Transform the model:  $X^* = PX \& y^* = Py$ .

Step 3. Do GLS; that is, OLS with the transformed variables.

<u>Key step to do GLS</u>: Step 1, getting the transformation matrix:  $\mathbf{P} = \mathbf{\Omega}^{-1/2}$ .

<u>Technical detail</u>: If we relax the CLM assumptions (A2) and (A4), as we did in Lecture 7-a, we only have asymptotic properties for GLS:

- Consistency - "well behaved data."

– Asymptotic distribution under usual assumptions.

(easy for heteroscedasticity, complicated for autocorrelation.) – Wald tests and *F*-tests with usual asymptotic  $\chi^2$  distributions.

# (Weighted) GLS: Pure Heteroscedasticity

Step 1. Find the transformation matrix  $\mathbf{P} = \mathbf{\Omega}^{-1/2}$  for:

(A3') 
$$Var[\varepsilon] = \Sigma = \sigma^2 \Omega = \sigma^2 \begin{bmatrix} \omega_1 & 0 & \dots & 0 \\ 0 & \omega_2 & \dots & 0 \\ 0 & 0 & & 0 \\ 0 & 0 & \dots & \omega_T \end{bmatrix}$$

$$\mathbf{\Omega}^{-1/2} = \mathbf{P} = \begin{bmatrix} 1/\sqrt{\omega_1} & 0 & \dots & 0 \\ 0 & 1/\sqrt{\omega_2} & \dots & 0 \\ 0 & 0 & \dots & 0 \\ 0 & 0 & \dots & 1/\sqrt{\omega_T} \end{bmatrix}$$

Step 2. Now, transform y & X:

$$\mathbf{y}^{*} = \mathbf{P}\mathbf{y} = \begin{bmatrix} 1/\sqrt{\omega_{1}} & 0 & \dots & 0 \\ 0 & 1/\sqrt{\omega_{2}} & \dots & 0 \\ 0 & 0 & \dots & 0 \\ 0 & 0 & \dots & 1/\sqrt{\omega_{T}} \end{bmatrix} * \begin{bmatrix} y_{1} \\ y_{2} \\ \vdots \\ y_{T} \end{bmatrix} = \begin{bmatrix} y_{1}/\sqrt{\omega_{1}} \\ y_{2}/\sqrt{\omega_{2}} \\ \vdots \\ y_{T}/\sqrt{\omega_{T}} \end{bmatrix}$$

• Each observation of y,  $y_i$ , is divided by  $\sqrt{\omega_i}$ . Similar transformation occurs with **X**:

$$\mathbf{X}^{*} = \mathbf{P}\mathbf{X} = \begin{bmatrix} 1/\sqrt{\omega_{1}} & 0 & \dots & 0 \\ 0 & 1/\sqrt{\omega_{2}} & \dots & 0 \\ 0 & 0 & \dots & 0 \\ 0 & 0 & \dots & 1/\sqrt{\omega_{T}} \end{bmatrix}^{*} \begin{bmatrix} 1 & x_{21} & \dots & x_{k1} \\ 1 & x_{22} & \dots & x_{k2} \\ \vdots & \vdots & \dots & \vdots \\ 1 & x_{2T} & \dots & x_{kT} \end{bmatrix} = \begin{bmatrix} 1/\sqrt{\omega_{1}} & x_{21}/\sqrt{\omega_{1}} & \dots & x_{k1}/\sqrt{\omega_{1}} \\ 1/\sqrt{\omega_{2}} & x_{22}/\sqrt{\omega_{2}} & \dots & x_{k2}/\sqrt{\omega_{2}} \\ \vdots & \vdots & \dots & \vdots \\ 1/\sqrt{\omega_{T}} & x_{2T}/\sqrt{\omega_{T}} & \dots & x_{kT}/\sqrt{\omega_{T}} \end{bmatrix}$$

Step 3. Do GLS (OLS with the transformed variables):  $\mathbf{b}_{\text{GLS}} = \mathbf{b}^* = (\mathbf{X}^* \mathbf{X}^*)^{-1} \mathbf{X}^* \mathbf{y}^* = (\mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{X})^{-1} \mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{y}$ 

• In the case of heteroscedasticity, GLS is also called *Weighted Least Squares* (WLS): Think of  $1/\sqrt{\omega_i}$ . as weights. The GLS estimator is:

$$\mathbf{b}_{\text{GLS}} = (\mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{X})^{-1} (\mathbf{X}' \mathbf{\Omega}^{-1} \mathbf{y}) = \left( \sum_{i=1}^{T} \frac{1}{\omega_i} x_i x_i' \right)^{-1} \left( \sum_{i=1}^{T} \frac{1}{\omega_i} x_i y_i \right)$$

Observations with lower (bigger) variances -i.e., lower (bigger)  $\omega_i$ - are given higher (lower) weights in the sums: More precise observations, more weight!

The GLS variance is given by:

$$\hat{\sigma}_{GLS}^2 = \frac{\sum_{i=1}^{T} \left(\frac{y_i - x'_i \mathbf{b}_{GLS}}{\omega_i}\right)^2}{T - K}$$

**Example**: Last Lecture, we found that squared market returns  $(r_{m,t} - r_f)^2$  influence the heteroscedasticity in DIS returns. Suppose we assume: (A3')  $\sigma_t^2 = (r_{m,t} - r_f)^2$ .

Steps for GLS:

**Step 1**. Find transformation matrix, **P**, with *i*<sup>th</sup> diagonal element:  $1/\sqrt{\sigma_i^2}$ 

Step 2. Transform model: Each y<sub>i</sub> and x<sub>i</sub> is divided ("weighted") by  $\sigma_t = \operatorname{sqrt}[(r_{m,t} - r_f)^2].$ 

Step 3. Do GLS, that is, OLS with transformed variables.

 $T \leq - \text{length}(\text{dis } x)$ Mkt RF 2 <- Mkt RF^2 # (A3') y w <- dis x/sqrt(Mkt RF 2) # transformed  $y = y^*$  $x_{0} < -matrix(1,T,1)$ xx w <- cbind(x0, Mkt RF, SMB, HML)/sqrt(Mkt RF2) # transformed  $X = X^*$ fit dis wls  $\leq - lm(y \ w \sim xx \ w)$ # GLS > summary(fit dis wls) Call:  $lm(formula = y_w \sim xx_w)$ **Residuals:** 30 Max Min 1Q Median -59.399 -0.891 0.316 1.503 77.434 Coefficients: Estimate Std. Error t value Pr(>|t|)-0.006607 0.001586 -4.165 3.59e-05 \*\*\* XX\_W xx wMkt RF 1.588057 0.334771 4.744 2.66e-06 \*\*\*  $\Rightarrow$  OLS b: **1.26056** xx wSMB -0.200423 0.067498 -2.969 0.00311 \*\* ⇒ OLS b: -0.028993 xx wHML -0.042032 0.072821 -0.577 0.56404  $\Rightarrow$  OLS b: **0.174545** ---Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' '1 Residual standard error: 7.984 on 566 degrees of freedom Multiple R-squared: 0.09078, Adjusted R-squared: 0.08435 F-statistic: 14.13 on 4 and 566 DF, p-value: 5.366e-11

<u>Conclusion</u>: Quite different results, including a change in sign in HML, from positive & significant at the 10% level (OLS) to negative & not significant (GLS) and change in significance in SMB, from not significant (OLS) to very significant (GLS).

#### **GLS: First-order Autocorrelation Case**

We assume an AR(1) process for the  $\varepsilon_t$ :

 $\varepsilon_t = \rho \varepsilon_{t-1} + u_t$ ,  $u_t$ : non-autocorrelated error ~ D(0,  $\sigma_u^2$ )

We need to find the transformation matrix  $\mathbf{P} = \mathbf{\Omega}^{-1/2}$  for:

(A3') 
$$Var[\boldsymbol{\varepsilon}] = \boldsymbol{\Sigma} = \begin{bmatrix} \sigma^2 & \sigma_{12} & \cdots & \sigma_{1T} \\ \sigma_{21} & \sigma^2 & \cdots & \sigma_{2T} \\ \vdots & \vdots & \vdots & \vdots \\ \sigma_{T1} & \sigma_{T2} & \cdots & \sigma^2 \end{bmatrix},$$

which we will decompose into  $\Sigma = \sigma^2 \Omega$  (our goal: get  $\mathbf{P} = \Omega^{-1/2}$ )

Notation: We use  $\gamma_l$  to denote a (auto-) *covariance* between two observations separated by *l* periods. For example,

when l = 1:  $\gamma_1 = \sigma_{21} = \sigma_{32} = \dots = \sigma_{T(T-1)} = \operatorname{Cov}[\varepsilon_t, \varepsilon_{t-1}] = \operatorname{E}[\varepsilon_t \varepsilon_{t-1}]$ when l = 2:  $\gamma_2 = \sigma_{31} = \sigma_{42} = \dots = \sigma_{T(T-2)} = \operatorname{Cov}[\varepsilon_t, \varepsilon_{t-2}] = \operatorname{E}[\varepsilon_t \varepsilon_{t-2}]$ 

 $\gamma_l$  measures how two errors separated in time by *l periods* covary.

When l = 0, we get the variance. That is,  $\gamma_0 = \sigma_{\varepsilon}^2 = E[\varepsilon_t \ \varepsilon_t].$ 

• Then, we can write (A3') as:

$$\boldsymbol{\Sigma} = \begin{bmatrix} \sigma^2 & \sigma_{12} & \cdots & \sigma_{1T} \\ \sigma_{21} & \sigma^2 & \cdots & \sigma_{2T} \\ \vdots & \vdots & \vdots & \vdots \\ \sigma_{T1} & \sigma_{T2} & \cdots & \sigma^2 \end{bmatrix} = \begin{bmatrix} \gamma_0 & \gamma_1 & \cdots & \gamma_{T-1} \\ \gamma_1 & \gamma_0 & \cdots & \gamma_{T-2} \\ \vdots & \vdots & \vdots & \vdots \\ \gamma_{T-1} & \gamma_{T-2} & \cdots & \gamma_0 \end{bmatrix}.$$

<u>Remark</u>: Eventually decompose  $\Sigma = \sigma^2 \Omega$ , since we need  $\mathbf{P} = \Omega^{-1/2}$ )

• Steps for GLS:

Step 1. To find the transformation matrix **P**, we need to derive the implied (A3') based on the AR(1) process for  $\varepsilon_t$ :

(1) Find diagonal elements of  $\Omega$ :  $\gamma_0 \gamma_0 = \text{Var}[\varepsilon_t] = \sigma_{\varepsilon}^2$ Given the AR(1) process:

$$\varepsilon_t = \rho \ \varepsilon_{t-1} + u_t$$

We take the variance on both sides of the AR(1) equation (recall that  $u_t$  is uncorrelated with anything). Then,

 $\operatorname{Var}[\varepsilon_t] = \rho^2 \operatorname{Var}[\varepsilon_{t-1}] + \operatorname{Var}[u_t] \qquad (\operatorname{Var}[\varepsilon_t] = \operatorname{Var}[\varepsilon_{t-1}] = \sigma_{\varepsilon}^2)$ 

–we need to assume 
$$|\rho| < 1$$
.

Using the above  $\gamma_l$  notation, we have

$$\gamma_0 = \sigma_t^2 = \mathbb{E}[\varepsilon_t \ \varepsilon_t] = \operatorname{Var}[\varepsilon_t] = \sigma_\varepsilon^2 = \frac{\sigma_u^2}{(1-\rho^2)}$$

Now, we have all the diagonal elements of  $\Sigma$ :  $\gamma_0 = \frac{\sigma_u^2}{(1-\rho^2)}$ 

(2) Find off-diagonal elements of  $\Omega$ :  $\gamma_l = E[\varepsilon_i \ \varepsilon_j]$ , where l = i - j: Using the definition of covariance, we compute all  $\gamma_l$ 

$$\sigma_{ij} = \gamma_l = \operatorname{Cov}[\varepsilon_i, \varepsilon_j] = \operatorname{E}[\varepsilon_i \ \varepsilon_j], \qquad \text{where } l = i - j$$

$$\begin{aligned} \gamma_1 &= Cov[\varepsilon_t, \varepsilon_{t-1}] = E[(\rho \varepsilon_{t-1} + u_t) \varepsilon_{t-1}] \\ &= \rho E[\varepsilon_{t-1} \varepsilon_{t-1}] + E[u_t \varepsilon_{t-1}] \\ &= \rho Var[\varepsilon_{t-1}] + 0 \\ &= \rho \sigma_{\varepsilon}^2 \\ &= \rho \gamma_0 \end{aligned}$$

$$\begin{aligned} \gamma_2 &= Cov[\varepsilon_t, \varepsilon_{t-2}] = E[(\rho \varepsilon_{t-1} + u_t) \varepsilon_{t-2}] \\ &= \rho E[\varepsilon_{t-1} \varepsilon_{t-2}] + E[u_t \varepsilon_{t-2}] \\ &= \rho Cov[\varepsilon_t, \varepsilon_{t-1}] \\ &= \rho \gamma_1 \\ &= \rho^2 \gamma_0 \end{aligned}$$

$$\begin{split} \gamma_{3} &= Cov[\varepsilon_{t}, \varepsilon_{t-3}] = E[(\rho\varepsilon_{t-1} + u_{t}) \varepsilon_{t-3}] \\ &= \rho E[\varepsilon_{t-1} \varepsilon_{t-3}] + E[u_{t} \varepsilon_{t-3}] \\ &= \rho Cov[\varepsilon_{t}, \varepsilon_{t-2}] = \rho \gamma_{2} \\ &= \rho^{2} \gamma_{1} \\ &= \rho^{3} \gamma_{0} \\ \vdots \\ \gamma_{l} &= Cov[\varepsilon_{t}, \varepsilon_{t-l}] = \rho^{l} \gamma_{0} \end{split}$$

Then,

$$\boldsymbol{\Sigma} = \begin{bmatrix} \gamma_0 & \rho \gamma_0 & \cdots & \rho^{T-1} \gamma_0 \\ \rho \gamma_0 & \gamma_0 & \cdots & \rho^{T-2} \gamma_0 \\ \vdots & \vdots & \vdots & \vdots \\ \rho^{T-1} \gamma_0 & \rho^{T-2} \gamma_0 & \cdots & \gamma_0 \end{bmatrix}.$$

<u>Note</u>: We take  $\gamma_0$  out of the matrix. It becomes  $\sigma^2$  in the decomposition of  $\Sigma$  into  $\sigma^2 \Omega$ .

Recall we defined  $\gamma_0 = \sigma_{\varepsilon}^2 = \frac{\sigma_u^2}{(1-\rho^2)}$ . Then, we decompose  $\Sigma$  into  $\gamma_0 \ \Omega$ :

$$\Rightarrow \sigma_{\varepsilon}^2 = \frac{\sigma_u^2}{(1-\rho^2)}$$

(A3') 
$$\Sigma = \sigma^2 \Omega = \left(\frac{\sigma_u^2}{1 - \rho^2}\right) \begin{bmatrix} 1 & \rho & \rho^2 & \cdots & \rho^{T-1} \\ \rho & 1 & \rho & \cdots & \rho^{T-2} \\ \rho^2 & \rho & 1 & \cdots & \rho^{T-3} \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ \rho^{T-1} & \rho^{T-2} & \rho^{T-3} & \cdots & 1 \end{bmatrix}$$

• Now, we get the transformation matrix  $\mathbf{P} = \mathbf{\Omega}^{-1/2}$ : c

$$\mathbf{P} = \mathbf{\Omega}^{-1/2} = \begin{bmatrix} \sqrt{1 - \rho^2} & 0 & 0 & \dots & 0 \\ -\rho & 1 & 0 & \dots & 0 \\ 0 & -\rho & 1 & \dots & 0 \\ \dots & \dots & \dots & \dots & \dots \\ 0 & 0 & 0 & -\rho & 0 \end{bmatrix}$$

Step 2. With  $\mathbf{P} = \mathbf{\Omega}^{-1/2}$ , we transform the data ( $\mathbf{y} \& \mathbf{X}$ ) to do GLS:

$$\mathbf{P} \, \mathbf{y} = \begin{bmatrix} \sqrt{1 - \rho^2} & 0 & 0 & \dots & 0 \\ -\rho & 1 & 0 & \dots & 0 \\ 0 & -\rho & 1 & \dots & 0 \\ \dots & \dots & \dots & \dots & \dots \\ 0 & 0 & 0 & -\rho & 0 \end{bmatrix} * \begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ \vdots \\ y_T \end{bmatrix}$$

$$\mathbf{y}^{*} = \mathbf{P} \, \mathbf{y} = \begin{pmatrix} (\sqrt{1 - \rho^{2}}) y_{1} \\ y_{2} - \rho y_{1} \\ y_{3} - \rho y_{2} \\ \dots \\ y_{T} - \rho_{T-1} \end{pmatrix}$$
$$\mathbf{P} \, \mathbf{x}_{k} = \begin{bmatrix} \sqrt{1 - \rho^{2}} & 0 & 0 & \dots & 0 \\ -\rho & 1 & 0 & \dots & 0 \\ 0 & -\rho & 1 & \dots & 0 \\ \dots & \dots & \dots & \dots & \dots \\ 0 & 0 & 0 & -\rho & 0 \end{bmatrix} * \begin{bmatrix} x_{k1} \\ x_{k2} \\ x_{k3} \\ \vdots \\ x_{k7} \end{bmatrix}$$

$$\boldsymbol{x}_{k}^{*} = \mathbf{P} \, \boldsymbol{x}_{k} = \begin{pmatrix} \left(\sqrt{1-\rho^{2}}\right) x_{k1} \\ x_{k2} - \rho \, x_{k1} \\ x_{k3} - \rho \, x_{k2} \\ \dots \\ x_{T} - \rho \, x_{T-1} \end{pmatrix}$$

**Step 3**. Do GLS: OLS with transformed data. In (A3') we assume  $\rho$  known. In practice, we need to estimate it.

## **GLS:** The Autoregressive Transformation

With AR models, sometimes it is easier to transform the data by taking pseudo differences.

• For the AR(1) model, we multiply the DGP by  $\rho$  and subtract it from it. That is,

Now, we have the errors,  $u_t$ , which are uncorrelated. We can do OLS with *pseudo differences*.

Note:  $y_t^* = y_t - \rho y_{t-1} \& x_t^* = x_t - \rho x_{t-1}$  are pseudo differences.

# FGLS: Unknown Ω

The problem with GLS is that  $\Omega$  is unknown. For example, in the AR(1) case,  $\rho$  is unknown.

Solution: Estimate  $\Omega$ .  $\Rightarrow$  *Feasible GLS* (FGLS).

In general, there are two approaches for GLS

(1) Two-step, or *Feasible estimation*: - First, estimate  $\Omega$  first.

- Second, do GLS.

Similar logic to HAC procedures: We do not need to estimate  $\Omega$ , difficult with *T* observations. We estimate  $(1/T)\mathbf{X'}\Omega^{-1}\mathbf{X}$ .

-Nice asymptotic properties for FGLS estimator. Not longer BLUE

(2) ML estimation of  $\beta$ ,  $\sigma^2$ , and  $\Omega$  at the same time (joint estimation of all parameters). With some exceptions, rare in practice.

# FGLS: Specification of $\Omega$

•  $\Omega$  must be specified first.

•  $\Omega$  is generally specified (modeled) in terms of a few parameters. Thus,  $\Omega = \Omega(\theta)$  for some small parameter vector  $\theta$ . Then, we need to estimate  $\theta$ .

#### **Examples**:

(1)  $\operatorname{Var}[\varepsilon_i | \mathbf{X}] = \sigma^2 f(\mathbf{\gamma}' \mathbf{z}_i)$ . Variance a function of  $\mathbf{\gamma}$  and some variable  $\mathbf{z}_i$  (say, market volatility, firm size, country dummy, etc). In general, f is an exponential to make sure the variance is positive.

(2)  $\varepsilon_i$  with AR(1) process. We have already derived  $\sigma^2 \Omega$  as a function of  $\rho$ .

<u>Technical note</u>: To achieve full efficiency, we do not need an *efficient* estimate of the parameters in  $\Omega$ , only a *consistent* one.

Py.

#### **FGLS: Estimation – Steps**

Steps for FGLS:

**Step 1**. Estimate the model proposed in (A3'). Get  $\hat{\sigma}_i^2 \& \hat{\sigma}_{ij}$ .

Step 2. Find transformation matrix, **P**, using the estimated  $\hat{\sigma}_i^2 \& \hat{\sigma}_{ij}$ .

**3**. Using **P** from Step 2, transform model:  $X^* = PX$ 

**4**. Do FGLS, that is, OLS with  $X^* \& y^*$ .

**Example:** In the pure heteroscedasticity case (**P** is diagonal):

1. Estimate the model proposed in (A3'). Get  $\hat{\sigma}_i^2$ .

- **2**. Find transformation matrix, **P**, with  $i^{\text{th}}$  diagonal element:  $1/\hat{\sigma}_i$
- **3**. Transform model (each  $y_i$  and  $x_i$  is divided ("weighted") by  $\hat{\sigma}_i$ ):

$$y_i^* = y_i / \hat{\sigma}_i$$
$$x_{k,i}^* = x_{k,i} / \hat{\sigma}_i$$

4. Do FGLS, that is, OLS with transformed variables.

#### FGLS Estimation: Heteroscesdasticity

**Example**: Last lecture, we found that  $(r_{m,t} - r_f)^2 \& (SMB_t)^2$  are drivers of the heteroscedasticity in DIS returns: Suppose we assume: (A3')  $\sigma_t^2 = \gamma_0 + \gamma_1 (r_{m,t} - r_f)^2 + \gamma_3 (SMB_t)^2$ 

Steps for FGLS:
1. Use OLS squared residuals to estimate (A3'):
fit\_dis\_ff3 <- lm(dis\_x ~ Mkt\_RF + SMB + HML)</li>
e\_dis <- fit\_dis\_ff3\$residuals</li>
e\_dis2 <- e\_dis^2</li>
fit\_dis2 <- lm(e\_dis2 ~ Mkt\_RF2 + SMB2)</li>
summary(fit\_dis2)
var\_dis2 <- fit\_dis2\$fitted</li>
# Estimated variance vector, with elements \$\bar{\bar{o}}\_i^2\$.

**2**. Find transformation matrix, **P**, with  $i^{th}$  diagonal element:  $1/\hat{\sigma}_i$ w\_fgls <- sqrt(var\_dis2) #  $1/\hat{\sigma}_i$ 

**3.** Transform model: Each  $y_i$  and  $x_i$  is divided ("weighted") by  $\hat{\sigma}_i$ . $y_fw <- dis_x/w_fgls$ # transformed y $xx_fw <- cbind(x0, Mkt_RF, SMB, HML)/w_fgls$ # transformed X

4. Do GLS, that is, OLS with transformed variables. fit\_dis\_fgls <-  $lm(y_fw \sim xx_fw - 1)$  > summary(fit dis fgls) Coefficients: Estimate Std. Error t value Pr(>|t|)xx fw -0.003097 0.002696 -1.149 0.251 xx fwMkt RF 1.208067 0.073344 16.471 <2e-16 \*\*\* xx fwSMB  $-0.043761 \ 0.105280 \ -0.416 \ 0.678$ xx fwHML 0.125125 0.100853 1.241 0.215  $\Rightarrow$  not longer significant at 10%. ---Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 0.9998 on 566 degrees of freedom Multiple R-squared: 0.3413, Adjusted R-squared: 0.3366 F-statistic: 73.31 on 4 and 566 DF, p-value: < 2.2e-16

	b <sub>OLS</sub>	SE	b <sub>GLS</sub>	SE	b <sub>FGLS</sub>	SE
Intercept	0.00417	0.00279	-0.00661	0.00159	-0.00310	0.00270
Mkt_RF	1.26056	0.06380	1.58806	0.33477	1.20807	0.07334
SMB	-0.02899	0.09461	-0.20042	0.06750	-0.04376	0.10528
HML	0.17455	0.09444	-0.04203	0.07282	0.12513	0.10085

• Comparing OLS, GLS & FGLS results:

• Comments:

- The GLS estimates are quite different than OLS estimates (remember OLS is unbiased and consistent). Very likely the assumed functional form in (A3') was not a good one.

- The FGLS results are similar to the OLS, as expected, if model is OK. FGLS is likely a more precise estimator (HML is not longer significant at 10%).

## FGLS Estimation: AR(1) Case – Cochrane-Orcutt

In the AR(1) case, it is easier to estimate the model in *pseudo differences*:

$$y_t^* = \mathbf{X}_t^* \beta + u_t$$
  

$$y_t - \rho y_{t-1} = (\mathbf{X}_t - \rho \mathbf{X}_{t-1})^* \beta + \varepsilon_t - \rho \varepsilon_{t-1}$$
  

$$\Rightarrow y_t = \rho y_{t-1} + \mathbf{X}_t' \beta - \mathbf{X}_{t-1}' \rho \beta + u_t$$

We have a linear model, but it is nonlinear in parameters. OLS is not possible, but non-linear estimation is possible.

Note: We can do a regression:

 $y_t = \delta_1 y_{t-1} + \mathbf{X}_t' \boldsymbol{\delta}_2 - \mathbf{X}_{t-1}' \boldsymbol{\delta}_3 + u_t$ 

OLS will estimate  $\delta_1$ ,  $\delta_2$ , &  $\delta_3$ . To get  $\rho$  &  $\beta$ , we need a restriction:  $\delta_1 * \delta_2 = -\delta_3$ 

d rss  $\leq$  abs(rss 1 - rss)

i <- i+1

Before today's computer power, Cochrane–Orcutt's (1949) iterative procedure was an ingenious way to do this estimation.

• Steps for Cochrane-Orcutt: (1) Do OLS in (A1) model:  $y = X \beta + \varepsilon$ .  $\Rightarrow$  Get residuals, *e*, & RSS. (2) Estimate  $\rho$  with a regression of  $\boldsymbol{e}_t$  against  $\boldsymbol{e}_{t-1} \Rightarrow \text{Get } \hat{\rho}$  (the estimator of  $\rho$ ). (3) FGLS Step. Use  $\hat{\rho}$  to transform the model to get  $y^*$  and  $X^*$ . Do OLS with  $y^*$  and  $X^*$  $\Rightarrow$  get **b** to estimate  $\beta$ . Get residuals,  $e^* = y - X b$ , and new RSS. Go back to (1). (4) Iterate until convergence, usually achieved when the difference in RSS of two consecutive iterations is lower than some tolerance level, say .0001. Then, stop when  $RSS_i - RSS_{i-1} < .0001$ . **Example**: Cochrane-Orcutt in R # C.O. function requires Y, X (with constant), OLS b. c.o.proc <- function(Y,X,b 0,tol) {  $T \leq - length(Y)$ e <- Y - X%\*%b 0 # OLS residuals rss <- sum( $e^2$ ) # Initial RSS of model, RSS9 rss  $1 \leq rss$ # RSS 1 will be used to reset RSS after each iteration d rss = rss# initialize d rss: difference between RSSi & RSSi-1 e2 <- e[-1] # adjust sample size for et e3 <- e[-T]# adjust sample size for et-1 ols  $e0 \le lm(e2 \sim e3 - 1)$ # OLS to estimate rho rho <- ols e0(coeff[1]) # initial value for rho,  $\rho_0$ i<-1 while (d rss > tol) { # tolerance of do loop. Stop when diff in RSS < tol rss <- rss 1# RSS at iter (i-1) YY <- Y[2:T] - rho \* Y[1:(T-1)] # pseudo-diff Y XX <- X[2:T, ] - rho \* X[1:(T-1), ] # pseudo-diff X  $ols_yx \le lm(YY \sim XX - 1)$ # adjust if constant included in X b <- ols yx\$coef # updated OLS b at iteration i # b[1] <- b[1]/(1-rho) # If constant not pseudo-differenced remove tag # e1 <- Y - X%\*%b # updated residuals at iteration i e2 <- e1[-1]# adjust sample size for updated et e3 <- e1[-T] # adjust sample size for updated e\_t-1 (lagged et) # updated regression to value for rho at iteration i ols  $e1 <-lm(e2 \sim e3 - 1)$ rho <- ols e1 coeff[1] # updated value of rho at iteration i,  $\rho_i$ rss  $1 \le sum(e1^2)$ # updated value of RSS at iteration i, RSS<sub>i</sub>

# diff in RSS (RSS<sub>i</sub> - RSS<sub>i-1</sub>)

```
}
result <-list()
result$Cochrane-Orc.Proc <- summary(ols_yx)
result$rho.regression <- summary(ols_e1)
# result$Corrected.b_1 <- b[1]
result$Iterations < -i-1
return(result)
}</pre>
```

**Example**: In the model for **Mexican interest rates** (iMX), we suspect an AR(1) in the residuals:

```
i_{MX,t} = \beta_0 + \beta_1 i_{US,t} + \beta_2 e_t + \beta_3 mx_I t + \beta_4 mx_y t + \varepsilon_t\varepsilon_t = \rho \varepsilon_{t-1} + u_t
```

Cochrane-Orcutt estimation.
y <- mx\_i\_1</li>
T\_mx <- length(mx\_i\_1)</li>
xx\_i <- cbind(us\_i\_1, e\_mx, mx\_I, mx\_y)</li>
x0 <- matrix(1,T\_mx,1)</li>
X <- cbind(x0,xx\_i)</li>
fit\_i <- lm(mx\_i\_1 ~ us\_i\_1 + e\_mx + mx\_I + mx\_y)</li>
b\_i <-fit\_i\$coefficients</li>
> summary(fit\_i)

# X matrix

# extract coefficients from lm

	Estimate	Std. Error	t value $Pr(> t )$
(Intercept)	0.04022	0.01506	2.671 0.00834 **
us_i_1	0.85886	0.31211	2.752 0.00661 **
e_mx	-0.01064	0.02130	-0.499 0.61812
mx_I	3.34581	0.19439	<b>17.212</b> < 2e-16 ***
mx_y	-0.49851	0.73717	-0.676 0.49985

> c.o.proc(y,X,b,.0001)
\$Cochrane.Orcutt.Proc
Call:
lm(formula = YY ~ XX - 1)

Residuals: Min 1Q Median

-0.69251 -0.02118 -0.01099 0.00538 0.49403

3Q

Max

Coefficients:

Coefficients:

	Estimate	Std. Error	t value	Pr(> t )	
XX	0.16639	0.07289	2.283	0.0238 *	
XXus_i_1	1.23038	0.76520	1.608	0.1098	$\Rightarrow$ not longer significant at 5% level.

XXe mx -0.00535 0.01073 -0.499 0.6187 XXmx I 0.41608 0.27260 **1.526** 0.1289  $\Rightarrow$  not longer significant at 5% level. XXmx y -0.44990 0.53096 -0.847 0.3981 Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 0.09678 on 160 degrees of freedom Multiple R-squared: 0.1082, Adjusted R-squared: 0.08038 F-statistic: 3.884 on 5 and 160 DF, p-value: 0.002381 \$rho e3 0.8830857  $\Rightarrow$  very high autocorrelation. \$Corrected.b 1 XX 0.1663884  $\Rightarrow$  Constant corrected if X does not include a constant **\$Number.Iteractions** [1] 10  $\Rightarrow$  algorithm converged in 10 iterations.

<u>Conclusion</u>: Quite high the autocorrelation in the residuals ( $\rho = 0.8830857$ ), which has a big effect on the results. Once we account for the autocorrelation in the residuals, U.S. interest rates and Mexican Inflation rates are not longer significant drivers of Mexican interest rates. The model needs to be reformulated. ¶

# **GLS: General Remarks**

GLS is great (BLUE) if we know  $\Omega$ . Very rare situation.

It needs the specification of  $\Omega$  –i.e., the functional form of autocorrelation and heteroscedasticity.

If the specification is bad  $\Rightarrow$  estimates are biased.

In general, GLS is used for larger samples, because more parameters need to be estimated.

Feasible GLS is not BLUE (unlike GLS); but, it is consistent and asymptotically more efficient than OLS.

We use GLS for inference and/or efficiency. OLS is still unbiased and consistent.

OLS and GLS estimates will be different due to sampling error. But, if they are very different, then it is likely that some other CLM assumption is violated.

# Lecture 8 - Time Series

## **Time Series: Introduction**

A time series  $y_t$  is a (stochastic) process observed in sequence over time,  $t = 1, ..., T \implies \mathbf{Y}_t = \{y_1, y_2, y_3, ..., y_t\}.$ 

**Examples**: IBM monthly stock prices from **1973:January** to **2020:September** (plot below); or USD/GBP daily exchange rates from February 15, 1923 to March 19, 1938.



<u>R Note</u>: There are different ways to do the above plot in R: - Using plot.ts, creating a timeseries object in R: ts\_ibm <- ts(x\_ibm,start=c(1973,1),frequency=12) # the function ts creates a timeseries object, start = starting year,

plot.ts(ts\_ibm,xlab="Time",ylab="IBM price", main="Time Series: IBM Stock Price")

- Using R package ggplot2 library(ggplot2) ggplot(data= SFX\_da, aes(x = x\_date1, y = x\_ibm)) + geom\_line() + labs(x = "Date", y = "IBM price", title = "Time Series: IBM ", subtitle = "Monthly: 1973-2020")

## **Time Series: Introduction – Types**

Usually, time series models are separated into two categories:

– Univariate ( $y_t \in R$ , it is a scalar)

**Example:** We are interested in the behavior of IBM stock prices as function of its past.

 $\Rightarrow$  Primary model: Autoregressions (ARs).

- Multivariate ( $y_t \in R^m$ , it is a vector-valued)

**Example:** We are interested in the joint behavior of IBM stock and bond prices as function of their joint past.

 $\Rightarrow$  Primary model: Vector autoregressions (VARs). ¶

## **Time Series: Introduction – Dependence**

Given the sequential nature of  $y_t$ , we expect  $y_t$  and  $y_{t-1}$  to be dependent. This is the main feature of time series: dependence. It creates statistical problems.

In classical statistics, we usually assume we observe several *i.i.d.* realizations of  $y_t$ . We use  $\bar{y}$  to estimate the mean.

With several independent realizations we are able to sample over the entire probability space and obtain a "good" –i.e., consistent or close to the population mean– estimator of the mean.

But, if the samples are highly dependent, then it is likely that  $Y_t$  is concentrated over a small part of the probability space. Then, the sample mean will not converge to the mean as the sample size grows.

<u>Technical note</u>: With dependent observations, the classical results (based on LLN & CLT) are not to valid. New assumptions and tools are needed: stationarity, ergodicity, CLT for martingale difference sequences (MDS CLT).

Roughly speaking, *stationarity* requires constant moments for  $y_t$ ; ergodicity requires that the dependence is short-lived, eventually  $y_t$  has only a small influence on  $y_{t+k}$ , when k is relatively large.

*Ergodicity* describes a situation where the expectation of a random variable can be replaced by the time series expectation.

An *MDS* is a discrete-time martingale with mean zero. In particular, its increments,  $\varepsilon_t$ 's, are uncorrelated with any function of the available dataset at time t. To these  $\varepsilon_t$ 's we will apply a CLT.

The amount of dependence in  $Y_t$  determines the 'quality' of the estimator. There are several ways to measure the dependence. The most common measure: Covariance.

$$Cov(Y_t, Y_{t+k}) = E[(Y_t - \mu)(Y_{t+k} - \mu)]$$

<u>Note</u>: When  $\mu = 0$ , then  $Cov(Yt, Y_{t+k}) = E[Y_t Y_{t+k}]$ 

## **Time Series: Introduction – Forecasting**

In a time series model, we describe how  $y_t$  depends on past  $y_t$ 's. That is, the information set is  $I_t = \{y_{t-1}, y_{t-2}, y_{t-3}, ....\}$ 

The purpose of building a time series model: Forecasting.

We estimate time series models to forecast out-of-sample. For example, the *l-step ahead* forecast:

$$\hat{\mathbf{y}}_{\mathrm{T+l}} = \mathrm{E}_{\mathrm{t}}[y_{t+l}|I_{t-1}]$$

In the 1970s it was found that very simple time series models out-forecasted very sophisticated (big) economic models. This finding represented a big shock to the big multivariate models that were very popular then. It forced a re-evaluation of these big models.

## **Time Series: Introduction – White Noise**

In general, we assume the error term,  $\varepsilon_t$ , is uncorrelated with everything, with mean 0 and constant variance,  $\sigma^2$ . We call a process like this a *white noise* (WN) *process*.

We denote a WN process as

 $\varepsilon_t \sim WN(0, \sigma^2)$ 

The WN is a very simple example of a stochastic process. We think of a white noise process as the basic building block of all time series. It can be written as:

 $\mathbf{z}_t = \sigma \ u_t, \qquad u_t \sim i.i.d \ (0, 1) \qquad \qquad \Rightarrow \mathbf{z}_t \sim WN(0, \sigma^2)$ 

The  $z_t$ 's are random shocks, with no dependence over time, representing unpredictable events. It represents a model of news.

<u>Technical note</u>: There may be dependence in the higher order moments of  $\varepsilon_t$ . For example,  $E[\varepsilon_s^2 \ \varepsilon_t^2] = E[\varepsilon_s^2] * E[\varepsilon_t^2]$ . If we assume  $\varepsilon_t$  is *i.i.d.*, this dependence is excluded.

## **Time Series: Introduction – Conditionality**

We make a key distinction: *Conditional* vs *Unconditional* moments. In time series we model the *conditional mean* as a function of its past, for example in an AR(1) process, we have:

$$y_t = \alpha + \beta y_{t-1} + \varepsilon_t.$$

Then, the *conditional mean* forecast at time *t*, conditioning on information at time *I*<sub>t-1</sub>, is:

$$\mathbf{E}_{t}[y_{t}|I_{t-1}] = \mathbf{E}_{t}[y_{t}] = \alpha + \beta y_{t-1}$$

Notice that the *unconditional mean* is given by:

$$E[y_t] = \alpha + \beta E[y_{t-1}] = \frac{\alpha}{1-\beta} = \mu = \text{constant} \qquad (\beta \neq 1)$$

The conditional mean is time varying; the unconditional mean is not!

Key distinction: Conditional vs. Unconditional moments.

## **Time Series: Introduction – AR and MA models**

Two popular models for  $E_t[y_t|I_{t-1}]$ :

- Autoregressive (AR): An AR process models  $E_t[y_t|I_{t-1}]$  with lagged dependent variables:  $E_t[y_t|I_{t-1}] = f(y_{t-1}, y_{t-2}, y_{t-3}, ...)$ Example: AR(1) process,  $y_t = \alpha + \beta y_{t-1} + \varepsilon_t$ .

- Moving Average (MA): An MA process models  $E_t[y_t|I_{t-1}]$ :with lagged errors,  $\varepsilon_t$ :  $E_t[y_t|I_{t-1}] = f(\varepsilon_{t-1}, \varepsilon_{t-2}, \varepsilon_{t-3}, ...)$ Example: MA(1) process,  $y_t = \mu + \theta_1 \varepsilon_{t-1} + \varepsilon_t$ .

- There is a third model, ARMA, that combines lagged dependent variables and lagged errors.

• We want to select an appropriate time series model to forecast  $y_t$ . In this class, we will use linear model, with choices: AR(p), MA(q) or ARMA(p, q).

Steps for forecasting:
(1) Identify the appropriate model. That is, determine p, q.
(2) Estimate the model.
(3) Test the model.
(4) Forecast.

## **Time Series: CLM Revisited & New Assumptions**

With autocorrelated data, we get dependent observations. For example, with autocorrelated errors:

 $\varepsilon_t = \rho \varepsilon_{t-1} + u_t$ , with  $u_t \sim WN(0, \sigma^2)$ , the independence assumption (A2) is violated. The LLN and the CLT cannot be easily applied in this context. We need new tools.

We introduce the concepts of *stationarity* and *ergodicity*. The ergodic theorem will give us a counterpart to the LLN.

To get asymptotic distributions, we also need a CLT for dependent variables, using new technical concepts: mixing and stationarity. Or we can rely on a new CLT: The *martingale difference sequence CLT*.

• We will not cover these technical points in detail.

# **Time Series – Stationarity**

Consider the joint probability distribution of the collection of RVs:

$$F(y_{t_1}, y_{t_2}, \dots, y_{t_T}) = F(Y_{t_1} \le y_{t_1}, Y_{t_2} \le y_{t_2}, \dots, Y_{t_T} \le y_{t_T})$$

To do statistical analysis with dependent observations, we need some extra assumptions. We need some form of invariance on the structure of the time series.

If the distribution F is changing with every observation, estimation and inference become very difficult.

Stationarity is an invariant property: the statistical characteristics of the time series do not change over time.

There different definitions of stationarity, they differ in how strong is the invariance of the distribution over time.

We say that a process is stationary of

 $\begin{array}{ll} I^{st} \text{ order if } & F(y_{t_1}) = F(y_{t_{1+k}}) & \text{for any } t_1, k \\ 2^{nd} \text{ order if } & F(y_{t_1}, y_{t_2}) = F(y_{t_{1+k}}, y_{t_{2+k}}) & \text{for any } t_1, t_2, k \\ N^{th} \text{-order if } & F(y_{t_1}, \dots, y_{t_T}) = F(y_{t_{1+k}}, \dots, y_{t_{T+k}}) & \text{for any } t_1, \dots, t_T, k \end{array}$ 

 $N^{th}$ -order stationarity is a strong assumption (& difficult to verify in practice).  $2^{nd}$  order stationarity is weaker: only consider mean and covariance (easier to verify in practice). 2<sup>nd</sup> order stationarity is also called Weak stationarity or Covariance stationarity.

Moments describe a distribution. We calculate moments as usual:

 $E[Y_t] = \mu$  $Var(Y_t) = \sigma^2 = E[(Y_t - \mu)^2]$  $\operatorname{Cov}(Y_{t_1}, Y_{t_2}) = E[(Y_{t_1} - \mu)(Y_{t_2} - \mu)] = \gamma(t_1 - t_2)$ 

# **Time Series – Stationarity, Autocovariances & Autocorrelations**

 $Cov(Y_{t_1}, Y_{t_2}) = \gamma(t_1 - t_2)$  is called the *auto-covariance function*. It measures how y<sub>t</sub>, measured at time  $t_1$ , and  $y_t$ , measured at time  $t_2$ , covary.

<u>Notes</u>:  $\gamma(t_1 - t_2)$  is a function of  $k = t_1 - t_2$  $\gamma(0)$  is the variance.

The autocovariance function is symmetric. That is,

 $\gamma(t_1 - t_2) = Cov(Y_{t_1}, Y_{t_2}) = Cov(Y_{t_2}, Y_{t_1}) = \gamma(t_1 - t_2)$ 

Autocovariances are unit dependent. We will have different values if we calculate the autocovariance for IBM returns in % terms or in decimal terms.

<u>Remark</u>: The autocovariance measures the (linear) dependence between the  $y_t$ 's separated by k periods.

From the autocovariances, we derive the autocorrelations:

$$\operatorname{Corr}(Y_{t_1}, Y_{t_2}) = \rho(Y_{t_1}, Y_{t_2}) = \frac{\gamma(t_1 - t_2)}{\sigma_{t_1} \sigma_{t_2}} = \frac{\gamma(t_1 - t_2)}{\gamma(0)}$$

the last step takes assumes:  $\sigma_{t_1} = \sigma_{t_2} = \sqrt{\gamma(0)}$ 

 $\operatorname{Corr}(Y_{t_1}, Y_{t_2}) = \rho(Y_{t_1}, Y_{t_2})$  is called the *auto-correlation function* (ACF), –think of it as a function of  $k = t_1 - t_2$ . The ACF is also symmetric.

Unlike autocovoriances, autocorrelations are not unit dependent. It is easier to compare dependencies across different time series.

Stationarity requires all these moments to be independent of time. If the moments are time dependent, we say the series is non-stationary.

For strictly stationary process (constant moments), we need:

because  $F(y_{t_1}) = F(y_{t_{1+k}}) \Rightarrow \qquad \mu_{t_1} = \mu_{t_{1+k}} = \mu$  $\sigma_{t_1} = \sigma_{t_{1+k}} = \sigma$ 

Then,

$$F(y_{t_1}, y_{t_2}) = F(y_{t_{1+k}}, y_{t_{2+k}}) \Rightarrow Cov(y_{t_1}, y_{t_2}) = Cov(y_{t_{1+k}}, y_{t_{2+k}})$$
  

$$\Rightarrow \rho(t_1, t_2) = \rho(t_1 + k, t_2 + k)$$
  
Let  $t_1 = t - k \& t_2 = t$   

$$\Rightarrow \rho(t_1, t_2) = \rho(t - k, t) = \rho(t, t - k) = \rho(k) = \rho_k$$

The correlation between any two RVs depends on the time difference. Given the symmetry, we have  $\rho(k) = \rho(-k)$ .

**Example:** Informally, we check if in any two periods separated by k observations, we have similar means, variances and covariances. That is,

$$\mu_{t_1} = \mu_{t_{1+k}} = \mu$$
  

$$\sigma_{t_1} = \sigma_{t_{1+k}} = \sigma$$
  

$$Cov(y_{t_1}, y_{t_2}) = Cov(y_{t_{1+k}}, y_{t_{2+k}})$$

Time Series: IBM Monthly Returns Period: 1973 - 2023



## **Time Series – Weak Stationarity**

A Covariance stationary process (or 2nd -order weakly stationary) has:

- constant mean
- constant variance
- covariance function depends on time difference between RVs.

That is,  $Z_t$  is covariance stationary if:

 $E(Z_t) = \text{constant} = \mu$ Var(Z\_t) = constant =  $\sigma$ Cov(Z\_{t\_1}, Z\_{t\_2}) = E[(Z\_{t\_1} - \mu\_{t\_1})(Z\_{t\_2} - \mu\_{t\_2})] = \gamma(t\_1 - t\_2) = f(t\_1 - t\_2)

<u>Remark</u>: Covariance stationarity is only concerned with the covariance of a process, only the mean, variance and covariance are time-invariant. *N*<sup>th</sup>-order stationarity is stronger and assumes that the whole distribution is invariant over time.

**Example**: Stationary time series. Assume  $\varepsilon_t \sim WN(0, \sigma^2)$ .  $y_t = \phi \ y_{t-1} + \varepsilon_t$  (AR(1) process)

#### • Mean

Taking expectations on both sides --or applying the expectations operator (E[.])--:

 $E[y_t] = \phi \ E[y_{t-1}] + E[\varepsilon_t]$   $\mu = \phi \ \mu + 0$  $\Rightarrow E[y_t] = \mu = 0 \qquad (assuming \ \phi \neq 1)$ 

#### • Variance

Computing the variance --or applying the variance operator (Var[.])-- on both sides:

 $Var[y_t] = \gamma(0) = \phi^2 Var[y_{t-1}] + Var[\varepsilon_t]$  $\gamma(0) = \sigma^2/(1 - \phi^2)$ (assuming |  $\phi$  |< 1)

Autocovariances

$$\gamma(1) = \operatorname{Cov}[y_t, y_{t-1}] = \operatorname{E}[y_t y_{t-1}] = \operatorname{E}[(\phi \ y_{t-1} + \varepsilon_t) \ y_{t-1}] \\ = \operatorname{E}[\phi \ y_{t-1} \ y_{t-1}] + \operatorname{E}[\varepsilon_t \ y_{t-1}] \\ = \phi \ \operatorname{E}[y_{t-1}^2] + 0 \\ = \phi \ \operatorname{Var}[y_{t-1}] \\ = \phi \ \gamma(0) \\ = \phi \ [\sigma^2/(1 - \phi^2)] \\ \gamma(2) = \operatorname{Cov}[y_t, y_{t-2}] = \operatorname{E}[y_t \ y_{t-2}]] = \operatorname{E}[(\phi \ y_{t-1} + \varepsilon_t) \ y_{t-2}] \\ = \operatorname{E}[\phi \ y_{t-1} \ y_{t-2}] + \operatorname{E}[\varepsilon_t \ y_{t-2}] \\ = \phi \ \operatorname{E}[y_{t-1} \ y_{t-2}] + \operatorname{E}[\varepsilon_t \ y_{t-2}] \\ = \phi \ \operatorname{E}[y_{t-1} \ y_{t-2}] + 0 \\ = \phi \ \operatorname{E}[y_{t-1} \ y_{t-2}] \\ = \phi \ \operatorname{Cov}[y_{t-1}, y_{t-2}] \\ = \phi \ \operatorname{Cov}[y_{t-1}, y_{t-2}] \\ = \phi \ \phi \ \gamma(0) \\ = \phi^2 \ \gamma(0) \\ = \phi^2 \ [\sigma^2/(1 - \phi^2)] \\ \end{cases}$$

• • •

$$\gamma(k) = \operatorname{Cov}[y_t, y_{t-k}] = \phi^k \gamma(0)$$

 $\Rightarrow$  If  $|\phi| < 1$ , the process is covariance stationary: mean, variance and covariance are constant.

<u>Remark</u>: To establish stationarity, we need to impose conditions on the AR parameters. (Conditions on parameters are not needed for MA processes; MA processes are always stationary.)

Note: From the autocovariance function, we can derive the auto-correlation function:

$$\rho(k) = \frac{\gamma(k)}{\gamma(0)} = \frac{\phi^k \gamma(0)}{\gamma(0)} = \phi^k$$

If  $|\phi| < 1$ , both the autocovariance function & ACF show exponential decay.

**Example**: Non-stationary time series. Assume  $\varepsilon_t \sim WN(0, \sigma^2)$ .  $y_t = \mu + y_{t-1} + \varepsilon_t$  (Random Walk with drift process)

Doing backward substitution:

$$y_{t} = \mu + (\mu + y_{t-2} + \varepsilon_{t-1}) + \varepsilon_{t}$$
  
= 2\*\mu + y\_{t-2} + \varepsilon\_{t} + \varepsilon\_{t-1} + \

• Mean & Variance

$$E[y_t] = \mu t + y_0$$
  
Var[y\_t] =  $\gamma(0) = \sum_{j=0}^{t-1} \sigma^2 = \sigma^2 t$ 

 $\Rightarrow$  the process is non-stationary; that is, moments are time dependent.

## **Stationary Series – Examples**

**Examples**: Assume  $\varepsilon_t \sim WN(0, \sigma^2)$ .  $y_t = 0.08 + \varepsilon_t + 0.4 \varepsilon_{t-1}$  -MA(1) process  $y_t = 0.13 y_{t-1} + \varepsilon_t$  -AR(1) process



Changes in the JPY/USD exchange rate  $(e_{f,t})$  is an example of a stationary series.

# **Non-Stationary Series – Examples**

**Examples**: Assume  $\varepsilon_t \sim WN(0, \sigma^2)$ .  $y_t = \mu \ t + \varphi_1 y_{t-1} + \varphi_2 y_{t-2} + \varepsilon_t$  $y_t = \mu + y_{t-1} + \varepsilon_t$ 

AR(2) with deterministic trendRandom Walk with drift



The level of the JPY/USD exchange rate  $(S_t)$  is an example of a non-stationary series.

## **Time Series – Stationarity: Remarks**

The main characteristic of time series is that observations are dependent.

To analyze time series, however, we need to assume that some features of the series are not changing. If we have non-stationary series (say, mean or variance are changing with each observation), it is not possible to make inferences.

Stationarity is an invariant property: the statistical characteristics of the time series do not vary over time.

If IBM is weak stationary, then, the returns of IBM may change month to month or year to year, but the average return and the variance in two equal-length time intervals will be more or less the same.

In the long run, say 100-200 years, the stationarity assumption may not be realistic. After all, technological change has affected the return of IBM over the long run. But, in the short-run, stationarity seems likely to hold.

In general, time series analysis is done under the stationarity assumption.

# **Time Series – Ergodicity**

We want to estimate the mean of the process  $\{Z_t\}$ ,  $\mu(Z_t)$ . But, we need to distinguishing between ensemble average (with *m* observations) and *time average* (with *T* observations):

- Ensemble Average:  $\overline{\overline{z}} = \frac{\sum_{i=1}^{m} Z_i}{m}$  Time Series Average:  $\overline{\overline{z}} = \frac{\sum_{t=1}^{T} Z_t}{T}$

Question: Which estimator is the most appropriate?

A: Ensemble Average. But, it is impossible to calculate. We only observe one Zt, with dependent observations.

Question: Under which circumstances we can use the time average (with only one realization of  $\{Z_t\}$ ? Is the time average an unbiased and consistent estimator of the mean? The Ergodic Theorem gives us the answer.

• Intuition behind Ergodicity:

We go to a casino to play a game with 20% return, but on average, one gambler out of 100 goes bankrupt. If 100 gamblers play the game, there is a 99% chance of winning and getting a 20% return. This is the *ensemble scenario*. Suppose that gambler 35 is the one that goes bankrupt. Gambler 36 is not affected by the bankruptcy of gamble 35.

Suppose now that instead of 100 gamblers you play the game 100 times. This is the *time series* scenario. You win 20% every day until day 35 when you go bankrupt. There is no day 36 for you (dependence at work!).

Result: The probability of success from the group (ensemble scenario) does not apply to one person (time series scenario).

Ergodicity describes a situation where the ensemble scenario outcome applies to the time series scenario.

• With dependent observation, we cannot use the LLN used before. The ergodicity theorem plays the role of the LLN with dependent observations.

The formal definition of ergodicity is complex and is seldom used in time series analysis. One consequence of ergodicity is the ergodic theorem, which is extremely useful in time series.

It states that if  $Z_t \mbox{ is an ergodic stochastic process then }$ 

$$\frac{1}{T}\sum_{t=1}^{T}g(Z_t) \xrightarrow{a.s} \mathbb{E}[g(Z)]$$

for any function g(.). And, for any time shift k

$$\frac{1}{T} \sum_{t=1} g(Z_{t_1+k}, Z_{t_2+k}, \dots, Z_{t_{\tau}+k}) \xrightarrow{a.s} \mathbb{E}[g(Z_{t_1}, Z_{t_2}, \dots, Z_{t_{\tau}}))]$$

where a.s. means *almost sure convergence*, a strong form of convergence.

Definition: A covariance-stationary process is *ergodic* for the **mean** if  $\bar{z} \xrightarrow{p} E[Z_t] = \mu$ 

This result needs the variance of  $\overline{z}$  to collapse to 0. It can be shown that the var $[\overline{z}]$  can be written as a function of the autocorrelations,  $\rho_k$ :

$$\begin{aligned} \operatorname{var}[\overline{z}] &= \operatorname{var}[(z_1 + z_2 + \dots + z_T)/T] \\ &= \{\operatorname{var}[z_1] + \operatorname{var}[z_2] + \dots + \operatorname{var}[z_T] \\ &+ 2\operatorname{cov}[z_1, z_2] + 2\operatorname{cov}[z_1, z_3] + \dots + 2\operatorname{cov}[z_1, z_T] \\ &+ 2\operatorname{cov}[z_2, z_3] + 2\operatorname{cov}[z_2, z_4] + \dots + 2\operatorname{cov}[z_2, z_T] \\ &+ 2\operatorname{cov}[z_3, z_4] + 2\operatorname{cov}[z_3, z_5] + \dots + 2\operatorname{cov}[z_3, z_T] \\ &\dots \\ &+ 2\operatorname{cov}[z_{T-1}, z_T]\}/T^2 \\ &= \frac{\gamma_0}{T^2} \{T\rho_0 + 2(T-1)\rho_1 + 2(T-2)\rho_2 + \dots + 2\rho_{T-1}\} \\ &= \frac{\gamma_0}{T^2} \{T\rho_0 + 2\sum_{k=1}^{T-1}(T-k)\rho_k\} \end{aligned}$$

Recalling that  $\rho_k = \rho_{-k}$ , then

$$\operatorname{var}[\overline{z}] = \frac{\gamma_0}{T^2} \sum_{k=-(T-1)}^{T-1} (T - |k|) \rho_k$$
$$= \frac{\gamma_0}{T} \sum_{k=-(T-1)}^{T-1} \left(1 - \frac{|k|}{T}\right) \rho_k$$

**Theorem**: A sufficient condition for ergodicity for the mean is that the autocorrelations  $\rho_k$  between two observations, say  $(y_{t_i}, y_{t_j})$ ,  $\rho(t_i, t_j) = \rho_{t_i - t_j}$ , go to zero as  $t_i \& t_j$  grow further apart.

Condition for ergodicity:  $\rho_k \to 0$ , as  $k \to \infty$ 

# **Time Series – Lag Operator**

Define the operator *L* as  $L^{k} z_{t} = z_{t-k}$ .

It is usually called *Lag operator*. But it can produce lagged or forward variables (for negative values of *k*). For example:

$$L^{-3} z_t = z_{t+3}.$$

Also note that if *c* is a constant  $\Rightarrow L c = c$ .

Sometimes the notation for L when working as a lag operator is B (backshift operator), and when working as a forward operator is F.

Important application: Differencing

$$\Delta z_t = (1 - L) z_t = z_t - z_{t-1}.$$
  
$$\Delta^2 z_t = (1 - L)^2 z_t = z_t - 2z_{t-1} + z_{t-2}.$$

#### **Time Series – Useful Result: Geometric Series**

The function  $f(x) = (1 - x)^{-1}$  can be written as an infinite geometric series (use a Maclaurin series around c=0):

$$f(x) = \frac{1}{1-x} = 1 + x + x^2 + x^3 + x^4 + \ldots = \sum_{n=0}^{\infty} x^n$$

If we multiply f(x) by a constant, *a*:

$$\sum_{n=0}^{\infty} ax^n = \frac{a}{1-x} \to \sum_{n=1}^{\infty} ax^n = a\left(\frac{1}{1-x} - 1\right)$$

**Example:** In Finance we have many applications of the above results.

- A stock price, P, equals the discounted some of all futures dividends. Assume dividends are constant, d, and the discount rate is r. Then:

$$P = \sum_{t=1}^{\infty} \frac{d}{(1+r)^t} = d\left(\frac{1}{1-\frac{1}{1+r}} - 1\right) = d\left(\frac{1}{\frac{1+r-1}{1+r}} - 1\right) = \frac{d}{r}$$
  
where  $x = \frac{1}{1+r}$ .

We will use this geometric series result when, under certain conditions, we invert a lag polynomial (say,  $\theta(L)$ ) to convert an AR (MA) process into an infinite MA (AR) process.

**Example:** Suppose we have an MA(1) process:

 $y_t = \mu + \theta_1 \varepsilon_{t-1} + \varepsilon_t = \mu + \theta(L) \varepsilon_t,$ 

with

$$\theta(L) = (1 + \theta_1 L)$$
 ( $\theta(L)$ : lag polynomial)

Recall,

$$f(x) = \frac{1}{1-x} = 1 + x + x^2 + x^3 + x^4 + \ldots = \sum_{n=0}^{\infty} x^n$$

Let  $x = -\theta_1 L$ . Then,

$$\theta(L)^{-1} = \frac{1}{1 - (-\theta_1 L)} = 1 + (-\theta_1 L) + (-\theta_1 L)^2 + (-\theta_1 L)^3 + (-\theta_1 L)^4 + \dots$$
$$= \sum_{n=0}^{\infty} (-\theta_1 L)^n = 1 - \theta_1 L + \theta_1^2 L^2 - \theta_1^3 L^3 + \theta_1^4 L^4 + \dots$$
That is, we get an AR( $\infty$ ), by multiplying both sides by  $\theta(L)^{-1}$ :

 $\theta(L)^{-1} y_t = \theta(L)^{-1} \mu + \varepsilon_t = \mu^* + \varepsilon_t$ 

Or

$$\theta(L)^{-1} y_t = y_t - \theta_1 y_{t-1} + \theta_1^2 y_{t-2} - \theta_1^3 y_{t-3} + \theta_1^4 y_{t-4} + \dots = \mu^* + \varepsilon_t$$

Solving for yt:

$$y_{t} = \mu^{*} + \theta_{1} y_{t-1} - \theta_{1}^{2} y_{t-2} + \theta_{1}^{3} y_{t-3} - \theta_{1}^{4} y_{t-4} + \dots + \varepsilon_{t} \cdot \P$$

#### **Moving Average Process**

An MA process models  $E[y_t|I_{t-1}]$  with lagged error terms. An MA(q) model involves q lags. We keep the white noise assumption for  $\varepsilon_t$ :  $\varepsilon_t \sim WN(0, \sigma^2)$ 

**Example**: A linear MA(q) model:

 $y_t = \mu + \theta_1 \varepsilon_{t-1} + \theta_2 \varepsilon_{t-2} + \dots + \theta_q \varepsilon_{t-q} + \varepsilon_t = \mu + \theta(L) \varepsilon_t,$ where  $\theta(L) = 1 + \theta_1 L + \theta_2 L^2 + \theta_3 L^3 + \dots + \theta_q L^q.$ 

In time series, the constant does not affect the properties of AR and MA process. It is usually removed (think of the data analyze as demeaned). Thus, in this situation we say "without loss of generalization", we assume  $\mu=0$ .

#### **Moving Average Process – Stationarity**

To check if an MA(q) process is stationary, we check the moments (assume  $\mu = 0$ ).

$$y_t = \varepsilon_t + \theta_1 \varepsilon_{t-1} + \theta_2 \varepsilon_{t-2} + \dots + \theta_q \varepsilon_{t-q}$$
  
$$y_{t-1} = \varepsilon_{t-1} + \theta_1 \varepsilon_{t-2} + \theta_2 \varepsilon_{t-3} + \dots + \theta_{q-1} \varepsilon_{t-q} + \theta_q \varepsilon_{t-(q+1)}$$

• Mean

$$\mathbf{E}[y_t] = \mathbf{E}[\varepsilon_t] + \theta_1 \mathbf{E}[\varepsilon_{t-1}] + \theta_2 \mathbf{E}[\varepsilon_{t-2}] + \dots + \theta_q \mathbf{E}[\varepsilon_{t-q}] = 0$$

• Variance

$$\operatorname{Var}[y_t] = \operatorname{Var}[\varepsilon_t] + \theta_1^2 \operatorname{Var}[\varepsilon_{t-1}] + \theta_2^2 \operatorname{Var}[\varepsilon_{t-2}] + \dots + \theta_q^2 \operatorname{Var}[\varepsilon_{t-q}]$$
  
=  $(1 + \theta_1^2 + \theta_2^2 + \dots + \theta_q^2) \sigma^2$ .

To get a positive variance, we require  $(1 + \theta_1^2 + \theta_2^2 + ... + \theta_q^2) > 0$ .

Autocovariances

$$y_{t} = \varepsilon_{t} + \theta_{1} \varepsilon_{t-1} + \theta_{2} \varepsilon_{t-2} + \theta_{3} \varepsilon_{t-3} + \dots + \theta_{q} \varepsilon_{t-q}$$

$$y_{t-1} = \varepsilon_{t-1} + \theta_{1} \varepsilon_{t-2} + \theta_{2} \varepsilon_{t-3} + \dots + \theta_{q} \varepsilon_{t-q} + \theta_{q} \varepsilon_{t-(q+1)}$$

$$\gamma(1) = \operatorname{Cov}[y_{t}, y_{t-1}] = \operatorname{E}[y_{t} y_{t-1}]$$

$$= \operatorname{E}[(\varepsilon_{t} + \theta_{1} \varepsilon_{t-1} + \theta_{2} \varepsilon_{t-2} + \dots + \theta_{q} \varepsilon_{t-q})^{*}(\varepsilon_{t-1} + \theta_{1} \varepsilon_{t-2} + \theta_{2} \varepsilon_{t-3} + \dots + \theta_{q} \varepsilon_{t-(q+1)})]$$

$$= \operatorname{E}[\varepsilon_{t} \varepsilon_{t-1}] + \theta_{1} \operatorname{E}[\varepsilon_{t} \varepsilon_{t-2}] + \theta_{2} \operatorname{E}[\varepsilon_{t} \varepsilon_{t-3}] + \dots$$

$$\begin{split} &+ \theta_1 \operatorname{E}[\varepsilon_{t-1}\varepsilon_{t-1}] + \theta_1^2 \operatorname{E}[\varepsilon_{t-1}\varepsilon_{t-2}] + \theta_1 \theta_2 \operatorname{E}[\varepsilon_{t-1}\varepsilon_{t-3}] + \dots \\ &+ \theta_2 \operatorname{E}[\varepsilon_{t-2}\varepsilon_{t-1}] + \theta_2 \theta_1 \operatorname{E}[\varepsilon_{t-2}\varepsilon_{t-2}] + \theta_2 \theta_1 \operatorname{E}[\varepsilon_{t-2}\varepsilon_{t-3}] + \dots \\ &\dots \\ &+ \theta_q \operatorname{E}[\varepsilon_{t-q}\varepsilon_{t-1}] + \theta_q \theta_1 \operatorname{E}[\varepsilon_{t-q}\varepsilon_{t-2}] + \dots + \theta_q \theta_{q-1} \operatorname{E}[\varepsilon_{t-q}\varepsilon_{t-q}] \\ &+ \theta_q^2 \operatorname{E}[\varepsilon_{t-q}\varepsilon_{t-(q+1)}] \\ &= \theta_1 \sigma^2 + \theta_2 \theta_1 \sigma^2 + \theta_3 \theta_2 \sigma^2 + \dots + \theta_q \theta_{q-1} \sigma^2 + 0 \\ &= \sigma^2 \sum_{j=1}^q \theta_j \theta_{j-1} \qquad (\text{where } \theta_0 = 1) \end{split}$$

We can also derive  $\gamma(1)$  without computing the expectation of the cross products of errors. It is easier to look at the sum of  $E[y_t \ \epsilon_{t-j}]$ 's:

$$\begin{aligned} \gamma(1) &= \mathbb{E}[y_t \ y_{t-1}] \\ &= \mathbb{E}[y_t \ * (\varepsilon_{t-1} + \ \theta_1 \ \varepsilon_{t-2} + \ \theta_2 \ \varepsilon_{t-3} + \dots + \ \theta_{q-1} \ \varepsilon_{t-q} + \ \theta_q \ \varepsilon_{t-(q+1)})] \\ &= \mathbb{E}[y_t \ \varepsilon_{t-1}] + \theta_1 \ \mathbb{E}[y_t \ \varepsilon_{t-2}] + \theta_2 \ \mathbb{E}[y_t \ \varepsilon_{t-3}] + \dots + \theta_{q-1} \ \mathbb{E}[y_t \ \varepsilon_{t-q}] + \theta_q \ \mathbb{E}[y_t \ \varepsilon_{t-q}] \\ &= \theta_1 \ \sigma^2 + \theta_2 \ \theta_1 \ \sigma^2 + \theta_3 \ \theta_2 \ \sigma^2 + \dots + \theta_q \ \theta_{q-1} \ \sigma^2 + 0 \\ &= \sigma^2 \sum_{j=1}^q \theta_j \ \theta_{j-1} \end{aligned}$$
(where  $\theta_0 = 1$ )

We continue with the derivations of the  $\gamma(k)$  function. It is easier to derive it by rewriting yt & **Y**t-2:

$$y_{t} = \varepsilon_{t} + \theta_{1} \varepsilon_{t-1} + \theta_{2} \varepsilon_{t-2} + \theta_{3} \varepsilon_{t-3} + \dots + \theta_{q} \varepsilon_{t-q}$$

$$y_{t-2} = \varepsilon_{t-2} + \theta_{1} \varepsilon_{t-3} + \theta_{2} \varepsilon_{t-4} + \theta_{3} \varepsilon_{t-5} + \dots + \theta_{q} \varepsilon_{t-(q+2)}$$

$$\gamma(2) = \operatorname{Cov}[y_{t}, y_{t-2}] = \operatorname{E}[y_{t} \ y_{t-2}]$$

$$= \operatorname{E}[y_{t} \ \varepsilon_{t-2}] + \theta_{1} \operatorname{E}[y_{t} \ \varepsilon_{t-3}] + \theta_{2} \operatorname{E}[y_{t} \ \varepsilon_{t-4}] + \dots + \theta_{q} \operatorname{E}[y_{t} \ \varepsilon_{t-q-2}]$$

$$= \theta_{2} \ \sigma^{2} + \theta_{3} \ \theta_{1} \ \sigma^{2} + \theta_{4} \ \theta_{2} \ \sigma^{2} + \dots + \theta_{q} \ \theta_{q-2} \ \sigma^{2} + 0$$

$$= \sigma^{2} \sum_{j=2}^{q} \theta_{j} \theta_{j-2} \qquad (\text{where } \theta_{0} = 1)$$

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$$\gamma(q) = \operatorname{E}[y_t \ y_{t-q}]] =$$

$$= \operatorname{E}[\varepsilon_t \ y_{t-q}] + \theta_1 \operatorname{E}[\varepsilon_{t-1} \ y_{t-q}] + \theta_2 \operatorname{E}[\varepsilon_{t-2} \ y_{t-q}] + \dots + \theta_q \operatorname{E}[\varepsilon_{t-q} \ y_{t-q}]$$

$$= \theta_q \ \sigma^2$$

$$= \sigma^2 \sum_{j=q}^q \theta_j \theta_{j-q} \qquad (\text{where } \theta_0 = 1)$$

In general, for the *k* autocovariance:

$$egin{aligned} &\gamma(k) = \sigma^2 \sum_{j=k}^q heta_j heta_{j-k} & ext{for } |k| \leq q \ &\gamma(k) = 0 & ext{for } |k| > q \end{aligned}$$

<u>Remark</u>: After lag q, the autocovariances are 0.

#### • Autocorrelations

From the autocovariances, we define the autocorrelations, by dividing the autocorrelations by *γ*(0):

$$\rho(q) = \frac{\sigma^2 \sum_{j=q}^{q} \theta_j \theta_{j\cdot q}}{(1+\theta_1^2+\theta_2^2+\ldots+\theta_q^2)\sigma^2} = \frac{\sum_{j=q}^{q} \theta_j \theta_{j\cdot q}}{(1+\theta_1^2+\theta_2^2+\ldots+\theta_q^2)} \quad (\theta_0=1)$$

In general, for the *k* autocorrelation function (ACF):

$$\rho(k) = \frac{\sum_{j=q}^{q} \theta_{j} \theta_{j-q}}{(1+\theta_{1}^{2}+\theta_{2}^{2}+\ldots+\theta_{q}^{2})} \quad \text{for } |k| \le q$$
  

$$\rho(k) = 0 \quad \text{for } |k| > q$$

<u>Remark</u>: After lag q, the ACF is 0.

• It can be shown that for  $\varepsilon_t$  with same distribution (say, normal) the ACF are non-unique. For example, for the MA(1) processes:

 $y_t = \varepsilon_t + 0.5 \varepsilon_{t-1} \qquad \Rightarrow \rho(1) = \theta_1 / (1 + \theta_1^2) = 0.4$  $y_t = \varepsilon_t + 2 \varepsilon_{t-1} \qquad \Rightarrow \rho(1) = \theta_1 / (1 + \theta_1^2) = 0.4$ 

• It is easy to verify that the sums  $\sum_{j=k}^{q} \theta_{j} \theta_{j-k}$  are finite. Then, mean, variance and covariance are constant.

 $\Rightarrow$  MA(q) is always stationary.

#### **Moving Average Process – Invertibility**

As mentioned above, it is possible that different time-series processes produce the same ACF.

**Example**: Two MA(1) produce the same  $\gamma(k)$ :  $y_t = \varepsilon_t + 0.2 \ \varepsilon_{t-1}, \qquad \varepsilon_t \sim i.i.d. \ N(0, 25)$  $z_t = v_t + 5 \ v_{t-1}, \qquad v_t \sim i.i.d. \ N(0, 1)$ 

We only observe the time series,  $y_t$  or  $z_t$ , and not the noise,  $\varepsilon_t$  or  $v_t$ , thus, we cannot distinguish between the models. Thus, we select only one of them. Which one? We select the model with an AR( $\infty$ ) representation.

Assuming  $\theta(L) \neq 0$ , we can invert  $\theta(L)$ . Then, by inverting  $\theta(L)$ , an MA(q) process generates an AR process:

$$y_t = \mu + \theta(L) \varepsilon_t \implies \theta(L)^{-1} y_t = \Pi(L) y_t = \mu^* + \varepsilon_t$$

Then, we have an infinite sum polynomial on  $\theta L$ . (Recall the geometric series result.) That is, we convert an MA(q) into an AR( $\infty$ ):

$$\sum_{j=0}^{\infty} \pi_j(L) y_t = \mu^* + \varepsilon_t$$

We need to make sure that  $\Pi(L) = \theta(L)^{-1}$  is defined: We require  $\theta(L) \neq 0$ . When this condition is met, we can write  $\varepsilon_t$  as a causal function of  $y_t$ . We say the MA is *invertible*. For this to hold, we require:

$$\sum_{j=0}^{\infty} |\pi_j(L)| < \infty$$

<u>Technical note</u>: An invertible MA(q) is typically required to have roots of the lag polynomial equation  $\theta(z) = 0$  greater than one in absolute value ("*outside the unit circle*"). In the MA(1) case, we require  $|\theta_1| < 1$ .

In the previous example, we select the model with  $\theta_1 = 0.2$ .

## Moving Average Process – MA(1)

**Example**:  $y_t = \theta_1 \varepsilon_{t-1} + \varepsilon_t = \mu + \theta(L) \varepsilon_t$ , with  $\theta(L) = (1 + \theta_1 L)$ 

#### • Moments

$$E[y_{t}] = 0$$
  

$$Var[y_{t}] = \gamma(0) = \sigma^{2} + \theta_{1}^{2} \sigma^{2} = \sigma^{2} (1 + \theta_{1}^{2})$$
  

$$Cov[y_{t}, y_{t-1}] = \gamma(1) = E[y_{t} y_{t-1}]$$
  

$$= E[(\theta_{1} \varepsilon_{t-1} + \varepsilon_{t}) * (\theta_{1} \varepsilon_{t-2} + \varepsilon_{t-1})]$$
  

$$= \theta_{1} \sigma^{2}$$
  

$$Cov[y_{t}, y_{t-2}] = \gamma(2) E[y_{t} y_{t-2}]$$
  

$$= E[(\theta_{1} \varepsilon_{t-1} + \varepsilon_{t}) * (\theta_{1} \varepsilon_{t-3} + \varepsilon_{t-2})]$$
  

$$= 0$$
  

$$\vdots$$
  

$$\gamma(k) = E[y_{t} y_{t-k}] = E[(\theta_{1} \varepsilon_{t-1} + \varepsilon_{t})) * (\theta_{1} \varepsilon_{t-(k+1)} + \varepsilon_{t-k})])] = 0 \quad (\text{for } k > 1)$$

That is, for |k| > 1,  $\gamma(k) = 0$ .

To get the ACF, we divide the autocovariances by  $\gamma(0)$ . Then, the autocorrelation function (ACF):

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$$\rho(1) = \gamma(1)/\gamma(0) = \theta_1 \sigma^2/\sigma^2 (1 + \theta_1^2) = \theta_1 / (1 + \theta_1^2)$$
  
$$\rho(k) = \gamma(k)/\gamma(0) = 0 \qquad (\text{for } k > 1)$$

<u>Remark</u>: The autocovariance function is zero after lag 1. Similarly, the ACF is also zero after lag 1.

Note that  $|\rho(1)| \leq 0.5$ .

When	$\theta_1 = 0.5$	$\Rightarrow \rho(1) = 0.4.$	
	$\theta_1 = -0.9$	$\Rightarrow \rho(1) = -0.497238.$	
	$\theta_1 = -2$	$\Rightarrow \rho(1) = -0.4.$	(same $\rho(1)$ for $\theta_1 \& 1/\theta_1$ )

#### Moving Average Process – MA(1): Simulations

We simulate and plot three MA(1) processes, with standard normal  $\varepsilon_t$ -i.e.,  $\sigma = 1$ :

$$y_t = \varepsilon_t + 0.5 \varepsilon_{t-1}$$
  

$$y_t = \varepsilon_t - 0.9 \varepsilon_{t-1}$$
  

$$y_t = \varepsilon_t - 2 \varepsilon_{t-1}$$

<u>R Note</u>: We use the *arima.sim* function to simulate the behavior of different ARIMA models (or ARMA, by setting the orden of Integration (I) equal to 0). Below we plot differen MA(1) process. On the first panel, we use the script below to plot  $y_t = \varepsilon_t + 0.5 \varepsilon_{t-1}$  with 100 simulations. The other panels are straightforward to get.

> plot(arima.sim(list(order=c(0,0,1), ma=0.5), n=100), ylab="ACF", main=(expression(MA(1)~~~theta==+.5)))



<u>Note</u>: The process  $\theta_1 > 0$  is smoother than the ones with  $\theta_1 < 0$ .

Below, we compute and plot the ACF for the 3 simulated process, using the acf R function.

1)  $y_t = \varepsilon_t + 0.5 \varepsilon_{t-1}$ sim ma1 5 <- arima.sim(list(order=c(0,0,1), ma=0.5), n=200) acf ma1 5 <- acf(sim ma1 5, main=(expression(MA(1)~~~theta==+.5)))  $> acf_ma1 5$ Autocorrelations of series 'sim mal 6', by lag 1.000 0.438 0.069 0.014 0.103 0.173 0.107 0.015 -0.080 -0.054 0.011 -0.006 0.041 0.000 -0.094 -0.147 -0.129 -0.082 -0.150 -0.196 -0.251 -0.235 -0.021 0.110



2)  $y_t = \varepsilon_t - 0.9 \varepsilon_{t-1}$ sim ma1 9 <- arima.sim(list(order=c(0,0,1), ma=-0.9), n=200) acf ma1 9 <- acf(sim ma1 5, main=(expression(MA(1)~~~theta==-0.9))) > acf mal 9Autocorrelations of series 'sim\_ma1\_9', by lag 1.000 -0.584 0.093 0.061 -0.132 0.147 -0.181 0.122 -0.013 -0.023 0.014 -0.012 0.092 -0.199 0.193 -0.155 0.143 -0.107 0.014 0.174 -0.244 0.196 -0.154 0.105



3)  $y_t = \varepsilon_t - 2 \varepsilon_{t-1}$ sim ma1 2 <- arima.sim(list(order=c(0,0,1), ma=-2), n=200) acf mal 2 <- acf(sim mal 2, main=(expression(MA(1)~~~theta==-2))) > acf mal 2Autocorrelations of series 'sim\_ma1\_2', by lag 1.000 -0.524 0.150 -0.064 0.006 -0.014 0.022 -0.070 0.068 -0.015 -0.002 0.054 -0.121 0.055 -0.029 0.026 -0.054 0.121 -0.156 0.106 -0.009 0.037 -0.080 0.104



• Invertibility: If  $|\theta_1| < 1$ , we can write  $(1 + \theta_1 L)^{-1} y_t + \mu^* = \varepsilon_t$ Or expanding

$$(1 - \theta_1 L + \theta_1^2 L^2 + \dots + \theta_1^j L^j + \dots) y_t + \mu^* = \mu^* + \sum_{i=1}^{\infty} \pi_i(L) y_t = \varepsilon_t$$

That is,  $\pi_i = (-\theta_1)^i$ .

The simulated process with  $\theta_1 = -2$  is non-invertible, the infinite sum of  $\pi_i$  would explode. We would select the MA(1) with  $\theta_1 = -.5$ .

#### Moving Average Process – MA(2)

**Example**:  $y_t = \mu + \theta_2 \epsilon_{t-2} + \theta_1 \epsilon_{t-1} + \epsilon_t = \mu + \theta(L) \epsilon_t$ , with  $\theta(L) = (1 + \theta_1 L + \theta_2 L^2).$ 

• Moments

$$E (Y_{t}) = \mu$$

$$\gamma_{k} = \begin{cases} \sigma^{2} (1 + \theta_{1}^{2} + \theta_{2}^{2}), & k = 0 \\ - \theta_{1} \sigma^{2} (1 - \theta_{2}), & |k| = 1 \\ - \theta_{2} \sigma^{2}, & |k| = 2 \\ 0, & |k| > 2 \end{cases}$$

<u>Remark</u>: The autocovariance function is zero after lag 2. Similarly, the ACF is also zero after lag 2.

– Invertibility: The roots of  $\lambda^2 - \theta_1 \lambda - \theta_2 = 0$  all lie inside the <u>unit circle</u>. It can be shown the invertibility condition for an MA(2) process is:

 $\theta_1 + \theta_2 < 1$   $\theta_1 - \theta_2 < 1$  $-1 < \theta_2 < 1.$ 

### **Moving Average Process – Estimation**

MA processes are more complicated to estimate. In particular, there are nonlinearities. Consider an MA(1):

$$y_t = \theta \ \varepsilon_{t-1} + \varepsilon_t, \qquad \varepsilon_t \sim WN.$$

We cannot do OLS, since we do not observe  $\varepsilon_{t-1}$ . But, based on the ACF, we can estimate  $\theta$ .

• The auto-correlation is  $\rho_1 = \theta/(1+\theta^2)$ . Then, we can use the method of moments (MM), which sets the theoretical moments equal to the sample moments and, then, solve for parameters of interest. In the MA(1) case, the theoretical formula for  $\rho_1$  is:

$$\rho_1 = \theta/(1 + \theta^2).$$

Then, we use the estimated  $\rho_1$ ,  $r_1$ , to estimate  $\theta$ :

$$r_1 = \frac{\widehat{\theta}}{(1+\widehat{\theta}^2)} \quad \Rightarrow \qquad \widehat{\theta} = \frac{1\pm \sqrt{1-4r_1^2}}{2r_1}$$

A nonlinear solution and difficult to solve.

• Alternatively, if  $|\theta| < 1$ , we can invert the MA(1) process. Then, based on the AR representation, we can try finding  $a \in (-1; 1)$ ,

$$\varepsilon_t(a) = y_t + ay_{t-1} + a^2 y_{t-2} + \dots$$

and look (numerically) for the least-square estimator

 $\hat{\boldsymbol{\theta}} = \arg \min_{\boldsymbol{\theta}} \{ \mathbf{S}(\boldsymbol{y}; \boldsymbol{\theta}) = \sum_{t=1}^{T} \varepsilon_t(\boldsymbol{a})^2 \}$ where  $\boldsymbol{a}^t = \theta_1^t$ .

## **The Wold Decomposition**

**Theorem -** Wold (1938).

Any covariance stationary  $\{y_t\}$  has infinite order, moving-average representation:

$$y_t = S_t + \kappa_t$$

where

 $\kappa_t$  is a deterministic term –i.e., completely predictable. For example,  $\kappa_t = \mu$  or a linear combination of past (known) values of  $\kappa_t$ .

$$\begin{split} S_t &= \sum_{j=0}^{\infty} \psi_j \, \varepsilon_{t-j} & (=\psi(L)\varepsilon_t, \text{ with } \psi(L) = \text{ infinite lag polynomial}) \\ \sum_{j=0}^{\infty} \psi_j^2 &< \infty & (\text{assumption for the stability of polynomial, "square summability"}) \\ \psi_j \text{ only depend on } j & (\text{weights of innovations are not time dependent}) \\ \psi_0 &= 1 & (\text{a convenient assumption}) \\ \varepsilon_t &\sim \text{WN}(0, \sigma^2) & (\varepsilon_t \text{ independent and uncorrelated with } S_t) \end{split}$$

Thus,  $y_t$  is a linear combination of innovations over time plus a deterministic part.

• A stationary process can be decomposed into a sum of two parts, one represented as an  $MA(\infty)$  and the other a deterministic "trend."

**Example**: Let  $x_t = y_t - \kappa_t$ . ( $x_t = MA(\infty)$  part) Then, check moments:

$$\begin{split} E[x_{t}] &= E[y_{t} - \kappa_{t}] = \sum_{j=0}^{\infty} \psi_{j} E[\varepsilon_{t-j}] = 0. \\ E[x_{t}^{2}] &= \sum_{j=0}^{\infty} \psi_{j}^{2} E[\varepsilon_{t-j}^{2}] = \sigma^{2} \sum_{j=0}^{\infty} \psi_{j}^{2} < \infty. \\ E[x_{t}, x_{t-j}] &= E[(\varepsilon_{t} + \psi_{1}\varepsilon_{t-1} + \psi_{2}\varepsilon_{t-2} + \dots)(\varepsilon_{t-j} + \psi_{1}\varepsilon_{t-j-1} + \psi_{2}\varepsilon_{t-j-2} + \dots) \\ &= \sigma^{2} (\psi_{j} + \psi_{1}\psi_{j+1} + \psi_{2}\psi_{j+2} + \dots) = \sigma^{2} \sum_{k=0}^{\infty} \psi_{k}\psi_{k+2} \end{split}$$

 $\{x_t\}$  is a covariance stationary process.

Remark: This old theorem is the backbone of time series analysis. We will approximate the Wold infinite lag polynomial  $\psi(L)$  with a ratio of two finite lag polynomials. This approximation is the basis of ARMA modeling.

#### **Autoregressive (AR) Process**

We model the conditional expectation of  $y_t$ ,  $E[y_t|I_{t-1}]$ , as a function of its past history. We assume  $\varepsilon_t$  follows a WN(0,  $\sigma^2$ ).

The most common models are AR models. An AR(1) model involves a single lag, while an AR(p) model involves p lags. Then, the AR(p) process is given by:

 $y_t = \mu + \phi_1 y_{t-1} + \phi_2 y_{t-2} + ... + \phi_p y_{t-p} + \varepsilon_t, \qquad \varepsilon_t \sim WN.$ 

Using the lag operator, we write the AR(p) process:

 $\phi(L) y_t = \varepsilon_t$ 

with

$$\boldsymbol{\phi}(L) = 1 - \boldsymbol{\phi}_1 L - \boldsymbol{\phi}_2 L^2 - \dots - \boldsymbol{\phi}_p L^p$$

• We can look at an AR(*p*) process:

 $y_t = \mu + \phi_1 \ y_{t-1} + \phi_2 \ y_{t-2} + \dots + \phi_p \ y_{t-p} + \varepsilon_t,$ 

as a stochastic (linear) difference equation (SDE). With difference equations we try to get a solution –i.e., given some initial conditions/history, we know the value of  $y_t$  for any t– and, then, we study its characteristics (stability, long-run value, etc.).

The solution to a difference equation can be written as a sum of two solutions:

1) Homogeneous equation (the part that only depends on the  $y_t$ 's):

 $y_t = \phi_1 y_{t-1} + \phi_2 y_{t-2} + \dots + \phi_p y_{t-p}$  (set  $\mu + \varepsilon_t = 0$ ) 2) A particular solution to the equation.

Once we get a solution, we study its stability. We want a stable one.

• We get a solution to the simple case, the AR(1) process.

$$y_t = \mu + \phi_1 y_{t-1} + \varepsilon_t, \qquad \varepsilon_t \sim WN.$$

We use the backward substitution method:

$$\begin{split} y_t &= \mu + \phi_1 \left( \mu + \phi_1 \, y_{t-2} + \varepsilon_{t-1} \right) + \varepsilon_t \\ &= \mu \left( 1 + \phi_1 \right) + \phi_1^2 \, y_{t-2} + \varepsilon_t + \phi_1 \, \varepsilon_{t-1} \\ &= \mu \left( 1 + \phi_1 \right) + \phi_1^2 \left( \mu + \phi_1 \, y_{t-3} + \varepsilon_{t-2} \right) + \varepsilon_t + \phi_1 \, \varepsilon_{t-1} \\ &= \mu \left( 1 + \phi_1 + \phi_1^2 \right) + \phi_1^3 \, y_{t-3} + \varepsilon_t + \phi_1 \, \varepsilon_{t-1} + \phi_1^2 \, \varepsilon_{t-2} \\ & \vdots \\ &\Rightarrow y_t &= \mu \left( 1 + \phi_1 + \phi_1^2 + \dots + \phi_1^{t-1} \right) + \sum_{j=0}^{t-1} \phi_j^j \varepsilon_{t-j} + \phi_1^t \, y_0 \end{split}$$

The solution is a function of *t*, the whole sequence  $\varepsilon_t$ ,  $\varepsilon_{t-1}$ , ...,  $\varepsilon_1$  and the initial condition  $y_0$ . The effect of  $y_0$  "dies out" if  $|\phi_1| < 1$ .

• The stability of the solution is crucial. With a stable solution,  $Y_t$  does not explode. This is good: We need well defined moments.

It turns out that the stability of the equation depends on the solution to the homogenous equation. In the AR(1) case:

$$y_t = \phi_1 y_{t-1}$$
  
with solution  $y_t = \phi_1^t y_0$ 

If  $|\phi_1| < 1$ ,  $y_t$  never explodes, as  $t \to \infty$ . In this case, in the solution to the AR(1) process, the effect of  $y_0$  "dies out" as  $t \to \infty$ .

• We can analyze the stability from the point of view of the *roots*, *z*, of the *characteristic* equation of the AR(*p*) process,  $\phi(L) = 0$ .

For the AR(1) process

 $\phi(z) = 1 - \phi_1 z = 0 \qquad \Rightarrow |z| = 1/|\phi_1| > 1.$ 

That is, the AR(1) process is stable if the root of  $\phi(z)$  is greater than one (also said as "*the roots lie outside the unit circle*").

This result generalizes to AR(p) process. For example, for the AR(3) process

 $y_t = \phi_1 y_{t-1} + \phi_2 y_{t-2} + \phi_3 y_{t-3} + \varepsilon_t,$  $\phi(z) = 1 - \phi_1 z - \phi_2 z^2 - \phi_3 z^3 \implies \text{the roots, } z_1, z_2 \& z_3, \text{ should lie outside the unit circle.}$ 

For an AR(*p*), we need the roots of  $\phi(z)$  to be outside the unit circle.

For the AR(2),  $y_t = \phi_1 y_{t-1} + \phi_2 y_{t-2}$ .

We need the roots of  $\phi(z)$  to be outside the unit circle.

The characteristic polynomial of the AR(2) can be written as:

 $\phi(z) = 1 - (\lambda_1 + \lambda_2) z - \lambda_1 \lambda_2 z^2 = (1 - \lambda_1 z) (1 - \lambda_2 z) = 0$ where  $\phi_1 = \lambda_1 + \lambda_2$ , and  $\phi_2 = \lambda_1 \lambda_2$ . ( $\lambda_1 \& \lambda_2$  are *eigenvalues* or *characteristic roots*.) If  $|\lambda_1| < 1$ , and  $|\lambda_2| < 1$ , the roots lie *outside the unit root*  $\Rightarrow$  stationary.

Then, some implications for  $\phi_1 \& \phi_2$ :

$$\begin{aligned} |\lambda_1 + \lambda_2| < 2 \quad \Rightarrow |\phi_1| < 2 \\ |\lambda_1, \lambda_2| < 1 \quad \Rightarrow |\phi_2| < 1 \end{aligned}$$

• Summary:

We say the process is globally (asymptotically) stable if the solution of the associated homogenous equation tends to 0, as  $t \rightarrow \infty$ .

#### Theorem

A necessary and sufficient condition for global asymptotical stability of a  $p^{\text{th}}$  order deterministic difference equation with constant coefficients is that all roots of the associated lag polynomial equation  $\phi(z)=0$  have *moduli* strictly more than 1.

For the case of real roots, moduli means "absolute values."

## AR(1) Process – Stationarity & ACF

An AR(1) model:

 $y_t = \phi_1 y_{t-1} + \varepsilon_t, \quad \varepsilon_t \sim WN.$ 

Recall that in a previous example, under the stationarity condition  $|\phi_1| < 1$ , we derived the moments:

$$E[y_t] = \mu = 0 \qquad (assuming \phi_1 \neq 1)$$
  

$$Var[y_t] = \gamma(0) = \sigma^2 / (1 - \phi_1^2) \qquad (assuming |\phi_1| < 1)$$
  

$$\gamma(1) = E[y_t y_{t-1}] = E[(\phi_1 y_{t-1} + \varepsilon_t) * y_{t-1}] = \phi_1 \gamma(0)$$
  

$$\gamma(2) = E[y_t y_{t-2}] = E[(\phi_1 y_{t-1} + \varepsilon_t) * y_{t-2}]$$
  

$$= \phi_1 E[y_{t-1} y_{t-2}] = \phi_1 \gamma(1) = \phi_1^2 \gamma(0)$$
  

$$\gamma(3) = E[y_t y_{t-3}] = E[(\phi_1 y_{t-1} + \varepsilon_t) * y_{t-3}]$$
  

$$= \phi_1 E[y_{t-1} y_{t-3}] = \phi_1 \gamma(2) = \phi_1^3 \gamma(0)$$
  

$$\vdots$$
  

$$\gamma(k) = \phi_1 \gamma(k-1) = \phi_1^k \gamma(0)$$

Now, we derive the autocorrelation:  $\rho(t_1, t_2) = \frac{\gamma(t_1 - t_2)}{\sigma_{t_1}\sigma_{t_2}}$ If the process is stationary  $(\sigma_t = \sigma_{t-1} = \sqrt{\gamma(0)})$   $\rho(1) = \rho(t, t-1) = \frac{\gamma(1)}{\sigma_t \sigma_{t-1}} = \frac{\gamma(1)}{\gamma(0)} = \phi_1$   $\rho(2) = \frac{\gamma(2)}{\gamma(0)} = \phi_1^2$ :  $\rho(k) = \frac{\gamma(k)}{\gamma(0)} = \phi_1^k$ 

<u>Remark</u>: The ACF decays with *k*.

Note that when  $\phi_1 = 1$ , the AR(1) is non-stationary,  $\rho(k) = 1$ , for all k. The present and the past are always correlated!

• Again, when  $|\phi_1| < 1$ , the autocorrelations do not explode as k increases. There is an exponential decay towards zero.

Note:

- when  $0 < \phi_1 < 1 \implies$  All autocorrelations are positive. - when  $-1 < \phi_1 < 0 \implies$  The sign of  $\rho(k)$  shows an alternating pattern beginning with a negative value.

The plot of  $\rho(k)$  against k, is called *autocorrelogram*. We also plot  $\rho(0)$ , which is 1.

## AR(1) Process – Stationarity & ACF: Simulations

We simulate and plot three AR(1) processes, with standard normal  $\varepsilon_t$ -i.e.,  $\sigma = 1$ :

$$y_t = 0.5 y_{t-1} + \varepsilon_t$$
  
 $y_t = -0.9 y_{t-1} + \varepsilon_t$   
 $y_t = 2 y_{t-1} + \varepsilon_t$ 

To simulate ARMA process, we use the *arima.sim* R function. Below, we start with the plot of the first AR(1) process:

 $y_t = 0.5 y_{t-1} + \varepsilon_t$  with 200 simulations.

> plot(arima.sim(list(order = c(1,0,0), ar = 0.5), n=200), ylab="Simulated Series", main=(expression(AR(1)~~~phi==+.5)))





<u>Note</u>: The process  $\theta_1 > 0$  is smoother than the ones with  $\theta_1 < 0$ . The process with  $|\theta_1| > 1$ , explodes!

Below, we compute and plot the ACF for the two stable simulated process. 1)  $y_t = 0.5 y_{t-1} + \varepsilon_t$ 

sim\_ar1\_5 <- arima.sim(list(order = c(1,0,0), ar = 0.5), n=200) acf\_ar1\_5 <- acf(sim\_ar1\_5, main=(expression(AR(1)~~~phi==+.5))) > acf\_ar1\_5

Autocorrelations of series 'sim mal 5', by lag 1.000 -0.860 0.720 -0.551 0.427 -0.330 0.258 -0.205 0.183 -0.202 0.209 -0.218 0.213 -0.216 -0.026 -0.106 -0.123 0.009 -0.009 -0.004 -0.012 -0.015 0.000 -0.008 AR(1)  $\phi = -0.9$ 



2)  $y_t = -0.9 y_{t-1} + \varepsilon_t$ sim\_ar1\_9 <- arima.sim(list(order=c(1,0,0), ar = -0.9), n=200) acf\_ar1\_9 <- acf(sim\_ar1\_9, main=(expression(AR(1)~~~phi==-.9))) > acf\_ar1\_9

Autocorrelations of series 'sim mal 9', by lag 1.000 -0.584 0.093 0.061 -0.132 0.147 -0.181 0.122 -0.013 -0.023 0.014 -0.012 0.092 -0.199 0.193 -0.155 0.143 -0.107 0.014 0.174 -0.244 0.196 -0.154 0.105

#### AR(1) Process – Stationarity & ACF: Real Data

**Example:** A process with  $|\phi_1| < 1$  (actually, 0.065) is the monthly changes in the USD/GBP exchange rate. Below we plot its corresponding ACF:



Below we plot the monthly changes in the USD/GBP exchange rate. Stationary series do not look smooth:



**Example:** A process with  $\phi_1 \approx 1$  (actually, 0.99) is the nominal USD/GBP exchange rate. Below, we plot the ACF, it is not 1 all the time, but its decay is very slow (after 30 months, it is still .40 correlated!):



Below we plot the nominal USD/GBP exchange rate. Stationary series look smooth, smooth enough that you can clearly spot trends:



USD/GBP Exchange Rate: Monthly Rates (1971-2020)

# AR(1) Process – Stationarity & ACF

An AR(2) model:

$$y_t = \mu + \phi_1 y_{t-1} + \phi_2 y_{t-2} + \varepsilon_t, \quad \varepsilon_t \sim WN.$$

• Moments:

$$E[y_t] = \mu/(1 - \phi_1 - \phi_2) = 0 \qquad (assuming\phi_1 + \phi_2 \neq 1) Var[y_t] = \sigma^2/(1 - \phi_1^2 - \phi_2^2) \qquad (assuming\phi_1^2 + \phi_2^2 < 1)$$

• Autocovariance function

$$\begin{split} \gamma(k) &= \operatorname{Cov}[y_t, y_{t-k}] = \operatorname{E}[(\phi_1 \ y_{t-1} + \phi_2 \ y_{t-2} + \varepsilon_t) \ y_{t-k}] \\ &= \phi_1 \ \operatorname{E}[y_{t-1} \ y_{t-k}] + \phi_2 \ \operatorname{E}[y_{t-2} \ y_{t-k}] + \operatorname{E}[\varepsilon_t \ y_{t-k}] \\ &= \phi_1 \ \gamma(k-1) + \phi_2 \ \gamma(k-2) + \operatorname{E}[\varepsilon_t \ y_{t-k}] \end{split}$$

• We have a recursive formula:

$$\begin{aligned} &(k=0) \quad \gamma(0) = \phi_1 \, \gamma(-1) + \phi_2 \, \gamma(-2) + \mathrm{E}[\varepsilon_t \, y_t] \\ &= \phi_1 \, \gamma(1) + \phi_2 \, \gamma(2) + \sigma^2 \\ &(k=1) \quad \gamma(1) = \phi_{1^1} \, \gamma(0) + \phi_2 \, \gamma(1) + \mathrm{E}[\varepsilon_t \, y_{t-1}] \\ &= \phi_1 \, \gamma(0) + \phi_2 \, \gamma(1) \end{aligned}$$

$$\Rightarrow \gamma(1) = [\phi_1/(1 - \phi_2)] \gamma(0)$$
(k=2)  $\gamma(2) = \phi_1 \gamma(1) + \phi_2 \gamma(0) + E[\varepsilon_t y_{t-2}]$ 

$$= \phi_1 \gamma(1) + \phi_2 \gamma(0)$$

$$\Rightarrow \gamma(2) = [\phi_1^2 \gamma(0)/(1 - \phi_2)] + \phi_2 \gamma(0)$$

$$= [\phi_1^2/(1 - \phi_2) + \phi_2] \gamma(0)$$
Replacing  $\gamma(1)$  and  $\gamma(2)$  back to  $\gamma(0)$ :
$$\gamma(0) = [\phi_1^2/(1 - \phi_2)] \gamma(0) + [\phi_2 \phi_1^2/(1 - \phi_2) + \phi_2^2] \gamma(0) + \sigma^2$$

$$= \frac{\sigma^2(1 - \phi_2)}{(1 - \phi_2) - \phi_1^2(1 + \phi_2) + \phi_2^2(1 - \phi_2)} \Rightarrow |\phi_2| < 1$$

• Dividing the previous formulas by  $\gamma(0)$ , we get the ACF:

$$\begin{split} \rho(k) &= \gamma(k)/\gamma(0) = \phi_1 \,\rho(k-1) + \phi_2 \,\rho(k-2) + \mathrm{E}[\varepsilon_t \, y_{t-k}]/\gamma(0) \\ (k=0) \quad \rho(0) &= 1 \\ (k=1) \quad \rho(1) &= \phi_1 \,/(1-\phi_2) \\ (k=2) \quad \rho(2) &= \phi_1 \,\rho(1) + \phi_2 \,\rho(0) = \phi_1^2/(1-\phi_2) + \phi_2 \\ (k=3) \quad \rho(3) &= \phi_1 \,\rho(2) + \phi_2 \,\rho(1) = \\ &= \phi_1^3/(1-\phi_2) + \phi_1 \,\phi_2 + \phi_2 \phi_1 \,/(1-\phi_2) \end{split}$$

<u>Remark</u>: Again, we see exponential decay in the ACF.

From the work above, we need:  $\phi_1 + \phi_2 \neq 1$ 

$$\phi_1 + \phi_2 \neq 1.$$
  
 $\phi_1^2 + \phi_2^2 < 1.$   
 $|\phi_2| < 1.$ 

#### AR(p) Process – VAR(1) Representation

With AR process with more lags than the AR(1) process, it is complicated to determine stationarity by looking at the  $\phi$ 's coefficients.

Stationarity conditions can be derived in a simplified way by rewriting an AR(p) as AR(1) process. For example, the AR(2) process:

 $y_t = \mu + \phi_1 y_{t-1} + \phi_2 y_{t-2} + \varepsilon_t \quad \Rightarrow (1 - \phi_1 L - \phi_2 L^2) y_t = \mu + \varepsilon_t$ can be written in matrix form as an AR(1):

$$\begin{bmatrix} y_t \\ y_{t-1} \end{bmatrix} = \begin{bmatrix} \mu \\ 0 \end{bmatrix} + \begin{bmatrix} \phi_1 & \phi_2 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} y_{t-1} \\ y_{t-2} \end{bmatrix} + \begin{bmatrix} \varepsilon_t \\ 0 \end{bmatrix} \implies \tilde{y}_t = \tilde{\mu} + A \tilde{y}_{t-1} + \tilde{\varepsilon}_t$$

• The AR(2) in matrix AR(1) form is called Vector AR(1) or VAR(1).

• We can derived a matrix lag polynomial A(L):  $\tilde{y}_t = \tilde{\mu} + A\tilde{y}_{t-1} + \tilde{\varepsilon}_t \implies A(L)\tilde{y}_t = [I - AL] \tilde{y}_t = \tilde{\varepsilon}_t.$ 

#### AR(2) Process – VAR(1) & Stationarity

If A(L) is invertible we can write an  $MA(\infty)$  representation:

$$\tilde{y}_t = \tilde{\mu} + A \tilde{y}_{t-1} + \tilde{\varepsilon}_t \qquad \Rightarrow \tilde{y}_t = [I - AL]^{-1} \tilde{\varepsilon}_t$$

Note: Recall the expansion:

Checking that [I - AL] is not singular, same as checking that  $A^j$  does not explode. The stability of the system (solution) can be determined by the *eigenvalues* of **A**. That is, get the  $\lambda_i$ 's and check if  $|\lambda_i| < 1$  for all *i*.

$$A = \begin{bmatrix} \phi_1 & \phi_2 \\ 1 & 0 \end{bmatrix} \Rightarrow |A - \lambda I| = \det \begin{bmatrix} \phi_1 & -\lambda & \phi_2 \\ 1 & -\lambda \end{bmatrix} = -(\phi_1 - \lambda)\lambda - \phi_2$$
$$= \phi_2 - \phi_1\lambda + \lambda^2$$
  
• Solution to quadratic equation:  $\lambda_i = \frac{\phi_1 \pm \sqrt{\phi_1^2 - 4\phi_2}}{2}$ 

<u>Stability and stationary</u>:  $|\lambda_i| < 1$ .  $\Rightarrow$  roots of  $\phi(z)$  *outside unit circle*. For the AR(2) process, we have already derived some relations between  $\lambda_i$ 's and  $\phi_i$ 's:

$$\begin{array}{l} \lambda_1 \lambda_2 = \phi_2 \quad \Rightarrow |\lambda_1 \lambda_2| = |\phi_2| < 1 \\ \lambda_1 + \lambda_2 = \phi_1 \quad \Rightarrow |\lambda_1 + \lambda_2| = |\phi_1| < 2 \end{array}$$

• We derived autocovariance function,  $\gamma(k)$ , before, getting a recursive formula. Let's write the first autocovariances:

 $\begin{array}{l} (k=0) \quad \gamma(0) = \phi_1 \ \gamma(1) + \phi_2 \ \gamma(2) + \sigma^2 \\ (k=1) \quad \gamma(1) = [\phi_1/(1 - \phi_2)] \ \gamma(0) \\ (k=2) \quad \gamma(2) = [\phi_1^2/(1 - \phi_2) + \phi_2] \ \gamma(0) \end{array}$ 

With  $|\phi_2| < 1$ , we get well defined  $\gamma(1)$ ,  $\gamma(2)$  &  $\gamma(0)$ .

The VAR(1) has a nice property: The VAR(1) is Markov -i.e., forecasts depend only on today's data.

It looks complicated, but it is straightforward to apply the VAR formulation to any AR(p) processes. We can also use the same eigenvalue conditions to check the stationarity of AR(p) processes.

#### **AR Process – Stationarity & Ergodicity**

**Theorem**: The linear AR(p) process is strictly stationary and ergodic if and only if the roots of  $\phi(L)$  are  $|r_i| > 1$  for all j, where  $|r_i|$  is the modulus of the complex number  $r_i$ .

<u>Note</u>: If one of the  $r_j$ 's equals 1,  $\phi(L)$  (&  $y_t$ ) has a unit root –i.e.,  $\phi(1)=0$ . This is a special case of *non-stationarity*.

Recall  $\phi(L)^{-1}$  produces an infinite sum on the  $\varepsilon_{t-j}$ 's. If this sum does not explode, we say the process is *stable*.

# AR Process – Dynamic Multiplier & IRF

If the process is stable, we can calculate  $\frac{\delta y_t}{\delta \varepsilon_{t-1}}$ 

 $\frac{\delta y_t}{\delta \varepsilon_{t-j}}$  = How much  $y_t$  is affected today by an innovation t - jj periods ago,  $\varepsilon_{t-j}$ . When expressed as a function of *j*, we call this *dynamic multiplier*. Accumulated over time it is the *impulse response function (IRF)*.

The *dynamic multiplier* measurers the effect of an innovation,  $\varepsilon_t$ , (economist like to call the  $\varepsilon_t$ 's, "*shocks*") on subsequent values of  $y_t$ : That is, the first derivative on the "Wold representation" – i.e., a stationary process represented as an MA process:

$$\frac{\delta y_{t+j}}{\delta \varepsilon_t} = \frac{\delta y_t}{\delta \varepsilon_0} = \psi_j.$$

where  $\psi_i$ 's are the coefficient of MA representation.

For an AR(1) process:

$$\frac{\delta y_{t+j}}{\delta \varepsilon_t} = \frac{\delta y_t}{\delta \varepsilon_0} = \Phi^j.$$

That is, the dynamic multiplier for any linear stochastic difference equation (SDE) depends only on the length of time j, not on time t.

• The *impulse-response function* (IRF) is an accumulation of the sequence of dynamic multipliers, as a function of time from the one time change in the innovation,  $\varepsilon_t$ .

Usually, IRFs are represented with a graph, that measures the effect of the innovation,  $\varepsilon_t$ , on  $y_t$  over time:

$$\frac{\delta y_{t+j}}{\delta \varepsilon_t} + \frac{\delta y_{t+j+1}}{\delta \varepsilon_t} + \frac{\delta y_{t+j+2}}{\delta \varepsilon_t} + \dots = \psi_j + \psi_{j+1} + \psi_{j+2} + \dots$$

Once we estimate the AR, MA or ARMA coefficients, we draw an IRF.



**Example**: AR(1) process:

 $y_t = \mu + \phi_1 y_{t-1} + \varepsilon_t,$   $\varepsilon_t \sim WN.$ The AR(1) is stable if  $|\phi_1| < 1$   $\Rightarrow$  stationarity condition.

We invert the AR(1) to get an MA( $\infty$ ):  $1/(1 - \phi_1) = \sum_{j=0}^{\infty} \phi_1^{j}$ 

Then,

$$y_{t} = \mu^{*} + \phi_{1} \varepsilon_{t-1} + \phi_{1}^{2} \varepsilon_{t-2} + \phi_{1}^{3} \varepsilon_{t-3} + \phi_{1}^{4} \varepsilon_{t-4} + \dots + \varepsilon_{t}.$$

Under the stationarity condition, we calculate the dynamic multiplier:  $\delta y_{t+1} / \delta \varepsilon_{t-j} = \phi_1^{j}$ 

Accumulated over time, after J periods, the effect of shock  $\varepsilon_t$  at t+J is: IRF(at t+J) =  $\sum_{i=0}^{J-1} \phi_i^{j}$ 

Suppose  $\phi_1 = 0.40$ . Then,  $\delta y_t / \delta \varepsilon_{t-1} = \phi_1 = 0.40$   $\delta y_t / \delta \varepsilon_{t-2} = \phi_1 = 0.40^2$ :  $\delta y_t / \delta \varepsilon_{t-J} = \phi_1 = 0.40^J$ 

After 5 periods, the accumulated effect of a shock today is: IRF(at t+5) =  $0.40 + 0.40^2 + 0.40^3 + 0.40^4 + 0.40^5 = 0.65984$ .

### **AR Process – Causality**

The AR(p) model:  $\phi(L)y_t = \mu + \varepsilon_t, \quad \varepsilon_t \sim WN.$ 

where  $\phi(L) = 1 - \phi_1 L^1 - L^2 \phi_2 - \dots - \phi_p L^p$ 

Then,  $y_t = \phi(L)^{-1}(\mu + \varepsilon_t)$ ,  $\Rightarrow$  an MA( $\infty$ ) process!

But, we need to make sure that we can invert the polynomial  $\phi(L)$ .

When  $\phi(L) \neq 0$ , we say the process  $y_t$  is *causal* (strictly speaking, a *causal function of*  $\{\varepsilon_t\}$ ).

<u>Definition</u>: A linear process  $\{y_t\}$  is *causal* if there is a

<u>Definition</u>: A linear process  $\{y_t\}$  is *causal* if there is a  $\psi(L) = 1 + \psi_1 L + \psi_2 L^2 + \cdots$ with  $\sum_{j=0}^{\infty} |\psi_j(L)| < \infty$ with  $y_t = \psi(L)\varepsilon_t$ .

**Example**: AR(1) process:

 $\phi(L)y_t = \mu + \varepsilon_t$ , where  $\phi(L) = 1 - \phi_1 L$ 

Then, yt is causal if and only if:

 $|\phi_1| < 1$  (same condition as stationarity) or

the root  $r_1$  of the polynomial  $\phi(z) = 1 - \phi_1 z$  satisfies  $|r_1| > 1$ .

Question: How do we calculate the 's coefficients for an AR(p)? A: Matching coefficients ( $\mu = 0$ ):

$$Y_t = \frac{1}{(1-\phi_{1L})} \varepsilon_t \stackrel{|\phi_1|<1}{\cong} \sum_{i=0}^{\infty} \phi 1^i L^i \varepsilon_t$$
$$= (1+\phi_{1L}+\phi_{1}^2 L^2 + \cdots) \varepsilon_t \qquad \Rightarrow \psi_i = \phi_i^i, \quad i \ge 0$$

#### **AR Process – Estimation and Properties**

We go back to the general AR(p). Define

$$\boldsymbol{x}_t = \begin{pmatrix} 1 & y_{t-1} & y_{t-2} & \dots & y_{t-p} \end{pmatrix}$$
$$\boldsymbol{\beta} = (\mu & \phi_1 & \phi_2 & \dots & \phi_p)$$

Then the model can be written as

$$y_t = x_t' \boldsymbol{\beta} + \varepsilon_t$$

The OLS estimator is

$$\mathbf{b} = (X'X)^{-1}X'y$$

• Properties:

– Using the Ergodic Theorem, OLS estimator is consistent.

– Using the MDS CLT, OLS estimator is asymptotically normal.

 $\Rightarrow$  asymptotic inference is the same.

The asymptotic covariance matrix is estimated just as in the cross-section case: The sandwich estimator.

#### **ARMA Process**

A combination of AR(p) and MA(q) processes produces an ARMA(p, q) process:

$$y_{t} = \mu + \phi_{1}y_{t-1} + \phi_{2}y_{t-2} + \dots + \phi_{p}y_{t-p} + \varepsilon_{t} - \theta_{1}\varepsilon_{t-1} - \theta_{2}\varepsilon_{t-2} - \dots - \theta_{q}\varepsilon_{t-q}$$
$$= \mu + \sum_{i=1}^{p} \phi_{i}y_{t-i} - \sum_{i=1}^{q} \theta_{i}L^{i}\varepsilon_{t} + \varepsilon_{t}$$
$$\Rightarrow \phi(L)y_{t} = \mu + \theta(L)\varepsilon_{t}$$

Usually, we insist that  $\phi(L) \neq 0$ ,  $\theta(L) \neq 0$  & that the polynomials  $\phi(L)$ ,  $\theta(L)$  have no *common factors*. This implies it is not a lower order ARMA model.

## **ARMA Process – Common Factors**

It is possible to reduce the order of an ARMA structure if the  $\phi(L)$  and  $\theta(L)$  lag polynomials have *common factors*.

**Example**: Suppose we have the following ARMA(2, 3) model

 $\phi(L)y_t = \theta(L)\varepsilon_t$ 

with

$$\phi(L) = 1 - .6L + .3L^2$$
  

$$\theta(L) = 1 - 1.4L + .9L^2 - .3L^3 = (1 - .6L + .3L^2)(1 - L)$$

This model simplifies to:  $y_t = (1 - L)\varepsilon_t \implies \text{an MA}(1) \text{ process.} \P$ 

## **ARMA Process – Representation**

An ARMA process can be rewritten as:

- Pure AR Representation:  $\Pi(L)(y_t \mu) = \varepsilon_t \Rightarrow \Pi(L) = \frac{\phi_p(L)}{\theta_q(L)}$
- Pure MA Representation:  $(y_t \mu) = \Psi(L)\varepsilon_t \Rightarrow \Psi(L) = \frac{\theta_q(L)}{\phi_p(L)}$
- Special ARMA(p, q) cases: -p = 0: MA(q) -q = 0: AR(p).

# **ARMA: Stationarity, Causality and Invertibility**

**Theorem:** If  $\phi(L)$  and  $\theta(L)$  have no common factors, a (unique) *stationary* solution to  $\phi(L)y_t = \theta(L)\varepsilon_t$  exists if and only if

$$|z|\leq 1\Rightarrow\phi(z)=1-\phi_1\,z-\phi_2z^2-\ldots-\phi_pz^p\neq 0.$$

This ARMA(*p*, *q*) model is causal if and only if  
$$|z| \le 1 \Rightarrow \phi(z) = 1 - \phi_1 z - \phi_2 z^2 - \dots - \phi_p z^p \ne 0.$$

This ARMA(p, q) model is invertible if and only if

$$|z| \le 1 \Rightarrow \theta(z) = 1 + \theta_1 z - \theta_2 z^2 + \ldots + \theta_p z^p \neq 0.$$

<u>Note</u>: Real data cannot be *exactly* modeled using a finite number of parameters. We choose p, q to create a good approximated model.

# Lecture 9 – ARIMA Models – Identification & Estimation

### **ARMA Process**

We defined the ARMA(p, q) model:  $\phi(L)(y_t - \mu) = \theta(L)\varepsilon_t$ 

The mean does not affect the order of the ARMA. Then, if  $\mu \neq 0$ , we demean the data:  $x_t = y_t - \mu$ .

Then,  $\phi(L)x_t = \theta(L)\varepsilon_t \implies x_t$  is a *demeaned* ARMA process.

- In this lecture, we will study:
- Identification of *p*, *q*.
- Estimation of ARMA(p, q)
- Non-stationarity of  $x_t$ .
- Differentiation issues ARIMA(p, d, q)
- Seasonal behavior SARIMA(p, d, q)s

## **Autocovariance Function (Again)**

We define the autocovariance function:  $\gamma(t - j) = E[y_t y_{t-j}]$ 

For an AR(*p*) process, WLOG with  $\mu$ =0 (or demeaned  $y_t$ ), we get:

$$\gamma(t-j) = E[(\phi_1 y_{t-1} y_{t-j} + \phi_2 y_{t-2} y_{t-j} + \dots + \phi_p y_{t-p} y_{t-j} + \varepsilon_t y_{t-j})]$$
  
=  $\phi_1 \gamma(j-1) + \phi_2 \gamma(j-2) + \dots + \phi_p \gamma(j-p)$ 

<u>Notation</u>:  $\gamma(k)$  or  $\gamma_k$  are commonly used. Sometimes,  $\gamma(k)$  is referred as "*covariance at lag k*."

The  $\gamma(t-j)$  determine a system of equations:

$$\begin{split} \gamma(0) &= E[y_t \ y_t] = \phi_1 \gamma(1) + \phi_2 \gamma(2) + \phi_3 \gamma(3) + \dots + \phi_p \gamma(p) + \sigma^2 \\ \gamma(1) &= E[y_t \ y_{t-1}] = \phi_1 \gamma(0) + \phi_2 \gamma(1) + \phi_3 \gamma(2) + \dots + \phi_p \gamma(p-1) \\ \gamma(2) &= E[y_t \ y_{t-2}] = \phi_1 \gamma(1) + \phi_2 \gamma(0) + \phi_3 \gamma(1) + \dots + \phi_p \gamma(p-2) \vdots \\ \vdots &\vdots &\vdots &\vdots \end{split}$$

This a pxp system of equations. Using linear algebra, we can write the system as:

 $\Gamma \phi = \gamma$ where  $\Gamma$  is a *pxp* matrix of autocovariances, with  $\gamma(0)$  on the diagonal;  $\phi$  is the *px*1 vector of AR(*p*) coefficients; and  $\gamma$  is the *px*1 vector of  $\gamma(k)$  autocovariances

**Example**: AR(1) model:

 $y_t = \phi_1 y_{t-1} + \varepsilon_t, \quad \varepsilon_t \sim WN.$ 

Then, the autocovariance function is:

$$\gamma(0) = E[y_t \ y_t] = \operatorname{Var}[y_t] = \sigma^2 / (1 - \phi_1^2)$$
  

$$\gamma(1) = E[y_t \ y_{t-1}] = E[(\phi_1 \ y_{t-1} + \varepsilon_t) * \ y_{t-1}] = \phi_1 \ \gamma(0)$$
  

$$\gamma(2) = E[y_t \ y_{t-2}] = E[(\phi_1 \ y_{t-1} + \varepsilon_t) * \ y_{t-2}] = \phi_1 \ \gamma(1) = \phi_1^2 \ \gamma(0)$$
  

$$\gamma(3) = E[y_t \ y_{t-3}] = E[(\phi_1 \ y_{t-1} + \varepsilon_t) * \ y_{t-3}] = \phi_1 \ E[y_{t-1} \ y_{t-3}]] = \phi_1^3 \ \gamma(0)$$
  

$$\gamma(k) = \phi_1 \ \gamma(k-1) = \phi_1^k \ \gamma(0) \implies \operatorname{If} |\phi_1| < 1, \text{ exponential decay.}$$

••••

.....

Under stationarity, moments are constant. That is,

 $\operatorname{Var}[y_t] = \operatorname{Var}[y_{t-1}] = \sqrt{\gamma(0)}. \P$ 

**Example**: MA(1) process:

 $y_t = \theta_1 \varepsilon_{t-1} + \varepsilon_t, \qquad \varepsilon_t \sim WN.$ 

Then, the autocovariance function is:

 $\gamma(0) = \sigma^{2} + \theta_{1}^{2} \sigma^{2} = \sigma^{2} (1 + \theta_{1}^{2})$   $\gamma(1) = E[y_{t} \ y_{t-1}] = E[(\theta_{1} \ \varepsilon_{t-1} + \varepsilon_{t}) (\theta_{1} \ \varepsilon_{t-2} + \varepsilon_{t-1})] = \theta_{1} \sigma^{2}$  $\gamma(k) = E[y_{t} \ y_{t-k}] = E[(\theta_{1} \ \varepsilon_{t-1} + \varepsilon_{t}) (\theta_{1} \ \varepsilon_{t-k-1} + \varepsilon_{t-k})] = 0 \quad (\text{for } k > 1)$ 

That is, for |k| > 1,  $\gamma(k) = 0$ .

**Example:** ARMA(1,1) process:  

$$y_t = \phi_1 y_{t-1} + \theta_1 \varepsilon_{t-1} + \varepsilon_t, \qquad \varepsilon_t \sim WN.$$

$$\begin{split} \gamma(k) &= E[y_t \ y_{t-k}] = E[\{\phi_1 y_{t-1} + \theta_1 \ \varepsilon_{t-1} + \varepsilon_t\} \ y_{t-k}] \\ &= \phi_1 E[y_{t-1} \ y_{t-k}] + E[\varepsilon_t \ y_{t-k}] + \theta_1 E[\varepsilon_{t-1} \ y_{t-k}] \\ &= \phi_1 \gamma(k-1) + E[\varepsilon_t \ y_{t-k}] + \theta_1 E[\varepsilon_{t-1} \ y_{t-k}] \\ \gamma(0) &= \phi_1 \gamma(-1) + \underbrace{E[\varepsilon_t \ y_t]}_{\sigma^2} + \theta_1 E\left[\varepsilon_{t-1} \ \underbrace{y_t}_{\phi_1 y_{t-1} + \varepsilon_t + \theta_1 \varepsilon_{t-1}}\right] \\ &= \phi_1 \gamma(1) + \sigma^2 + \theta_1 E\left[\varepsilon_{t-1}(\phi_1 \ \underbrace{y_{t-1}}_{\phi_1 y_{t-2} + \varepsilon_{t-1} + \theta_1 \varepsilon_{t-2}}\right] \\ &= \phi_1 \gamma(1) + \sigma^2 + \theta_1 (\phi_1 \ \sigma^2 + \theta_1 \sigma^2) \end{split}$$

• Similarly,

for 
$$k = 1$$
,  
 $\gamma(1) = \phi_1 \gamma(0) + E[\varepsilon_t y_{t-1}] + \theta_1 E[\varepsilon_{t-1} y_{t-1}]$   
 $= \phi_1 \gamma(0) + \theta_1 E[\varepsilon_{t-1} \{\phi_1 y_{t-2} + \theta_1 \varepsilon_{t-2} + \varepsilon_{t-1}\}]$   
 $= \phi_1 \gamma(0) + \theta_1 \gamma(1)$ 

For k = 2,  $\gamma(2) = \phi_1 \gamma(1) + E[\varepsilon_t y_{t-2}] + \theta_1 E[\varepsilon_{t-1} y_{t-2}]$ 

$$= \phi_1 \gamma(1) + \theta_1 E[\varepsilon_{t-1} \{ \phi_1 y_{t-3} + \theta_1 \varepsilon_{t-3} + \varepsilon_{t-2} \}] \\ = \phi_1 \gamma(1)$$

For a general k,

 $\begin{aligned} \gamma(k) &= \phi_1 \gamma(k-1) \\ &= \phi_1^{k-1} \gamma(1), \quad k > 1 \qquad \Rightarrow \text{If } |\phi_1| < 1, \text{ exponential decay.} \end{aligned}$ 

• Two equations for  $\gamma(0)$  and  $\gamma(1)$ :  $\gamma(0) = \phi_1 \gamma(1) + \sigma^2 + \theta_1 (\phi_1 \sigma^2 + \theta_1 \sigma^2)$   $\gamma(1) = \phi_1 \gamma(0) + \theta_1 \gamma(1)$ Solving for  $\gamma(0) \& \gamma(1)$ :

$$\gamma(0) = \sigma^{2} \frac{1 + \theta_{1}^{2} + 2\phi_{1}\theta_{1}}{1 - \phi_{1}^{2}}$$
  

$$\gamma(1) = \sigma^{2} \frac{(1 + \phi_{1}\theta_{1}) * (\phi_{1} + \theta_{1})}{1 - \phi_{1}^{2}}$$
  

$$\vdots$$
  

$$\gamma(k) = \phi_{1}^{k-1}\gamma(1), \quad k > 1.$$

<u>Note</u>: If stationary, ARMA(1,1) & AR(1) show exponential decay. Difficult to distinguish one from the other through autocovariances.

#### **Autocorrelation Function (ACF)**

Now, we define the autocorrelation function (ACF):

$$\rho(k) = \frac{\gamma(k)}{\gamma(0)} = \frac{\text{covariance at lag } k}{\text{variance}}$$

The ACF lies between -1 and +1, with  $\rho(0) = 1$ .

Dividing the autocovariance system by  $\gamma(0)$ , we get:

Γ	$\rho(0)$	$\rho(1)$	•••	$\rho(p-1)$	[ <b>φ</b> <sub>1</sub> ]		$\left[\rho(1)\right]$	1
I	$\rho(1)$	ho(0)	•••	$\rho(p-2)$	$\phi_2$	_	$\rho(2)$	
I	:	:	•••	÷	:	_	:	
L	$\rho(p-1)$	$\rho(p-2)$		$\rho(0)$	$\left\lfloor \phi_{p} \right\rfloor$		$\rho(p)$	

Or using linear algebra:

$$\mathbf{P} \boldsymbol{\phi} = \boldsymbol{\rho}$$

These are "Yule-Walker" equations, which can be solved numerically.

#### **Autocorrelation Function (ACF) – Estimation & Correlogram** • Estimation:

Easy: Use sample moments to estimate  $\gamma(k)$  and plug in formula:

$$r_k = \hat{\rho}_k = \frac{\sum (Y_t - \bar{Y})(Y_{t+k} - \bar{Y})}{\sum (Y_t - \bar{Y})^2}$$

Then, we plug the  $\hat{\rho}_k$  in the Yule-Walker equations and solve for  $\phi$ :  $\hat{\mathbf{P}} \phi = \hat{\rho}$ 

The sample *correlogram* is the plot of the ACF against k. As the ACF lies between -1 and +1, the correlogram also lies between these values.

#### • Distribution:

For a linear, stationary process, with large *T*, the distribution of the sample ACF,  $r_k = \hat{\rho}_k$  is approximately normal with:

 $\mathbf{r} \xrightarrow{d} N(\mathbf{\rho}, \mathbf{V}/T),$  V is the covariance matrix.

Under H<sub>0</sub>:  $\rho_k = 0$  for all k > 1.  $\mathbf{r} \xrightarrow{d} \mathbf{N}(\mathbf{0}, \mathbf{I}/T) \implies \operatorname{Var}[\mathbf{r}(k)] = 1/T$ .

Under H<sub>0</sub>:  $\rho_k = 0$  for all k, the SE =  $1/\sqrt{T} \implies 95\%$  C.I.:  $0 \pm 1.96 * 1/\sqrt{T}$ 

Then, for a white noise sequence, approximately 95% of the sample ACFs should be within the above C.I. limits.

<u>Note</u>: The SE =  $1/\sqrt{T}$  are sometimes referred as *Bartlett's SE*.

**Example**: Sample ACF for an AR(1) process: Under stationarity (constant moments, in particular,  $\operatorname{Var}[y_t] = \operatorname{Var}[y_{t-1}] = \gamma(0)$ ):  $\rho(k) = \frac{\gamma(k)}{\gamma(0)} = \phi_1^k \qquad k = 0, 1, 2, \dots$ 

If  $|\phi_1| < 1$ , the ACF will show exponential decay.

Suppose  $\phi_1 = 0.4$ . Then,  $\rho(0) = 1$   $\rho(1) = 0.4$   $\rho(2) = 0.4^2 = 0.16$   $\rho(3) = 0.4^3 = 0.064$   $\rho(4) = 0.4^4 = 0.0256$ :  $\rho(k) = 0.4^k$ 

• We simulate an AR(1) series with with  $\phi_1 = 0.4$ , using the R function *arima.sim*.

sim\_ar1\_04 <- arima.sim(list(order=c(1,0,0), ar=0.4), n=200) #simulate AR(1) series
plot(sim\_ar1\_04, ylab="Simulated Series", main=(expression(AR(1):~~~phi==0.4)))
acf(sim\_ar1\_04) #plot ACF for sim series</pre>



**Example:** Sample ACF for an MA(1) process:  $\rho(0) = 1$   $\rho(k) = \theta_1/(1+\theta_1^2)$ , for k = 1, -1  $\rho(k) = 0$  for |k| > 1. After k = 1 –i.e., one lag– the ACF dies out.

Suppose  $\theta_1 = 0.5$ . Then,  $\rho(0) = 1$   $\rho(1) = 0.4$  $\rho(k) = 0$  for |k| > 1.

• We simulate an MA(1) series with  $\phi_1=0.4$ 

sim\_ma1\_05 <- arima.sim(list(order=c(0,0,1), ma=0.5), n=200) #simulate MA(1) series
plot(sim\_ma1\_05, ylab="Simulated Series", main=(expression(MA(1):~~~theta==0.5)))
acf(sim\_ma1\_05) #plot ACF for sim series</pre>



**Example**: Sample ACF for an MA(q) process:  $\gamma(0) = E[y_t \ y_t]] = \sigma^2 (1 + \theta_1^2 + \theta_2^2 + ... + \theta_q^2)$   $\gamma(1) = E[y_t \ y_{t-1}] = \sigma^2 (\theta_1 + \theta_2 \theta_1 + \theta_3 \theta_2 + ... + \theta_q \theta_{q-1})$   $\gamma(2) = E[y_t \ y_{t-2}]] = \sigma^2 (\theta_2 + \theta_3 \theta_1 + ... + \theta_q \theta_{q-2})$  $\gamma(q) = \theta_q$ 

In general,

$$\gamma(k) = \sigma^2 \sum_{j=k}^{q} \theta_j \theta_{j-k} \qquad k \le q \qquad \text{(with } \theta_0 = 0 \\ = 0 \qquad \text{otherwise.}$$

1).

Then,

$$\rho(k) = \frac{\sum_{j=k}^{q} \theta_{j} \theta_{j-k}}{(1+\theta_{1}^{2}+\theta_{2}^{2}+\dots+\theta_{q}^{2})} \qquad k \le q$$
  
= 0 otherwise

For an MA(3):

$$y_t = \mu + \varepsilon_t + \theta_1 \varepsilon_{t-1} + \theta_2 \varepsilon_{t-2} + \theta_3 \varepsilon_{t-3}$$

Then,

$$\rho(0) = 1 \rho(1) = \frac{\theta_1 + \theta_2 \theta_1 + \theta_3 \theta_2}{(1 + \theta_1^2 + \theta_2^2 + \theta_3^2)}$$

= 0

$$\rho(2) = \frac{\theta_2 + \theta_3 \theta_1}{(1 + \theta_1^2 + \theta_2^2 + \theta_3^2)}$$

$$\rho(3) = \frac{\theta_3}{(1 + \theta_1^2 + \theta_2^2 + \theta_3^2)}$$

$$\rho(k) = 0 \qquad \text{for } |k| > 3.$$

Suppose  $\theta_1 = 0.5$ ;  $\theta_2 = 0.4$ ;  $\theta_3 = 0.2$ .

Then,

Plot of simulated series and ACF > sim\_ma3\_05 <- arima.sim(list(order=c(0,0,3), ma=c(0.5, 0.4, 0.1)), n=200) # sim MA(3)





From the autocovariances, we get

$$\gamma(0) = \sigma^{2} \frac{1+\theta_{1}^{2}+2\phi_{1}\theta_{1}}{1-\phi_{1}^{2}}$$

$$\gamma(1) = \sigma^{2} \frac{(1+\phi_{1}\theta_{1})*(\phi_{1}+\theta_{1})}{1-\phi_{1}^{2}}$$

$$\gamma(k) = \phi_{1}\gamma(k-1) = \phi_{1}^{k-1}\sigma^{2} \frac{(1+\phi_{1}\theta_{1})*(\phi_{1}+\theta_{1})}{1-\phi_{1}^{2}}$$

Then,

$$\rho(k) = \phi_1^{k-1} \frac{(1+\phi_1\,\theta_1)*(\phi_1+\theta_1)}{1+\theta_1^2+2\phi_1\theta_1}$$

 $\Rightarrow$  If  $|\phi_1| < 1$ , exponential decay. Similar pattern to AR(1).

- The ACF for an ARMA(1,1):  $\rho(k) = \phi_1^{k-1} \frac{(1+\phi_1\theta_1) * (\phi_1+\theta_1)}{1+\theta_1^2 + 2\phi_1\theta_1}$
- Suppose  $\phi_1 = 0.4, \theta_1 = 0.5$ . Then,  $\rho(0) = 1$   $\rho(1) = \frac{(1+0.4*0.5)*(0.4+0.5)}{1+0.5^2+2*0.4*0.5} = 0.6545$   $\rho(2) = 0.4*\frac{(1+0.4*0.5)*(0.4+0.5)}{1+0.5^2+2*0.4*0.5} = 0.2618$   $\rho(3) = 0.4^2*\frac{(1+0.4*0.5)*(0.4+0.5)}{1+0.5^2+2*0.4*0.5} = 0.0233$ :  $\rho(k) = 0.4^{k-1}*\frac{(1+0.4*0.5)*(0.4+0.5)}{1+0.5^2+2*0.4*0.5}$

Plot of simulated series ARMA (1,1) and ACF > sim\_arma11 <- arima.sim(list(order=c(1,0,1), ar=0.4, ma=0.5), n=200) # sim ARMA(1,1)



**Example: US Monthly Returns** (1871 – 2020, *T*=1,795) Sh da <- read.csv("C://Financial Econometrics/Shiller 2020data.csv", head=TRUE, sep=",")  $x P \le Sh da P$  $x D \le Sh da$  $T \leq - length(x P)$  $\ln p \le \log(x P[-1]/x P[-T])$  $\ln d \le \log(x D[-1]/x D[-T])$  $acf p \leq acf(lr p)$ # acf: R function that estimates the ACF > acf pAutocorrelations of series 'lr p', by lag 1.000 0.279 0.004 -0.043 0.017 0.074 0.039 0.039 0.044 0.035 0.034 0.022 -0.010 -0.059 -0.058 -0.056 0.009 0.033 0.047 -0.040 -0.087 -0.090 -0.029 0.005 24 25 0.003 -0.013 -0.058 -0.018 -0.005 0.026 0.011 0.000 0.020

 $SE(r_k) = 1/sqrt(T) = 1/sqrt(1,795) = .0236. \implies 95\% \text{ C.I.} \pm 2* 0.0236$ 



<u>Note</u>: With the exception of first correlation, correlations are small. However, many are significant, not strange result when *T* is large.  $\P$ 

**Example: US Monthly Changes in Stock Dividends** (1871 – 2020, *T*=1,795) > acf dAutocorrelations of series 'lr d', by lag **1.000 0.462** 0.516 0.432 0.444 0.326 0.442 0.288 0.283 0.265 0.202 0.168 0.142 0.100 0.122 0.123 0.085 0.045 0.026 -0.013 0.001 -0.029 -0.049 -0.077 -0.038 -0.100 -0.095 -0.055 -0.081 -0.092 -0.034 -0.063 -0.089

High correlations and significant even after 32 months!



Note: Correlations are positive for almost 1.5 years, then correlations become negative.

#### **ACF – Joint Significance Tests**

Recall the Ljung-Box (LB) statistic as:

$$LB = T(T+2) \sum_{k=1}^{m} (\frac{\hat{\rho}_{k}^{2}}{(T-k)})$$

The LB test can be used to determine if the first *m* sample ACFs are jointly equal to zero. Under H<sub>0</sub>:  $\rho_1 = \rho_2 = ... = \rho_m = 0$ , the LB has an asymptotic  $\chi^2(m)$  distribution.

**Example**: LB test with 20 lags for US Monthly Returns and Changes in Dividends (1871 – 2020) > Pox test(lr, p, lag=20, type= "Liung Pox")

> Box.test(lr\_p, lag=20, type= "Ljung-Box")

Box-Ljung test

data:  $lr_p$ X-squared = 208.02, df = 20, p-value < 2.2e-16  $\Rightarrow$  Reject first 20 correlations.

 $\Rightarrow$  Reject H<sub>0</sub> at 5% level. Joint significant

> Box.test(lr d, lag=20, type= "Ljung-Box")

Box-Ljung test

data: lr d

X-squared = 2762.7, df = 20, p-value < 2.2e-16  $\Rightarrow$  Reject H<sub>0</sub> at 5% level. Joint significant first 20 correlations. ¶

## **Partial ACF (PACF)**

The ACF gives us a lot of information about the order of the dependence when the series we analyze follows a MA process: The ACF is zero after q lags for an MA(q) process.

If the series we analyze, however, follows an ARMA or AR, the ACF alone tells us little about the orders of dependence: We only observe an exponential decay.

We introduce a new function that behaves like the ACF of MA models, but for AR models, namely, the *partial autocorrelation function* (PACF).

The PACF is similar to the ACF. It measures correlation between observations that are k time periods apart, after controlling for correlations at intermediate lags.

Intuition: Suppose we have an AR(1):

 $y_t = \phi_1 y_{t-1} + \varepsilon_t.$ 

Then,

 $\gamma(2) = \phi_1^2 \gamma(0)$ 

The correlation between  $y_t$  and  $y_{t-2}$  is not zero, as it would be for an MA(1), because  $y_t$  is dependent on  $y_{t-2}$  through  $y_{t-1}$ .

Suppose we break this chain of dependence by removing ("partialing out") the effect  $y_{t-1}$ . Then, we consider the correlation between  $[y_t - \phi_1 y_{t-1}]$  and  $[y_{t-2} - \phi_1 y_{t-1}]$  –i.e, the correlation between  $y_t$  and  $y_{t-2}$  with the linear dependence of each on  $y_{t-1}$  removed:

 $\gamma(2) = \operatorname{Cov}(y_t - \phi_1 y_{t-1}, y_{t-2} - \phi_1 y_{t-1}) = \operatorname{Cov}(\varepsilon_t, y_{t-2} - \phi_1 y_{t-1}) = 0.$ 

Similarly,

$$\gamma(k) = \operatorname{Cov}(\varepsilon_t, y_{t-k} - \phi_1 y_{t-1}) = 0 \quad \text{for all } k > 1.$$

<u>Definition</u>: The PACF of a stationary time series  $\{y_t\}$  is  $\phi_{11} = \text{Corr}(y_t, y_{t-1}) = \rho(1)$  $\phi_{hh} = \text{Corr}(y_t - \text{E}[y_t|I_{t-1}], y_{t-h} - \text{E}[y_{t-h}|I_{t-1}])$  for h = 2, 3, ...

This removes the linear effects of  $y_{t-1}, y_{t-2}, \dots, y_{t-h}$ .

The PACF  $\Phi_{hh}$  is also the last coefficient in the best linear prediction of  $y_t$  given  $y_{t-1}, y_{t-2}, ..., y_{t-h}$ .

Estimation by Yule-Walker equation, using sample estimates:

 $\widehat{\boldsymbol{\phi}}_{h} = [\widehat{\boldsymbol{R}}]^{-1}\widehat{\boldsymbol{\gamma}}(k) \implies \text{a recursive system,}$ where  $\boldsymbol{\phi}_{h} = (\phi_{h1}, \phi_{h2}, ..., \phi_{hh})$  and  $\boldsymbol{R}$  is the (hxh) correlation matrix.

A recursive algorithm, Durbin-Levinson, can be used. Also OLS can be used.

## Partial ACF – AR(p)

**Example**: AR(*p*) process:

 $y_{t} = \mu + \phi_{1}y_{t-1} + \phi_{2}y_{t-2} + \dots + \phi_{p}y_{t-p} + \varepsilon_{t}$   $E[y_{t}|I_{t-1}] = \mu + \phi_{1}y_{t-1} + \phi_{2}y_{t-2} + \dots + \phi_{p}y_{t-h-1}$  $E[y_{t-h}|I_{t-1}] = \mu + \phi_{1}y_{t-h-1} + \phi_{2}y_{t-h-2} + \dots + \phi_{p}y_{t-1}$  Then,

 $\Rightarrow$  After the *p*<sup>th</sup> *PACF*, all remaining PACF are 0 for AR(*p*) processes.

The plot of the PACF is called the *partial correlogram*.

<u>R Note</u>: The R function *pacf* computes the PACF.

**Example**: We simulate an AR(2) process:  $y_t = \mu + \phi_1 y_{t-1} + \phi_2 y_{t-2} + \varepsilon_t$ 

sim\_ar2 <- arima.sim(list(order=c(1,0,0), ar=c(0.5, 0.3)), n=200) #simulate AR(2) series
plot(sim\_ar2, ylab="Simulated Series", main=(expression(AR(2))))
pacf\_ar2 <- pacf(sim\_ar2)</pre>

• Print PACF > pacf ar2 0.558 0.286 0.038 0.103 -0.010 0.009 0.111 0.060 -0.021 -0.076 0.016  $-0.086 - 0.139 \quad 0.100 \quad 0.061 - 0.156 \quad 0.078 - 0.103 \quad 0.043 - 0.075 \quad 0.104 \quad 0.024 \quad 0.061$ -0.038 -0.100 -0.095 -0.055 -0.081 -0.092 -0.034 -0.063 -0.089

 $SE(r_k) \approx 1/sqrt(200) = .0707 \implies 95\%$  C.I.:  $0 \pm 1.96 * 0.0707$ 

Plot of simulated series and PACF

> plot(sim\_ar2, ylab="Simulated Series", main=(expression(AR(2))))
>pacf\_ar2 <- pacf(sim\_ar2)</pre>







<u>Note</u>: The PACF can be calculated by *h* regressions, each one with *h* lags. The *hh* coefficient is the  $h^{\text{th}}$  order PACF.

> ar(sim\_ar2, order.max=1, method = "ols")
Coefficients:

# 1

**0.5586** Intercept: -0.008403 (0.0761) Order selected 1 sigma<sup>2</sup> estimated as 1.152

> ar(sim\_ar2, order.max=2, method = "ols") Coefficients: 1 2 0.3974 0.2869 Intercept: -0.009847 (0.07326) Order selected 2 sigma^2 estimated as 1.063. ¶

# Partial ACF – MA(q)

Following the analogy that PACF for AR processes behaves like an ACF for MA processes, we will see exponential decay (*"tails off"*) in the partial correlogram for MA process. Similar pattern will also occur for ARMA(p, q) process.

**Example**: We simulate an MA(1) process with  $\theta_1 = 0.5$ . sim\_ma1 <- arima.sim(list(order=c(0,0,1), ma=0.5), n=200) > pacf(sim\_ma1)





# Partial ACF – ARMA(p,q)

For an ARMA processes, we will see exponential decay ("tails off") in the partial correlogram.

**Example**: We simulate an ARMA(1) process with  $\phi_1 = 0.4 \& \theta_1 = 0.5$ . sim\_arma11 <- arima.sim(list(order=c(1,0,1), ar=0.4, ma=0.5), n=200) > pacf(sim arma11)



## **Partial ACF – Examples**

Partial autocorrelations of series 'lr\_p', by lag

**0.278** -0.081 -0.026 0.041 0.058 0.002 0.038 0.032 0.016 0.022 0.009 -0.023 -0.057 -0.032 -0.045 0.027 0.017 0.037 -0.059 -0.051 -0.050 0.005 0.006 0.004 -0.005 -0.051 0.014 -0.007 0.037 0.008 0.018

 $SE(r_k) = 1/sqrt(1,795) = .0236$   $\Rightarrow 95\%$  C.I.:  $0 \pm 1.96 * 0.0236$ 

 $> pacf(lr_p)$ 



<u>Note</u>: With the exception of the first partial correlation, partial correlations are small, though, again, some are significant.  $\P$ 

**Example**: US Monthly Stock Dividends (1871 – 2020, *T* = 1,795) pacf\_d <- pacf(lr\_d) > pacf\_d

```
Partial autocorrelations of series 'lr d', by lag
 1
                   4
                                            8
                                                 9
                                                       10
       2
             3
                         5
                               6
                                      7
                                                            11
0.462 0.385 0.160 0.150 -0.033 0.189 -0.054 -0.056 0.027 -0.082 -0.019
12
       13
             14
                   15
                         16
                                     18
                                            19
                                                 20
                                                       21
                                                             22
                               17
-0.023 -0.057 -0.032 -0.045 0.027 0.017 0.037 -0.059 -0.051 -0.050 0.005
  23
       24
             25
                         27
                                      29
                                            30
                   26
                               28
                                                 31
                                                       32
-0.041 0.050 -0.036 -0.030 0.091 0.006 -0.017 0.044 -0.002 -0.042
```

Higher partial correlations than for stock returns.

 $> pacf(lr_d)$ 



Note: Partial correlations are positive for almost 6 lags, then become small.
## **Non-Stationary Time Series Models**

The ACF is as a rough indicator of whether a trend is present in a series. A slow decay in ACF is indicative of highly correlated data, which suggests a true unit root process, or a trend stationary process.

Formal tests can help to determine whether a system contains a trend and whether the trend is deterministic or stochastic (unit root). Popular tests are the ADF(Augmented Dickey-Fuller), PP( Phillips-Perron) and KPSS (Kwiatkowski-Phillips-Schmidt-Shin).

We will analyze two situations faced in ARMA models: (1) Deterministic trend – Simple model:  $y_t = \alpha + \beta t + \varepsilon_t$ . - Solution: Detrending -i.e., regress yt on t. Then, keep residuals for further modeling.

(2) Stochastic trend – Simple model:  $y_t = c + y_{t-1} + \varepsilon_t$ . - Solution: Differencing –i.e., apply  $\Delta = (1 - L)$  operator to y<sub>t</sub>. Then, use  $\Delta y_t$  for further modeling.

**Example**: Below, we plot two series with a clear trend: U.S. Monthly Prices and U.S. Dividends (1871 - 2020).



#### Monthly U.S. Stock Price

### Non-Stationary Time Series Models – Deterministic Trend

Suppose we have the following model:

$$y_t = \alpha + \beta t + \varepsilon_t. \qquad \Rightarrow \Delta y_t = y_t - y_{t-1}$$

 $\{y_t\}$  shows only temporary departures, given by the  $\varepsilon_t$ 's, from trend line  $\alpha + \beta t$ . This type of model is a *trend stationary* (TS) model.

We take first differences in the TS model:

$$\Delta y_t = y_t - y_{t-1} = \alpha + \beta t + \varepsilon_t - (\alpha + \beta (t-1) + \varepsilon_{t-1}) = \beta + \varepsilon_t - \varepsilon_{t-1}$$

Taking expectations:

 $E[\Delta y_t] = \beta \implies y_t$  shows constant change over time.

If a series is TS –i.e., it has a deterministic time trend–, then we simply regress  $y_t$  on an intercept and a time trend (t = 1, 2, ..., T) and save the residuals:

$$e_t = y_t - \hat{\alpha} - \hat{\beta} t$$

The residuals are the *detrended*  $y_t$  series (= $y_t$  without the influence of t).

If  $y_t$  is stochastic, we do not necessarily get stationary series, by detrending.

Many economic series exhibit "exponential trend/growth". They grow over time like an exponential function over time instead of a linear function. In this cases, it is common to work with logs

 $\ln(y_t) = \alpha + \beta t + \varepsilon_t. \qquad (\Rightarrow y_t = e^{\alpha + \beta t + \varepsilon_t})$ 

We take first differences in the exponential trend/growth model:

 $\Delta \ln(y_t) = \ln(y_t) - \ln(y_{t-1}) = \alpha + \beta t + \varepsilon_t - \alpha - \beta (t-1) - \varepsilon_{t-1}$  $= \beta + \varepsilon_t - \varepsilon_{t-1}$  $\Rightarrow \text{The average growth rate is: } E[\Delta \ln(y_t)] = \beta$ 

• We can have ARMA models, with more complex trend structure:

 $y_t = \alpha + \phi_1 y_{t-1} + \ldots + \phi_p y_{t-p} + \beta_1 t + \beta_2 t^2 + \ldots + \beta_k t^k + \varepsilon_t.$ 

In these cases, in general, the estimation of ARMA involves two steps, both with OLS. For example for the case of AR(p) with a trend and quadratic trend components:

(1) Detrend  $y_t$ : regress  $y_t$  against a constant, t, and  $t^2$ .  $\Rightarrow$  get the residuals (= $y_t$  without the influence of t).

(2) Estimate AR(p): Use residuals to estimate the AR(p) model.

Note: This 2-step method is usually called Frish-Waugh method.

Simulated Example 1: We simulate an AR(1) series with a trend:  $y_t = 0.3 + 0.2 y_{t-1} + 0.05 t + \varepsilon_t.$ 

```
T sim <- 200
                                      # Length of simulation
y_sim \le matrix(0, T_sim, 1)
                                      # vector to accumulate simulated data
                                      # Draw T_sim normally distributed errors
u \leq rnorm(T sim, sd = 1)
mu <- 0.3
                                      # Constant
phi1 <- 0.2
                                      # Change to create different AR(1) patterns
mu t <- .05
                                      # Trend coefficient
y sim[1] <- mu/(1 - phi1)
                                      # Initial observation (= to unconditional mean & t=0)
t <- 2
                                      # Time index for observations
while (t \leq T_sim) {
y_sim[t] = mu + phi1 * y_sim[t-1] + mu_t * t + u[t] # y_sim simulated autocorrelated values
t < -t + 1
}
y det t \leq y sim[2: T sim]
plot(y det t, type="l", col = "blue", main = "Simulated Series with a Deterministic Trend")
# Detrend series
T sim1 <- length(y det t)
trend <- c(1:T \text{ sim1})
fit det t <- lm(y det t \sim trend)
y det t filt <- fit det t$residuals
                                                     # Filtered series
plot(y det t filt, type="l", main = "Detrended Simulated Series")
```

• Below, we plot the simulated AR(1) series (blue) and the detrended simulated series (red).



<u>Remark</u>: There is no longer a trend, only the AR(1) component remains in the detrended series.

Simulated Example 2: Now, in the previous simulated example, we add a quadratic trend:  $y_t = 0.3 + 0.2 y_{t-1} + 0.05 t + 0.003 t^2 + \varepsilon_t$ .

mu\_t2 <- .003 # Trend square coefficient t <- 2 # Time index for observations while (t <= T\_sim) { y\_sim[t] = mu + phi1 \* y\_sim[t-1] + mu\_t \* t + u[t]# y\_sim simulated autocorrelated values t <- t + 1 } y\_det\_t <- y\_sim[2: T\_sim] # Detrend series with only a linear trend trend <- c(1:(T\_sim1-1))</pre>

 $\begin{array}{l} \text{fit\_det\_t <- lm(y\_det\_t \sim trend)} \\ \text{y\_det\_t\_filt <- fit\_det\_t$residuals} & \# \text{ Filtered series} \end{array}$ 

• Below, we plot the simulated AR(1) series (blue) and the detrended series from the above regression, which only involves a constant (violet).

plot(y\_det\_t, type="l", col = "blue", main = "Simulated Series with a Deterministic Trend")



plot(y det t filt, type="l", main = "Detrended Simulated Series")



<u>Remark</u>: We made a mistake, we detrended a series with a linear and a quadratic trend, using a model with only a linear trend. As observe above, an unexpected deterministic (U-shape) patterns shows up in the detrended series –i.e., the residuals. We need to detrend using an appropriate model, with a linear and quadratic trends. This is what we do below:

## Detrend series with a linear & Quadratic trends
trend2 <- trend^2fit\_det\_t <- lm(y\_det\_t ~ trend + trend2)
y\_det\_t\_filt <- fit\_det\_t\$residuals # Filtered series
plot(y\_det\_t\_filt, type="l", col = "violet", main="Detrended Simulated Series")</pre>

• Below, we plot the detrended simulated series with a linear and quadratic trends (red).



<u>Remark</u>: A series with a quadratic trend, needs to be detrended with a quadratic trend, otherwise extra patterns (U-shape, in this case) appear. Once we use an appropriate detrending model, we use the detrended series –i.e., the residuals– for furthering (ARMA) modeling.

<b>Example:</b> We detrend U.S. Stock Prices	
$T \leq - length(x_P)$	# length of series
trend $\leq$ - c(1:T)	# create trend

det  $P \leq -lm(x P \sim trend)$ # regression to get detrended e detrend P <- det P\$residuals plot(detrend P, type="l", col="blue", ylab ="Detrended U.S. Prices", xlab ="Time") title("Detrended U.S. Stock Prices")



• Not very appealing series. We still see trends and an extra U-shape pattern shows up in detrended series. Now, we detrend U.S. Stock Prices adding a squared trend.

trend2 <- trend^2 det P2 <-  $lm(x P \sim trend + trend2)$ # regression to get detrended e detrend P2 <- det P2\$residuals plot(detrend P2, type="l", col="blue", ylab ="Detrended U.S. Prices", xlab ="Time")





• Still, trends and an extra pattern are still observed. It is possible that there is exponential growth in original series. Then, we detrend Log U.S. Stock Prices adding, first, a linear trend and. Second, both a linear and square trends.

 $1 P \leq \log(x P)$ det  $1P \leq -lm(1 P \sim trend)$ # regression to get detrended e detrend 1P <- det 1P\$residuals plot(detrend lP, type="l", col="blue", ylab ="Detrended Log U.S. Prices", xlab ="Time") title("Detrended Log U.S. Stock Prices")

det\_lP2 <- lm(l\_P ~ trend + trend2) # regression to get detrended e det\_lP2 <- det\_lP2\$residuals plot(det\_lP2, type="l", col="blue", ylab ="Det Log U.S. Prices", xlab ="Time") title("Detrended Log U.S. Stock Prices with linear and quadratic trends")



<u>Remark</u>: The second detrended series, with linear and quadratic trends looks better, but we still see trends in the graph, and, thus, evidence of a time dependent mean. ¶

## Non-Stationary Time Series Models – Stochastic Trend

The more modern approach is to consider trends in time series as a variable.

A variable trend exists when a trend changes in an unpredictable way. Therefore, it is considered as *stochastic*.

Recall the AR(1) model:  $y_t = c + \phi_1 y_{t-1} + \varepsilon_t$ .

As long as  $|\phi| < 1$ , everything is fine, we have a stationary AR(1) process: OLS is consistent, t-stats are asymptotically normal, etc.

Now consider the extreme case where  $\phi_1 = 1$ ,  $\Rightarrow y_t = c + y_{t-1} + \varepsilon_t$ .

Where is the (stochastic) trend? No *t* term.

Let us replace recursively the lag of  $y_t$  on the right-hand side:

$$y_t = \mu + y_{t-1} + \varepsilon_t$$
  
=  $\mu + (\mu + y_{t-2} + \varepsilon_{t-1}) + \varepsilon_t$   
...  
=  $y_0 + t \mu + \sum_{j=0}^t \varepsilon_{t-j}$ 

A constant (y<sub>0</sub>), a determinist trend (t  $\mu$ ) and an accumulation of errors over time ( $\sum_{j=0}^{t} \varepsilon_{t-j}$ ) appear in the recursive formulation. This is what we call a "random walk with drift". The series grows with *t*.

Each  $\varepsilon_t$  shock represents a shift in the intercept. All values of  $\{\varepsilon_t\}$  have a 1 as coefficient  $\Rightarrow$  each shock never vanishes (permanent).

We remove the trend by differencing  $y_t \Rightarrow \Delta y_t = (1 - L) y_t = \mu + \varepsilon_t$ 

<u>Note</u>: Applying the (1 - L) operator to a time series is called *differencing* 

**Example:** We difference **U.S. Stock Prices**, using the *diff* R function: diff\_P <- diff(x\_P)

> plot(diff\_P,type="l", col="blue", ylab ="Differenced U.S. Stock Prices", xlab ="Time")
> title("Differenced U.S. Stock Prices")



<u>Remark</u>: The trend is gone from the graph. ¶

•  $y_t$  is said to have a *stochastic trend* (ST), since each  $\varepsilon_t$  shock gives a permanent and random change in the conditional mean of the series.

For these situations, we use Autoregressive Integrated Moving Average (ARIMA) models.

Question: Deterministic or Stochastic Trend?

They appear similar: Both lead to growth over time. The difference is how we think of  $\varepsilon_t$ . Should a shock today affect  $y_{t+1}$ ?

- TS:  $y_{t+1} = c + \beta (t+1) + \varepsilon_{t+1}$ - ST:  $y_{t+1} = c + y_t + \varepsilon_{t+1} = c + [c + y_{t-1} + \varepsilon_t] + \varepsilon_{t+1}$  $\Rightarrow \varepsilon_t$  affects  $y_{t+1}$ . (In fact, the shock will have a permanent impact.)

#### ARIMA(*p*, *d*, *q*) Models

For p, d,  $q \ge 0$ , we say that a time series  $\{y_t\}$  is an ARIMA (p, d, q) process if  $w_t = \Delta^d y_t = (1 - L)^d y_t$  is ARMA(p, q). That is,

$$\phi(L)(1-L)^d y_t = \theta(L) \varepsilon_t$$

Applying the (1 - L) operator to a time series is called *differencing*.

<u>Notation</u>: If  $y_t$  is non-stationary, but  $\Delta^d y_t$  is stationary, then  $y_t$  is *integrated* of order d, or I(d). A time series with *unit root* is I(1), typical of asset prices. A stationary time series is I(0), typical of log changes of asset prices (returns).

#### **Examples**:

Example 1: RW:  $y_t = y_{t-1} + \varepsilon_t$ . y<sub>t</sub> is non-stationary, but  $(1 - L) y_t = \varepsilon_t \Rightarrow$  white noise! Now,  $y_t \sim ARIMA(0,1,0)$ .

Example 2: AR(1) with time trend:  $y_t = \mu + \delta t + \phi_1 y_{t-1} + \varepsilon_t$ .  $y_t$  is non-stationary, but  $w_t = (1 - L) y_t = \mu + \delta t + \phi_1 y_{t-1} + \varepsilon_t - (\mu + \delta (t-1) + \phi_1 y_{t-2} + \varepsilon_{t-1})$   $= \delta + \phi_1 w_{t-1} + \varepsilon_t - \varepsilon_{t-1}$ Now,  $y_t \sim \text{ARIMA}(1,1,1)$ .

We call both process first difference stationary.

Note:

- Example 1: Differencing a series with a unit root in the AR part of the model reduces the AR order.

- Example 2: Differencing can introduce an extra MA structure. We introduced non-invertibility. This happens when we difference a TS series. Detrending should be used in these cases.

• In practice:	
A root near 1 of the AR polynomial	$\Rightarrow$ differencing
A root near 1 of the MA polynomial	$\Rightarrow$ over-differencing

• In general, we have the following results:

- Too little differencing: not stationary.

- Too much differencing: extra dependence introduced.

• Finding the right *d* is crucial. For identifying preliminary values of *d*:

- Use a time plot.

- Check for slowly decaying (persistent) ACF/PACF.

### **ARIMA Models: Unit Roots 1?**

**Example 1**: Monthly Stock Price levels (1871-2020) acf\_P <- acf(x\_P)

> acf\_P Autocorrelations of series 'x\_p', by lag

1.000 0.992 0.984 0.977 0.971 0.966 0.961 0.954 0.946 0.938 0.931 0.924  $0.917 \ 0.911 \ 0.904 \ 0.897 \ 0.891 \ 0.884 \ 0.877 \ 0.871 \ 0.865 \ 0.860 \ 0.854 \ 0.848$ 0.841 0.834 0.827 0.821 0.815 0.809 0.803 0.797 0.790



Very high autocorrelations. Looks like  $\phi_1 \approx 1$ .

#### **Example 2: Monthly Interest Rates** (1871-2020)

acf  $i \leq acf(x i)$ > acf i Autocorrelations of series 'x i', by lag 1.000 0.996 0.990 0.985 0.980 0.975 0.970 0.965 0.960 0.956 0.951 0.946 0.940 0.934 0.929 0.924 0.919 0.915 0.912 0.908 0.904 0.901 0.899 0.896 0.894 0.891 0.889 0.887 0.884 0.882 0.879 0.877 0.874



Very high autocorrelations. Looks like  $\phi_1 \approx 1$ .

#### **ARIMA Models – Random Walk**

A *random walk* (RW) is defined as a process where the current value of a variable is composed of the past value plus an error term defined as a white noise (a normal variable with zero mean and variance one).

A Random Walk is an ARIMA(0,1,0) process  $y_t = y_{t-1} + \varepsilon_t \implies \Delta y_t = (1-L)y_t = \varepsilon_t, \qquad \varepsilon_t \sim WN(0, \sigma^2).$ 

Popular model. Used to explain the behavior of financial assets, unpredictable movements (Brownian motions, drunk persons).

It is a special case (limiting) of an AR(1) process: a *unit-root* process.

Implication:  $E[y_{t+1}|I_t] = y_t \implies \Delta y_t$  is absolutely random.

Thus, a RW is nonstationary, and its variance increases with t.

<b>Examples</b> : Two simulated RW	
T_sim <- 200	# Sample size for simulation
u <- rnorm(200)	<pre># Draw T_sim normally distributed errors</pre>
y_sim <- matrix(0,T_sim,1)	# Vector to collect simulated data
phi <- 1	# Change to create different correlation patterns

a <- 2 # Time index for observations mu <- 0 # RW Drift (mu = 0, no drift) while (a <= T\_sim) { y\_sim[a] = mu + rho \* y\_sim[a-1] + u[a] # y\_sim simulated autocorrelated values a <- a + 1 }

plot(y\_sim, type="l", col="blue", ylab ="Simulated Series", xlab ="Time") title("Simulated RW Series with no drift")



<u>Remark</u>: The (stochastic trends) are clear in both graphs.

### **ARIMA Models – Random Walk with Drift**

Change in  $y_t$  is partially deterministic ( $\mu$ ) and partially stochastic.

$$y_t - y_{t-1} = \Delta y_t = \mu + \varepsilon_t$$

It can also be written as

 $y_t = y_0 + t \mu + \sum_{j=0}^t \varepsilon_{t-j}$  $\Rightarrow \varepsilon_t \text{ has a permanent effect on the mean of } y_t.$ 

Recall the difference between conditional and unconditional forecasts:

 $E[y_t] = y_0 + t \mu$  (Unconditional forecast)  $E[y_{t+s}|y_t] = y_t + s \mu$  (Conditional forecast)

### **ARIMA Models: Box-Jenkins**

An effective procedure for building empirical time series models is the Box-Jenkins approach, which consists of three stages:

(1) Identification or Model specification (of ARIMA order)

(2) **Estimation** of identified ARM(p, q) structure.

(3) **Diagnostics testing**: Checking that residuals are white noise.

Two main approaches to (1) Identification.

- Correlation approach: Mainly based on ACF & PACF.

1) Make sure data is stationary –check a time plot. If not, differentiate.

2) Using ACF & PACF, guess small values for *p* & *q*.

- *Information criteria*: Very common situation: The order choice not clear from looking at ACF & PACF. Then, use AIC (or AICc), BIC, or HQIC (Hannan and Quinn (1979)). This is the usual (& easier) approach.

Value parsimony. When in doubt, keep it simple (KISS).

## **ARIMA Models: Identification – ACF & PACF**

Basic tools: sample ACF and sample PACF.

- ACF identifies order of MA: Non-zero at lag q; zero for lags > q.
- PACF identifies order of AR: Non-zero at lag p; zero for lags > p.
- All other cases, try ARMA(p, q) with p > 0 and q > 0.

<u>Summary</u>: For p > 0 and q > 0.

	AR(p)	MA(q)	ARMA(p,q)
ACF	Tails off	0 after lag q	Tails off
PACF	0 after lag p	Tails off	Tails off

<u>Note</u>: Ideally, "Tails off" is exponential decay. In practice, in these cases, we may see a lot of non-zero values for the ACF and PACF.

## **ARIMA Models: Identification – AR(1)**





# **ARIMA Models: Identification – MA(1)**

**ARIMA Models: Identification – ARMA(1,1)** 





























## **ARIMA Models: Identification – Examples Example 1: Monthly US Returns** (1871 - 2020).



Note: ARMA(1,1), MA(1), AR(2)?

**Example 2: Monthly US Dividend Changes** (1871 - 2020).



Note: Not clear: Maybe long a ARMA(p,q) or needs differencing?



**Example 3**: Monthly Log Changes in Oil Prices (1973 - 2020).

<u>Note</u>: MA(1), AR(4)? ¶

**Example 4**: Monthly Log Changes in Gold (1973 - 2020).





Note: No clear ARMA structure.

## **ARIMA Model: Identification – IC**

In general, it is not easy to identify an ARMA model using the ACF and PACF. It is common to rely on information criteria (IC).

IC's are equal to the estimated variance or the log-likelihood function plus a penalty factor, that depends on *k*. Many IC's:

- Akaike Information Criterion (AIC) AIC = -2 \* (ln L - k) = -2 ln L + 2 \* k $\Rightarrow$  if normality AIC =  $T * \ln(\mathbf{e}'\mathbf{e}/T) + 2*k$  (+constants)

- Bayes-Schwarz Information Criterion (BIC or SBIC) BIC = -2 \* ln  $L - \ln(T)$  \* k $\Rightarrow$  if normality AIC =  $T * \ln(\mathbf{e}^{*}\mathbf{e}/T) + \ln(T) * k$  (+constants)

- Hannan-Quinn (HQIC) HQIC =  $-2*(\ln L - k [\ln(\ln(T))]$  $\Rightarrow$  if normality AIC =  $T * \ln(\mathbf{e}^*\mathbf{e}/T) + 2 k [\ln(\ln(T))]$  (+constants)

It is very common to compute the IC's under normality (it is the default setting in R and almost all other packages). Recall that under normality, we write the Likelihood function as:

$$\ln L = -\frac{T}{2}\ln(2\pi\sigma^2) - \frac{1}{2\sigma^2}\underbrace{S(\phi p, \theta_q, \mu)}_{Errors\,SS} = -\frac{T}{2}\ln(2\pi\sigma^2) - \frac{1}{2\sigma^2}T\sigma^2\ln L =$$
$$= -\frac{T}{2}\ln\sigma^2 - \underbrace{\frac{T}{2}(1+\ln 2\pi)}_{\text{constant}}$$

Since we compare different ARIMA models, using the same data, the constants play no role in our decision. They can be ignored. Then,

- AIC =  $T * \ln(\hat{\sigma}^2) + 2 * k$ - BIC =  $T * \ln(\hat{\sigma}^2) + \ln(T) * k$ - HQIC =  $T * \ln(\hat{\sigma}^2) + 2 * k * [\ln(\ln(T))]$  The goal of these criteria is to provide us with an easy way of comparing alternative model specifications, by ranking them.

<u>General Rule</u>: The lower the IC, the better the model. For the previous IC's, then choose model to AIC<sub>J</sub>, BIC<sub>J</sub>, or HQIC.

## **ARIMA Model: Identification – Remarks**

Some remarks about IC's:

- IC's are not test statistics. They do not test a model.
- They are used for ranking. The raw value tends to be ignored.
- They have two components: a goodness of fit component -based on lnL- and a model
- complexity component –the penalty based on *k*.
- Different penalties, different IC's.
- Some authors scale the IC's by T. Since raw values tend to be irrelevant, this is not an issue.

We would like these statistics -i.e., the IC's- to have good properties. For example, if the true model is being considered among many, we want the IC to select it. This can be done on average (unbiased) or as *T* increases (consistent).

Some results regarding AIC and BIC.

- AIC and Adjusted  $R^2$  are not consistent.

- AIC is conservative –i.e., it tends to over-fit:  $k_{AIC}$  too large models.

- In time series, AIC selects the model that minimizes the out-of-sample one-step ahead forecast MSE.

- BIC is more parsimonious than AIC. It penalizes the inclusion of parameters more ( $k_{BIC} \le k_{AIC}$ ).

- BIC is consistent in autoregressive models.

- No agreement which criteria is better.

## **ARIMA Model: Identification – Small Sample Modifications**

• There are modifications of IC to get better finite sample behavior, a popular one is AIC corrected, *AICc*, statistic:

$$AICc = T \ln \hat{\sigma}^2 + \frac{2k(k+1)}{T-k-1}$$

AICc converges to AIC as T gets large. Using AICc is not a bad idea.

For AR(*p*) models, other AR-specific criteria are possible: Akaike's final prediction error (FPE), Akaike's BIC, Parzen's CAT.

Hannan and Rissannen's (1982) *minic (=Minimum IC)*: Calculate the BIC for different p's (estimated first) and different q's. Select the best model –i.e., lowest BIC.

Note: Box, Jenkins, and Reinsel (1994) proposed using the AIC above.

<u>**R** Note</u>: The R function auto.arima uses *AICc* to select p, q; d is selected using a formal unit root test (KPSS).

## **ARIMA Model: Identification – In practice**

**Example**: We compute, for **monthly US Returns (1871 - 2020)**, annan and Rissannen (1982)'s minic, based on AIC.

Lags	MA 0	MA 1	MA 2	MA 3	MA 4	MA 5
AR 0	-6403.59	-6552.94	-6552.69	-6554.27	-6552.88	-6557.37
AR 1	-6545.22	-6552.23	-6551.86	-6552.42	-6552.64	-6561.48
AR 2	-6554.76	-6553.28	-6554.85	-6554.35	-6564.32	-6559.48
AR 3	-6553.94	-6552.53	-6554.44	-6552.33	-6550.36	-6558.52
AR 4	-6554.98	-6559.83	-6559.92	-6558.94	-6554.1	-6558.16
AR 5	-6558.81	-6558.65	-6557.45	-6555.78	-6558.66	-6556.06

**Minimum Information Criterion** 

<u>Note</u>: Best Model is ARMA(2,4); other potential candidates: ARMA(1,5), ARMA(4,2), ARMA (5,0).

<u>R Note</u>: R has a couple of functions that select automatically the "best" ARIMA model: *armaselect* (using package *caschrono*) minimizes BIC and *auto.arima* (using package *forecast*) minimizes AIC, AICc (default) or BIC.

> armaselect(lr\_p)

# shows the best 10 models according to BIC

p q sbc [1,] 2 0 -11644.79 [2,] 1 0 -11641.53 [3,] 3 0 -11637.71 [4,] 4 0 -11632.43 [5,] 5 0 -11629.95 [6,] 2 1 -11627.42 [7,] 6 0 -11621.70 [8,] 1 3 -11620.18 [9,] 3 1 -11619.93 [10,] 2 2 -11619.44

> auto.arima(lr\_p, ic="bic", trace=TRUE)
function approximates models.

# ic="BIC".

Fitting models using approximations to speed things up...

ARIMA(2,0,2) with non-zero mean : -6519.957ARIMA(0,0,0) with non-zero mean : -6392.599ARIMA(1,0,0) with non-zero mean : -6527.879ARIMA(0,0,1) with non-zero mean : -6536.548ARIMA(0,0,0) with zero mean : -6385.246ARIMA(1,0,1) with non-zero mean : -6529.358ARIMA(0,0,2) with non-zero mean : -6530.806ARIMA(1,0,2) with non-zero mean : -6523.415ARIMA(0,0,1) with zero mean : -6534.284

Now re-fitting the best model(s) without approximations...

ARIMA(0,0,1) with non-zero mean : -6536.463

> auto.arima(lr\_p, ic="bic", max.p=5, max.q = 5, trace=TRUE) # approximates models. Series: lr\_p ARIMA(0,0,1) with non-zero mean Coefficients: mal mean 0.2880 0.0037 s.e. 0.0218 0.0012 sigma^2 estimated as 0.001523: log likelihood=3279.47 AIC=-6552.94 AICc=-6552.93 BIC=-6536.46

Note: The function auto.arima does not try a lot of models, it tries to keep the  $p+q \le 5$ .

<u>Remark</u>: Do not take the results from auto.arima or armaselect or minic as the final model. We still need to check the residuals are WN.

• Script in R to select model using arima function. # set max order for AR part: p-1 p <- 6 # set max order for Ma part: q-1 q <- 6  $npq \leq p^{*}q$ aic  $m \le matrix(0, nrow = npq, ncol=3)$ # matrix collects p, q, AIC: AIC in last column i <- 0 k <- 1 while (j < p) { i <- 0 while (i < q)mod  $j \leq arima(lr p, order=c(i,0,j))$ # fit arima(p,0,q) process aic\_m[k,] <- cbind(i, j, mod j\$aic) # extract aic from arima fit model i < -i + 1

## **ARIMA Model: Identification – Final Remarks**

There is no agreement on which criteria is best. The AIC is the most popular, but others are also used.

Asymptotically, the BIC is consistent –i.e., it selects the true model if, among other assumptions, the true model is among the candidate models considered.

The AIC is not consistent, generally producing too large a model, but is more efficient –i.e., when the true model is not in the candidate model set the AIC asymptotically chooses whichever model minimizes the MSE/MSPE.

## **ARIMA Process – Estimation**

We assume:

- The model order (d, p and q) is known. Make sure yt is I(0).
- The data has zero mean ( $\mu$ =0). If this is not reasonable, demean y .

Fit a zero-mean ARMA model to the demeaned  $y_t$ :

 $\phi(L)(y_t - \bar{y}) = \theta(L)\varepsilon_t$ 

Several ways to estimate an ARMA(p, q) model:

- 1) *Maximun Likelihood Esimation* (MLE). Assume a distribution, usually a normal distribution, and, then, do ML.
- 2) Yule-Walker for ARMA(p,q). Method of moments. Not efficient.
- 3) Innovations algorithm for MA(q).
- 4) Hannan-Rissanen algorithm for ARMA(p, q).

## **ARIMA Process – Estimation: MLE**

#### Steps:

1) Assume a distribution for the errors. Typically, *.i.i.d.* normal, say:

$$\varepsilon_t \sim i.i.d. \ N(0,\sigma^2)$$
  
 $\Rightarrow pdf: f(\varepsilon_t) = \frac{1}{\sqrt{2\pi}} \exp\left[-\frac{\varepsilon_t^2}{2\sigma^2}\right]$ 

2) Write down the joint pdf for  $\boldsymbol{\varepsilon}$ :  $f(\varepsilon_1, ..., \varepsilon_T) = f(\varepsilon_1) ... f(\varepsilon_T)$ 

<u>Note</u>: We are not writing the joint pdf in terms of the  $y_t$ 's, as a multiplication of the marginal pdfs because of the dependency in  $y_t$ .

3) Get  $\varepsilon_t$ . For the general stationary ARMA(p,q) model:

 $\varepsilon_t = y_t - \phi_1 y_{t-1} - \dots - \phi_p y_{t-p} - \theta_1 \varepsilon_{t-1} - \dots - \theta_q \varepsilon_{t-q}$ (if  $\mu \neq 0$ , demean  $y_t$ .)

4) The joint pdf for 
$$\{\varepsilon_1, ..., \varepsilon_T\}$$
 is:  

$$\mathcal{L} = f(\varepsilon_1, \cdots, \varepsilon_T | \mu, \phi, \theta, \sigma^2) = (2\pi\sigma^2)^{-T/2} \exp\left\{-\frac{1}{2\sigma^2} \sum_{t=1}^n \varepsilon_t^2\right\}$$

5) Let  $Y = (y_1, y_2, ..., y_T)$ . With an AR(p, q) model, we need p and q initial lags for  $y_t$  and  $\varepsilon_t$ . We assume that initial conditions  $Y_* = (y_0, y_{-1}, ..., y_{-p+1})^{'}$  and  $\varepsilon_* = (\varepsilon_0, \varepsilon_{-1}, ..., \varepsilon_{-q+1})^{'}$  are known.

6) The conditional log-likelihood function is given by

 $\mathcal{L} = \ln L\left(\mu, \phi, \theta, \sigma^{2}\right) = -\frac{T}{2}\ln(2\pi\sigma^{2}) - \frac{S_{*}(\mu, \phi, \theta)}{2\sigma^{2}}$ 

where  $S_*(\mu, \phi, \theta) = \sum_{t=1}^n \varepsilon_t^2(\mu, \phi, \theta | Y, Y_*, \varepsilon_*)$  is the conditional sum of squares (SS).

<u>Note</u>: Usual Initial conditions:  $y_* = \bar{y}$  and  $\varepsilon_* = E[\varepsilon_t] = 0$ .

• Numerical optimization problem, where initial values (y\*) matter.

**Example**: AR(1) process:

$$y_t = \phi_1 y_{t-1} + \varepsilon_t, \qquad \varepsilon_t \stackrel{i.i.d.}{\sim} N(0, \sigma^2).$$

- Write down the joint likelihood for  $\varepsilon_t$ 

$$\mathcal{L} = f(\varepsilon_1, \cdots, \varepsilon_n) = (2\pi\sigma)^{-n/2} \exp\left\{-\frac{1}{2\sigma^2} \sum_{t=1}^n \varepsilon_t^2\right\}$$

First, we need to solve for  $\varepsilon_t$ :

 $\begin{array}{ll} Y_1 = \phi_1 Y_0 + \varepsilon_1 & \rightarrow \text{Let's take } Y_0 = 0 \\ Y_2 = \phi_1 Y_1 + \varepsilon_2 & \Rightarrow \varepsilon_2 = Y_2 - \phi_1 Y_1 \\ Y_3 = \phi_1 Y_2 + \varepsilon_3 & \Rightarrow \varepsilon_3 = Y_3 - \phi_1 Y_2 \\ \vdots \\ Y_n = \phi_1 Y_{n-1} + \varepsilon_n \Rightarrow \varepsilon_n = Y_n - \phi_1 Y_{n-1} \end{array}$ 

<u>Technical note</u>: The joint likelihood is in terms of  $\varepsilon_t$ . We want to change the joint from  $\varepsilon_t$  to  $\mathbf{y}_t$ , for this, we need the Jacobian  $|\mathbf{J}|$ .

$$|J| = \begin{vmatrix} \frac{\partial \varepsilon_2}{\partial Y_2} & \frac{\partial \varepsilon_2}{\partial Y_3} & \cdots & \frac{\partial \varepsilon_2}{\partial Y_n} \\ \vdots & \vdots & \ddots & \vdots \\ \frac{\partial \varepsilon_n}{\partial Y_2} & \frac{\partial \varepsilon_n}{\partial Y_3} & \cdots & \frac{\partial \varepsilon_n}{\partial Y_n} \end{vmatrix} = \begin{vmatrix} 1 & 0 & \cdots & 0 \\ -\phi_1 & 1 & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & 1 \end{vmatrix} = 1$$

Then,

$$f(Y_2, \dots, Y_n | Y_1) = f(\varepsilon_2, \dots, \varepsilon_n) |J| = f(\varepsilon_2, \dots, \varepsilon_n)$$

- Then, the likelihood function can be written as

$$\begin{split} \mathcal{L}(\phi_1, \sigma_a^2) &= f(Y_1, \cdots, Y_n) = f(Y_1) f(Y_2, \cdots, Y_n | Y_1) = f(Y_1) f(\varepsilon_2, \cdots, \varepsilon_n) \\ &= \left(\frac{1}{2\pi\gamma_0}\right)^{1/2} e^{-\frac{(Y_1 - 0)^2}{2\gamma_0}} \left(\frac{1}{2\pi\sigma^2}\right)^{(T-1)/2} e^{-\frac{1}{2\sigma^2} \sum_{t=2}^{T} (Y_t - \phi_1 Y_{t-1})^2}, \\ &\text{where } Y_1 \sim N\left(0, \gamma_0 = \frac{\sigma^2}{1 - \phi_1^2}\right). \end{split}$$
Then,

Then

$$\mathcal{L}(\phi_1, \sigma^2) = \frac{\sqrt{1 - \phi_1^2}}{\left(2\pi\sigma^2\right)^{n/2}} \exp\left\{-\frac{1}{2\sigma^2} \left[\sum_{t=2}^n (Y_t - \phi_1 Y_{t-1})^2 + (1 - \phi_1^2)Y_1^2\right]\right\}$$

- Then, the log likelihood function:

$$L = ln \mathcal{L}(\phi_{I}, \sigma^{2}) = -\frac{n}{2} ln 2 \pi - \frac{n}{2} ln \sigma^{2} - \frac{1}{2} ln (1 - \phi_{I}^{2}) - \frac{1}{2\sigma^{2}} \underbrace{\left[ \underbrace{\sum_{t=2}^{n} (Y_{t} - \phi_{I}Y_{t-1})^{2}}_{S_{*}(\phi_{I})} + (1 - \phi_{I}^{2})Y_{1}^{2} \right]}_{S(\phi_{I})}$$

where  $S^*(\phi_1)$  is the conditional SS and  $S(\phi_1)$  is the unconditional SS.

• F.o.c.'s:

$$\frac{\partial L(\phi_1,\sigma^2)}{\partial \phi_1} = 0$$
$$\frac{\partial L(\phi_1,\sigma^2)}{\partial \sigma} = 0$$

Note:

- If we neglect  $ln(1-\phi_1^2)$ , then MLE = Conditional LSE.  $\max_{I} L(\phi_{1}, \sigma^{2}) = \min S(\phi_{1}).$ - If we neglect both  $ln(1-\phi_1^2)$  and  $(1-\phi_1^2)Y_1^2$ , then  $\max_{\phi} L(\phi_1, \sigma^2) = \min S(\phi_1) \cdot \P$ 

## **ARIMA Process – Estimation: Yule-Walker**

Yule-Walker for AR(p): Regress yt against yt-1, yt-2,..., yt-p Yule-Walker for ARMA(p, q): Method of moments. Not efficient.

**Example**: For an AR(*p*), we the Yule-Walker equations are

Γ	$\rho(0)$	$\rho(1)$	•••	$\rho(p-1)$	<b>[</b> \$\phi_1]		$\rho(1)$	I
I	$\rho(1)$	ho(0)	•••	$\rho(p-2)$	$\phi_2$	_	$\rho(2)$	
I	:	:	•••	÷	1	_	:	
L	$\rho(p-1)$	$\rho(p-2)$	•••	$\rho(0)$	$\left\lfloor \phi_{p} \right\rfloor$		$\rho(p)$	

Method of Moments (MM) Estimation: Equate sample moments to population moments, and solve the equation. In this case, we use:

$$E(Y_t) = \frac{1}{T} \sum_{t=1}^{T} Y_t \Rightarrow \mu = \bar{Y}$$
  

$$E[(Y_t - \mu)(Y_{t-k} - \mu)) = \frac{1}{T} \sum_{t=1}^{T} (Y_t - \mu)(Y_{t-k} - \mu)) \Rightarrow \gamma_k = \hat{\gamma}_k \quad (\& \rho_k = \hat{\rho}_k)$$

• Then, the Yule-Walker estimator for  $\phi$  is given by solving

$$\begin{bmatrix} 1 & \hat{\rho}(1) & \cdots & \hat{\rho}(p-1) \\ \hat{\rho}(1) & 1 & \cdots & \hat{\rho}(p-2) \\ \vdots & \vdots & \cdots & \vdots \\ \hat{\rho}(p-1) & \hat{\rho}(p-2) & \cdots & 1 \end{bmatrix} \begin{bmatrix} \hat{\phi}_1 \\ \hat{\phi}_2 \\ \vdots \\ \phi_p \end{bmatrix} = \begin{bmatrix} \hat{\rho}(1) \\ \hat{\rho}(2) \\ \vdots \\ \hat{\rho}(p) \end{bmatrix}$$
$$\Rightarrow \hat{\phi} = \hat{R}_p^{-1} \hat{\rho}_p \P$$

<u>Note</u>: If  $\hat{\gamma}_0 > 0$ , then,  $\hat{\Gamma}_m$  is nonsingular.

• If {*Y*<sub>t</sub>} is an AR(*p*) process,  

$$\hat{\phi} \xrightarrow{d} N\left(\phi, \frac{\sigma^2}{T}\Gamma_p^{-1}\right)$$

$$\hat{\phi}_{kk} \xrightarrow{d} N\left(0, \frac{1}{T}\right) \text{ for } k > p$$
The set of the part of th

• Thus, we can use the sample PACF to test for AR order, and we can calculate approximated C.I. for  $\phi$ .

• Distribution:

If  $\overline{y_t}$  is an AR(p) process, and T is large,  $\sqrt{T}(\hat{\phi} - \phi) \stackrel{approx.}{\sim} N(0, \hat{\sigma}^2 \hat{\Gamma}_p^{-1})$ 

100(1- $\alpha$ )% approximate C.I. for  $\phi_j$  is  $\hat{\phi}_j \pm z_{\alpha/2} \frac{\hat{\sigma}}{\sqrt{T}} (\hat{\Gamma}_p^{-1})_{jj}^{1/2}$ 

<u>Note</u>: The Yule-Walker algorithm requires  $\Gamma^{-1}$ .

• For AR(p). The Levinson-Durbin (LD) algorithm avoids  $\Gamma^{-1}$ . It is a recursive linear algebra prediction algorithm. It takes advantage that  $\Gamma$  is a symmetric matrix, with a constant diagonal (Toeplitz matrix). Use LD replacing  $\gamma$  with  $\hat{\gamma}$ .

Side effect of LD: automatic calculation of PACF and MSPE.

**Example 1**: AR(1) (MM) estimation:  $y_t = \phi_1 y_{t-1} + \varepsilon_t$ It is known that  $\rho_l = \phi_1$ . Then, the MME of  $\phi_1$  is • Also,  $\sigma^2$  is unknown:  $\gamma_0 = \frac{\sigma^2}{(1-\phi_1^2)} \Rightarrow \hat{\sigma}^2 = \hat{\gamma}_0 \left(1 - \widehat{\phi_1}^2\right). \P$  **Example 2**: Suppose we suspect an AR(3). We have estimated  $\hat{\rho}_1$ ,  $\hat{\rho}_2$ , and  $\hat{\rho}_3$ . Then,

Example 2. Suppose we suspect an AA(3): we have estimated  $p_1, p_2, a$   $\begin{bmatrix} 1 & \hat{p}_1 & \hat{p}_2 \\ \hat{p}_1 & 1 & \hat{p}_1 \\ \hat{p}_2 & \hat{p}_1 & 1 \end{bmatrix} \begin{bmatrix} \phi_1 \\ \phi_2 \\ \phi_3 \end{bmatrix} = \begin{bmatrix} \hat{p}_1 \\ \hat{p}_2 \\ \hat{p}_3 \end{bmatrix}$ Suppose we get:  $\hat{p}_1 = 0.5, \hat{p}_2 = 0.4$ , and  $\hat{p}_3 = -0.3$ . Then, solving for  $\boldsymbol{\phi}$ :  $\begin{bmatrix} \hat{\phi}_1 \\ \hat{\phi}_2 \\ \hat{\phi}_2 \end{bmatrix} = \begin{bmatrix} 1 & 0.5 & 0.4 \\ 0.5 & 1 & 0.5 \\ 0.4 & 0.5 & 1 \end{bmatrix}^{-1} \begin{bmatrix} 0.5 \\ 0.4 \\ -0.3 \end{bmatrix} = \begin{bmatrix} 0.555 \\ 0.511 \\ -0.777 \end{bmatrix}$ 

• Solving system with R: Rho <- matrix(c(1, 0.5, 0.4, 0.5, 1, 0.5, 0.4, 0.5, 1), nrow=3)  $r \le c(.5, 0.4, -0.3)$ solve(Rho)%\*%r.¶

**Example**: MA(1) process with MM estimation:

 $y_t = \varepsilon_t - \theta_1 \varepsilon_{t-1}$ Again using the autocorrelation of the series at lag 1,

$$\rho_{1} = -\frac{\theta_{1}}{(1+\theta_{1}^{2})} = \hat{\rho}_{1}$$
  
$$\theta_{1}^{2} \hat{\rho}_{1} + \theta_{1} + \hat{\rho}_{1} = 0$$
  
$$\hat{\theta}_{1,2} = \frac{-1\pm\sqrt{1-4\hat{\rho}_{1}^{2}}}{2\hat{\rho}_{1}}$$

• Choose the root satisfying the invertibility condition. For real roots:  $1 - 4\hat{\rho}_1^2 \ge 0 \implies 0.25 \ge \hat{\rho}_1^2 \implies -0.5 \le \hat{\rho}_1 \le 0.5$ 

If  $\hat{\rho}_1 = \pm 0.5$ , unique real roots but non-invertible.

If  $|\hat{\rho}_1| < 0.5$ , unique real roots and invertible.  $\Rightarrow$  We keep this one.

• Remarks

- The MMEs for MA and ARMA models are complicated.

- In general, regardless of AR, MA or ARMA models, the MMEs are sensitive to rounding errors. They are usually used to provide initial estimates needed for a more efficient nonlinear estimation method.

- The moment estimators are not recommended for final estimation results and should not be used if the process is close to being nonstationary or noninvertible.

## **ARIMA Process – Estimation: Yule-Walker – Remarks**

The MM estimations for MA and ARMA models are complicated.

In general, regardless of AR, MA or ARMA models, the MMEs are sensitive to rounding errors.

They are usually used to provide initial estimates needed for a more efficient nonlinear estimation method.

The moment estimators are not recommended for final estimation results and should not be used if the process is close to being nonstationary or noninvertible.

### **ARIMA Process – Estimation: Hannan-Rissanen**

Hannan-Rissanen algorithm for ARMA(p,q)

Steps:

Estimate high-order AR.
 Use Step (1) to estimate (unobserved) noise ε<sub>t</sub>
 Regress y<sub>t</sub> against y<sub>t-1</sub>, y<sub>t-2</sub>, ..., y<sub>t-p</sub>, ê<sub>t-1</sub>, ..., ê<sub>t-q</sub>

4. Get new estimates of  $\varepsilon_t$ . Repeat Step (3).

Example: We estimate a ARIMA(0,0,1) model for S&P 500 historical returns, using the arima
function, part of the R forecast package.
> arima(lr\_p, order=c(0,0,1), method="ML") #ML estimation method
Call:
arima(x = lr\_p, order = c(0, 0, 1), method = "ML")
Coefficients:
 mal intercept
 0.2880 0.0037
s.e. 0.0218 0.0012
sigma^2 estimated as 0.001522: log likelihood = 3279.47, aic = -6552.94. ¶

<u>Note</u>: Model was selected by ACF/PACF and confirmed with *auto.arima* function. Not a lot of structure in stock returns.

**Example**: We use auto.arima function to estimate a model for **DIS**, **GE**, and **IBM** returns.

```
> auto.arima(lr_dis)
Coefficients:
    ar1 mean
    0.0538 0.0072
s.e. 0.0419 0.0038
sigma^2 estimated as 0.007462: log likelihood=588.13
AIC=-1170.25 AICc=-1170.21 BIC=-1157.22
> auto.arima(lr_ge)
Coefficients:
    ar1 ma1
    0.0592 -0.9848
s.e. 0.0428 0.0096
sigma^2 estimated as 0.005591: log likelihood=667.5
```

Note: For both **DIS** & **GE returns**, we observe low AR(1) coefficient, and not significant.

> auto.arima(lr\_ibm)
Series: lr\_ibm
ARIMA(0,0,0) with zero mean
sigma^2 estimated as 0.005126: log likelihood=694.13
AIC=-1386.26 AICc=-1386.25 BIC=-1381.91
sigma^2 estimated as 0.001522: log likelihood = 3279.47, aic = -6552.94.

<u>Note</u>: Unpredictable! In general, we do not find a lot of structure in stock returns; autocorrelations die out very quickly. This result is expected, given the Efficient Markets Hypothesis. ¶

Example: We use auto.arima function to estimate a model for changes in oil prices.
> auto.arima(lr\_oil)
Series: lr\_oil
ARIMA(4,0,0) with zero mean
Coefficients:
 ar1 ar2 ar3 ar4
 0.2950 -0.1024 -0.0570 -0.0984
s.e. 0.0521 0.0543 0.0551 0.0539

sigma<sup>2</sup> estimated as 0.008913: log likelihood=344.52 AIC=-679.04 AICc=-678.87 BIC=-659.55

<u>Note</u>: AR(4)  $\Rightarrow$  significant autocorrelation in changes in oil prices, but mainly decaying at .30.

Example: We use auto.arima function to estimate a model for monthly U.S. interest long rates (1871 – 2020). > auto.arima(x\_i) Series: x\_i ARIMA(0,1,2) Coefficients: ma1 ma2 0.4012 -0.0957 s.e. 0.0236 0.0238 sigma^2 estimated as 0.02719: log likelihood=690.02 AIC=-1374.04 AICc=-1374.03 BIC=-1357.56

Note: We need to differentiate interest rates to get a stationary MA(2) model.

## **ARIMA Process – Diagnostic Tests**

Once the model is estimated, we run diagnostic tests. Usually, we check for extra-AR structure in the mean. We check visual plots of residuals, ACFs, and the distribution of residuals. More formally, we compute the LB test on the residuals. If we find extra-AR structure, we increase p and/or q.

<u>**R** Note</u>: If we use *arima()* or *auto.arima()* functions, we can use the function *checkresiduals()* to do the plots and testing for us.

**Example:** We check the MA(1) model for U.S. historical stock returns > fit\_arima\_lr\_p <- arima(lr\_p, order=c(0,0,1), method="ML") > checkresiduals(fit\_arima\_lr\_p)

Ljung-Box test

data: Residuals from ARIMA(0,0,1) with non-zero mean  $Q^* = 18.579$ , df = 8, p-value = 0.01728  $\Rightarrow$  There seems to be more AR structure





<u>R Note</u>: We check stationarity/invertibility too -i.e., if the roots are inside the unit circle. In this case, an MA model, stationarity is not an issue (MA are stationary), but invertibility is. We use the R function *autoplot*, part of the forecast package. Be aware that autoplot plots the inverse roots, not the roots; in this case, a stationary AR (or invertible MA) process will have the inverse roots inside the unit circle.

> autoplot(fit\_arima\_lr\_p)



<u>Note</u>: The inverse root is inside the unit circle and are real: invertible MA(1).  $\P$ 

**Example:** We change the model for **U.S. stock returns**. We estimate an ARIMA(1,0,5). > fit\_arima\_lr\_p15 <- arima(lr\_p, order=c(1,0,5)) > fit\_arima\_lr\_p15

Coefficients:

ar1ma1ma2ma3ma4ma5intercept0.7077-0.4071-0.1965-0.06710.03380.08070.0035s.e.0.10390.10580.03920.02630.02560.02500.0014

sigma<sup>2</sup> estimated as 0.001502: log likelihood = 3278.2, aic = -6540.4

> checkresiduals(fit\_arima\_lr\_p15)
 Ljung-Box test

data: Residuals from ARIMA(1,0,5) with non-zero mean  $Q^* = 1.7047$ , df = 3, p-value = 0.6359  $\Rightarrow$  The joint 10 lag autocorrelation not significant.

Model df: 7. Total lags used: 10



Note: We still see some small autocorrelations different from 0.

We check the stationarity and invertibility of ARIMA(1,0,5) model > autoplot(fit\_arima\_lr\_p15)



<u>Note</u>: All inverse roots inside the unit circle: stationary and invertible. Notice that we have some roots on the MA part that are imaginary. ¶

**Example:** We check the fit of the ARIMA model for **U.S. long interest rates** > fit\_arima\_i <- auto.arima(x\_i) ARIMA(0,1,2) Coefficients:

ma1 ma2 0.4012 -0.0957 s.e. 0.0236 0.0238

sigma<sup>2</sup> estimated as 0.02719: log likelihood=690.02 AIC=-1374.04 AICc=-1374.03 BIC=-1357.56

> checkresiduals(fit\_arima\_i)

Ljung-Box test data: Residuals from ARIMA(0,1,2)  $Q^* = 34.029$ , df = 8, p-value = 4.014e-05  $\Rightarrow$  Again, more AR or MA structure needed





<u>Note</u>: We still see some large autocorrelations.  $\Rightarrow$  change model (usually, increase *p* and/or *q*). But, we may in the presence of a series with regime change. We may need to focus on 2<sup>nd</sup> regime (post 1950s).

We check the invertibility of ARIMA(0,1,2) model > autoplot(fit\_arima\_i)



<u>Note</u>: All inverse roots are inside the unit circle. MA process is invertible. Notice that all roots are real.  $\P$ 

**Example:** We check the fit of the ARIMA(4,0,0) model selected by auto.arima for changes in **Oil Prices.** 

fit\_arima\_oil<- auto.arima(lr\_oil) > fit\_arima\_oil Series: lr\_oil ARIMA(4,0,0) with zero mean

Coefficients: ar1 ar2 ar3 ar4 0.295 -0.102 -0.057 -0.098 s.e. 0.052 0.054 0.055 0.054

sigma<sup>2</sup> estimated as 0.00891: log likelihood=344.52 AIC=-679.04 AICc=-678.87 BIC=-659.55

> checkresiduals(fit\_arima\_oil)

Ljung-Box test

data: Residuals from ARIMA(4,0,0) with zero mean  $Q^* = 2.72$ , df = 9, p-value = 0.84  $\Rightarrow$  No significant joint AR structure

Model df: 4. Total lags used: 10



Note: Nothing significant. Happy with fit. Ready to forecast.

We check the stationarity of AR(4) model > autoplot(fit arima oil)



Note: All (inverse) roots inside the unit circle -we have imaginary roots.

## Non-Stationarity in Variance

Stationarity in mean does not imply stationarity in variance. However, non-stationarity in mean implies non-stationarity in variance.

If the mean function is time dependent:

- 1. The variance,  $Var(y_t)$  is time dependent.
- 2. Var[y<sub>t</sub>] is unbounded as  $t \rightarrow \infty$ .

3. Autocovariance functions and ACFs are also time dependent.

4. If *t* is large with respect to the initial value  $y_0$ , then  $\rho_k \approx 1$ .

• It is common to use *variance stabilizing* transformations: Find a function G(.) so that the transformed series  $G(y_t)$  has a constant variance. Very popular transformation:

#### 1) Log transformation:

$$G(Y_t) = \log(Y_t)$$

**Example**: We log transform the monthly variable Total U.S. Vehicle Sales data (1976: Jan – 2020: Sep):

ts\_car <- ts(x\_car,start=c(1976,1),frequency=12) plot.ts(ts\_car,xlab="Time",ylab="div", main="Total U.S. Vehicle Sales")



 $l_car <- \log(ts_car)$ 

> plot.ts(l\_car,xlab="Time",ylab="div", main="Log Total U.S. Vehicle Sales")library(tseries)



<u>Note</u>: The volatility is significantly reduced by the log transformation.  $\P$ 

## 2) Box-Cox transformation:

$$G(Y_t) = \frac{Y_t^{\lambda} - 1}{\lambda}$$

where  $\lambda > 0$ , usually between 0 and 2 (it can be estimated too). When  $\lambda=1$ , we have a linear  $y_t$ ; when  $\lambda \to 0$ , we have a log transformation for  $y_t$ .

**Example**: We do a Box-Cox transformation of the monthly variable Total U.S. Vehicle Sales data (1976: Jan – 2020: Sep), setting  $\lambda = 0.75$ : lambda <- 0.75 b\_cox\_car <- (ts\_car^lambda - 1)/lambda > plot.ts(b\_cox\_car, xlab="Time",ylab="cars", main=" Box-Cox Total U.S. Vehicle Sales")



<u>Note</u>: Again, we see a reduced volatility. But, different  $\lambda$ s will have a different impact on volatility. ¶

Remarks:

- Variance stabilizing transformation is only done for positive series, usually for nominal series (say, in USD total retail sales or units, like Total U.S. vehicle sales).

- If a series has negative values, then, we need to add each value with a positive number so that all the values in the series are positive.

- Then, we can search for any need for transformation.

- It should be performed before any other analysis, such as differencing.

- Not only stabilize the variance, but we tend to find that it also improves the approximation of the distribution by the Normal distribution.

## **Seasonal Time Series**

In time series, seasonal patterns ("seasonalities") can show up in two forms: additive and multiplicative.

- Additive: The seasonal variation is independent of the level.

- Multiplicative: The seasonal variation is a function of the level.



<u>Note</u>: In the multiplicative case, the amplitude of the seasonal pattern is changing over time, while in the additive the amplitude is constant.

**Examples:** We simulate the two seasonal patterns, additive and multiplicative, with trend and no trend.

A. With trend


B. With no trend



• In the presence of seasonal patterns, we proceed to do seasonal adjustments to remove these predictable influences, which can blur both the true underlying movement in the series, as well as certain non-seasonal characteristics which may be of interest to analysts.

The type of adjustment depends on how we view the seasonal pattern: Deterministic or Stochastic.

Similar to the situation where the series had a trend, once we determine the nature of the seasonal pattern, we filter the series –i.e., we remove the seasonal patter- to conduct further ARIMA modeling.

When we work with a nominal series (not changes, say, USD total retail sales or total units sold), it is common to first apply a variance stabilizing transformation to the data, usually using logs.

#### **Seasonal Time Series – Types**

Two types of seasonal behavior: - **Deterministic** – Usual treatment: Build a deterministic function,  $f(t) = f(t + k \times s), \qquad k = 0, \pm 1, \pm 2, \cdots$ 

We can include seasonal (means) dummies, for example, monthly or quarterly dummies. (This is the approach in Brooks' Chapter 10).

Instead of dummies, trigonometric functions (sum of cosine curves) can be used. A linear time trend is often included in both cases.

-Stochastic – Usual treatment: SARIMA model. For example:

$$y_t = \theta_0 + \varphi_1 \, y_{t-s} + \, \varepsilon_t + \Theta_1 \varepsilon_{t-s}$$

or

$$(1 - \Phi_1 L^s) y_t = (1 - \Theta_1 L^s) \varepsilon_t$$

where *s* the seasonal periodicity –associated with the frequency– of  $y_t$ . For quarterly data, s = 4; monthly, s = 12; daily, s = 7, etc.

# Seasonal Time Series – Finding Seasonality with Visual Patterns

The raw series along with the ACF and PACF can be used to discover seasonal patterns.



<u>Signs</u>: Periodic repetitive wave pattern in ACF, repetition of significant ACFs, PACFs after *s* periods.

• We simulate an ARMA(1,1) with a December seasonal pattern, typical of retail sales with a significant Christmas spike.





Suppose  $y_t$  has monthly frequency and we suspect that in every December  $y_t$  increases. – For the additive model, we can regress  $y_t$  against a constant and a December dummy,  $\mathbf{D}_t$ :  $y_t = \mu + \mathbf{D}_t \boldsymbol{\mu}_s + \varepsilon_t$ 

For the multiplicative model, we can regress  $y_t$  against a constant and a December dummy,  $\mathbf{D}_t$ :, interacting with a trend:

$$y_t = \mu + \boldsymbol{D}_t \boldsymbol{\mu}_s * t + \varepsilon_t$$

The residuals of this regressions,  $e_t$ , –i.e.,  $e_t$  = filtered  $y_t$ , free of "monthly seasonal effects"– are used for further ARMA modeling.

**Example:** We simulate an AR(1) series, with a multiplicative December seasonal behavior.

 $y_t = \mu + \phi_1 y_{t-1} + \boldsymbol{D}_t \boldsymbol{\mu}_s * t + \varepsilon_t$ Seas  $12 \le \operatorname{rep}(c(0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 1), (\operatorname{length}(y \ sim)/12+1))$ # Create Oct dummy T sim <- 500 u <- rnorm(T sim, sd=0.75) # Draw T sim normally distributed errors  $y_sim \le matrix(0, T_sim, 1)$ # vector to accumulate simulated data phi1 <- 0.2 # Change to create different correlation patterns k <- 12 # Seasonal Periodicity a <- k+1 # Time index for observations mu <- 0.2 mu s <- .02 while (a  $\leq T \sin$ ) {  $y \sin[a] = mu + phi1 * y \sin[a-1] + Seas 12[a] * mu s * a + u[a]$ #y sim simulated autocorrelated values a <- a + 1 } y seas  $\leq y sim[(k+1):T sim]$ plot(y seas, type="l", main="Simulated Deterministic Seasonality")

We plot simulated series, ACF, & PACF.



Residual standard error: 0.8209 on 484 degrees of freedom Multiple R-squared: 0.7929, Adjusted R-squared: 0.7917 F-statistic: 617.8 on 3 and 484 DF, p-value: < 2.2e-16

• We plot the detrended simulated series, along with the ACF and PACF.



The strong December seasonal pattern is gone from the detrended series. We run an ARIMA(1,0,0):

> fit\_y\_seas\_ar1 <- arima(y\_seas\_filt, order=c(1,0,0))
Call:
arima(x = y\_seas\_filt, order = c(1, 0, 0))</pre>

Coefficients: ar1 intercept 0.1785 -0.0001  $\Rightarrow$  Very close to phi1 = 0.20 s.e. 0.0446 0.0443

sigma<sup>2</sup> estimated as 0.6471: log likelihood = -586.26, aic = 1178.51

y seas filt 2 <- fit seas det ar1\$residuals # Extract Residuals

plot(y\_seas\_filt\_2,type="l", main="AR(1) Residuals") acf(y\_seas\_filt\_2, main="ACF: AR(1) Residuals") pacf(y\_seas\_filt\_2, main="ACF: AR(1) Residuals")



There is no seasonality pattern in the residuals.

**Example:** We model **log changes in real estate prices in the LA market**,  $y_t$ . First, we run a regression to remove (*filter*) the monthly effects from  $y_t$ . Then, we model  $y_t$  as an ARMA(p, q) process.

RE\_da <- read.csv(" https://www.bauer.uh.edu/rsusmel/4397/Real\_Estate\_2019.csv", head=TRUE, sep=",") x\_la <- RE\_da\$LA\_c zz <- x\_la T <- length(zz) plot(x\_la, type="l", main="Changes in Log Real Estate Prices in LA")



#### Changes in Log Real Estate Prices in LA

We look at the ACF & PACF for LA > acf(x\_la) > pacf(x\_la)



<u>Note</u>: ACF shows highly autocorrelated data, with some seasonal pattern (there is a periodic decreasing wave).

• We define monthly dummies. Then, we regress x la against the monthly dummies. # Create January dummy  $Feb1 \le rep(c(1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0), (length(zz)/12+1))$ Mar1 <- rep(c(0, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0), (length(zz)/12+1))# Create March dummy Apr1 <- rep(c(0, 0, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0), (length(zz)/12+1)) # Create April dummy # Create May dummy May1 <- rep(c(0, 0, 0, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0), (length(zz)/12+1))Jun1 <- rep(c(0, 0, 0, 0, 1, 0, 0, 0, 0, 0, 0, 0), (length(zz)/12+1)) # Create June dummy  $Jul1 \le rep(c(0, 0, 0, 0, 0, 1, 0, 0, 0, 0, 0, 0, 0), (length(zz)/12+1))$ # Create Jul dummy Aug1 <- rep(c(0, 0, 0, 0, 0, 0, 1, 0, 0, 0, 0, 0), (length(zz)/12+1)) # Create Aug dummy # Create Sep dummy Sep1  $\leq$  rep(c(0, 0, 0, 0, 0, 0, 0, 1, 0, 0, 0, 0), (length(zz)/12+1))  $Oct1 \le rep(c(0, 0, 0, 0, 0, 0, 0, 0, 1, 0, 0, 0), (length(zz)/12+1))$ # Create Oct dummy Nov1 <- rep(c(0, 0, 0, 0, 0, 0, 0, 0, 0, 1, 0, 0), (length(zz)/12+1))# Create Oct dummy  $Dec1 \le rep(c(0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 1, 0), (length(zz)/12+1))$ # Create Oct dummy seas1 <- cbind(Feb1, Mar1, Apr1, May1, Jun1, Jul1, Aug1, Sep1, Oct1, Nov1, Dec1) seas  $\leq$  seas1[1:T,] x la fit sea  $\leq - lm(x la \sim seas)$ # Regress x la against constant + seasonal dummies > summary(x la fit sea)

Coefficients:

	Estim	ate Std.	Error t value P	r(> t )
(Intercept)	-0.0014063	0.0020125	-0.699 0.48515	7
seasFeb1	0.0006752	0.0028223	0.239 0.81107	9
seasMar1	0.0049095	0.0028223	1.740 0.08283	8.
seasApr1	0.0090903	0.0028223	<b>3.221</b> 0.00140	0 **
seasMay1	0.0104159	0.0028223	<b>3.691</b> 0.00026	0 ***

seasJun1	0.0103464	0.0028223	<b>3.666</b> 0.000285 <sup>3</sup>	***
seasJul1	0.0080593	0.0028223	<b>2.856</b> 0.004557 <sup>3</sup>	**
seasAug1	0.0062247	0.0028223	2.206 0.028080 <sup>3</sup>	*
seasSep1	0.0032244	0.0028223	1.142 0.254055	
seasOct1	0.0011967	0.0028461	0.420 0.674421	
seasNov1	-0.0006218	0.0028461	-0.218 0.827181	
seasDec1	-0.0009031	0.0028461	-0.317 0.751195	
Signif. codes:	0 '***' 0.001	<b>***</b> 0.01 <b>**</b>	0.05 '.' 0.1 ' ' 1	

Note: Returns -i.e., home prices- are higher from April to August.

Now, we model et, the filtered LA series

x_la_filt <- x_la_fit_sea\$residuals	# residuals, $e_t = filtered x_la series$
fit_ar_la_filt <- auto.arima(x_la_filt)	# use auto.arima to look for a good model
> fit_ar_la_filt	

Series: x\_la\_filt ARIMA(2,0,1) with zero mean

Coefficients: ar1 ar2 ma1 0.0987 0.7737 0.7245 s.e. 0.0963 0.0866 0.1136

sigma<sup>2</sup> estimated as 1.668e-05: log likelihood=1453.66 AIC=-2899.33 AICc=-2899.21 BIC=-2883.83

> checkresiduals(fit ar la filt)

Ljung-Box test

data: Residuals from ARIMA(2,0,1) with zero mean  $Q^* = 13.5$ , df = 7, p-value = 0.06083  $\Rightarrow$  Reject H<sub>0</sub> at 5% lever. But, judgement call is OK.

Model df: 3. Total lags used: 10

We check residual plots.



<u>Note</u>: ACF shows some small, but significant autocorrelations, but the seasonal (wave) pattern is no longer there.

Finally, we check the stationarity & the invertibility of the ARIMA(2,0,1) process.



Note: All inverse roots inside the unit circle (& real): stationarity and invertibility.

# **Seasonal Time Series – SARIMA**

For stochastic seasonality, we use the Seasonal ARIMA model. In general, we have the SARIMA(P, D, Q)<sub>s</sub>:

$$\Phi_P(L^s)(1-L^s)^D y_t = \theta_0 + \Theta_Q(L^s)\varepsilon_t$$

where  $\theta_0$  is constant and

$$\Phi_P(L^s) = 1 - \Phi_1 L^s - \Phi_2 L^{2s} - \dots - \Phi_P L^{sP} \\ \Theta_O(L^s) = 1 + \Theta_1 L^s + \Theta_2 L^{2s} + \dots + \Theta_O L^{sQ}$$

**Example 1**: SARIMA(0,0,1)<sub>12</sub>= SMA(1)<sub>12</sub>  $y_t = \theta_0 + \varepsilon_t + \Theta_1 \varepsilon_{t-12}$ 

- Invertibility Condition:  $|\Theta_1| < 1$ .

$$- \operatorname{E}[y_t] = \theta_0.$$
  

$$- \operatorname{Var}(y_t) = (1 + \theta_1^2)\sigma^2$$
  

$$- \operatorname{ACF}: \rho_k = \begin{cases} \frac{\theta_1}{1 + \theta_1^2}, & |k| = 12\\ 0, & \text{otherwise} \end{cases} \Rightarrow \operatorname{ACF} \text{ non-zero at seasonal lags } 12, 24,...$$

Example 2: SARIMA(1,0,0)<sub>12</sub> = SAR(1)<sub>12</sub> (1 -  $\Phi_1 L^{12}$ ) $y_t = \theta_0 + \varepsilon_t$ The process is

The process is

$$y_t = \theta_0 + \Phi_1 y_{t-12} + \varepsilon_t$$

- This is a simple seasonal AR model.

- Stationarity Condition: 
$$|\Phi_1| < 1$$
.  
 $- E[Y_t] = \frac{\theta_0}{1 - \Phi_1}$   
 $- Var(Y_t) = \frac{\sigma^2}{1 - {\Phi_1}^2}$   
 $- ACF: \ \rho_{12k} = {\Phi_1}^k, \qquad k = 0, \ \pm 1, \ \pm 2, \cdots$   
When  $\Phi_1 = 1$ , the series is non-stationary. ¶

• Now, we put together the seasonal behavior and the ARMA behavior. That is, we have the multiplicative SARIMA model  $(p,d,q) \ge (P,D,Q)_s$ 

**Example 1**: ARIMA(0,0,1) x (0,0,1)<sub>12</sub> (usually, with monthly data):  $y_t = (1 + \theta_1 L)(1 + \Theta L^{12})\varepsilon_t$ Then, the process is  $y_t = \varepsilon_t + \theta_1 \varepsilon_{t-1} + \varepsilon_{t-12} + \theta_1 \Theta \varepsilon_{t-12}$ . ¶

**Example 2**: Suppose p = Q = 1 and P = q = 0, with s=4, then, we have an ARIMA(1,0,0) x (0,0,1)<sub>4</sub> (usually, with quarterly data):

 $(1-\phi_1 L) y_t = (1+\Theta L^4)\varepsilon_t$ 

Then, the process is

 $y_t = \phi_1 y_{t-1} + \varepsilon_t + \Theta \varepsilon_{t-4}. \P$ 

In general, we the multiplicative SARIMA model  $(p,d,q) \ge (P,D,Q)_s$  is written as:  $\Phi(L)\phi(L)y_t = \theta(L)\Theta(L)\varepsilon_t$ 

where  $\phi(L)$  is the AR lag polynomial,  $\theta(L)$  is the MA lag polynomial,  $\phi(L)$  is the seasonal AR lag polynomial, and  $\theta(L)$  is the seasonal MA lag polynomial.

**Example**: We model with a SARIMA model for **U.S. vehicle sales**. First, we look at the raw data:

Car\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/TOTALNSA.csv", head=TRUE, sep=",")





Note: ACF shows a highly autocorrelated data, with some clear seasonal wave pattern.

• Then, we plot the data and, then, log transform the data: ts\_car <- ts(x\_car,start=c(1976,1),frequency=12) plot.ts(ts\_car, xlab="Time", ylab="div", main="Total U.S. Vehicle Sales")



l\_car <- log(ts\_car)
> plot.ts(l\_car,xlab="Time",ylab="div", main="Log Total U.S. Vehicle Sales")library(tseries)



<u>R Note</u>: R has a function, *decompose*, that decomposes the data in trend, seasonal and random (unexplained): comp\_lcar <- decompose(l\_car) > plot(comp\_lcar)



Decomposition of additive time series

• Question: Should we try deterministic seasonalities?

No clear trend in data. We regress  $l_car$  against monthly dummies:  $zz \le 1$  car

seasd <- cbind(Jan1, Feb1, Mar1, Apr1, May1, Jun1, Jul1, Aug1, Sep1, Oct1, Nov1)

seas\_d <- seasd[1:length(zz),]</pre>

 $fit\_car\_det <- lm(l\_car ~ seas\_d)$ 

> summary(fit\_car\_det)

Coefficients:

```
EstimateStd. Error t value Pr(>|t|)
(Intercept)
            7.03020 0.02564 274.171 < 2e-16 ***
seas dJan1
            0.08235 0.03626 2.271 0.023551 *
seas dFeb1
            -0.09854 0.03606 -2.733 0.006494 **
seas dMar1 0.01462 0.03606 0.406 0.685259
seas dApr1 0.19884 0.03606 5.514 5.51e-08 ***
seas dMay1 0.11396 0.03606 3.160 0.001668 **
seas dJun1 0.20192 0.03606 5.599 3.47e-08 ***
seas dJul1
            0.17824 0.03606 4.943 1.04e-06 ***
seas dAug1 0.12804 0.03606 3.551 0.000419 ***
seas dSep1 0.14824 0.03606 4.111 4.57e-05 ***
seas dOct1
            0.06599 0.03606 1.830 0.067813.
seas dNov1 0.07014 0.03626 1.934 0.053638.
```

• Check ACF and PACF

res\_car\_det <- fit\_car\_det\$residuals
acf(res\_car\_det)
pacf(res\_car\_det)</pre>





• Now, we use auto.arima to check for best SARIMA model:

> fit\_lcar <- auto.arima(l\_car, trace=TRUE, ic="bic")</pre>

Fitting models using approximations to speed things up...

ARIMA(2,0,2)(1,1,1)[12] with drift	: -1049.585
ARIMA(0,0,0)(0,1,0)[12] with drift	: -609.8308
ARIMA(1,0,0)(1,1,0)[12] with drift	: -928.3348
ARIMA(0,0,1)(0,1,1)[12] with drift	: -780.978
ARIMA(2,0,2)(0,1,2)[12] with drift	: -1072.605
ARIMA(2,0,2)(1,1,2)[12] with drift	: -1055.059
ARIMA(1,0,2)(0,1,2)[12] with drift	: -1080.563
ARIMA(0,0,2)(0,1,2)[12] with drift	: -905.0785
ARIMA(1,0,1)(0,1,2)[12] with drift	: -1081.598
Now re-fitting the best model(s) withou	t approximations.

ARIMA(1,0,1)(0,1,2)[12] :-1132.208

Best model: ARIMA(1,0,1)(0,1,2)[12]

• Check estimated best SARIMA model and check its residuals: > fit\_lcar Series: 1\_car ARIMA(1,0,1)(0,1,2)[12]

Coefficients: ar1 ma1 sma1 sma2 **0.9539 -0.5113 -0.5921 -0.2099** s.e. 0.0163 0.0509 0.0464 0.0442

sigma<sup>2</sup> estimated as 0.006296: log likelihood=581.76 AIC=-1153.52 AICc=-1153.41 BIC=-1132.21

> checkresiduals(fit\_lcar)

Ljung-Box test

data: Residuals from ARIMA(1,0,1)(0,1,2)[12]  $Q^* = 44.006$ , df = 20, p-value = 0.001502

Model df: 4. Total lags used: 24





<u>Note</u>: ACF shows small and significant autocorrelation, but the seasonal pattern is gone. More lags maybe needed. ¶

#### **Forecasting**

One of the most important objectives in time series analysis is to forecast its future values. It is the primary objective of ARIMA modeling.

Two types of forecasts.

- In sample (prediction): The expected value of the RV (in-sample), given the estimates of the parameters.

- Out of sample (forecasting): The value of a future RV that is not observed by the sample.

To evaluate forecasts, we can use in-sample estimation to learn about the order of the ARMA(p,q) model and then use the model to forecast. We do the in-sample estimation keeping a hold-out sample. We use the hold-out sample to validate the selected ARMA model.

Any forecasts needs an information set,  $I_T$ . This includes data, models and/or assumptions available at time *T*. The forecasts will be conditional on  $I_T$ .

The variable to forecast  $Y_{T+\ell}$  is a RV. It can be fully characterized by a pdf.

In general, it is difficult to get the pdf for the forecast. In practice, we get a point estimate (the forecast) and a C.I.

Notation:

- Forecast for  $T + \ell$  made at  $T: \hat{Y}_{T+\ell}, \hat{Y}_{T+\ell|T}, \hat{Y}_{T}(\ell)$ .
- $T + \ell$  forecast error:  $e_{T+\ell} = e_T(\ell) = Y_{T+\ell} \hat{Y}_{T+\ell}$
- Mean squared error (MSE):  $MSE(e_{T+\ell}) = E[Y_{T+\ell} \hat{Y}_{T+\ell}]^2$

To get a point estimate,  $\pi \pi Y = T + \ell$ , we need a cost function to judge various alternatives. This cost function is call *loss function*. Since we are working with forecast, we work with a expected loss function.

A popular loss functions is the MSE, which is quadratic and symmetric. We can use asymmetric functions, for example, functions that penalize positive errors more than negative errors.

If we use the MSE as the loss function, we look for  $\hat{Y}_{T+l}$ , which minimizes it. That is,

$$\min E\left[e_{T+\ell}^2\right] = E\left[(Y_{T+\ell} - \hat{Y}_{T+\ell})^2\right] = E\left[Y_{T+\ell}^2 - 2Y_{T+\ell}\hat{Y}_{T+\ell} + \hat{Y}_{T+\ell}^2\right]$$

Then, f.o.c. implies:

$$E[-2Y_{T+\ell} + 2\hat{Y}_{T+\ell}] = 0 \qquad \Rightarrow E[Y_{T+\ell}] = \hat{Y}_{T+\ell}.$$

The optimal point forecast under MSE is the (conditional) mean:

$$\widehat{Y}_{T+\ell} = \mathbb{E}[Y_{T+\ell}|I_T]$$

Different loss functions lead to different optimal forecast. For example, for the MAE, the optimal point forecast is the median.

The computation of  $E[Y_{T+\ell} | I_T]$  depends on the distribution of  $\{\varepsilon_t\}$ . If  $\{\varepsilon_t\} \sim WN$ , then  $E[\varepsilon_{T+\ell} | I_T] = 0$ , which greatly simplifies computations, especially in the linear model.

Then, for an ARMA(p, q) stationary process (with a Wold representation), the minimum MSE linear forecast (best linear predictor) of  $Y_{T+\ell}$ , conditioning on I<sub>T</sub> is:

 $Y_{T+\ell} = \theta_0 + \Psi_l \varepsilon_{T+\ell} + \Psi_{l+1} \varepsilon_{T+\ell-1} + \cdots$ 

### **Forecasting Steps for ARMA Models**

The usual process has the following steps:

- ARIMA model: - Estimation (Evaluation in-sample) - Forecast  $Y_t = \phi Y_{t-1} + \varepsilon_t$   $\hat{\phi}$  (Estimate of  $\phi$ )  $\Rightarrow \hat{Y}_t = \hat{\phi} Y_{t-1}$  (Prediction)  $\hat{Y}_{t+1} = \hat{\phi} \hat{Y}_t$  (Forecast)

(Evaluation out-of-sample)

We observe the time series:  $I_T = \{Y_1, Y_2, ..., Y_T\}$ .

- At time T, we want to forecast:  $Y_{T+1}$ ,  $Y_{T+2}$ , ...,  $Y_{T+\ell}$ .
- T: The forecast origin.
- $\ell$ : Forecast horizon
- $\hat{Y}_T(\ell)$ :  $\ell$ -step ahead forecast = Forecasted value  $Y_{T+\ell}$

Use the conditional expectation of  $Y_{T+\ell}$ , given the observed sample.  $\hat{Y}_{T+\ell} = E[Y_{T+\ell}|Y_T, Y_{T-1}, \dots, Y_1]$ 

**Example:** One-step ahead forecast:  $\hat{Y}_{T+1} = E[Y_{T+1}|Y_T, Y_{T-1}, \dots, Y_1]$ .

Forecast accuracy to be measured by MSE

 $\Rightarrow$  conditional expectation, best forecast.

# Forecasting From MA(q) Models

The stationary MA(q) model for  $Y_t$  is

 $Y_t = \mu + \varepsilon_t + \theta_1 \varepsilon_{t-1} + \dots + \theta_q \varepsilon_{t-q}$ 

 $Y_{T+1} = \mu + \varepsilon_{T+1} + \theta_1 \varepsilon_T + \dots + \theta_q \varepsilon_{T-q+1}$  $Y_{T+2} = \mu + \varepsilon_{T+2} + \theta_1 \varepsilon_{T+1} + \dots + \theta_q \varepsilon_{T-q+2}$ 

Then, assuming we have the data up to time T ( $Y_1$ ,  $Y_2$ , ...,  $Y_T$ ,  $\varepsilon_1$ ,  $\varepsilon_2$ , ...,  $\varepsilon_T$ ) and parameter constancy, we produce at time *T l*-step ahead forecasts using:

÷

 $Y_{T+l} = \mu + \varepsilon_{T+l} + \theta_1 \varepsilon_{T+l-1} + \dots + \theta_a \varepsilon_{T+l-a}$  (l > 2)

Now, we take conditional expectations:

 $\hat{Y}_{T+l} = E[Y_{T+l}|I_T] = \mu + E[\varepsilon_{T+l}|I_T] + \theta_1 E[\varepsilon_{T+l-1}|I_T] + \dots + \theta_q E[\varepsilon_{T+l-q}|I_T]$ 

Note: The forecasts are a linear combination of forecast and past errors.

Some of the errors are know at time  $T: \varepsilon_1 = \hat{\varepsilon}_1, \varepsilon_2 = \hat{\varepsilon}_2, ..., \varepsilon_T = \hat{\varepsilon}_T$ , the rest are unknown. Thus,  $E[\varepsilon_{T+l}|I_T] = 0$  for l > 1.

**Example:** For an MA(2) we have:  $\hat{Y}_{T+1} = \mu + \mathbb{E}[\varepsilon_{T+1}|I_T] + \theta_1 \mathbb{E}[\varepsilon_T|I_T] + \theta_2 \mathbb{E}[\varepsilon_{T-1}|I_T]$ 

$$\begin{aligned} \hat{Y}_{T+2} &= \mu + \mathbb{E}[\varepsilon_{T+2}|I_T] + \theta_1 \mathbb{E}[\varepsilon_{T+1}|I_T] + \theta_2 \mathbb{E}[\varepsilon_T|I_T] \\ \hat{Y}_{T+3} &= \mu + \mathbb{E}[\varepsilon_{T+3}|I_T] + \theta_1 \mathbb{E}[\varepsilon_{T+2}|I_T] + \theta_2 \mathbb{E}[\varepsilon_{T+1}|I_T] \end{aligned}$$

At time T=t, we know  $\varepsilon_t$  and  $\varepsilon_{t-1}$ . Set  $\mathbb{E}[\varepsilon_{t+j}|I_t]=0$  for j > 1. Then,  $\hat{Y}_{t+1} = \mu + \theta_1 \mathbb{E}[\varepsilon_t|I_t] + \theta_2 \mathbb{E}[\varepsilon_{t-1}|It] = \mu + \theta_1 \hat{\varepsilon}_t + \theta_2 \hat{\varepsilon}_{t-1}$   $\hat{Y}_{t+2} = \mu + \theta_2 \mathbb{E}[\varepsilon_t|It] = \mu + \theta_2 \hat{\varepsilon}_t$   $\hat{Y}_{t+3} = \mu$  $\hat{Y}_{t+1} = \mu$  for l > 2

 $\Rightarrow$  MA(2) memory of 2 periods. For l > 2, all forecast are constant (=  $\mu$ ).

The example generalizes: An MA(q) process has a memory of only q periods. All forecasts beyond q revert to the unconditional mean,  $\mu$ .

Example: An industrial firm uses an MA(2) to forecast sales. The estimated MA(2) model is:

 $\hat{Y}_T = 2.2 + 0.4 \varepsilon_{T-1} + 0.2 \varepsilon_{T-2}$ 

At time T=t, the firms know  $\varepsilon_t = 1.42$  and  $\varepsilon_{t-1} = -0.91$ .

Then, the first three forecast are:

$$\begin{split} \hat{Y}_{t+1} &= 2.2 + 0.4 * \mathbf{1.42} + 0.2 * (-0.91) = 2.586 \\ \hat{Y}_{t+2} &= 2.2 + 0.2 * (\mathbf{1.42}) = 2.484 \\ \hat{Y}_{t+3} &= 2.2 \qquad \qquad (\Rightarrow \hat{Y}_{t+l} = 2.2 \text{ for } l > 3.) \end{split}$$

Later, the firm observes:  $Y_{t+1} = 4.77$ ,  $Y_{t+2} = 3.15 \& Y_{t+3} = 1.85$ . Then, the MSE:

MSE =  $\frac{1}{3} * [(4.77 - 2.586)^2 + (3.15 - 2.484)^2 + (1.85 - 2.2)^2] = 1.779.$ 

**Example:** We fit an MA(1) to the U.S. stock returns (T=1,975):

```
library(tseries)
library(forecast)
fit_p_ts <- arima(lr_p, order=c(0,0,1))  #fit an MA(1) model
fcast_p <- forecast(fit_p_ts, h=4)  #produce 4-step ahead forecasts
> fit_p_ts
> fcast_p
Coefficients:
    mal intercept
    0.2888    0.0037
s.e. 0.0218    0.0012
```

sigma<sup>2</sup> estimated as 0.001522: log likelihood = 3275.83, aic = -6545.67

```
> fcast_p
Point Forecast Lo 80 Hi 80 Lo 95 Hi 95
1796 0.012570813 -0.03742238 0.06256401 -0.06388718 0.08902881
```

1797	<b>0.003689524</b> -0.04834634 0.05572539 -0.07589247 0.08327152
1700	0.002(00524.0.04024(24.0.05572520.0.07500247.0.0022715

**0.003689524** -0.04834634 0.05572539 -0.07589247 0.08327152

1799 **0.003689524** -0.04834634 0.05572539 -0.07589247 0.08327152

<u>Remark</u>: After the first forecast, the MA(1) process generates constant forecasts.

### Forecasting From AR(p) Models

The stationary AR(p) model for  $Y_t$  is

 $Y_t = \mu + \phi_1 Y_{t-1} + \phi_2 Y_{t-2} + \dots + \phi_p Y_{t-p} + \varepsilon_t$ 

Then, assuming we have the data up to time  $T(Y_1, Y_2, ..., Y_T)$  and parameter constancy, we produce at time *T l*-step ahead forecasts using:

$$Y_{T+1} = \mu + \phi_1 Y_T + \phi_2 Y_{T-1} + \dots + \phi_p Y_{T-p+1} + \varepsilon_{T+1}$$
  

$$Y_{T+2} = \mu + \phi_1 Y_{T+1} + \phi_2 Y_T + \dots + \phi_p Y_{T-p+2} + \varepsilon_{T+2}$$
  

$$\vdots$$
  

$$Y_{T+l} = \mu + \phi_1 Y_{T+l-1} + \phi_2 Y_{T+l-2} + \dots + \phi_p Y_{T+l-p} + \varepsilon_{t+l} \qquad (l > 2)$$

Now, we take conditional expectations:

$$\widehat{Y}_{T+l} = E[Y_{T+l}|I_{T}] = \mu + \phi_{1}E[Y_{T+l-1}|I_{T}] + \phi_{2}E[Y_{T+l-2}|I_{T}] + \dots + \phi_{p}E[Y_{T+l-p}|I_{T}]$$

Note that  $E[Y_{T+l-j}|I_T]$  are also forecasts. The forecasts  $\hat{Y}_{T+l}$  is a linear combination of past forecast.

**Example:** AR(2) model for  $Y_{t+l}$  is  $Y_{t+l} = \mu + \phi_1 Y_{t+l-1} + \phi_2 Y_{t+l-2} + \varepsilon_{t+l}$ 

Then, taking conditional expectations at time T=t, we get the forecasts:

$$\begin{split} \hat{Y}_{t+1} &= \mu + \phi_1 Y_t + \phi_2 Y_{t-1} \\ \hat{Y}_{t+2} &= \mu + \phi_1 \hat{Y}_{t+1} + \phi_2 Y_t \\ \hat{Y}_{t+3} &= \mu + \phi_1 \hat{Y}_{t+2} + \phi_2 \hat{Y}_{t+1} \\ \vdots \\ \hat{Y}_{t+l} &= \mu + \phi_1 \hat{Y}_{t+l-1} + \phi_2 \hat{Y}_{T+l-1} \end{split}$$

AR-based forecasts are autocorrelated, they have long memory!

**Example:** An industrial firm uses an AR(2) to forecast sales. The estimated AR(2) model is:

$$\hat{Y}_T = 0.7 + 0.51 y_{T-1} + 0.1 y_{T-2}$$

At time T=t, the firms know  $Y_t = 3$  and  $Y_{t-1} = 3.52$ . Then, the first three forecast are:

$$\hat{Y}_{t+1} = 0.7 + 0.51 * 3 + 0.1 * 3.52 = 2.582$$
  
 $\hat{Y}_{t+2} = 0.7 + 0.51 * 2.582 + 0.1 * 3 = 2.317$ 

 $\hat{Y}_{t+3} = 0.7 + 0.51 * 2.317 + 0.1 * 2.582 = 2.140$ 

Later, the firm observes:  $Y_{t+1} = 4.77$ ,  $Y_{t+2} = 3.15 \& Y_{t+3} = 1.85$ . Then, the MSE: MSE  $= \frac{1}{3} * [(4.77 - 2.582)^2 + (3.15 - 2.317)^2 + (1.85 - 2.140)^2] = 1.855$ .

**Example:** We fit an AR(4) to the changes in Oil Prices (T=346): fit\_oil\_ts <- arima(lr\_oil, order=c(4,0,0)) fcast\_oil <- forecast(fit\_oil\_ts, h=12) > fit\_oil\_ts

Coefficients:

ar1 ar2 ar3 ar4 intercept 0.2946 -0.1027 -0.0571 -0.0983 0.0017 s.e. 0.0521 0.0543 0.0551 0.0539 0.0051

sigma<sup>2</sup> estimated as 0.008812: log likelihood = 344.57, aic = -677.14

> fcast\_oil Point Forecast Lo 80 Hi 80 Lo 95 Hi 95 365 -5.425015e-02 -0.1745546 0.0660543 -0.2382399 0.1297396 366 -1.578754e-02 -0.1412048 0.1096297 -0.2075966 0.1760216 367 2.455760e-03 -0.1229760 0.1278875 -0.1893755 0.1942871 368 1.356917e-02 -0.1123501 0.1394884 -0.1790077 0.2061460 369 1.160479e-02 -0.1154462 0.1386558 -0.1827029 0.2059125 370 5.060891e-03 -0.1221954 0.1323172 -0.1895608 0.1996826 371 9.059104e-04 -0.1263511 0.1281629 -0.1937169 0.1955287

<u>Note</u>: You can extract the point forecasts from the forecast function using \$mean. That is, fcast\_oil\$mean extracts the whole vector of forecasts.

• We plot the 12 forecasts:

> plot(fcast\_oil)



<u>Remark</u>: Different from the MA(1) forecasts, the AR(1) process generates non-constant forecasts.  $\P$ 

## Forecasting From ARMA(p,q) Models

The stationary ARMA model for  $Y_t$  is

 $Y_t = \theta_0 + \phi_1 Y_{t-1} + \dots + \phi_p Y_{t-p} + \varepsilon_t + \theta_1 \varepsilon_{t-1} + \dots + \theta_q \varepsilon_{t-q}$ 

Assume that we have data  $Y_1, Y_2, ..., Y_T$ ;  $\varepsilon_1 = \hat{\varepsilon}_1, \varepsilon_2 = \hat{\varepsilon}_2, ..., \varepsilon_T = \hat{\varepsilon}_T$ . We want to forecast  $Y_{T+\ell}$ . Then,

$$Y_{T+\ell} = \theta_0 + \phi_1 Y_{T+\ell-1} + \dots + \phi_p Y_{T+\ell-p} + \varepsilon_{T+\ell} + \theta_1 \varepsilon_{T+\ell-1} + \dots + \theta_q \varepsilon_{T+\ell-q}$$

Taking expectations:

$$\hat{Y}_{T+l} = \theta_0 + \phi_1 \hat{Y}_{T+l-1} + \dots + \phi_p \hat{Y}_{T+l-p} + E[\varepsilon_{T+\ell} | I_T] + \theta_1 E[\varepsilon_{T+\ell-1} | I_T] + \dots + \theta_q E[\varepsilon_{T+\ell-q} | I_T]$$

<u>Remark</u>: An ARMA forecasting is a combination of past  $\hat{Y}_{T+l-i}$  forecasts and observed past  $\hat{\varepsilon}_{t+l-i}$ .

**Example:** An industrial firm uses an ARMA(1,2) to forecast sales. The estimated ARMA(1,2) model is:

 $\hat{Y}_T = 1.9 + 0.32 y_{T-1} + 0.25 \varepsilon_{T-1} - 0.1 \varepsilon_{T-2}$ 

At time *T*=*t*, the firm knows:  $Y_t = 3$ ,  $\varepsilon_t = 0.81$ , &  $\varepsilon_{t-1} = 0.47$ .

Then, the first three forecast are:

$$\hat{Y}_{t+1} = 1.9 + 0.32 * 3 + 0.25 * (0.81) - 0.1 * 0.47 = 2.981$$
  
 $\hat{Y}_{t+2} = 1.9 + 0.32 * 2.981 - 0.1 * 0.81 = 2.770$   
 $\hat{Y}_{t+3} = 1.9 + 0.32 * 2.770 = 2.786$ 

Later, the firm observes:  $Y_{t+1} = 4.77$ ,  $Y_{t+2} = 3.15$ , &  $Y_{t+3} = 1.85$ . Then, the MSE:

$$MSE = \frac{1}{3} * \left[ (4.77 - 2.981)^2 + (3.15 - 2.770)^2 + (1.85 - 2.786)^2 \right] = 1.407.$$

Alternatively, we can forecast considering the Wold representation:

$$Y_{T+\ell} = \mu + \Psi(B)\varepsilon_t = \theta_0 + \frac{\theta_q(B)}{\phi_p(B)}\varepsilon_t = \mu + \varepsilon_{T+\ell} + \Psi_1\varepsilon_{T+\ell-1} + \Psi_2\varepsilon_{T+\ell-2} + \dots + \Psi_\ell\varepsilon_T + \dots$$

Taking the expectation of  $Y_{T+\ell}$ , we have

 $\hat{Y}_{T+\ell} = E(Y_{T+\ell}|Y_T, Y_{T-1}, \cdots, Y_1) = \mu + \Psi_{\ell}\varepsilon_T + \Psi_{\ell+1}\varepsilon_{T-1} + \cdots$ where

$$E(\varepsilon_{T+j}|Y_T,\cdots,Y_1) = \begin{cases} 0, & j > 0\\ \varepsilon_{T+j}, & j \le 0 \end{cases}$$

Then, we define the forecast error:

$$e_{T}(\ell) = Y_{T+\ell} - \hat{Y}_{T+\ell} = \varepsilon_{T+\ell} + \Psi_{1}\varepsilon_{T+\ell-1} + \dots + \Psi_{\ell-1}\varepsilon_{T+1}$$
$$= \sum_{i=0}^{\ell-1} \Psi_{i}\varepsilon_{T+\ell-i}$$

The forecast error is:  $e_T(\ell) = \sum_{i=0}^{\ell-1} \Psi_i \varepsilon_{T+\ell-i}$ 

<u>Note</u>: The expectation of the forecast error:  $E[e_T(\ell)] = 0$  $\Rightarrow$  we say the forecast is *unbiased*.

• The variance of the forecast error:

$$Var(e_{T}(\ell)) = Var\left(\sum_{i=0}^{\ell-1} \Psi_{i}\varepsilon_{T+\ell-i}\right) = \sigma^{2}\sum_{i=0}^{\ell-1} \Psi_{i}^{2}$$

、

**Example 1:** One-step ahead forecast (l = 1).

$$Y_{T+1} = \mu + \varepsilon_{T+1} + \Psi_1 \varepsilon_T + \Psi_2 \varepsilon_{T-1} + \cdots$$
  

$$\hat{Y}_{T+1} = \mu + \Psi_1 \varepsilon_T + \Psi_2 \varepsilon_{T-1} + \cdots$$
  

$$e_T(1) = Y_{T+1} - \hat{Y}_{T+1} = \varepsilon_{T+1}$$
  

$$Var(e_T(1)) = \sigma^2 . \P$$

**Example 2:** One-step ahead forecast (
$$\ell = 2$$
).

$$Y_{T+2} = \mu + \varepsilon_{T+2} + \Psi_1 \varepsilon_{T+1} + \Psi_2 \varepsilon_T + \cdots \hat{Y}_{T+2}$$
  
=  $\mu + \Psi_2 \varepsilon_T + \cdots e_T(2)$   
=  $Y_{T+2} - \hat{Y}_{T+2} = \varepsilon_{T+2} + \Psi_1 \varepsilon_{T+1} Var(e_T(2)) = \sigma^2 * (1 + \Psi_1^2)$ 

Note:

$$\lim_{\ell \to \infty} Y_T(\ell) = \mu$$
$$\lim_{\ell \to \infty} Var[e_T(\ell)] = \gamma_0 < \infty$$

Recall that the Wold representation depends on an infinite number of parameters, but, in practice, they decay rapidly. Then, as we forecast into the future, the forecasts are not very interesting (unconditional forecasts!).

That is why ARMA (or ARIMA) forecasting is useful only for short-term forecasting.

# Forecasting From ARMA(p,q) Models: C.I.

A 100(1-  $\alpha$ )% prediction interval for  $Y_{T+\ell}$  ( $\ell$ -steps ahead) is

$$\hat{Y}_{T}(\ell) \pm \frac{z_{\alpha}}{2} \sqrt{Var(e_{T}(\ell))}$$

$$\hat{Y}_{T}(\ell) \pm \frac{z_{\alpha/2}}{2} \sigma \sqrt{\sum_{i=0}^{\ell-1} \Psi_{i}^{2}}$$

**Example:** 95% C.I. for the 2-step-ahead forecast:  $\hat{Y}_T(2) \pm 1.96 \sigma \sqrt{1 + \Psi_1^2}$  When computing prediction intervals from data, we substitute estimates for parameters, giving approximate prediction intervals.  $\P$ 

<u>Note</u>: Since  $\Psi'_i$ 's are RV, MSE[ $\varepsilon_{T+\ell}$ ] = MSE[ $e_{T+\ell}$ ] =  $\sigma^2 \sum_{i=0}^{\ell-1} \Psi_i^2$ 

```
Example: We fit an ARMA(4, 5), as selected by the function auto.arima, to changes in monthly
U.S. earnings (1871 – 2020):
x E \leq Sh da E
T \leq - \text{length}(x | E)
\ln e \le \log(x E[-1]/x E[-T])
fit e \le auto.arima(lr e)
> auto.arima(lr e)
Series: lr e
ARIMA(4,0,5) with non-zero mean
Coefficients:
                                               ma3
     ar1
          ar2
                    ar3
                           ar4
                                ma1
                                        ma2
                                                      ma4
   0.3541 0.9786 0.2530 -0.6381 0.2943 -0.6794 -0.5720 0.1787
s.e. 0.0414 0.0466 0.0414 0.0363 0.0455 0.0400 0.0465 0.0362
     ma5 mean
   -0.1498 0.0032
s.e. 0.0286 0.0008
sigma<sup>2</sup> estimated as 0.0005759: log likelihood=4140.46
AIC=-8258.91 AICc=-8258.76 BIC=-8198.52
• We forecast 20 periods ahead
> fcast e <- forecast(fit e, h=20)
                                        # h=number of step-ahead forecasts
> fcast e
  Point Forecast
                   Lo 80
                              Hi 80
                                           Lo 95
                                                     Hi 95
1791 -0.054521445 -0.08527728 -0.023765608 -0.10155844 -0.007484451
1792 -0.048064225 -0.08471860 -0.011409845 -0.10412226 0.007993811
1793 -0.032702992 -0.07280271 0.007396723 -0.09403021 0.028624230
1794 -0.030680456 -0.07365723 0.012296320 -0.09640776 0.035046851
1795 -0.017583413 -0.06228564 0.027118816 -0.08594957 0.050782746
1796 -0.013681751 -0.05882105 0.031457550 -0.08271635 0.055352853
1797 -0.008775187 -0.05458154 0.037031165 -0.07882996 0.061279583
1798 -0.001197077 -0.04705319 0.044659034 -0.07132795 0.068933794
1799 -0.001083388 -0.04698821 0.044821436 -0.07128876 0.069121982
1800 0.005124015 -0.04078796 0.051035988 -0.06509229 0.075340318
1801
      0.006219195 -0.03973961 0.052178005 -0.06406874 0.076507130
1802 0.007874051 -0.03809120 0.053839304 -0.06242374 0.078171840
      0.011029600 -0.03506469 0.057123889 -0.05946553 0.081524732
1803
      0.010082045 -0.03611076 0.056274848 -0.06056375 0.080727841
1804
```

<u>Note</u>: You can extract the point forecasts from the forecast function using \$mean. That is, fcast\_e\$mean extracts the whole vector of forecasts.

• We plot the forecast and the C.I. > plot(fcast\_e, type="l", include = 24, main = "Changes in Earings: Forecast 2020:Oct - 2021:Jun") #We include the last 24 observations along the forecast.



# Forecasting From ARMA(p,q) Models - Updating

Suppose we have T observations at time t=T. We have a good ARMA model for  $Y_t$ . We obtain the forecast for  $Y_{T+1}$ ,  $Y_{T+2}$ , etc.

• At t = T + 1, we observe  $Y_{T+1}$ . Now, we update our forecasts using the original value of  $Y_{T+1}$  and the forecasted value of it.

The forecast error is:  $e_T(\ell) = Y_{T+\ell} - \hat{Y}_T(\ell) = \sum_{i=0}^{\ell-1} \Psi_i \varepsilon_{T+\ell-i}$ 

The forecast error associated with  $\hat{Y}_{T-1}(\ell + 1)$  is:

$$e_{T-1}(\ell+1) = Y_{T-1+\ell+1} - Y_{T-1}(\ell+1)$$
  
=  $\sum_{i=0}^{\ell} \Psi_i \varepsilon_{T-1+\ell+1-i} = \sum_{i=0}^{\ell} \Psi_i \varepsilon_{T+\ell-i}$   
=  $\sum_{i=0}^{\ell-1} \Psi_i \varepsilon_{T+\ell-i} + \Psi_\ell \varepsilon_T = e_T(\ell) + \Psi_\ell \varepsilon_T$ 

• Then,

$$\begin{split} e_{T-1}(\ell+1) &= Y_{T+\ell} - \hat{Y}_{T-1}(\ell+1) = Y_{T+\ell} - \hat{Y}_{T}(\ell) + \Psi_{\ell} \varepsilon_{T} \\ \hat{Y}_{T}(\ell) &= \hat{Y}_{T-1}(\ell+1) + \Psi_{\ell} \varepsilon_{T} \\ \hat{Y}_{T}(\ell) &= \hat{Y}_{T-1}(\ell+1) + \Psi_{\ell} \{Y_{T} - \hat{Y}_{T-1}(1)\} \\ \hat{Y}_{T+1}(\ell) &= \hat{Y}_{T}(\ell+1) + \Psi_{\ell} \{Y_{T+1} - \hat{Y}_{T}(1)\} \end{split}$$

Example: 
$$\ell = 1, T = 100.$$
  
 $\hat{Y}_{101}(1) = \hat{Y}_{100}(2) + \Psi_1 \{Y_{101} - \hat{Y}_{100}(1)\}.$ 

Forecasting From ARMA(p,q) Models - Remarks

In general, we need a large *T*. Better estimates and it is possible to check for model stability and check forecasting ability of model by withholding data.

Seasonal patterns also need large T. Usually, you need 4 to 5 seasons to get reasonable estimates.

Parsimonious models are very important. Easier to compute and interpret models and forecasts. Forecasts are less sensitive to deviations between parameters and estimates.

# **Forecasting From Simple Models: ES**

Industrial companies, with a lot of inputs and outputs, want quick and inexpensive forecasts. Easy to fully automate. In general, they only use past observations of the series to forecat. That is, we use past  $Y_t$  to forecast future  $Y_t$ 's, which in the literature is usually referred as the "*level's* forecasts."

Exponential Smoothing Models (ES) fulfill these requirements.

In general, these models are limited and not optimal, especially compared with Box-Jenkins methods.

Goal of these models: Suppress the short-run fluctuation by smoothing the series. For this purpose, a weighted average of all previous values works well.

There are many ES models. We will go over the Simple Exponential Smoothing (SES) and Holt-Winter's Exponential Smoothing (HW ES).

# Simple Exponential Smoothing: SES

We "smooth" the series  $Y_t$  to produce a quick forecast,  $S_{t+1}$  called the "level's forecast." Smooth? The graph of  $S_t$  is less jagged than the graph of original series  $Y_t$ .

Observed time series at time T:  $Y_1$ ,  $Y_2$ , ...,  $Y_T$ .

The SES Model has only one equation, we only forecast the level:

 $S_t = \alpha Y_{t-1} + (1-\alpha)S_{t-1}$ 

where

-  $\alpha$ : The smoothing parameter,  $0 \le \alpha \le 1$ .

- *Y<sub>t</sub>*: Value of the observation at time t.
- *St*: Value of the smoothed observation at time t –i.e., the forecast.

The equation can also be written as an updating equation:

 $S_t = S_{t-1} + \alpha(Y_{t-1} - S_{t-1}) = S_{t-1} + \alpha * \text{ (past forecast error)}$ 

Note: The updating form of the SES model looks like an MA(1) model.

## **SES: Forecast and Updating**

From the updating equation for  $S_t$ :

$$S_t = S_{t-1} + \alpha (Y_{t-1} - S_{t-1})$$

we compute the forecast:

$$S_{t+1} = \alpha Y_t + (1 - \alpha)S_t = S_t + \alpha(Y_t - S_t)$$

That is, a simple updating forecast: last period forecast + adjustment.

For the next period, we have:

 $S_{t+2} = \alpha Y_{t+1} + (1-\alpha)S_{t+1} = \alpha S_{t+1} + (1-\alpha)S_{t+1} = S_{t+1}$ 

Then the  $\ell$ -step ahead forecast is:

 $S_{t+\ell} = S_{t+1} \implies$  A naive forecast!

<u>Note</u>: Similar to an MA(1) process, SES forecasts are not very interesting after  $\ell > 1$ .

**Example:** An industrial firm uses SES to forecast sales:

 $S_{t+1} = S_t + \alpha * (Y_t - S_t)$ 

The firm estimates  $\alpha = 0.25$ . The firm observes  $Y_t = 5$  and, last period's forecast,  $S_t = 3$ . Then, the forecast for time t+1 is:

 $S_{t+1} = 3 + 0.25 * (5 - 3) = 3.50$ 

The forecast for time *t*+1 and any period after time *t*+1, we have  $S_{t+\ell} = S_{t+1} = 3.50$  for  $\ell > 1$ .

Later, the firm observes:  $Y_{t+1} = 4.77$ ,  $Y_{t+2} = 3.15$ , &  $Y_{t+3} = 1.85$ . Then, the MSE:

$$MSE = \frac{1}{3} * [(4.77 - 3.50)^2 + (3.15 - 3.50)^2 + (1.85 - 3.50)^2] = 1.486.$$

<u>Note</u>: If  $\alpha = 0.75$ , then

 $S_{t+1} = 3 + 0.75 * (5 - 3) = 4.50$ 

A bigger  $\alpha$  gives more weight to the more recent observation –i.e.,  $Y_t$ .

#### **SES:** Exponential?

Question: Why Exponential?

For the observed time series  $\{Y_1, Y_2, ..., Y_T, Y_{T+1}\}$ , using backward substitution,  $S_{t+1} = \hat{Y}_t(1)$  can be expressed as a weighted sum of previous observations:

$$S_{t+1} = \alpha Y_t + (1 - \alpha)S_t = \alpha Y_t + (1 - \alpha)[\alpha Y_{t-1} + (1 - \alpha)S_{t-1}]$$
  
=  $\alpha Y_t + \alpha (1 - \alpha)Y_{t-1} + (1 - \alpha)^2 S_{t-1}$   
 $\Rightarrow \hat{Y}_t(1) = c_0 Y_t + c_1 Y_{t-1} + c_2 Y_{t-2} + \cdots$   
where  $c_i$ 's are the weights, with  
 $c_i = \alpha (1 - \alpha)^i$ ;  $i = 0, 1, ...; 0 \le \alpha \le 1$ .

We have decreasing weights, by a constant ratio for every unit increase in lag.

Then,

$$\begin{split} \hat{Y}_t(1) &= \alpha (1-\alpha)^0 Y_t + \alpha (1-\alpha)^1 Y_{t-1} + \alpha (1-\alpha)^2 Y_{t-2} + \cdots \\ \hat{Y}_t(1) &= \alpha Y_t + (1-\alpha) \hat{Y}_{t-1}(1) \quad \Rightarrow S_{t+1} = \alpha Y_t + S_t \end{split}$$

• Let's look at the weights:

$$c_i = \alpha (1 - \alpha)^i;$$
  $i = 0, 1, ...; 0 \le \alpha \le 1$ 

$c_i = \alpha (1-\alpha)^i$	$\alpha = 0.25$	$\alpha = 0.75$
<i>c</i> <sub>0</sub>	0.25	0.75
<i>c</i> <sub>1</sub>	0.25 * 0.75 = 0.1875	0.75 * 0.25 = 0.1875
<i>c</i> <sub>2</sub>	$.25 * 0.75^2 = 0.140625$	$0.75 * 0.25^2 = 0.046875$
<i>C</i> <sub>3</sub>	$.25 * 0.75^3 = 0.1054688$	$0.75 * 0.25^3 = 0.01171875$
C <sub>4</sub>	$.25 * 0.75^4 = 0.07910156$	$0.75 * 0.25^4 = 0.002929688$
:		
<i>c</i> <sub>12</sub>	$.25 * 0.75^{12} = 0.007919088$	$0.75 * 0.25^{12} = 4.470348e-08$

Decaying weights. Faster decay with greater  $\alpha$ , associated with faster learning: we give more weight to more recent observations.

We do not know  $\alpha$ ; we need to estimate it.

# SES: Selecting α

Choose  $\alpha$  between 0 and 1.

- If  $\alpha = 1$ , it becomes a naive model; if  $\alpha \approx 1$ , more weights are put on recent values. The model fully utilizes forecast errors.

- If  $\alpha$  is close to 0, distant values are given weights comparable to recent values. Set  $\alpha \approx 0$  when there are big random variations in  $Y_t.$ 

-  $\alpha$  is often selected as to minimize the MSE.

In empirical work,  $0.05 \le \alpha \le 0.3$  are used ( $\alpha \approx 1$  is used rarely).

Numerical Minimization Process:

- Take different  $\alpha$  values ranging between 0 and 1.

- Calculate 1-step-ahead forecast errors for each  $\alpha$ , where the forecast error is  $e_t = Y_t S_t$ .
- Calculate MSE for each case.

- Then, choose the  $\alpha$  which produces the minimum MSE:  $min_{\alpha} \sum_{t=1}^{n} e_t^2$ 

Time	$Y_t$	$S_{t+I} (\alpha = 0.10)$	$(Y_t - S_t)^2$
1	5	-	-
2	7	(0.1) <b>5</b> + $(0.9)$ <b>5</b> = <b>5</b>	4
3	6	(0.1)7 + (0.9)5 = 5.2	0.64
4	3	(0.1)6 + (0.9)5.2 = 5.28	5.1984
5	4	(0.1)3 + (0.9) <b>5.28</b> = 5.052	1.107
TOTAI	[]		10.945
$MSE = \frac{SSE}{n-1} = 2.74$			

**Example:** 

Calculate this for  $\alpha = 0.2, 0.3, \dots, 0.9, 1$  and compare the MSEs. Choose  $\alpha$  with minimum MSE.

Note:  $Y_{t=1} = 5$  is set as the initial value for the recursive equation.

# **SES:** Initial Values

We start forecasting at time 2. Since we have a recursive equation, we need an initial value for  $S_1$ (or  $Y_0$ ).

Approaches:

- Set  $S_1$  to  $Y_1$  is one method of initialization. Then,  $S_2 = Y_1$ .

- Take the average of the first p observations, say first 4 or 5 observations:  $Y_0 = \frac{\sum_{t=1}^{p} Y_t}{p}$ 

Use this average as an initial value  $S_1 = Y_0$ . Obviously, in the case our first prediction will be for time (p+1), which becomes:  $S_2$ .

- Estimate  $S_1$  (similar to the estimation of *a*).

# **SES:** Forecasting Examples

**Example 1:** We want to forecast log changes in U.S. monthly dividends (T=1796) using SES. First, we estimate the model using the R function HoltWinters(), which has as a special case SES: set beta=FALSE, gamma=FALSE. We use estimation period T=1750.

mod1 <- HoltWinters(lr d[1:1750], beta=FALSE, gamma=FALSE) > mod1Holt-Winters exponential smoothing without trend and without seasonal component. Call:

HoltWinters(x = lr d[1:1750], beta = FALSE, gamma = FALSE)



}
ses\_f <- sm[T1:T]
ses\_f
f\_error\_ses <- sm[T1:T] - y[T1:T] # forecast errors
MSE\_ses <- sum(f\_error\_ses^2)/h # MSE
plot(ses\_f, type="l", main ="SES Forecasts: Changes in Dividends")</pre>



•••

Note: Constant forecasts, but C.I. gets wider (as expected) with h. ¶

**Example 2:** We want to forecast **log monthly U.S. vehicles** (1976-2020, T=537) using SES. mod\_car <- HoltWinters(l\_car[1:512], beta=FALSE, gamma=FALSE)

> mod\_car Holt-Winters exponential smoothing without trend and without seasonal component. Call: HoltWinters(x = l\_car[1:512], beta = FALSE, gamma = FALSE) Smoothing parameters: alpha:  $0.4888382 \implies$  Estimated  $\alpha$ beta : FALSE gamma: FALSE Coefficients: [,1] a 7.315328

 Now, we do one-step ahead forecasting ses\_f\_c <- sm\_c[T1:T] f\_error\_c\_ses <- sm\_c[T1:T] - y[T1:T]</li>
 > plot(ses\_f\_c, type="l", main ="SES Forecasts: Log Vehicle Sales")

#### SES Forecasts: Log Vehicle Sales



> plot(f error c ses, type="l", main ="SES Forecasts Errors: Log Vehicle Sales")



#### **SES: Remarks**

Some computer programs automatically select the optimal  $\alpha$  using a line search method or nonlinear optimization techniques.

We have a recursive equation, we need initial values for  $S_1$ .

This model ignores trends or seasonalities. Not very realistic, especially for manufacturing facilities, retail sector, and warehouses. But, deterministic components,  $D_t$ , can be easily incorporated.

The model that incorporates both features is called *Holt-Winter's ES*.

# Holt-Winters (HW) Exponential Smoothing

Now, we introduce trend  $(T_t)$  & seasonality  $(I_t)$  factors. Since we also produce smooth forecasts for  $T_t \& I_t$ , this method is also called *triple exponential smoothing*.

The *h*-step ahead forecast is a combination of the smooth forecasts of  $S_t$  (Level),  $T_t$  (Trend) &  $I_{t+h-s}$  (Seasonal).

Both,  $T_t \& I_t$ , can be included as *additively* or *multiplicatively* factors. In this class, we consider an additive trend and the seasonal factor as additive or multiplicative, see Figure 9.1. We produce *h*-step ahead forecasts:

For the additive model: For the multiplicative model:  $\hat{Y}_t(h) = S_t + h T_t + I_{t+h-s}$  $\hat{Y}_t(h) = (S_t + h T_t) * I_{t+h-s}$ 

Note: Seasonal factor is multiplied in the *h*-step ahead forecast.

Figure 9.1 – Different Seasonality Models with Additive Trend





Additive Model: Additive seasonal variability with an additive trend.

Multiplicativ Model: Multiplicative seasonal variability with an additive trend.

# Holt-Winters (HW) ES: Additive Model

Additive model (additive trend and additive seasonality) *h*-step ahead forecast:

$$\hat{Y}_t(h) = S_t + h T_t + I_{t+h-s}$$

where *s* is the number of periods in seasonal cycles (s = 4 for quarters).

• Components:

- The level,  $S_t$ , is a weighted average of seasonal adjusted  $Y_t$  and the non-seasonal forecast  $(S_{t-1} + T_{t-1})$ :

$$S_t = \alpha (Y_t - I_{t-s}) + (1 - \alpha)(S_{t-1} + T_{t-1})$$

- The trend,  $T_t$ , is a weighted average of  $T_{t-1}$  and the change in  $S_t$ .  $T_t = \beta(S_t - S_{t-1}) + (1 - \beta)T_{t-1}$  - The seasonality is also a weighted average of seasonal index of *s* last year,  $I_{t-s}$ , and the current seasonal index  $(Y_{t-1} - S_{t-1} - T_{t-1})$ :

$$I_{t} = \gamma (Y_{t} - S_{t-1} - T_{t-1}) + (1 - \gamma)I_{t-s}$$

• Summary:

The additive model produces the following *h*-step ahead forecast:

 $\hat{Y}_t(h) = S_t + h T_t + I_{t+h-s}$ 

We use three equations:

 $S_{t} = \alpha (Y_{t} - I_{t-s}) + (1 - \alpha)(S_{t-1} + T_{t-1})$  $T_{t} = \beta (S_{t} - S_{t-1}) + (1 - \beta) T_{t-1}$  $I_{t} = \gamma (Y_{t} - S_{t-1} - T_{t-1}) + (1 - \gamma)I_{t-s}$ 

We have only three smoothing parameters:

 $\alpha$  = level coefficient  $\beta$  = trend coefficient  $\gamma$  = seasonality coefficient

#### Holt-Winters (HW) ES: Multiplicative Model

In the multiplicative seasonal case (with an additive trend), we have the *h*-step ahead forecast:  $\hat{Y}_t(h) = (S_t + h T_t) * I_{t+h-s}$ 

Details for multiplicative seasonality -i.e., Yt/It- and additive trend

- The forecast,  $S_t$ , now shows the average  $Y_t$  adjusted  $(\frac{Y_t}{I_{t-s}})$ .

- The trend, Tt, is a weighted average of Tt-1 and the change in St.

- The seasonality is also a weighted average of  $I_{t-S}$  and the  $Y_t/S_t$ 

Then, the model has three equations:

 $S_{t} = \alpha \frac{Y_{t-1}}{I_{t-s}} + (1 - \alpha) (S_{t-1} - T_{t-1})$   $T_{t} = \beta (S_{t} - S_{t-1}) + (1 - \beta) T_{t-1}$  $I_{t} = \gamma \frac{Y_{t}}{S_{t}} + (1 - \gamma) I_{t-s}$ 

We think of  $(Y_t / S_t)$  as capturing *seasonal effects*, where *s* represents the number of periods in the seasonal cycles. For example, s = 4, for quarterly data; s = 12, for monthly data;

Again We have only three parameters:

 $\alpha$  = smoothing parameter  $\beta$  = trend coefficient  $\gamma$  = seasonality coefficient

Question: How do we determine these 3 parameters?

- Ad-hoc method:  $\alpha$ ,  $\beta$  and  $\gamma$  can be chosen as value between  $0.02 < \alpha$ ,  $\gamma$ ,  $\beta < 0.2$ 

- Optimal method: Minimization of the MSE, as in SES.

**Example:** An industrial firm uses HW ES to forecast sales next three quarters (h = 1, 2 & 3; with s = 4):

$$\hat{Y}_{t}(h) = \hat{Y}_{t+h} = (S_{t} + h T_{t}) * I_{t+h-s}$$
with  $S_{t}, T_{t}, \& I_{t}$  factors given by:  
 $S_{t} = \alpha \frac{Y_{t}}{I_{t-s}} + (1 - \alpha) (S_{t-1} + T_{t-1})$   
 $T_{t} = \beta (S_{t} - S_{t-1}) + (1 - \beta) T_{t-1}$   
 $I_{t} = \gamma \frac{Y_{t}}{S_{t}} + (1 - \gamma) I_{t-s}$ 

The firm estimates:  $\alpha = 0.25$ ;  $\beta = 0.1$ ; and  $\gamma = 0.4$ . The firm observes  $Y_t = 5$ ; last quarter's smoothed forecasts:  $S_{t-1} = 3$ , &  $T_{t-1} = 1.2$ ; and last year's seasonal factors:  $I_{t-4} = 1.1$ ,  $I_{t-3} = 0.7$  &  $I_{t-2} = 1.2$ , &  $I_{t-2} = 0.8$ .

• Components forecasts:

 $S_t = 0.25 * \frac{5}{1.1} + (1 - 0.25) * (3 + 1.2) = 4.2864$   $T_t = 0.1 * (4.2864 - 3) + (1 - 0.1) * 1.2 = 1.2086$  $I_t = 0.4 * \frac{5}{4.2864} + (1 - 0.4) * 1.1 = 1.1266$ 

The forecast for h = 1 (next quarter) is:  $\hat{Y}_{t+1} = (4.2864 + 1.2086) * 0.7 = 4.8125$ 

The forecast for h = 2 & 3 are:  $\hat{Y}_{t+2} = (4.2864 + 2 * 1.2086) * 1.2 =$ 

 $\hat{Y}_{t+2} = (4.2864 + 2 * 1.2086) * 1.2 = 7.8475.$  $\hat{Y}_{t+3} = (4.2864 + 3 * 1.2086) * 0.8 = 6.1329.$ 

#### Holt-Winters (HW) ES: Initial Values

We have three recursive equations. That is, we need initial values for  $S_0$ ,  $T_0$  and  $I_{t-s}$ . To calculate initial values for the algorithm, we need at least one complete season of data to determine the initial estimates.

Like in the SES model, there are different approaches. Below, we present one approach for the multiplicative model:

- Initial values for S<sub>0</sub> and T<sub>0</sub>:

$$S_{0} = \frac{\sum_{t=1}^{S} Y_{t}}{sT_{0}}$$
  

$$T_{0} = \frac{1}{s} \left( \frac{Y_{s+1} - Y_{1}}{s} + \frac{Y_{s+2} - Y_{2}}{s} + \dots + \frac{Y_{s+s} - Y_{s}}{s} \right)$$
  
or 
$$T_{0} = \left[ \left\{ \sum_{t=1}^{S} Y_{t} / s \right\} - \left\{ \sum_{t=s+1}^{2s} Y_{t} / s \right\} \right] / s$$

- Initial values for *I*<sub>*t*-*s*</sub>.:

Assume we have *T* observation and quarterly seasonality (s=4): (1) Compute the averages of each of *T* years.

 $A_t = \sum_{i=1}^4 Y_{t,i}/4$ ,  $t = 1, 2, \dots, 6$  (yearly averages)

(2) Divide the observations by the appropriate yearly mean:  $Y_{t,i}/A_t$ .

(3) Is is formed by computing the average  $Y_{t,i}/A_t$  per year:

$$I_s = \sum_{i=1}^{T} Y_{t,s} / A_t$$
  $s = 1, 2, 3, 4$ 

## Holt-Winters (HW) ES: Damped Model

We can damp the trend as the forecast horizon increases, using a parameter  $\phi$ . For the multiplicative model, see Figure 9.2, we have:

$$S_{t} = \alpha \frac{Y_{t-1}}{I_{t-s}} + (1-\alpha)(S_{t-1} - \phi T_{t-1})T_{t} = \beta(S_{t} - S_{t-1}) + (1-\beta)T_{t-1}I_{t} = \gamma \frac{Y_{t}}{S_{t}} + (1-\gamma)I_{t-s}$$

Then, the *h*-step ahead forecast:

$$\hat{Y}_t(h) = \{S_t + (1 + \phi + \phi^2 + \dots + \phi^{2h-1})T_t\} * I_{t+h-s}$$

Figure 9.2 – Damped Additive Trend with Multiplicative Seasonality



This model is based on practice: It seems to work well for industrial outputs. Not a lot of theory or clear justification behind the damped trend.

### **HW ES Models – Different Types**

We have many variations:

- 1. No trend and additive seasonal variability.
- 2. Additive seasonal variability with an additive trend.
- 3. Multiplicative seasonal variability with an additive trend.
- 4. Multiplicative seasonal variability with a multiplicative trend.
- 5. Dampened trend with additive seasonal variability.
- 6. Multiplicative seasonal variability and dampened trend.
- Q: Which model should be used?

A: Select the type of model to fit based on the presence of

- Trend additive or multiplicative, dampened or not
- Seasonal variability additive or multiplicative

# HW ES: Example – Log U.S. Vehicles Sales

**Example:** We want to forecast log U.S. monthly vehicle sales with HW. We use the R function *HoltWinters*(). 1\_car\_18 <- 1\_car[1:512] 1\_car\_ts <- ts(1\_car\_18, start = c(1976, 1), frequency = 12) # convert lr\_d in a ts object hw\_d\_car <- HoltWinters(1\_car\_18, seasonal="additive") > hw\_d\_car Holt-Winters exponential smoothing with trend and additive seasonal component.

Call: HoltWinters(x = lr d ts, seasonal = "additive")

Smoothing parameters:	
alpha: <b>0.4355244</b>	$\Rightarrow$ Estimated smoothing parameter
beta : 0.009373815	$\Rightarrow$ Estimated trend parameter $\approx 0$ (no trend)
gamma:0.3446495	$\Rightarrow$ Estimated seasonal parameter
> hw_d_car	
Coefficients:	
[,1]	
a 7.177857555	$\Rightarrow$ forecast for level
b 0.0001100345	$\Rightarrow$ forecast for trend
s1 -0.075314457	$\Rightarrow$ forecast for seasonal month 1
s2 -0.084468361	$\Rightarrow$ forecast for seasonal month 2
s3 0.049447067	
s4 -0.273299309	
s5 -0.138251757	
s6 -0.026603921	
s7 -0.144953062	
s8 0.079214066	
s9 0.037899454	
s10 0.020477134	
s11 0.089309775	
s12 -0.012530316	
>plot(hw_d_car)	



```
• Now, we forecast one-step ahead forecasts
T last \leq- nrow(hw d car$fitted)
h <- 25
ses f hw <- matrix(0,h,1)
alpha <- 0.4355244
beta <- 0.009373815
gamma <- 0.3446495
y < -1 car
T \leq - length(l car)
sm \leq matrix(0,T,1)
Tr \leq matrix(0,T,1)
I \leq matrix(0,T,1)
T1 < T-h+1
a <- T1
sm[a-1] <- 7.177857555
Tr[a-1] <- -0.000309358
I[501:512] <- c(-0.075314457,-0.084468361,0.049447067,-0.273299309,-0.138251757, -
0.026603921, -0.144953062, 0.079214066, 0.037899454, 0.020477134, 0.089309775, -
0.012530316)
while (a \le T)
       sm[a] = alpha * y[a-1] + (1-alpha) * sm[a-1]
       Tr[a] = beta * (sm[a] - sm[a-1]) + (1 - beta) * Tr[a-1]
       I[a] = gamma * (y[a] - sm[a]) + (1 - gamma) * I[a - 12]
a <- a + 1
}
hh <- c(1:h)
car f hw <- sm[T1:T] + hh*Tr[T1:T] + I[T1:T]
car f hw
f error c hw<- car f hw - y[T1:T]
plot(car f hw, type="l", main ="SES Forecasts: Log Vehicle Sales")
```


MSE\_hw <- sum(f\_error\_c\_hw^2)/h > MSE\_hw [1] 0.01655964. ¶

### **HW ES: Remarks**

If a computer program selects  $\gamma = 0 = \beta$ , it has a lack of trend or seasonality. It implies a constant (deterministic) component. In this case, an ARIMA model with deterministic trend may be a more appropriate model.

- For HW ES, a seasonal weight near one implies that a non-seasonal model may be more appropriate.

We can model seasonalities as multiplicative or additive:

### **Evaluation of forecasts – Accuracy measures**

The mean squared error (*MSE*) and mean absolute error (*MAE*) are the most popular accuracy measures:

$$MSE = \frac{1}{m} \sum_{i=T+1}^{T+m} (\hat{y}_i - y_i)^2 = \frac{1}{m} \sum_{i=T+1}^{T+m} e_i^2$$
$$MAE = \frac{1}{m} \sum_{i=T+1}^{T+m} |\hat{y}_i - y_i| = \frac{1}{m} \sum_{i=T+1}^{T+m} |e_i|$$

where *m* is the number of out-of-sample forecasts.

But other measures are routinely used:

- Mean absolute percentage error  $(MAPE) = \frac{100}{T - (m-1)} \sum_{i=T+1}^{T+m} \left| \frac{\hat{y}_i - y_i}{y_i} \right|$ - Absolute  $MAPE (AMAPE) = \frac{100}{T - (m-1)} \sum_{i=T+1}^{T+m} \left| \frac{\hat{y}_i - y_i}{\hat{y}_i + y_i} \right|$ 

<u>Remark</u>: There is an asymmetry in MAPE, the level  $y_i$  matters.

- % correct sign predictions (PCSP) =  $\frac{1}{T-(m-1)} \sum_{i=T+1}^{T+m} z_i$ where  $z_i = 1$  if  $(\hat{y}_{i+l} * y_{i+l}) > 0$ = 0, otherwise. - % correct direction change predictions (PCDP)=  $\frac{1}{T-(m-1)} \sum_{i=T+1}^{T+m} z_i$ 

where  $z_i = 1$  if  $(\hat{y}_{i+l} - y_i) * (y_{i+l} - y_i) > 0$ = 0, otherwise.

<u>Remark</u>: We value forecasts with the right direction (sign) or forecast that can predict turning points. For stock investors, the sign matters!

MSE penalizes large errors more heavily than small errors, the sign prediction criterion, like MAE, does not penalize large errors more.

Example: We compute MSE and the % of correct direction change (PCDC) predictions for the
one-step forecasts for U.S. monthly vehicles sales based on the SES and HW ES models.
> MSE\_ses
[1] 0.027889
> MSE\_hw
[1] 0.01655964

```
We calculate PCDC with following script for HW and SES:

bb_hw <- (car_f_hw - y[(T1-1):(T-1)]) * (y[T1:T] - y[(T1-1):(T-1)])

indicator_hw <- ifelse(bb_hw > 0,1,0)  # ifelse ("if else") produces a 1 if condition is true

pcdc_hw <- sum(indicator_hw)/h

> indicator_hw

[1] 1 1 1 0 1 1 1 1 1 1 0 1 1 1 1 0 1 1 1 1 1 0 0 0

> pcdc_hw

[1] 0.76

bb_s <- (ses_f_c - y[(T1-1):(T-1)]) * (y[T1:T] - y[(T1-1):(T-1)])

indicator_s <- ifelse(bb_s > 0,1,0)

pcdc_s <- sum(indicator_s)/h

> indicator_s

[1] 1 0 1 1 1 1 1 1 1 1 1 1 0 1 1 1 0 1 1 0 0 0
```

<u>Note</u>: Same percentage of correct direction change (PCDC) predictions, but the sequence of correct predictions is not the same. ¶

### **Evaluation of forecasts – DM Test**

> pcdc\_s [1] 0.76.

To determine if one model predicts better than another, we define the loss differential between two forecasts:

$$d_t = g(e_t^{M1}) - g(e_t^{M2})$$

where g(.) is the forecasting loss function. M1 and M2 are two competing sets of forecasts – could be from models or something else.

We only need  $\{e_t^{M1}\}$  &  $\{e_t^{M2}\}$ , not the structure of M1 or M2. In this sense, this approach is "*model-free*."

Typical (symmetric) loss functions:  $g(e_t) = e_t^2 \& g(e_t) = |e_t|$ .

But other g(.)'s can be used:  $g(e_t) = exp(\lambda e_t^2) - \lambda e_t^2$  ( $\lambda > 0$ ).

Then, we test the null hypotheses of equal predictive accuracy:

H<sub>0</sub>: 
$$E[d_t] = 0$$
  
H<sub>1</sub>:  $E[d_t] = \mu \neq 0$ .

- Diebold and Mariano (1995) assume  $\{e_t^{M1}\}$  &  $\{e_t^{M2}\}$  is covariance stationarity and other regularity conditions (finite Var[dt], independence of forecasts after  $\ell$  periods) needed to apply CLT. Then,

$$\frac{\bar{d}-\mu}{\sqrt{Var[\bar{d}]/T}} \xrightarrow{d} N(0,1), \qquad \bar{d} = \frac{1}{m} \sum_{i=T+1}^{T+m} d_i$$

• Then, under H<sub>0</sub>, the DM test is a simple *z*-test:

$$DM = \frac{\tilde{d}}{\sqrt{\hat{V}ar[\tilde{d}]/T}} \xrightarrow{d} N(0,1)$$

where  $\hat{V}ar[\bar{d}]$  is a consistent estimator of the variance, usually based on sample autocovariances of  $d_i$ :

$$\widehat{V}ar[\overline{d}] = \gamma(0) + 2\sum_{j=k}^{\ell} \gamma(j)$$

There are some suggestion to calculate small sample modification of the DM test. For example, :  $DM^* = DM/\{[T+1-2 \ell + \ell (\ell-1)/T]/T\}^{1/2} \sim t_{T-1}.$ 

where  $\ell$ -step ahead forecast. If ARCH is suspected, replace  $\ell$  with  $[0.5 \sqrt{T}] + \ell$ .

<u>Note</u>: If  $\{e_t^{M1}\}$  &  $\{e_t^{M2}\}$  are perfectly correlated, the numerator and denominator of the DM test are both converging to 0 as  $T \rightarrow \infty$ .

 $\Rightarrow$  Avoid DM test when this situation is suspected (say, two nested models.) Though, in small samples, it is OK.

```
Code in R
dm.test <- function (e1, e2, h = 1, power = 2) {
d <- c(abs(e1))^power - c(abs(e2))^power
d.cov <- acf(d, na.action = na.omit, lag.max = h - 1, type = "covariance", plot = FALSE)$acf[, ,
1]
d.var <- sum(c(d.cov[1], 2 * d.cov[-1]))/length(d)
dv <- d.var  #max(1e-8,d.var)
if(dv > 0)
STATISTIC <- mean(d, na.rm = TRUE) / sqrt(dv)
else if(h==1)
```

```
stop("Variance of DM statistic is zero")
else
{
    warning("Variance is negative, using horizon h=1")
    return(dm.test(e1,e2,alternative,h=1,power))
}
    n <- length(d)
k <- ((n + 1 - 2*h + (h/n) * (h-1))/n)^(1/2)
STATISTIC <- STATISTIC * k
names(STATISTIC) <- "DM"
}</pre>
```

**Example**: We compare the SES and HW forecasts for the log of U.S. monthly vehicle sales. We use the *dm.test* function, part of the forecast package. library(forecast)

> dm.test(f\_error\_c\_ses, f\_error\_c\_hw, power=2)
 Diebold-Mariano Test
data: f\_error\_c\_sesf\_error\_c\_hw
DM = 1.6756, Forecast horizon = 1, Loss function power = 2, p-value = 0.1068
alternative hypothesis: two.sided

```
> dm.test(f_error_c_ses,f_error_c_hw, power=1)
    Diebold-Mariano Test
data: f_error_c_sesf_error_c_hw
DM = 1.94, Forecast horizon = 1, Loss function power = 1, p-value = 0.064
alternative hypothesis: two.sided
```

<u>Note</u>: Cannot reject H<sub>0</sub>: MSE<sub>SES</sub> = MSE<sub>HW</sub> at 5% level. ¶

### **Evaluation of forecasts – DM Test: Remarks**

The DM tests is routinely used. Its "model-free" approach has appeal. There are model-dependent tests, with more complicated asymptotic distributions.

The loss function does not need to be symmetric (like MSE).

The DM test is based on the notion of unconditional –i.e., on average over the whole sample-expected loss.

Following Morgan, Granger and Newbold (1977), the DM statistic can be calculated by regression of  $d_t$ , on an intercept, using NW SE. But, we can also condition on variables that may explain  $d_t$ . We move from an unconditional to a conditional expected loss perspective.

### **Combination of Forecasts**

Idea – from Bates & Granger (Operations Research Quarterly, 1969):

- We have different forecasts from R models:

 $\widehat{Y}_T^{M1}(\ell), \widehat{Y}_T^{M2}(\ell), \dots, \widehat{Y}_T^{MR}(\ell)$ 

Question: Why not combine them?  $\hat{Y}_T^{Comb}(\ell) = \omega_{M1}\hat{Y}_T^{M1}(\ell) + \omega_{M2}\hat{Y}_T^{M2}(\ell) + \dots + \omega_{MR}\hat{Y}_T^{MR}(\ell)$ 

Very common practice in economics, finance and politics, reported by the press as "consensus forecast." Usually, as a simple average.

Question: Advantage? Lower forecast variance. Diversification argument.

<u>Intuition</u>: Individual forecasts are each based on partial information sets (say, private information) or models.

The variance of the forecasts is:

$$Var[\hat{Y}_{T}^{Comb}(\ell)] = \sum_{j=1}^{R} (\omega_{Mj})^{2} Var[\hat{Y}_{T}^{Mj}(\ell)] + 2\sum_{j=1}^{R} \sum_{i=j+1}^{R} \omega_{Mj} \omega_{Mi} Co \operatorname{var}[\hat{Y}_{T}^{Mj}(\ell) \hat{Y}_{T}^{Mi}(\ell)]$$

Note: Ideally, we would like to have negatively correlated forecasts.

Assuming unbiased forecasts and uncorrelated errors,

$$Var[\hat{Y}_T^{Comb}(\ell)] = \sum_{j=1}^{K} (\omega_{Mj})^2 \sigma_j^2$$

**Example:** Simple average:  $\omega_j = 1/R$ . Then,  $Var[\hat{Y}_T^{Comb}(\ell)] = 1/R^2 \sum_{i=1}^R \sigma_i^2$ .

Example: We combine the SES and HW forecast of log US vehicles sales: f\_comb <- (ses\_f\_c + car\_f\_hw)/2 f\_error\_comb <- f\_comb - y[T1:T] > var(f\_comb) [1] 0.0178981 > var(car\_f\_hw) [1] 0.02042458 > var(ses\_f\_c)

# [1] 0.01823237. ¶

### **Combination of Forecasts – Optimal & Regression Weights**

We can derived optimal weights -i,e.,  $\omega_j$ 's that minimize the variance of the forecast. Under the uncorrelated assumption:

$$\omega_{Mj} *= \sigma_j^{-2} / \sum_{j=1}^R \sigma_j^{-2}$$

The  $\omega_j^*$ 's are inversely proportional to their variances.

In general, forecasts are biased and correlated. The correlations will appear in the above formula for the optimal weights. For the two forecasts case:

$$\omega_{Mj} *= (\sigma_1^2 - \sigma_{12})/(\sigma_1^2 + \sigma_2^2 - 2\sigma_{12}) = (\sigma_1^2 - \rho\sigma_1\sigma_2)/(\sigma_1^2 + \sigma_2^2 - 2\rho\sigma_1\sigma_2)$$

Ideally, we would like to have negatively correlated forecasts.

- Granger and Ramanathan (1984) used a regression method to combine forecasts.
- Regress the actual value on the forecasts. The estimated coefficients are the weights.

$$y_{T+\ell} = \beta_1 \hat{Y}_T^{M1}(\ell) + \beta_2 \hat{Y}_T^{M2}(\ell) + \dots + \beta_R \hat{Y}_T^{MR}(\ell) + \varepsilon_{T+\ell}$$

Should use a constrained regression

- Omit the constant
- Enforce non-negative coefficients.
- Constrain coefficients to sum to one

**Example:** We regress the SES and HW forecasts against the observed car sales to obtain optimal weights. We omit the constant  $\geq lm(u[T1:T] = aga f a + aga f b + aga$ 

```
> lm(y[T1:T] \sim ses_f_c + car_f_hw - 1)
```

```
Call:
lm(formula = y[T1:T] \sim ses_f_c + car_f_hw - 1)
```

Coefficients:  $ses_f_c car_f_hw$ -0.5426 1.5472

<u>Note</u>: Coefficients (weights) add up to 1. But, we see negative weights. In general, we use a constrained regression, forcing parameters to be between 0 and 1 (& non-negative). But, h=25 delivers not a lot of observations to do non-linear estimation. ¶

• Remarks:

- To get weights, we do not include a constant. Here, we are assuming unbiased forecasts. If the forecasts are biased, we include a constant.

- To account for potential correlation of errors, we can allow for ARMA residuals or include  $y_{T+l-1}$  in the regression.

- Time varying weights are also possible.

Question: Should weights matter? Two views:

- Simple averages outperform more complicated combination techniques.

- Sampling variability may affect weight estimates to the extent that the combination has a larger MSE.

### **Combination of Forecasts: Final Remarks**

• Since Bates and Granger (1969) and Granger and Ramanathan (1984), combination weights have generally been chosen to minimize a symmetric, squared-error loss function.

• But, asymmetric loss functions can also be used. Elliot and Timmermann (2004) allow for general loss functions (and distributions). They find that the optimal weights depend on higher order moments, such a skewness.

• It is also possible to forecast quantiles and combine them. We will not explore these issues in more detail in this class.

# Lecture 10 – Efficient Markets Hypothesis & Predictability

# **Efficient Markets Hypothesis (EMH)**

Questions: Can we predict stock returns? Can past information be used to build profitable trading rules in financial markets? In particular, can past return realizations tell us anything about expected future returns? Very old questions.

• The efficient markets hypothesis (EMH) is a first attempt to address the predictability issue.

• Earliest known version:

"When shares become publicly known in an open market, the value which they acquire there may be regarded as the judgement of the best intelligence concerning them." - George Gibson, *The Stock Exchanges of London, Paris and New York*, G. P. Putnman & Sons, New York, 1889.

• In 1900, Louis Bachelier, a French PhD student at the time, was the first to propose the *"Random Walk* Model" for security prices.

• Samuelson (1965)

"In an informationally efficient market, price changes must be unforecastable."

• Fama (1970)

"A market in which prices always *fully reflect* available information is '*efficient*'." If we have new information (a new earnings announcement) prices will adjust immediately (or very fast). Prices (significantly) jump with relevant information. But, they have to jump a proper amount, not too much (over-reaction) or not too little (under-reaction)!

• Grossman and Stiglitz (1980)

"There must be sufficient profit opportunities, i.e. inefficiencies, to compensate investors for the cost of trading and information-gathering."

The, under a frictionless world, it is impossible to have efficient prices (& EM). Only when all information gathering & trading costs are zero we can expect prices to fully reflect all available information.

But, if prices reflect fully and instantly all available information, who is going to gather information?

• Malkiel (1992)

"The market is said to be efficient with respect to some information set... implies that it is impossible to make *economic profits* by trading on the basis of [the information in that set]."

The first sentence of Malkiel's definition expands Fama's definition and suggests a test for efficiency useful in a laboratory.

The second sentence suggests a way to judge efficiency that can be used in empirical work. This is what is usually done in the finance literature.

**Example**: If Fund managers *outperform* the market consistently, then prices are not efficient with respect to their information set.

Many examples of "inefficiencies" with respect to some information sets.

The behavioral finance field has found that investors often show predictable and financially ruinous behavior (irrational?). Different causes: overreaction, overconfidence, loss aversion, herding, psychological accounting, miscalibration of probabilities, regret, etc.

**Examples:** Momentum strategies (buying past winners and selling past losers, under-reaction?) and Contrarian strategies (buying past losers and selling past winners, over-reaction?) achieve abnormal returns.

### • Lo (2004)

"... much of what behavioralists cite as counterexamples to economic rationality [...] are, in fact, consistent with an evolutionary model of individuals adapting to a changing environment." There is a time dimension. It takes time to adapt to new circumstances.

# **EMH: Versions**

Efficiency can only be defined with reference to a specific type of information. Fama (1970) defined three classes of information sets:

(a) Historical sequence of prices. This set gives Weak form EMH.

(b) Public records of companies and public forecasts regarding the future performance

possible actions. Sets (a) & (b) create the Semi-strong form EMH. (c) Private or inside information. Sets (a), (b) & (c) deliver the Strong form EMH.

• Violations:

and

- Technical traders devising profitable strategies (weak EMH)
- Reading a newspaper and devising a profitable trading strategy (semi-strong EMH)
- Corporate insiders making profitable trades (strong EMH).

Question: Can markets really be strong-form efficient? Very unlikely, plenty of examples of successful trading with private information: Jeffrey Skilling (Enron), Ivan Boesky & Michael Milken (junk bonds), Eugene Plotkin and David Pajcin (from Goldman Sachs, trading on M&A inside information), James McDermott Jr (Keefe, Bruytee & Woods, passed M&A tips to his mistress), Raj Rajaratnam (Galleon Group), Scott London (KPMG, passed tips from clients to a friend).

- Perfectly rational factors may account for violations of EMH:
  - Microstructure issues and trading costs.
  - Rewarding investors for bearing certain dynamic risks.
  - Time-varying expected returns due to changing conditions can generate predictability.

# **EMH: Joint Tests**

We are talking about economic profits, adjusting for risk and costs. Thus, a model for risk adjustment is needed. Results will be conditional on the underlying asset pricing model.

Fama (1991) remarks that tests of efficiency are *joint tests* of efficiency and some asset pricing model, or benchmark.

**Example**: Many benchmarks assume constant "normal" returns. This is easier to implement, but may not be correct. Thus, rejections of efficiency could be due to rejections of the benchmark.

Most tests suggest that if the security return (beyond the mean) cannot be forecasted, then market efficiency is not rejected.

**Example**: A wrong asset pricing model may reject efficiency. It would be easy to find (demeaned) returns to be forecastable if we use the wrong mean.

# **EMH: Expectations and Information Set**

The conditional expectation of the stochastic process  $X_{t+1}$ , conditioned on information set  $I_t$ , can be written as:

$$\mathbf{E}[\mathbf{X}_{t+1}|\mathbf{I}_t] = \mathbf{E}_t [\mathbf{X}_{t+1}]$$

Information set, I<sub>t</sub>: It describes what we know at time t. The usual assumption is that we do not forget anything. Over time, the information set increases: I<sub>t</sub> is contained in I<sub>t+1</sub>; I<sub>t+1</sub> is contained in I<sub>t+2</sub>, etc. That is, we have a sequence I<sub>0</sub>  $\subseteq$  I<sub>1</sub>  $\subseteq$  I<sub>2</sub>...  $\subseteq$  I<sub>t</sub>. In stochastic processes this sequence is called a "*filtration*," with notation { $\mathscr{F}_t$ }.

<u>Technical note</u>: We say a stochastic process  $\{X_t\}$  is *adapted* to a filtration  $\{\mathscr{F}_t\}$  if  $X_t$  is *measurable*  $\mathscr{F}_t$  for all t.

Measurable? The event of interest is in  $\mathcal{F}_t$ .

# **EMH: Random Prices**

Efficient market: A market where prices are random with respect to an information set (*"filtration"*), It.

Let the price of a security at time t be given by the expectation of some "fundamental value," V\*, conditional on  $I_t$ :

 $P_t = E[V^*|I_t] = E_t[V^*]$ 

The same equation holds one period ahead so that:  $P_{t+1} = E[\mathbf{V}^*|I_{t+1}] = E_{t+1}[\mathbf{V}^*]$ 

The expectation of the price change over the next period is:

 $E_t[P_{t+1} - P_t] = E_t[E_{t+1}[V^*] - E_t[V^*]] = 0$ since It is contained in It+1  $\Rightarrow$  Et[Et+1[V^\*]] = Et[V^\*] (by the law of IE).

<u>Remark</u>: Under efficiency, financial asset prices are unpredictable.

# EMH: Martingale & Fair Games

$$\begin{split} \textbf{Martingale:} \ A \ stochastic \ process \ P_t \ is \ a \ martingale \ if: \\ E[P_{t+1} \mid \Omega_t] = P_t \qquad (or \ E_t[P_{t+1}] = P_t) \\ where \ the \ information \ set \ is \ \Omega_t \ (what \ we \ know \ at \ time \ t, \ includes \ P_t). \end{split}$$

Submartingale:	If $E[P_{t+1} \mid \Omega_t] \ge P_t$ .	-Pt is a lower bound for $E_t[P_{t+1}]$
Supermartingale:	If $E[P_{t+1} \mid \Omega_t] \leq P_t$	-Pt is an upper bound for $E_t[P_{t+1}]$

Fair game model: A stochastic process  $r_t$  is a fair game if:

 $E[r_{t+1} | \Omega_t] = 0$  $\Rightarrow \text{ if } P_t \text{ is a martingale or pure random walk, } (P_{t+1} - P_t) \text{ is a fair game.}$ 

Note: Only referring to expected values!

The Martingale process can be setup as a special case of an AR(1) process:

 $p_t = \mu + \phi p_{t-1} + \varepsilon_t$ with  $\phi = 1$ ,  $\mu = 0$ , & E<sub>t</sub>[ $\varepsilon_{t+1}$ ] = 0. A non-stationary process. <u>Technical detail</u>: Martingale condition is neither a necessary nor a sufficient condition for *rational expectations* models of asset prices (LeRoy (1973), Lucas (1978)).

According to Lucas (1978), in markets where all investors have rational expectations, prices do fully reflect all available information and *marginal-utility weighted* prices follow martingales.

But, we consider the martingale as an important starting point.

# EMH: The Random Walk Hypothesis (RWH)

**Definition:** Random Walk (RW) A stochastic process  $p_t$  is a RW if:  $p_t = \mu + p_{t-1} + \varepsilon_t$  -where  $p_t = \ln(P_t)$ 

$$\Rightarrow r_t = \mu + \varepsilon_t = \Delta p_t$$

Assumptions about  $\varepsilon_t$ : Uncorrelated with past information, with constant mean (=0) & variance ( $\sigma^2$ ). That is,

$$\begin{split} \epsilon_t &\sim D \; (0, \, \sigma^2), \\ \text{with } E_t[\epsilon_{t+1}] = 0, \, E_t[\epsilon^2_{t+1}] = \sigma^2 \end{split}$$

If  $\mu \neq 0$ , the process is called a RW *with a drift*.

A RW with no drift is a martingale with structure for the error term,  $\varepsilon_t$ , uncorrelated, zero mean and constant variance.

• We start testing the EMH by assuming log returns,  $r_t$ , follow a RW with a drift. We called this "Random Walk Model":

 $\Rightarrow r_t = \Delta p_t = \mu + \varepsilon_t = \Delta p_t$ where  $\varepsilon_t \sim D (0, \sigma^2)$ .

• Different specifications for  $\varepsilon_t$  produce different testable hypothesis for the EMH-RW Model: - **RW1**:  $\varepsilon_t$  is *independent and identically distributed* (*i.i.d.*) ~ D(0,  $\sigma^2$ ). Not realistic. (Old tests: Cowles and Jones (1937)).

- **RW2**:  $\varepsilon_t$  is *independent* (allows for heteroskedasticity). Test using filter rules, technical analysis. (Alexander (1961, 1964), Fama (1965)).

- **RW3**:  $\varepsilon_t$  is *uncorrelated* (allows for dependence in higher moments). Test using autocorrelations, variance ratios, long horizon regressions.

### The RWH: Autocorrelations & ACF

Assume  $r_t$  is covariance stationary and ergodic. Then,

 $\gamma_k = \operatorname{cov}(r_t, r_{t-k})$  - Auto-covariance between times t & t - k  $\rho_k = \gamma_k / \gamma_0$ . is not time dependent. We estimate both statistics with  $\hat{\gamma}_k$  and  $\hat{\rho}_k$ . (Recall that  $\operatorname{Var}[r_t] = \gamma_0$ .)

• Under RW1 Hypothesis (and some assumptions)

$$\frac{\sqrt{T} \,\hat{\rho}_k \xrightarrow{a} N(0, 1)}{\operatorname{SE}[\hat{\rho}_k] = 1/\sqrt{T}}$$

 $\Rightarrow$ 

<u>Technical Note</u>: The sample correlation coefficients,  $\hat{\rho}_k$ , are negatively biased in finite samples. See Fuller (1976).

• To check autocorrelations up to order k, we use the ACF for  $r_t$ . Confidence Intervals can be easily approximated by  $\pm 2/\sqrt{T}$ .

**Example:** ACF with k = 24 lags for the monthly Equal- and Value-weighted (EW & VW, respectively) CRSP index returns from 1926:Jan – 2022:March (T = 1,155):

EMH\_da <- read.csv("http://www.bauer.uh.edu/rsusmel/4397/crsp\_ew\_vw\_m.csv", head=TRUE,sep=",")

lr_vw <- EMH_da\$vwretd	<pre># Value weighted CRSP returns (including distributions)</pre>
lr_ew <- EMH_da\$ewretd	# Equal weighted CRSP returns (including distributions)
T <- length(lr_vw)	
$SE_rho \le 1/sqrt(T)$	# Asymptotic SE for rho's:  rho  > 2 * SE => significant
> SE_rho	
[1] 0.02942449	#  rho  > 2 * SE => significant

## Monthly Value-Weighted  $acf y \leq acf(lr vw)$ > acf yAutocorrelations of series 'lr vw', by lag 1.000 -0.011 0.044 -0.183 0.140 -0.001 0.002 -0.010 0.121 -0.024 -0.003 -0.045 -0.002 0.045 0.009 -0.004 0.007 0.010 0.015 -0.010 -0.004 -0.005 0.051 -0.009 -0.015







<u>Conclusion for Monthly EW returns</u>: Again, a few significant autocorrelations, but, small in size.

**Example:** ACF with k = 24 lags for the daily Equal- and Value-weighted (EW & VW, respectively) CRSP index returns from 1926:Jan 1 – 2022:March 30 (T = 23,359):

EMH\_d\_da <- read.csv("http://www.bauer.uh.edu/rsusmel/4397/crsp\_ew\_vw\_d.csv", head=TRUE,sep=",")

lr\_vw\_d <- EMH\_d\_da\$vwretd</td># VW CRSP returns (including distributions)lr\_ew\_d <- EMH\_d\_da\$ewretd</td># EW CRSP returns (including distributions)T <- length(lr\_ew\_d)</td># EW CRSP returns (including distributions)SE\_rho <- 1/sqrt(T)</td># Asymptotic SE for rho's: |rho| > 2 \* SE => significant> SE\_rho# |rho| > 2 \* SE => significant

## Daily Value-Weighted acf  $y \le acf(lr vw d)$ > acf yAutocorrelations of series 'lr vw d', by lag 1.000 0.053 -0.027 0.001 0.018 0.007 -0.027 -0.003 0.002 0.019 0.014 0.010 0.023 -0.001 -0.001 -0.017 0.024 -0.010 -0.011 0.013 0.014 -0.008 -0.003 0.011 0.006



<u>Conclusion for daily VW returns</u>: There are many significant autocorrelations, with the exception of the first one, all very small.

## Daily Equal-Weighted acf  $y \leq -acf(lr ew d)$ > acf yAutocorrelations of series 'lr ew d', by lag 1.000 **0.198** 0.016 0.046 0.061 0.049 0.000 0.018 0.023 0.042 0.034 0.033 0.040 

0.015 0.014 0.004 0.021 0.007 0.008 0.033 0.025 0.003 0.009 0.023 0.007

Conclusion for daily EW rturns: Lots of significant autocorrelations, but, in general, small.

#### **The RWH: Autocorrelations Joint Tests**

We already know two tests to check for zero autocorrelation in a time series: Box-Pierce Q and Ljung-Box tests. We usually rely on the Ljung-Box (1978), LB, test, since it has better small sample properties.

-The Q & LB statistics test a joint hypothesis that the first *p* autocorrelations are zero:  $H_0: r_1 = ... = r_p = 0$ 

Under **RW1** and using the asymptotic distribution of  $\hat{\rho}_k$ :

$$Q = T \sum_{k=1}^{p} \hat{r}_{k}^{2} \xrightarrow{d} \chi_{p}^{2}.$$
  

$$LB = T * (T-2) * \sum_{k=1}^{p} \frac{\hat{r}_{k}^{2}}{T-k} \xrightarrow{d} \chi_{p}^{2}.$$

• Q & LB tests are widely use, but they have two main limitations:

(1) The test was developed under the independence (RW1) assumption.

If  $y_t$  shows dependence, such as heteroscedasticity, the asymptotic variance of  $\sqrt{T} \hat{\rho}$  is no longer I, but a non-diagonal matrix.

There are several proposals to "*robustify*" both Q & LB tests, see Diebold (1986), Robinson (1991), Lobato et al. (2001). The "robustified" Portmanteau statistic uses  $\tilde{\rho}_k$  instead of  $\rho_k$ :

$$\tilde{\rho}_{k} = \frac{\tilde{\gamma}_{k}}{\tau_{k}} = \frac{\sum_{t=k+1}^{T} (y_{t} - \bar{y})(y_{t-k} - \bar{y})}{\sum_{t=k+1}^{T} (y_{t} - \bar{y})^{2} (y_{t-k} - \bar{y})^{2}}$$

Thus, for Q we have:

$$Q^* = T \ \sum_{k=1}^p \tilde{\rho}_k^2 \xrightarrow{d} \chi_p^2.$$

(2) The selection of the number of autocorrelations *p* is arbitrary.

The traditional approach is to try different p values, say 3, 6 & 12. Another popular approach is to let the data "select" p, for example, using AIC or BIC, an approach sometimes referred as "*automatic selection*."

Escanciano and Lobato (2009) propose combining BIC's and AIC's penalties to select p in Q\* (BIC for small r and AIC for bigger r).

Note: It is common to reach different conclusion from Q and Q\*.

**Example:** Q and LB tests with p = 3 & 12 lags for the monthly EW & VW CRSP index returns from 1926:Jan – 2022:March (T = 1155):

• Q test for monthly VW

> Box.test(lr\_vw, lag = 4, type="Box-Pierce") Box-Pierce test
data: lr\_vw
X-squared = 22.812, df = 4, p-value = 0.000138
> Box.test(lr\_vw, lag = 12, type="Box-Pierce") Box-Pierce test
data: lr\_vw
X-squared = 34.696, df = 12, p-value = 0.0005234
• LB tests for monthly VW

> Box.test(lr\_vw, lag = 4, type="Ljung-Box")
Box-Ljung test
data: lr\_vw
X-squared = 22.891, df = 4, p-value = 0.0001332
> Box.test(lr\_vw, lag = 12, type="Ljung-Box")
Box-Ljung test
data: lr\_vw
X-squared = 34.87, df = 12, p-value = 0.0004912

Q\* tests with automatic lag selection. In R, the package *vrtest* has the Auto.Q function that computes this test. As always, you need to install *vrtest* first.

```
• Q* test for monthly VW
> Auto.Q(lr_vw, 12)
$Stat
[1] 3.059582
$Pvalue
```

```
[1] 0.08026232
```

<u>Conclusion for monthly VW returns</u>: Once we take into consideration potential heteroscedasticity in  $y_t$ , there is weak evidence for autocorrelation in monthly Value-weighted CRSP index returns.

• Q test for **monthly EW** > Box.test(lr\_ew, lag = 4, type="Box-Pierce") Box-Pierce test data: lr ew

X-squared = 61.607, df = 4, p-value = 1.333e-12

> Box.test(lr\_ew, lag = 12, type="Box-Pierce")
X-squared = 83.328, df = 12, p-value = 9.531e-13
• LB tests for monthly EW
> Box.test(lr\_ew, lag = 4, type="Ljung-Box")
X-squared = 61.793, df = 4, p-value = .218e-12
> Box.test(lr\_ew, lag = 12, type="Ljung-Box")
X-squared = 83.719, df = 12, p-value = 8.02e-13

• Q\* test for monthly EW library(vrtest)
> Auto.Q(lr\_ew, 12)
\$Stat
[1] 6.487553
\$Pvalue
[1] 0.01086324

<u>Conclusion for monthly EW returns</u>: Strong evidence for autocorrelation in monthly EW CRSP returns (the evidence was weaker, once we take into consideration potential heteroscedasticity in  $y_t$ , for monthly VW CRSP returns). That is, we reject the RW hypothesis for monthly EW CRSP returns.

**Example:** Q and LB tests with p = 5 & 20 lags for the daily Equal- and Value-weighted (EW & VW, respectively) CRSP index returns from 1926: Jan 1 – 2022 :March 30 (T = 25,359):

 $EMH\_d\_da <- read.csv("http://www.bauer.uh.edu/rsusmel/4397/crsp\_ew\_vw\_d.csv", head=TRUE, sep=",")$ 

lr\_vw\_d <- EMH\_d\_da\$vwretd # Value weighted CRSP returns (with distributions)
lr\_ew\_d <- EMH\_d\_da\$ewretd # Equal weighted CRSP returns (with distributions)
T <- length(lr\_ew\_d)
• Q tests for daily VW
> Box.test(lr\_vw\_d, lag = 5, type="Box-Pierce")
data: lr\_vw\_d
X-squared = 100.64, df = 5, p-value = 2.2e-16
> Box.test(lr\_vw\_d, lag = 20, type="Box-Pierce")
data: lr\_vw\_d
X-squared = 184.68, df = 20, p-value < 2.2e-16</pre>

```
• Q* test for daily VW (continuation)
> Auto.Q(y, 20) # Q* test automatic selection of p
$Stat
[1] 11.73454
$Pvalue
[1] 0.0006135076
```

• Q tests for daily EW > Box.test(lr\_ew\_d, lag = 5, type="Box-Pierce") data: lr\_ew\_d X-squared = 1213.3, df = 5, p-value = 2.2e-16 > Box.test(lr\_ew\_d, lag = 20, type="Ljung-Box") data: lr\_ew\_d X-squared = 1445.4, df = 20, p-value = 2.2e-16

• Q\* test for daily EW (continuation)
> Auto.Q(y, 40) # Q\* test automatic selection of p
\$Stat
[1] 235.7106
\$Pvalue
[1] 0

<u>Conclusion</u>: Strong evidence for autocorrelation in daily VW & EW CRSP returns. That is, we reject the uncorrelated returns hypothesis as implied by the RW hypothesis for daily VW & EW CRSP returns. ¶

### The RWH: Variance Ratio (VR) Test

<u>Intuition</u>: For all 3 RW hypotheses, the variance of RW increments is linear in the time interval. If the interval is twice as long, the variance must be twice as big. That is, the variance of monthly data should be 4 times bigger than the variance of weekly data. (Recall the log approximation rules for *i.i.d.* returns.)

If  $r_t$  is a covariance stationary process (constant first two moment, and covariance independent of time), then for the variance ratio of 2-period versus 1-period returns, VR(2):

$$VR(2) = \frac{Var[r_t(2)]}{2^*Var[r_t]} = \frac{Var[r_t + r_{t+1}]}{2^*Var[r_t]} =$$

$$=\frac{\operatorname{Var}[r_{t}] + \operatorname{Var}[r_{t+1}] + 2\operatorname{Cov}[r_{t}, r_{t+1}]}{2^{*}\operatorname{Var}[r_{t}]} = \frac{2\sigma^{2} + 2\gamma_{1}}{2\sigma^{2}} = 1 + \rho_{1}$$

where  $r_t(2) = r_t + r_{t+1}$ 

• Three cases:

 $\rho_1 = 0 \Rightarrow VR(2) = 1$  (True under **RW1**, random walk)  $\rho_1 > 0 \Rightarrow VR(2) > 1$  (mean aversion)  $\rho_1 < 0 \Rightarrow VR(2) < 1$  (mean reversion)

The intuition generalizes to longer horizons:

$$\operatorname{VR}(q) = \frac{\operatorname{Var}[r_t(q)]}{q \operatorname{Var}[r_t]} = 1 + 2 \operatorname{Var}[r_{t-1}] \sum_{k=1}^{q-1} (1 - \frac{k}{q}) \rho_k.$$

The VR(q) is a particular linear combination of the 1<sup>st</sup> (q - 1) autocorrelation coefficients (with linearly declining weights).

• Under **RW1**, we have  $H_0: VR(q) = 1$ .  $H_1: VR(q) \neq 1$ .

<u>Technical Note</u>: Under **RW2** and **RW3**, VR(q) = 1 provided  $1/T \Sigma_t Var[r_t] \rightarrow \overline{\sigma}^2 > 0$ 

we need this assumption, since some "fat-tailed" distributions do not have a well-defined second moment.

• To do any testing we need the sampling distribution of the VRs (estimated variance ratios) under  $H_0$ : VR(q) = 1. We use the statistic:

$$\frac{\sqrt{Tq}}{\sqrt{2*(q-1)}}(\widehat{\mathrm{VR}}(q)-1) \xrightarrow{a} \mathrm{N}(\mathbf{0},1)$$

This is Cochrane's (1988) VR test. The test rejects  $H_0$  –i.e., the RWH – if the above statistic is greater in absolute value than **1.96**.

For the special case of q = 2, we use  $\sqrt{T} (\widehat{\text{VR}}(2) - 1) \xrightarrow{a} \text{N}(\mathbf{0}, 1)$ 

• Var[ $r_t(q)$ ] is computed using the **MLE formulation**, that is, dividing by *T*, not by (*T* - 1) (or T minus degrees of freedom).

**Example**: We have monthly data from Jan 1973. Then, we compute

$$\operatorname{Var}[r_t] = \frac{\sum_{t=1}^{T} (r_t - \bar{r})^2}{T}$$
$$\operatorname{Var}[r_t(2)] = \frac{\sum_{t=1}^{T} (r_t(2) - 2*\bar{r})^2}{T}.$$

<u>Note</u>: Since the tests are asymptotic tests, in this case, relying on the Normal distribution, dividing by *T* or by (T - k) does not make any difference. ¶

•  $Var[r_t(q)]$  is computed using **non-overlapping returns**.

**Example**: We compute **non-overlapping bi-monthly returns**, using monthly data from Jan 1973.

(1) monthly returns:  $r_t$  is computed as usual. For the first return:

 $r_{t=Jan\,73} = \ln(P_{t=Jan\,31,\,73}) - \ln(P_{t=Jan\,1,\,73})$ 

(2) bi-monthly returns. The first three  $r_t(2)$  are computed as:

 $r_{t=Feb \ 73}(2) = r_{t=Feb \ 73} + r_{t=Jan \ 73}$   $r_{t=Apr \ 73}(2) = r_{t=Apr \ 73} + r_{t=Mar \ 73}$  $r_{t=June \ 73}(2) = r_{t=June \ 73} + r_{t=May \ 73}$ 

<u>Note</u>: We have "clean data," with no introduced serial correlation. But, we lose observations. If we have 1,000 monthly returns, using non-overlapping bi-monthly returns we end up with only 500 observations.  $\P$ 

**Example:** We check the RW Hypothesis, under RW3, for the monthly CRSP EW and VW Index returns. In R, the package *vrtest* has functions to compute the above mentioned VR tests.

• VR tests for <b>monthly VW</b>	
library(vrtest)	
kvec <- c(2,3,12)	#Vector with different q
y <- <b>lr_vw</b>	
> vr_1 <- VR.minus.1(y, kvec)	# Stat should be close to 0 if RW
> vr_1	
\$VR.auto	# VR with Automatic ("optimal) q selection
[1] 0.1954746	
\$Holding.Periods	
[1] 2 3 12	
\$VR.kvec	(VR - 1) stat for each q=kvec[i]
[1] 0.1007011 0.1187365 0.1212423	
<pre>&gt; sqrt(T*kvec)/sqrt(2*(kvec-1))*vr 1\$VR.k</pre>	twee # VR test for each $q=kvec[i] \sim N(0,1)$
[1] 3.422358 3.494666 3.043158	
• VR tests for <b>monthly EW</b>	
> y <- lr ew	
$>$ vr_1 <- VR.minus.1(y, kvec)	# Stat should be close to 0 if RW
> vr 1	
\$VR.auto	# VR with Automatic ("optimal) q selection
[1] 0.1954746	

\$Holding.Periods [1] 2 3 12

\$VR.kvec [1] 0.2043236 0.2789327 0.2180176 (VR-1) stat for each q=kvec[i]

> sqrt(T\*kvec)/sqrt(2\*(kvec-1))\*vr\_1\$VR.kvec # VR
[1] 6.943998 8.209583 5.472199

# VR test for each q=kvec[i] ~ N(0,1)

<u>Conclusion</u>: Using the VR test (with q = 2, 3, 12), we reject the RW Hypothesis  $\Rightarrow$  tests are greater in absolute value than **1.96**.

# The RWH: Variance Ratio (VR) Test - Issues

Several issues has been raised regarding the VR's tests. The main issues are:

(1) Choice of q. In the previous examples, we have arbitrarily selected q. Similar to the situation with the Q and LB tests, there are suggestions to automatically (or "optimally," according to some loss function) select q. Choi (1999) is one example of this approach, (the *vrtest* R package uses this approach in the *Auto*. VR test).

(2) Poor asymptotic approximation. In simulations, it is found that the asymptotic Normal distribution is a poor approximation to the small-sample distribution of the VR statistic. The usual solution is to use a bootstrap (Kim's (2009) bootstrap gives the p-value of the automatic VR test in the *Auto*. VR function).

**Example:** We use VR tests with automatic selection and a bootstrap to check the RW Hypothesis for the monthly CRSP EW and VW Index returns. Again, we use AutoBoot.test function in R package *vrtest*.

• Automatic VR tests for **monthly VW** y <- lr\_vw > AutoBoot.test(y, nboot=1000, wild="Normal", prob=c(0.025,0.975))

(Automatic variance ratio test statistic as in Choi (1999))
(1+ weighted sum of autocorrelation up to the optimal order)

\$CI.VRsum 2.5% 97.5% 0.8323731 1.1927214

```
• Automatic VR tests for monthly EW
y \le lr ew
> AutoBoot.test(y, nboot=1000, wild="Normal",
prob=c(0.025,0.975))
# Choi (1999)
$test.stat
                            (Automatic variance ratio test statistic as in Choi (1999))
[1] 4.173898
                            (1+ weighted sum of autocorrelation up to the optimal order)
$VRsum
[1] 1.382554
$pval
[1] 0.021
$CI.stat
  2.5%
          97.5%
-3.262026 3.359002
$CI.VRsum
  2.5%
         97.5%
0.7687769 1.2610106
```

<u>Conclusion</u>: Using the Automatic VR test and a bootstrap, we have strong evidence against the RW Hypothesis for EW, but weak for VW.  $\P$ 

### The RWH: VR Tests – LM's Modifications

Lo & MacKinlay (LM, 1988, 1989) propose modifications to the test:

- Allow for **overlapping returns**, and, thus, using more observations. But, overlapping returns will be autocorrelated, even if underlying process is not. We need to adjust for this feature.

- Use unbiased estimators of variances -i.e., divide by (T - df).

$$M_1(q) = \frac{\sqrt{3*T*q}}{\sqrt{2*(2q-1)*(q-1)}} (\overline{VR}(q) - 1) \xrightarrow{a} N(0, 1),$$

where  $\overline{VR}(q)$  is the VR statistic computed using overlapping returns.

- Allow for possible **heteroscedasticity** of returns (more realistic)

$$M_2(q) = \frac{(\overline{VR}(q)-1)}{\sqrt{\phi(q)}} \xrightarrow{a} N(0, 1),$$

where

$$\phi(q) = \sum_{j=1}^{q} \left[\frac{2(q-j)}{q}\right]^2 * \left\{\frac{\sum_{t=j+1}^{T} (r_t - \bar{r})^2 (r_{t-j} - \bar{r})^2}{[\sum_{t=1}^{T} (r_t - \bar{r})^2]^2}\right\}.$$

**Example:** We check the RW Hypothesis, under RW3, for the monthly CRSP EW and VW Index returns using the LM's tests: M1 and M2. Again, we use the R package *vrtest*.

```
Automatic VR tests for monthly EW
y <- lr_ew</li>
Lo.Mac(y, kvec)  # LM's tests M1 & M2 ~ asymptotic N(0,1)
$Stats
M1 M2
k=2 6.943998 2.5480302
k=3 6.359116 2.5009114
k=12 1.976326 0.9975538
```

<u>Conclusion</u>: Strong rejection of RW using M1, especially for q = 2, 3; but, using M2 test with q = 12, we cannot reject the RW Hypothesis. Consistent with previous result, stronger evidence for EW returns than for VW returns. ¶

# The RWH: VR & LM Tests – Issues

Several issues has been raised regarding the LM's tests:

(1) Poor asymptotic approximation. The asymptotic standard normal distribution provides a poor approximation to the small-sample distribution of the VR statistic. LM's tests tend to be biased and right-skewed, in finite samples.

• Proposed solutions:

- Alternative asymptotic distributions, as in Richardson and Stock (1989) or Chen and Deo (2006).

- Bootstrapping, as in Kim (2006) or Malliaropulos and Priestley (1999).

(2) Joint tests. The LM's tests are individual tests, where  $H_0$  is tested for a specific value of q. But, under  $H_0$ , VR(q) = 1, for all q. LM's tests ignore the joint nature of testing for the RW Hypothesis.

• Proposed solutions:

- **RS statistic**, a Wald Test, as proposed by Richardson and Smith (1993):

$$\operatorname{RS}(q) = T\left(\mathbf{VR} - \mathbf{\iota}\right)' \Phi^{-1} T\left(\mathbf{VR} - \mathbf{\iota}\right) \xrightarrow{d} \chi_q^2.$$

where VR is the  $(q \times 1)$  vector of q sample variance ratios,  $\mathbf{i}$  is the  $(q \times 1)$  unit vector, and  $\Phi$  is the covariance matrix of VR.

- QP statistic, a Wald Test based on a "power transformed" VR statistic, as proposed by Chen and Deo (2006). QP asymptotically follows a  $\chi_q^2$  distribution. This test is a one-sided test (H<sub>1</sub>:

VR(q) < 1 for all q.)

- CD statistic, a join test, as proposed by Chow and Denning (1993):

 $CD = \sqrt{T} \max_{1 \le i \le m} |M_2(q_i)|$ which follows a complex distribution, the studentized maximum modulus [SMM] distribution with *m* and *T* degrees of freedom (*m* is the number of k values). This SMM distribution is tabulated in Hahn and Hendrickson (1971) and Stoline and Ury (1979).

In general, we use the simulated critical values obtained by simulations as done by Chow and Denning themselves or a bootstrap as in Kim (2006).

**Example:** We check the monthly LM test results using a bootstrap instead of the asymptotic distribution. We use the *Boot.test* function in the R package vrtest, which provides two bootstrapped p-values: one for the LM statistic and the other one for the CD statistic.

• VR tests for <b>monthly VW</b>	
> y <- <b>lr_vw</b>	
> Lo.Mac(y, kvec)	# LM's tests M1 & M2
\$Stats	
M1 M2	
k=2 3.422358 1.7485059	
k=3 2.706957 1.4241521	
k=12 1.099060 0.6373211	
> Boot.test(y, kvec, nboot=1000, \$Holding.Period [1] 2 3 12	wild="Normal", prob=c(0.025,0.975)) #Kim's Bootstrap
\$LM.pval	(Bootstrap p-values for the Lo-MacKinlav M2 tests)
[1] 0.067 0.157 0.503	
> Lo.Mac(y, kvec) \$Stats	# LM's tests M1 & M2
M1 M2	
k=2 3.422358 1.7485059	
k=3 2.706957 1.4241521	
k=12 1.099060 0.6373211	

> Boot.test(y, kvec, nboot=1000, wild="Normal" prob=c(0.025,0.975)) #Kim's Bootstrap \$Holding.Period

[1] 2 3 12

\$LM.pval (Bootstrap p-values for the Lo-MacKinlay M2 tests) [1] 0.067 0.157 0.503 (Bootstrap p-value for the Chow-Denning test) \$CD.pval [1] 0.153 \$CI (C.I. for Lo-Mackinlay M2 tests from Bootstrap distr) 2.5% 97.5% k=2 -1.825961 1.827630 k=3 -1.847447 1.855263 k=12 -1.712367 2.152280 # RS Wald test > Wald(y, kvec) \$Holding.Period [1] 2 3 12 \$Wald.stat [1] 12.42735 \$Critical.Values 10 5 1 percent [1] 6.251389 7.814728 11.344867 > Chen.Deo(y, kvec) # QP Wald test \$Holding.Period [1] 2 3 12 \$VRsum [1] 0.07335402 \$OPn [1,] **3.154226** \$ChiSQ.Quantiles 1 2 5 10 20 percent [1] 11.344867 9.837409 7.814728 6.251389 4.641628 • VR tests for **monthly EW** > y < -lr ew> Lo.Mac(y, kvec) # LM's tests M1 & M2 **\$Stats** M1 M2 k=2 6.943998 2.5480302 k=3 6.359116 2.5009114 k=12 1.976326 0.9975538 > Boot.test(y, kvec, nboot=1000, wild="Normal", prob=c(0.025, 0.975))#Kim's Bootstrap

\$Holding.Period [1] 5 20 60 \$LM.pval (Bootstrap p-values for the Lo-MacKinlay M2 tests) [1] 0.001 0.004 0.279 \$CD.pval (Bootstrap p-value for the Chow-Denning test) [1] 0.017 \$LM.pval (Bootstrap p-values for the Lo-MacKinlay M2 tests) [1] 0.001 0.004 0.279 \$CD.pval (Bootstrap p-value for the Chow-Denning test) [1] **0.017** \$CI (C.I. for Lo-Mackinlay M2 tests from Bootstrap distr) 2.5% 97.5% k=2 -1.754012 1.708415 k=3 -1.710910 1.816157 k=12 1.563058 2.092434 # RS Wald test > Wald(y, kvec) \$Holding.Period [1] 2 3 12 \$Wald.stat [1] **52.68679** \$Critical.Values\_10\_5\_1\_percent [1] 6.251389 7.814728 11.344867 # QP Wald test > Chen.Deo(y, kvec)

Schen.Deo(y, kvec \$Holding.Period [1] 2 3 12

\$VRsum [1] 0.1442001

\$QPn [1,] **6.524497** 

\$ChiSQ.Quantiles\_1\_2\_5\_10\_20\_percent [1] **11.344867 9.837409 7.814728 6.251389 4.641628** 

<u>Conclusion</u>: Consistent with previous result, solid evidence for the RW for EW returns, but weak evidence (only the Wald test rejects  $H_0$ ) for VW returns. ¶

**Example:** We check the RW Hypothesis, under RW3, for the daily CRSP EW and VW Index returns.

• VR tests for **daily VW** kvec <- c(5, 20, 60)

#Vector with different q

y <- lr\_vw vr\_1 <- VR.minus.1(y, kvec) > vr\_1 \$VR.auto [1] 0.08049192

# Stat should be close to 0 if RW

(value of VR-1 with automatic selection of holding vectors)

\$Holding.Periods [1] 5 20 60

\$VR.kvec(the values of VR-1 for the chosen holding periods)[1] 0.060158750.111556930.16958754

> sqrt(T\*kvec)/sqrt(2\*(kvec-1))\*vr\_1\$VR.kvec # VR test for each q=kvec[i] (~ N(0,1) dist)
[1] 1.616329 2.750494 4.109789

> AutoBoot.test(y, nboot=300, wild="Normal", prob=c(0.025,0.975)) # Choi (1999)
\$test.stat
[1] 4.354851

\$VRsum [1] 1.080492

\$pval [1] 0.02333333

\$CI.stat 2.5% 97.5% -3.423189 4.067023 \$CI.VRsum 2.5% 97.5% 0.9483973 1.0656480

# LM's tests M1 & M2

> Boot.test(y, kvec, nboot=1000, wild="Normal", prob=c(0.025,0.975))#Kim's Bootstrap \$Holding.Period [1] 2 3 12

\$LM.pval (Bootstrap p-values for the Lo-MacKinlay M2 tests) [1] 0.06333333 0.08000000 0.07333333

\$CD.pval [1] 0.11333	(Bootstrap p-value for the Chow-Denning test)
\$CI 2.5% 97.5% k=5 -1.602225 2.333427 k=20 -1.594718 1.935643 k=60 -1.748524 1.782090	(C.I. for Lo-Mackinlay M2 tests from Bootstrap distrib)
> Wald(y, kvec) \$Holding.Period [1] 5 20 60	# RS Wald test
\$Wald.stat [1] 21.19834	
\$Critical.Values_10_5_1_percent [1] 6.251389 7.814728 11.344867	
> Chen.Deo(y, kvec) \$VRsum [1] 0.05863072	# QP Wald test
\$QPn [,1] [1,] <b>3.639522</b>	
\$ChiSQ.Quantiles_1_2_5_10_20_pe [1] <b>11.344867 9.837409 7.814728</b>	rcent 6.251389 4.641628. ¶

### **The RWH: Overall Evidence & Implications**

Tests results are based on CRSP value-weighted (VW) and equal weighted (EW) indices from 1925 & individual securities from 1962.

Daily, weekly and monthly returns from VW and EW indices show significant (positive) autocorrelation.

VR(q) > 1 statistics reject RW3 for EW index but not VW index. Market capitalization or size may be playing a role. Rejection of RW stronger for smaller firms. Their returns more serially correlated.

For individual securities,  $VR(q) \le 1$ , suggesting small and negative correlations (and not significant).

VR tests in other countries and financial markets. Tests also tend to reject the RWH, with stronger rejections for smaller markets and less liquid markets.

The rejection of the RWH does not necessarily imply a violation of the EMH.

Main implication: Theoretical pricing models should be able to explain the pattern of serial correlation.

Side Question: How can portfolios show VR(q) > 1 when individual securities show VR(q) < 1?

### **Predictability**

Traditional view pre 1980:

- CAPM is a good measure of risk
- Usual findings:
  - (a) Stock, bond and foreign exchange changes are not predictable
  - (b) Constant equity premium
- Market volatility does not change much through time

- Professional managers do not reliably outperform simple indices and passive portfolios once one corrects for risk

• Summary of State of the Art, late 1970s (Jensen, 1978):

"I believe there is no other proposition in economics which has more solid evidence supporting it than the Efficient Markets Hypothesis."

Modern view post-1980:

- Rejection of the RW Hypothesis.

- Stock returns are predictable.

Valuation ratios (D/P, E/P, B/M ratios)

Interest rates (term spread, short-long T-bill rates, etc.)

Decision of market participants (corporate financing, consumption).

Cross-sectional equity pricing.

Bond and foreign exchange returns are also predictable.

- Some funds seem to outperform simple indices, even after controlling for risk through market betas.

- New equilibrium (theory) models with time-varying equity premium.

# **Predictive Regressions**

Motivation:

1. Mounting evidence that stock and bond returns are predictable.

2. Q: Market inefficiency vs Rational variation in expected returns?

Economic questions:

1. Do the expected returns on bonds and stocks move together?

- 2. Do the same variables forecast bond and stock returns?
- 3. Is the variation in expected returns related to business cycles?

#### Setup:

Regress future returns,  $r_{t+\tau}$ , on variables  $x_t$  known at time t.

 $r_{t+\tau} = \boldsymbol{\mu}_t + \beta \boldsymbol{x}_t + \boldsymbol{\varepsilon}_{t+\tau}$ 

where  $\tau$  can be one month, one quarter, and one to four years.

# **Predictive Regressions – Fama-French (1989)**

One of the first papers to show a predictive pattern at different horizons. The setup of Fama and French (JFE, 1989):

(1)

-  $r_{t+\tau}$ : value- & equal-weighted market portfolios of NYSE; value-weighted corporate bond portfolios.

- x<sub>t</sub>variables:
  - Dividend yields,  $D_t/P_t$ : Add monthly dividends for the year preceding time t divided by the value of the portfolio at time t
  - Term Premium, *TERM<sub>t</sub>*: Long term government bond yield minus treasuries –see, Keim and Stambaugh (1986).
  - Default premium,  $DEF_t$ : AAA bond yields minus BAA bond yields –see, Keim and Stambaugh (1986).

<u>Sample</u>: Non-overlapping data for quarterly (T=244) & annual (T=61) data. For longer horizons (bi-annual+), overlapping observations.

• Findings:

-  $x_t$  variables work, especially  $D_t/P_t$  with high t-stats & high R<sup>2</sup> for forecast horizons beyond 1 year.

- (Conditional) Expected returns move with the predictors,  $x_t$ :

$$r_{t+\tau}] = \widehat{\boldsymbol{\mu}}_t + \widehat{\boldsymbol{\beta}} \boldsymbol{x}_t$$

That is, even with  $\mu_t = \mu$ , expected future returns are time-varying!

- Regression coefficients and R<sup>2</sup> increase with the forecast horizon.

• Interpretation of the Fama-French's slope estimate for  $D_t/P_t$  (similar of other financial ratios with  $P_t$  in the denominator):

- There is a positive relation between  $D_t/P_t$  and  $r_{t+\tau}$ . A high (low)  $D_t/P_t$  forecasts high (low) subsequent returns (higher  $P_{t+\tau}$ !). Since we tend to observe high  $D_t/P_t$  when  $P_t$  is low, we have evidence for **mean reversion** in stock prices.

- Let's look at the one-year  $D_t/P_t$  EW slope coefficient: 5.75. Then, a 1% increase in dividend increase expected (total) returns by 5.75% (an investor gets 1% dividend plus 4.75% extra return). **Big number**!

- Using the above 5.75 slope, we can derive an informal range for the expected 1-year return: In the past 40 years  $D_t/P_t$  ranged from 1% to 6%, ignoring  $\hat{\mu}_t$ , the range for  $E_t[r_{t+1y}]$  is {5.75% -

### 34%}. Very big!

• Interpretation of Fama-French's  $\mathbb{R}^2$  for  $D_t/P_t$  (again, similar interpretation for other ratios with  $P_t$  in the denominator):

-  $R^2$  are small, but they start to be worth paying attention to for horizons of 1-year ahead or longer. "Small" and "big" are relative term, remember that according to the RW the  $R^2$  should be 0! Then, any  $R^2 > 0$  is "interesting."

- For the EW returns,  $D_t/P_t$  predicts 7% of the variability of one-year ahead returns and 23% of the variability of 4-year ahead returns. These are results that, on average, can produce profitable investment strategies.

- Rational explanations for time-variation of expected return:
  - Time-varying risk aversion
  - Time-varying amount of risk
  - Parallel behavior explanation (investor sentiment).

<u>Remark</u>: We expect low prices –relative to  $D_t$ ,  $E_t$ ,  $Book_t$ – to be followed by high returns (high prices). Going back to the EMH, can we profit from this predictability?

<u>Note</u>: Another well-cited paper is Lamont (JF, 1998), who finds that other financial ratios also work as predictors: **dividends yield** & **earnings yield**. Lamont also find that the dividend payout ratio has cross-sectional predictive power.

**Example:** We use Shiller's data (1871:Jan - 2021:Dec) to redo the **monthly** predictive regressions of Fama-French (see FEc\_prog\_Pred for code and links to data).

	Independent Variable: Excess Returns at <i>t</i> +1 (1871-2021)						
	$r_t$	$D_t/P_t$	DY <sub>t</sub>	$E_t/P_t$	$D_t/E_t$	DFY <sub>t</sub>	DFR <sub>t</sub>
μ	0.00398 (0.0011)	0.00992 (0.0141)	0.01694 (0.0142)	0.02570 (0.0150)	-0.0183 (0.0195)	0.00979 (0.0025)	0.00435 (0.0030)
β	0.11256 (0.0234)	0.00095 (0.0025)	0.00218 (0.0025)	0.00410 (.0029)	-0.0041 (0.0035)	-0.0939 (0.0435)	0.11446 (0.2169)
R <sup>2</sup>	0.00731	0.0004	0.00001	0.0011	0.0008	0.0041	0.0002

<u>Findings</u>: With the exception of **lagged excess returns** and the **default yield spread** (AAA yield – BBB yield) nothing is significant.  $\P$ 

**Example:** We use Shiller's data (1925:Jan - 2021:Dec) to redo the **monthly** predictive regressions of Fama-French (see FEc\_prog\_Pred for code and links to data).

	Independent Variable: Excess Returns at <i>t</i> +1 (1871-2021)						
	r <sub>t</sub>	$D_t/P_t$	DY <sub>t</sub>	$E_t/P_t$	$D_t/E_t$	DFY <sub>t</sub>	DFR <sub>t</sub>
μ	0.00354 (0.0016)	0.02127 (0.0199)	0.02709 (0.0201)	0.02605 (0.0199)	-0.00118 (0.0277)	0.00724 (0.0030)	0.01038 (0.0025)
β	0.08548 (0.0294)	0.00296 (0.0034)	0.00396 (0.0034)	0.00424 (.0038)	-0.0009 (0.0049)	0.10056 (0.2165)	-0.0475 (0.0438)
R <sup>2</sup>	0.01267	0.0007	0.00117	0.00216	0.00002	0.00017	0.00104

<u>Findings</u>: Now, only **lagged excess returns** are significant. We just see "momentum" at work at the monthly level.  $\P$ 

# **Predictive Regressions: Methodological Issues**

• Data snooping. Are  $D_t/P_t$ ,  $TERM_t$ , Payout Ratios the only variables used in those regressions? The standard finance and economic databases used in academic and industry research (CRSP, Compustat, Refinivit) have thousands of potential predictors.

Recall Type I error: If we use 100 regressors, 5 will be significant at the 5% level!

• **Peso problem.** In the sample, we do not observe a "crash," which are very low probability events, but agents do compute that probability in the expectation. Then, on average, the sample average is biased!

• **Regime Change**. Always a potential problem. Maybe coefficients change with the business cycle, Fed policy, bull/bear markets, etc.

• **Endogeneity.** Regressors are only predetermined, but not exogenous. OLS slopes have a small bias (Stambaugh, 1986). Traditional OLS S.E. are likely not appropriate (Hodrick, 1992).

• Persistence of Financial Ratios. Valuation ratios are persistent and their innovations are correlated with returns, causing

- biased predictive coefficients: Stambaugh (1999)
- over-rejection by standard *t*-test: Cavanagh-Elliott-Stock (1995)

<u>Note</u>: These issues are less relevant for interest rates & recently proposed predictor variables (persistent, but less correlated with  $r_t$ ).

### **Predictive Regressions: Valuation Ratios – Persistence**

•  $D_t/P_t$  is persistent,  $D_t/P_t$  stays "high" or "low" for a long time. It moves around a constant mean (in red) & has no trend (stationary?).



There is some evidence for **mean reversion**, but it can take many years (decades?) to get back to the mean.

Given the persistence in  $D_t/P_t$ , the Fama-French results imply that we should also have persistence in the forecast of expected returns. That is, we have high (low) expected returns for a long time (decades?)!

Issue: How persistent is  $D_t/P_t$ ?

-  $D_t/P_t$  is likely to be persistent: it reflects long-run expectations.

- But, is  $D_t/P_t$  stationary? unit root? explosive?

To answer the above question, we compute the ACF for  $D_t/P_t$ . (Recall that a persistent series will show a slow decay in the ACF.)



The first order autocorrelation is **0.882**. Very persistent series! That is, next period dividend yield is very likely to be similar to this period.

There seems to be a relation (non-linear?) between  $D_t/P_t$  & the business cycle. We see big spikes in  $D_t/P_t$  when there is a recession (clear spike in the 1930s and in 2008-2009). Though these spikes are relatively short-lived (years, not decades).

Thus, expected returns vary with the business cycle (not a surprise): A big increase when there is a recession (risk is higher).

• Potential Problem with  $D_t$ : "too smooth" (measurement error?). The observed data may not be the "true" series of interest.

<u>Subtle point</u>: Since  $D_t$  is too smooth, all the predictability comes from  $P_t$ . What news affect more future stock prices (& returns): "Cash Flows news or Discount Rates news"? Discount rates news.

### **Predictive Regressions: Stambaugh Bias**

One econometric issue in Fama and French (1989): Regressors are only predetermined, but not exogenous.

• Start with predictive regression for returns,  $r_{t+1}$ :

 $\begin{aligned} r_{t+1} &= \alpha + \beta \; x_t + \; \varepsilon_{t+1} \\ x_t &: D_t / P_t & -\text{i.e., the dividend price ratio} \end{aligned}$ 

<u>Note</u>:  $x_t$  depends on the price at the beginning of t, the change of x at the end of t+1 reflects changes in price from t to t+1, as does  $r_{t+1}$ ;  $E[\varepsilon_{t+1}|x_{t+1}, x_t] \neq 0$ , more generally,  $E[\varepsilon_t|x_s, x_w] \neq 0$ , s < t < w.

Assumption (A2) is violated!

In addition,  $x_t$  is persistent. It can be modeled with an ARMA.

Stambaugh (1999) assumes that  $x_t$  follows an AR(1)  $x_t = \mu + \phi x_{t-1} + v_t$  (2) where  $v_t \& \varepsilon_t$  follow a multivariate N(0,  $\Sigma$ ), independent across t.

<u>Results</u>: b (OLS estimate) is biased upward, positively skewed, and has higher variance and kurtosis than the normal sampling distribution of the OLS estimator.

• Stambaugh bias:

 $E[b - \beta] = (\sigma \boldsymbol{\varepsilon} \nu / \sigma_{\nu}^{2}) E[\widehat{\boldsymbol{\phi}} - \boldsymbol{\phi}]$ 

It turns out  $\hat{\phi}$  has a downward bias and  $\sigma \epsilon \nu$  is negative

 $\Rightarrow$  b shows an upward bias. Conventional t-tests are misleading.

Finding: Correcting the bias weakens the predictability evidence.

Since conventional t-tests are misleading, there are many suggestions to check if the predictability of the very persistent valuation ratios remains after correcting for the bias.

One approach is Lewellen (2004): Adjust the OLS estimator under worst case scenario for persistence ( $\phi = 1$ ):

 $b_{adj} = b - (\sigma \epsilon \nu / \sigma_{\nu}^2) E[\hat{\phi} - 1]$ 

In practice, the estimated persistence is very close to one. The bias correction is small. Predictability survives:

- $D_t/P_t$  predicts market returns from 1946–2000 and sub-samples.
- B/M and  $E_t/P_t$  predict returns during the shorter sample 1963–2000.

Interesting Result: In a (1)-(2) framework NW SE are not reliable in small samples. Result from Hodrick (1992) & Kim and Nelson (1993).

### **Predictive Regressions: Long Horizon Returns (Aside)**

 $D_t/P_t$  and other ratios forecast excess returns on stocks. Regression coefficients and R<sup>2</sup> rise with the forecast horizon.

This is a result of the fact that the forecasting variable is **persistent**.

Model (1)-(2), assuming  $\alpha = \mu = 0$ .  $r_{t+1} = \alpha + \beta x_t + \varepsilon_{t+1}$ (1)  $\boldsymbol{x_t} = \boldsymbol{\mu} + \boldsymbol{\phi} \; \boldsymbol{x_{t-1}} + \boldsymbol{\nu_t} \tag{2}$ Now, we compound 2-period returns (with log returns, we add them):  $r_{t+2}(2) = \beta x_{t+1} + \varepsilon_{t+2} + \beta x_t + \varepsilon_{t+1}$ 

$$\begin{aligned} & = \beta x_{t+1} + \varepsilon_{t+2} + \beta x_t + \varepsilon_{t+1} \\ &= \beta (x_{t+1} + x_t) + \varepsilon_{t+2} + \varepsilon_{t+1} \\ &= \beta (\phi x_t + v_{t+1} + x_t) + \varepsilon_{t+2} + \varepsilon_{t+1} \\ &= \beta (1 + \phi) x_t + \beta v_{t+1} + \varepsilon_{t+2} + \varepsilon_{t+1} \\ &= \beta_2 x_t + \omega_{2t} \implies \beta_2 > \beta. \end{aligned}$$

• The previous result generalizes:

$$r_{t+k}(k) = \beta x_{t+k} + \varepsilon_{t+k} + \beta x_{t+k-1} + \varepsilon_{t+k-1} + \dots + \beta x_t + \varepsilon_{t+1}$$
  
=  $\beta (x_{t+k} + x_{t+k-1} + \dots + x_t) + \varepsilon_{t+k} + \dots + \varepsilon_{t+1}$   
=  $\beta (1 + \varphi + \varphi^2 + \dots + \varphi^k) x_t + \omega_{kt}$   
=  $\beta_k x_t + \omega_{kt} \implies \beta_k > \beta_{k-1}.$ 

The coefficient of the persistent ratio is increasing with the horizon of compounding returns.

Note: A more complicated derivation is needed for the increase in  $\mathbb{R}^2$ .

### **Predictive Regressions: More Predictors**

Lots of variables have been proposed as predictors. A short list:

- Book-to-market (b/mt-1), equity share in new issues (S, equist-1), and lagged returns, as in Baker and Wurgler (2000) (B-W, next slide).

- Cross-sectional premium (csp): The relative valuations of high- and low-beta stocks, as in Polk, Thompson, and Vuolteenaho (2006).

- *Net Equity Expansion* (ntis): The ratio of 12-month moving sums of net issues by NYSE listed stocks divided by the total end-of-year market capitalization, as in Boudoukh, et al. (2007).

- Long Term Yield (lty): Long-term government bond yields.

- Investment to Capital Ratio (i/k): The ratio of aggregate (private nonresidential fixed) investment to aggregate capital for the whole economy, as in Cochrane (1991).

- **Consumption, wealth, income ratio** (cay): Estimated from an equation from a model proposed by Lettau and Ludvigson (2001).

**Example:** We use the expanded Goyal and Welch data (1927 - 2021) to redo the **annual** predictive regressions of Baker-Wurgler, using S&P **excess returns** (see FEc\_prog\_Pred for code & data). Script for ik:

```
Pred_da_a <- read.csv("http://www.bauer.uh.edu/rsusmel/4397/goyal-welch-a_27.csv", head=TRUE, sep=",")
```

```
\label{eq:linear_sp} \begin{split} & || Value weighted S&P returns (with distributions) \\ & || k <- Pred_da_a \$ ik & # Investment-to-capital \\ & TA <- length(lr_sp) \\ & TI <- 21 \\ & y_a_i ik <- lr_sp[(TI+1):TA] - Rf_a[(TI+1):TA]/100 \\ & || k_a <- ik[TI:(TA-1)] \\ & fit_lag_y_i k <- lm(y_a_i k \sim ik_a) \\ & > summary(fit_lag_y_i k) \\ & Coefficients: \\ & Estimate Std. Error t value Pr(>|t|) \\ & (Intercept) & 0.23805 & 0.06206 & 3.836 & 0.000268 *** \end{split}
```

ik a	-0.07640	0.01747	-4.372 4.13e-05 ***
	0.07010	0.01/1/	

Independent Variable: Excess Returns at <i>t</i> +1 (1871-2021)							
$r_t$	$D_t/P_t$	DY <sub>t</sub>	$E_t/P_t$	$D_t/E_t$	DFY <sub>t</sub>	DFR <sub>t</sub>	
μ	-0.0087	0.02039	0.00511	-0.0152	-0.0212	0.04094	0.23805
----------------	----------	----------	----------	----------	----------	----------	----------
	(0.0096)	(0.0668)	(0.0626)	(0.0169)	(0.0214)	(0.0174)	(0.0621)
β	0.00234	0.00856	0.00501	0.00555	0.00023	-0.0027	-0.07640
	(0.1093)	(0.0195)	(0.0225)	(0.0119)	(0.0003)	(0.0008)	(0.0175)
R <sup>2</sup>	0.00001	0.0021	0.0005	0.0331	0.0046	0.1066	0.2121

<u>Findings</u>: Consistent with the previous table for VW returns, **equity share in new equity** is significant. We also run predictive regressions for the other variables mentioned above. **Investment-to-capital** (ik, starting in 1947) was very significant, with very high  $R^2$ . (Note: cay (starting in 1944) & csp (starting in 1937) were not significant).

# **Predictive Regressions: Way More Predictors**

With the advances in computer power, the success of finding predictors of future returns has continued almost exponentially. For example, using Machine Learning models (Neural Networks) we have:

- Gu, Kelly and Xiu (2020): **176 predictors**, grouped in **94 stock-level predictive characteristics** (Green et al. (2017)); **8 macroeconomic & financial variables predictors** (Welch and Goyal (2008)); and **74 industry dummies** (& even 94 \* 8 interaction terms!).

- Bianchi, Buchner and Tamoni (2021): **128 monthly macroeconomic and financial variables** (McCracken and Ng (2015)).

• Always keep in mind that the standard finance databases for research (CRSP & Compustat) have **over 1,000 potential predictors** (without counting interactions). It is always possible to find more predictors!

Question: Why not use them all?

#### **Predictive Regressions: In-sample vs Out-of-sample**

In a very well know paper, Goyal and Welch (2008) argue that the in-sample (IS) predictability seen in predictive regression, once evaluated out-of-sample (OOS), becomes very weak or just disappears.

- Setup of OOS Evaluation
- (1) Perform Q  $\tau$ -step-ahead forecasts using:

- Rolling predictive regressions, adding one observation at a time. That is, we obtain Q forecasts,  $\hat{r}_{t+\tau}$ .

- Use the mean of the rolling period at time t as the forecast. That is, we obtain Q forecasts,  $\bar{r}_t$ .
- (2) Get Q rolling forecast errors,  $e_A$ , & Q mean forecasts,  $e_N$ .

- (3) Compute  $MSE_A \& MSE_N$ .
- (4) Evaluate MSEs using the Diebold-Mariano test.
- An OOS R<sup>2</sup> can be computed as:

$$R_{OOS}^2 = 1 - \frac{MSE_A}{MSE_N}$$

with  $MSE_{A} = \sum_{t=1}^{Q} (r_{t+\tau} - \hat{r}_{t+\tau})^{2}$ 

$$MSE_N = \sum_{t=1}^Q (\boldsymbol{r}_{t+\tau} - \bar{r}_t)^2$$

<u>Note</u>: Goyal and Welch (2008) evaluate the MSEs using other tests, proposed by Clark and McCracken (2001) and McCracken's (2004) variation of the Diebold-Mariano test.

<u>Findings</u>: Very difficult to identify any robust predictor of excess stock returns. There are short time intervals of significant OOS predictability, but these "*pockets of predictability*" are surrounded by long periods of little or no predictability, see Lansing, LeRoy & Ma (2022).

**Example:** We use the expanded Goyal and Welch data (1927 - 2021) to compute their **annual** OOS R<sup>2</sup>, using rolling regressions starting in 1967, and perform Diebold-Mariano (DM) tests for significant differences of the forecasts (R script below for ik).

<u>Findings</u>: Consistent with the results of Goyal and Welch (2008), we do not find a lot of consistent predictability out of sample. In general, DM tests fail to reject  $H_0$  that the predictors do better than the unconditional mean in forecasting next year excess returns.

```
• R Code for ik (OOS rolling regressions)
yy <- y_a_ik
                                         # Dependent variable (y t+1) of rolling regression
                                         # Independent variable x_t
xx <- ik a
                                         # Initialize empty (a space to put forecasts errors)
Alles = NULL
                                         # Start of Rolling Sample
k for <-40
                                         # Counter for while loop
i \leq k for
TF \leq - length(yy)
while (i \leq TF-1) {
 y tp1 <- yy[1:i]
 x t <- xx[1:i]
 pred_reg <- lm (y_tp1 \sim x_t)
                                                 # OLS predictive regression
 b hat <- pred reg$coefficients
                                                 # Extract coefficient
 y_{hat} <- b_{hat}[1] + b_{hat}[2] * xx[i+1]
                                                 #t+1 forecast
 f e a \leq y hat - yy[i+1]
                                                 # t+1 forecast error for model
 f_e_n \le mean(y_tp1) - yy[i+1]
                                                 # t+1 forecast error for mean
 f 2e \leq c(f e a, f e n)
                                                 # Combine both forecast errors in a vector
 \overline{\text{Alles}} = r\overline{\text{bind}}(\overline{\text{Alles}}, \overline{f} 2e)
                                                 # accumulate forecast errors in rows (two columns)
 i <- i+1
}
```

# Checking accuracy of forecasts with OOS R^2 mse <- colSums(Alles^2)/(TF-k\_for) r2\_oos <- 1 - mse[1]/mse[2] > r2\_oos [1] 0.02177127 ⇒ Relative to IS results, big reduction in R<sup>2</sup>.

# Testing accuracy of forecasts with Diebold-Mariano
> dm.test(Alles[,1], Alles[,2], power=2)
 Diebold-Mariano Test
data: Alles[, 1]Alles[, 2]
DM = -0.12985, Forecast horizon = 1, Loss function power = 2, p-value = 0.8975
alternative hypothesis: two.sided

>dm.test(Alles[,1], Alles[,2], power=1) DM = -0.23874, Forecast horizon = 1, Loss function power = 1, p-value = 0.8128alternative hypothesis: two.sided. ¶

# **Predictive Regressions: Final Remarks**

There is a big and active literature on the predictability of stock returns, lately using ML/AI models. It has found lots of potential predictors of excess stock returns, for example, Gu, Kelly and Xiu (2020) use Neural Networks to discover **176 predictors** (with interaction terms, they almost use almost 1,000 predictors!)

Given the usual data mining results in large datasets, many of the discovered predictors are not "true predictors," but "false positive (FP) predictors." A lot of FP predictors will increase C.I. for forecasts.

We have a typical model selection problem. If we use the General-to-specific approach, the question is: How to reduce the GUM? Several proposals: optimize  $R_{OOS}^2$ , OOS SR, minimize FP predictors, etc.

Old question: Can we make money from these predictors? Not clear.

# Lecture 11 – Volatility Models

#### **Linear and Non-linear Models**

So far, we have focused on linear models. We have relied on Assumption (A1), where the relation between  $y_t \& X_t$  is given by:

$$y_t = X_t \beta + \varepsilon_t$$
,  $\varepsilon_t \sim i. i. d. D(0, \sigma^2)$ 

There are, however, many relationships in finance that are intrinsically non-linear: The payoffs to options are non-linear in some of the input variables, for example,  $S_t$ ; investors' willingness to trade off returns and risks are also non-linear; CEO compensation that depends on thresholds and with a big option component are also non-linear.

The textbook of Campbell *et al.* (1997) defines a non-linear data generating process as one where the current value of  $y_t$  is related non-linearly to current and previous values of the error term,  $\varepsilon_t$ :

$$y_t = f(\varepsilon_t, \varepsilon_{t-1}, \varepsilon_{t-2}, \ldots)$$

where  $\varepsilon_t$  is *i.i.d.* and *f* is a non-linear function.

A friendlier and slightly more specific definition of a non-linear model is given by the equation  $y_t = g(\varepsilon_t, \varepsilon_{t-1}, \varepsilon_{t-2}, ...) + \varepsilon_t \sigma^2(\varepsilon_t, \varepsilon_{t-1}, \varepsilon_{t-2}, ...)$ 

where g is a function of past error terms only, and 
$$\sigma^2$$
 can be interpreted as a variance term, since  
it is multiplied by the current value of the error.

Cases

- Non-linear in mean only: $g(\bullet) = \text{non-linear } \& \sigma^2(\bullet) = \sigma^2$ - Non-linear in variance only: $g(\bullet) = \text{linear } \& \sigma^2(\bullet) \neq \text{non-linear } g(\bullet)$ - Non-linear in mean and variance:both  $g(\bullet) \& \sigma^2(\bullet)$  are non-linear.

Most popular non-linear models in finance: The ARCH models, where we model a time-varying variance as a function of past  $\varepsilon_t$ 's.

#### **ARCH Models**

Until the early 1980s econometrics had focused almost solely on modeling the conditional means of time series, conditioning on information set at time t,  $I_t$ :

 $y_t = E[y_t | I_t] + \varepsilon_t, \quad \varepsilon_t \sim D(0, \sigma^2)$ Suppose we have an AR(1) process:

 $y_t = \alpha + \phi y_{t-1} + \varepsilon_t.$ The conditional mean is:

 $\mathrm{E}[y_{t+1}|\ I_t] = \mathrm{E}_{\mathrm{t}}[y_{t+1}] = \alpha + \phi \ y_t$ 

The unconditional mean and variance are:

$$E[y_t] = \frac{\alpha}{1 - \phi} \alpha / (1 - \phi) = \text{constant}$$
$$Var[y_t] = \frac{\sigma^2}{1 - \phi^2} = \text{constant}$$

Note: Conditional mean is time varying; unconditional mean is not!

Similar idea for the variance. For the AR(1) process, we have:

- Conditional variance:

 $\operatorname{Var}[y_{t+1}|I_{t}] = \operatorname{E}_{t}[(y_{t+1} - \operatorname{E}_{t}[y_{t+1}|I_{t}])^{2}] = \operatorname{E}_{t}[\varepsilon_{t+1}^{2}]$ 

- Unconditional variance:

$$Var[y_{t+1}] = E[(y_{t+1} - E[y_{t+1}])^2] = \frac{\sigma^2}{1 - \phi^2}$$

The unconditional variance measures overall uncertainty. In the AR(1) example, the information available at time t,  $I_t$ , plays no role, it is always a constant!

The conditional variance,  $Var[y_t|I_t]$ , is a better measure of uncertainty at time t. It is a function of information at time t,  $I_t$ .

Notation:  $E_t[Z_{t+1}] = E[Z_{t+1}|I_t]$ 

Summary:

- Unconditional variance measures the overall uncertainty.

- Conditional variance measures uncertainty at time *t*.

Remark: Conditional moments are time varying; unconditional moments are not!

# **ARCH Models: Stylized Facts of Asset Returns**

(1) *Thick tails*: Leptokurtic (thicker tails than Normal).

(2) Volatility clustering: "Large changes tend to be followed by large changes of either sign."

(3) *Leverage Effects*: Tendency for changes in stock prices to be negatively correlated with changes in volatility.

(4) *Non-trading Effects, Weekend Effects*: When a market is closed, information accumulates at a different rate to when it is open –for example, the weekend effect, where stock price volatility on Monday is not three times the volatility on Friday.

(5) *Expected events*: Volatility is high at regular times such as news announcements or other expected events, or even at certain times of day –for example, less volatile in the early afternoon.
(6) *Volatility and serial correlation*: Inverse relationship between the two.

(7) Co-movements in volatility: Volatility is positively correlated across markets/assets.

We need a model that accommodates all these (non-linear) facts.

Stylized facts (1) and (2) form the basis of Volatility (ARCH) Models.

• Easy to check leptokurtosis (Stylized Fact #1).

# Descriptive Statistics and Distribution for Monthly S&P500 Returns

	Statistic
Mean (%)	0. 0585 (p-value: 0.0004)
Standard Dev (%)	0.0449
Skewness	-0.7294
Excess Kurtosis	2. 6406
Jarque-Bera	216.15 ( <i>p-value</i> : <0.000001)



Note: Excess kurtosis greater than 0! Heavy tails are very common in financial time series.







Note: Periods with low changes, usually long, and periods of high changes, usually short. That is, volatility shows autocorrelation.

#### **ARCH Models: Engle (1982)**

We start with assumptions (A1) to (A5), but with a specific (A3'):

 $y_t = \gamma X_t + \varepsilon_t, \qquad \varepsilon_t \sim N(0, \sigma_t^2)$ (A3')  $\sigma_t^2 = Var_{t-1}[\varepsilon_t] = E_{t-1}[\varepsilon_t^2] = \omega + \alpha_1 \varepsilon_{t-1}^2 + \dots + \alpha_q \varepsilon_{t-q}^2$ which we can write, using the L operator, as:

$$\sigma_t^2 = \sum_{i=1}^q \alpha_i \varepsilon_{t-i}^2 = \omega + \alpha(L)\varepsilon^2$$

We can write the model in terms of an AR(q) for  $\varepsilon_t^2$ . Define  $v_t \equiv \varepsilon_t^2 - \sigma_t^2$ , -an error term for the variance.

Then,

 $\varepsilon_t^2 = \omega + \alpha(L)\varepsilon_t^2 + \nu_t$ 

Correlated  $\varepsilon_t^2$ 's: High (low) past  $\varepsilon_t^2$ 's produce a high (low)  $\varepsilon_t^2$  today.

The model

$$\sigma_t^2 = \sum_{i=1}^q \alpha_i \varepsilon_{t-i}^2 = \omega + \alpha(L)\varepsilon^2$$

is an AR(q) model for squared innovations,  $\varepsilon_t^2$ . We have the ARCH model: Auto-Regressive Conditional Heteroskedasticity.

The ARCH(q) model estimates the unobservable (*latent*) variance.

Non-negative constraints: Since we are dealing with a variance, we usually impose  $\omega > 0$  and  $\alpha_i > 0$ for all i.

Notation:  $E_{t-1}[\varepsilon_t^2] = E[\varepsilon_t^2 || I_{t-1}]$ 

<u>Useful result</u>: Since  $E[\varepsilon_t] = 0$ , then  $E_{t-1}[\varepsilon_t^2] = \sigma_t^2$ 

#### **ARCH Models: Unconditional Variance**

The unconditional variance is determined by:

 $\sigma^{2} = E[\sigma_{t}^{2}] = \omega + \sum_{i=1}^{q} \alpha_{i} E[\varepsilon_{t-i}^{2}] = \omega + \sum_{i=1}^{q} \alpha_{i} \sigma^{2}$ 

That is,

$$\sigma^2 = \frac{\omega}{1 - \sum_{i=1}^q \alpha_i}$$

To obtain a positive  $\sigma^2$ , we impose another restriction:  $(1 - \sum_{i=1}^{q} \alpha_i) > 0$ 

**Example:** ARCH(1)

$$Y_t = \beta X_t + \varepsilon_t, \qquad \varepsilon_t \sim N(0, \sigma_t^2)$$
  
$$\sigma_t^2 = \omega + \alpha_1 \varepsilon_{t-1}^2 \qquad \Rightarrow \sigma^2 = \frac{\omega}{1 - \alpha_1}$$

We need to impose restrictions:  $\omega > 0$ ,  $\alpha_1 > 0$ , &  $(1 - \alpha_1) > 0$ .

#### **ARCH Models: Leptokurtosis**

Errors may be serially uncorrelated, but they are not independent: There will be volatility clustering, which produces fat tails.

We want to calculate the kurtosis of the errors:

$$\kappa(\varepsilon_t) = E[\varepsilon_t^4] / E[\varepsilon_t^2]^2$$

We define standardized errors:  $z_t = \frac{\varepsilon_t}{\sigma_t}$ 

They have conditional mean zero and a time invariant conditional variance equal to 1. That is,  $z_t \sim N(0, 1)$ .

From the definition of  $z_t$  we have:  $\varepsilon_t = z_t \sigma_t$ 

Now, we compute the fourth (also central, since  $E[\varepsilon_t]=0$ ) moment:  $E[\varepsilon_t^4] = E[z_t^4]E[\sigma_t^4]$ 

Then, using Jensen's inequality:

 $E[\varepsilon_t^4] = E[z_t^4]E[\sigma_t^4] > E[z_t^4]E[\sigma_t^2]^2 = E[z_t^4]E[\varepsilon_t^2]^2$  $= 3 E[\varepsilon_t^2]^2$ 

$$\kappa(\varepsilon_t) = E[\varepsilon_t^4]/E[\varepsilon_t^2]^2 > 3.$$
  
where we have used the fact that since  $E[\varepsilon_t] = 0$ , then  $E[\varepsilon_t^2] = E[\sigma_t^2].$ 

<u>Technical point</u>: It can be shown that for an ARCH(1), the 4<sup>th</sup> moment for an ARCH(1):  $\kappa(\varepsilon_t) = \frac{3(1-\alpha^2)}{1-3\alpha^2}$  if  $3\alpha^2 < 1$ .

More convenient, but less intuitive, presentation of the ARCH(1) model:

$$y_t = \gamma X_t + \varepsilon_t$$
  

$$\varepsilon_t = \sigma_t v_t, \qquad v_t \sim D(0, 1)$$

that is,  $v_t$  is *i.i.d.* with mean 0, and  $Var[v_t]=1$ . Since  $v_t$  is *i.i.d.*, then:

$$E_{t-1}[\varepsilon_t^2] = E_{t-1}[\sigma_t^2 v_t^2] = E_{t-1}[\sigma_t^2] E_{t-1}[v_t^2] = \omega + \alpha_1 \varepsilon_{t-1}^2$$

which delivers the AR(1) representation for  $\varepsilon_t^2$ .

Also, if we assume  $v_t$  is normally distributed, then  $\varepsilon_t \sim N(0, \sigma_t^2)$ .

#### **GARCH Model: Bollerslev (1986)**

An early technique to determine q was to look at the ACF/PACF for squared returns,  $\varepsilon_t^2$ , which usually determined a very large q.

**Example:** We calculate the ACF and PACF for the squared of the **U.S. monthly stock returns** (1871-2020).



Squared U.S. Monthly Stock Returns (1871-2020)

<u>Note</u>: Highly autocorrelated squared returns. To accommodate the long autocorrelations, we use large q.

This result is not surprising,  $\sigma_t^2$  is a very persistent process. Persistent processes can be captured with an AR(*p*), where *p* is large. This is not efficient.

Following the idea of an ARMA process, we can use a more parsimonious representation of the ARCH model: The Generalized ARCH model or GARCH(q, p):

$$\sigma_t^2 = \omega + \sum_{i=1}^q \alpha_i \varepsilon_{t-j}^2 + \sum_{j=1}^p \beta_j \sigma_{t-j}^2$$
$$= \omega + \alpha(L)\varepsilon^2 + \beta(L)\sigma^2$$

which can be shown it is an ARMA(max(p,q), p) model for the squared innovations.

Popular GARCH model: GARCH(1,1):  $\sigma_{t+1}^2 = \omega + \alpha_1 \varepsilon_t^2 + \beta_1 \sigma_t^2$ with an unconditional variance:  $\operatorname{Var}[\varepsilon_t^2] = \sigma^2 = \frac{\omega}{1 - \alpha_1 - \beta_1}$   $\Rightarrow$  Restrictions:  $\omega > 0$ ,  $\alpha_1 > 0$ ,  $\beta_1 > 0$ ;  $(1 - \alpha_1 - \beta_1) > 0$ .

<u>Technical details</u>: This is *covariance stationary* if all the roots of  $\alpha(L) + \beta(L) = 1$ lie outside the unit circle. For the GARCH(1,1) this amounts to  $\alpha_1 + \beta_1 < 1$ .

Question: What should be the order of the GARCH Model? We should use enough lags to make sure the residuals do not have any more autocorrelation in the square residuals.

If the order of GARCH process is well determined, the ACF/PACF for  $\varepsilon_t^2$  should show no significant autocorrelations.

We can add lags until the tests for ARCH structure in the squared residuals, discussed later, are not longer significant.

• A GARCH(1,1) is a very good starting point.

# **GARCH-X**

In the GARCH-X model, exogenous variables are added to the conditional variance equation.

Consider the GARCH(1,1)-X model:

 $\sigma_t^2 = \omega + \alpha_1 \varepsilon_t^2 + \beta_1 \sigma_{t-1}^2 + \delta f(X_{t-l}, \theta),$ 

where  $f(X_t, \theta)$  is strictly positive for all *t*. Usually,  $X_t$ , is an observed economic variable or indicator, for example, a liquidity index, and f(.) is a non-linear transformation, which should be non-negative.

**Examples**: We can use 3-mo T-bill rates for modeling stock return volatility, or interest rate differentials between countries to model FX return volatility.

The US congressional budget office uses inflation in an ARCH(1) model for interest rate spreads.  $\P$ 

# **ARCH Estimation: MLE**

All of these models can be estimated by maximum likelihood. First we need to construct the sample likelihood.

Since we are dealing with dependent variables, we use the conditioning trick to get the joint distribution:

 $f(y_1, y_2, \dots, y_T; \theta) = f(y_1 | x_1; \theta) * f(y_2 | y_1, x_2, x_1; \theta) * f(y_3 | y_2, y_1, x_3, x_2, x_1; \theta) *$ \* ...\*  $f(y_T | y_{T-1}, \dots, y_1, x_{T-1}, \dots, x_1; \theta).$  Taking logs:

$$L = \log(f(y_1, y_2, \dots, y_T; \theta))) = \log(f(y_1|x_1; \theta)) + \log(f(y_2|y_1, x_2, x_1; \theta) + \dots + \log(f(y_T|y_{T-1}, \dots, y_1, x_{T-1}, \dots, x_1; \theta)))$$
  
=  $\sum_{t=1}^T \log(f(y_t|Y_{t-1}, X_t; \theta))$ 

We maximize this function with respect to the *k* mean parameters ( $\gamma$ ) and the *m* variance parameters ( $\omega$ ,  $\alpha$ ,  $\beta$ ).

**Example**: ARCH(1) model.

Mean equation:  $y_t = X_t \gamma + \varepsilon_t$ ,  $\varepsilon_t \sim N(0, \sigma_t^2)$ Variance equation:  $\sigma_t^2 = \omega + \alpha_1 \varepsilon_{t-1}^2$ 

We write the pdf for the normal distribution,

$$f(\varepsilon_t|\gamma,\omega,\alpha_1) = \frac{1}{\sqrt{2\pi\sigma_t^2}} \exp\left[-\frac{\varepsilon_t^2}{2\sigma_t^2}\right] = \frac{1}{\sqrt{2\pi\sigma_t^2}} \exp\left[-\frac{(y_t - X_t\gamma)^2}{2\sigma_t^2}\right]$$

We form the likelihood  $\mathcal{L}$  (the joint pdf):

$$\mathcal{L} = \prod_{t=1}^{T} \frac{1}{\sqrt{2\pi\sigma_t^2}} \exp\left(-\frac{\varepsilon_t^2}{2\sigma_t^2}\right) = (2\pi)^{-T/2} \prod_{t=1}^{T} \frac{1}{\sqrt{\sigma_t^2}} \exp\left(-\frac{\varepsilon_t^2}{2\sigma_t^2}\right)$$

We take logs to form the log likelihood,  $L = \log \mathcal{L}$ :

$$L = \sum_{t=1}^{T} \log(f_t) = -\frac{T}{2} \log(2\pi) - \frac{1}{2} \sum_{t=1}^{T} \log(\sigma_t^2) - \frac{1}{2} \sum_{t=1}^{T} \varepsilon^2_t / \sigma_t^2$$

Then, we maximize *L* with respect to  $\boldsymbol{\theta} = (\gamma, \omega, \alpha_1)$  the function *L*.

$$L = -\frac{T}{2}\log(2\pi) - \frac{1}{2}\sum_{t=1}^{T}\log(\omega + \alpha_{1}\varepsilon_{t-1}^{2}) - \frac{1}{2}\sum_{t=1}^{T}\varepsilon_{t}^{2}/(\omega + \alpha_{1}\varepsilon_{t-1}^{2})$$

Taking derivatives with respect to  $\theta = (\omega, \alpha_1, \gamma)$ , where  $\gamma$  is a vector of k mean parameters:

$$\frac{\partial L}{\partial \omega} = (-1/2) \sum_{t=1}^{T} 1/(\omega + \alpha_1 \varepsilon_{t-1}^2) - (-1/2) \sum_{t=1}^{T} \varepsilon_t^2 / (\omega + \alpha_1 \varepsilon_{t-1}^2)^2$$

$$\frac{\partial L}{\partial \alpha_1} = (-1/2) \sum_{t=1}^{T} \varepsilon_{t-1}^2 / (\omega + \alpha_1 \varepsilon_{t-1}^2) - (-1/2) \sum_{t=1}^{T} \varepsilon_t^2 \varepsilon_{t-1}^2 / (\omega + \alpha_1 \varepsilon_{t-1}^2)^2$$

$$\frac{\partial L}{\partial \gamma} = -\sum_{t=1}^{T} \mathbf{X}'_t \varepsilon_t / \sigma_t^2 \qquad (kx1 \text{ vector of derivatives})$$

We form the f.o.c.; that is, we write the first derivative vectors as  $\frac{\partial L}{\partial \theta}$  and, then, set it equal to 0:

$$\frac{\partial L}{\partial \theta} = S(y_t, \theta) = 0$$
 -a (k+2) system of equations

The vector of first derivatives is called the score vector,  $S(y_t, \theta)$ . Take the last f.o.c., the *k*x1 vector,  $\frac{\partial L}{\partial \gamma} = 0$ :

$$\frac{\partial L}{\partial \boldsymbol{\gamma}} = -\sum_{t=1}^{T} \boldsymbol{X}'_{t} \varepsilon_{t} / \sigma_{t,MLE}^{2} = \sum_{t=1}^{T} \boldsymbol{X}'_{t} (\boldsymbol{y}_{t} - \boldsymbol{X}_{t} \boldsymbol{\gamma}_{MLE}) / \sigma_{t,MLE}^{2} = 0$$
$$= \sum_{t=1}^{T} \frac{\boldsymbol{X}'_{t}}{\sigma_{t,MLE}} (\frac{\boldsymbol{y}_{t}}{\sigma_{t,MLE}} - \frac{\boldsymbol{X}_{t}}{\sigma_{t,MLE}} \boldsymbol{\gamma}_{MLE}) = 0$$

The last equation shows that MLE is GLS for the mean parameters,  $\gamma$ : each observation is weighted by the inverse of  $\sigma_{t,MLE}$ .

We have a (k+2) system. It is a non-linear system. The system is solved using numerical optimization (usually, with the Newton-Raphson method). ¶

<u>Technical Note</u>: If the conditional density for  $\varepsilon_t$  is well specified and  $\theta_0$  (the true parameter) belongs to the parameter space,  $\Omega$ , then

$$T^{\frac{1}{2}}(\widehat{\theta} - \theta_0) \rightarrow N(0, A_0^{-1}), \quad \text{where } A_0 = T^{-1} \sum_{t=1}^{I} \frac{\partial S_t(y_t, \theta_0)}{\partial \theta}$$

 $A_0$  is the matrix of second derivatives of the log likelihood, *L*. It is called the *Hessian*. In general, it is difficult to numerically compute and make sure it is positive definite (so it can be inverted), especially when the dimensions are big.

• There a lot of computational tricks to compute a Hessian that is invertible, the most popular algorithm is the Broyden–Fletcher–Goldfarb–Shanno, or "BFGS."

#### **ARCH Estimation: MLE – Standard Errors**

Under the correct specification assumption,  $A_0 = B_0$ , where

$$B_0 = T^{-1} \sum_{t=1}^{T} E[S_t(y_t, \theta_0), S_t(y_t, \theta_0)']$$

We estimate A<sub>0</sub> and B<sub>0</sub> by replacing  $\theta_0$  by its estimated MLE value,  $\theta_{MLE}$ .

The estimator  $B_0$  has a computational advantage over  $A_0$ . Only first derivatives are needed. But  $A_0 = B_0$  only if the distribution is correctly specified. This is very difficult to know in practice.

Common practice in empirical studies: Assume the necessary regularity conditions are satisfied.

# **ARCH Estimation: Numerical Optimization**

In general, we have a  $(k+m \ge k+m)$  system; *k* mean parameters and *m* variance parameters. But, it is a non-linear system. We use *numerical optimization*, which are methods that search over the parameter space looking for the values that maximize the log likelihood function.

In R, the function *optim* does numerical optimization. It minimizes any non-linear function. It needs as inputs:

- Initial values for the parameters,  $\theta_0$ .
- Function to be minimized (includes the GARCH process).
- Data used.
- Other optional inputs: Choice of method, hessian calculated, etc.

**Example:** *optim*(theta0, log\_lik\_garch11, data=z, method="BFGS", hessian=TRUE) theta0 = initial values log\_lik\_garch11 = function to be minimized. ¶

• Initial values:

- Numerical optimization needs initial values for  $\theta$ , say  $\theta_0$ . It is very common to find that the optimization is sensitive to the initial values. It is a good practice to try different sets of initial values.

We want to avoid selecting a local maximum:

• Initial values (continuation):

- Numerical optimization needs initial values for  $\theta$ , say  $\theta_0$ . It is very common to find that the optimization is sensitive to the initial values. It is a good practice to try different sets of initial values.

We want to avoid selecting a local maximum:



Figure 9.2 The problem of local optima in maximum likelihood estimation

- Given the autoregressive structure in  $\sigma_t^2$ , and sometimes we have AR(p) in the mean, we need to make assumptions about  $\sigma_0$  and the  $\varepsilon_0$ , ...,  $\varepsilon_q$  (and  $\varepsilon_0$ ,  $\varepsilon_1$ , ...,  $\varepsilon_p$  if we assume an AR(p) process for the mean).

<u>Usual assumptions</u>:  $\sigma_0$  = unconditional SD;  $\varepsilon_0 = \varepsilon_1 = ... = \varepsilon_p = 0$ .

- Alternatively, we can take  $\sigma_0$  (and  $\epsilon_0$ ,  $\epsilon_1$ , ...,  $\epsilon_p$ ) as parameters to be estimated (it can be computationally more intensive and estimation can lose power.)

# **ARCH Estimation: MLE – Example (in R)**

Log likelihood of AR(1)-GARCH(1,1) Model: log lik garch11 <- function(theta, data) {  $mu \le theta[1]$ ;  $rho1 \le theta[2]$ ;  $omega \le abs(theta[3])$ ;  $alpha1 \le abs(theta[4])$ ;  $beta1 \ge abs(theta[4])$ ;  $beta1 \ge$ abs(theta[5]); chk0 <- (1 - alpha1 - beta1) $r \leq ts(data)$  $n \leq - length(r)$  $u \leq vector(length=n); u \leq ts(u)$ u[1] = 0for (t in 2:n) $\{u[t] = r[t] - mu - rho1*r[t-1]\}$ # this setup allows for ARMA in mean  $h \le vector(length=n); h \le ts(h)$ h[1] = omega/chk0# set initial value for h[t] series if (chk0==0) {h[1]=.000001} # check to avoid dividing by 0 for (t in 2:n)  $\{h[t] = abs(omega + alpha1*(u[t-1]^2) + beta1*h[t-1])\}$ if  $(h[t] == 0) \{h[t] = .00001\}$ # check to avoid log(0) return(-1\*sum(- 0.5 \* log(abs(h[2:n])) - 0.5 \* (u[2:n]^2)/abs(h[2:n]))) } # I use optim to minimize a function, to maximize multiply by -1

**Example 1**: GARCH(1,1) model for **changes in CHF/USD**. We will use R function *optim* (*mln* can also be used) to maximize the likelihood function.

PPP da <read.csv("https://www.bauer.uh.edu/rsusmel/4397/ppp\_2020\_m.csv",head=TRUE,sep=",") x chf <- PPP da\$CHF USD # CHF/USD 1971-2020 monthly data  $T \leq - length(x chf)$  $z \le \log(x \operatorname{chf}[-1]/x \operatorname{chf}[-T])$ theta0 = c(-0.002, 0.026, 0.001, 0.19, 0.71)# initial values ml 2 <- optim(theta0, log lik garch11, data=z, method="BFGS", hessian=TRUE)  $\log L g_{11} \le \log lik garch_{11}(ml 2\par, z)$ # value of log likelihood logL g11 ml 2\$par # estimated parameters I Var m2 <- ml 2\$hessian eigen(I Var m2) # check if Hessian is pd. sqrt(diag(solve(I Var m2))) # parameters SE chf usd  $\leq$ - ts(z, frequency=12, start=c(1971,1)) plot.ts(chf usd) # time series plot of data # Log likelihood value  $> \log L g 11$ [1]-1745.197 # Extract from ml 2 function parameters > ml 2\$par  $[1] -0.0021051742 \ 0.0260003610 \ 0.00012375 \ 0.1900276519 \ 0.710\overline{0}718082$ 

>I Var m2 <- ml 2\$hessian # Extract Hessian (matrix of 2nd derivatives) > eigen(I Var m2) # Check if Hessian is pd to invert. eigen() decomposition # Eigenvalues: if positives => Hessian is pd \$values [1] 1.687400e+08 6.954454e+05 7.200084e+03 5.120984e+02 2.537958e+02 \$vectors [,1] [,3] [,2] [,4] [,5] [1,] 4.265907e-05 9.999960e-01 -0.0011397586 0.0018331957 -0.0018541203 [2,] -3.333961e-06 -2.188159e-03 -0.0010048203 0.9769058449 -0.2136566699 [3,] 9.999998e-01 -4.223001e-05 -0.0003544245 0.0001291633 0.0005770707 [4,] -3.599974e-06 -1.702277e-03 -0.8603563865 -0.1097470278 -0.4977344477 [5,] -6.893837e-04 6.416141e-04 -0.5096905472 0.1833226197 0.8405994743 > sqrt(diag(solve(I Var m2))) # Invert Hessian: Parameters Var on diag [1] 1.203690e-03 4.419049e-02 7.749756e-05 5.014454e-02 3.955411e-02 > t stats <- ml 2\$par/sqrt(diag(solve(I Var m2))) >t stats [1] -1.7489333 0.5883701 1.5967743 3.7895984 17.9519078 Summary for CHF/USD changes

• *T*: 562 (January 1971 - July 2020, monthly).

The estimated model for  $e_{f,t}$  is given by:  $e_{f,t} = -0.00211 + 0.02600 e_{f,t-1},$  (.0012) (0.044)  $\sigma_t^2 = 0.00012 + 0.19003 \varepsilon_{t-1}^2 + 0.71007 \sigma_{t-1}^2.$   $(0.00096)^* (0.050)^* (0.040)^*$ Unconditional  $\sigma = 0.00012 / (1 - 0.19003 - 0.71007) = 0.001201201$  Log likelihood: 1745.197

<u>Note</u>:  $\alpha_1 + \beta_1 = .90 < 1.$  (Persistent.) ¶



**Example 2**: Using Robert Shiller's monthly data set for the S&P 500 (1871:Jan - 2020:Aug, T=1,795), we estimate an AR(1)-GARCH(1,1) model:

 $\begin{aligned} \mathbf{r}_{t} &= [\log(\mathbf{P}_{t}) - \log(\mathbf{P}_{t-1})] = \mathbf{a}_{0} + \mathbf{a}_{1} \mathbf{r}_{t-1} + \varepsilon_{t}, \qquad \varepsilon_{t} \mid I_{t-1} \sim \mathrm{N}(0, \sigma_{t}^{2}) \\ \sigma_{t}^{2} &= \boldsymbol{\omega} + \boldsymbol{\alpha}_{1} \varepsilon_{t-1}^{2} + \boldsymbol{\beta}_{1} \sigma_{t-1}^{2} \end{aligned}$ 

The estimated model for st is given by:

$$\begin{aligned} \mathbf{r}_{t} &= & \mathbf{0.338} & + & \mathbf{0.278} \ \mathbf{r}_{t-1}, \\ & (.08)^{*} & (0.025)^{*} \\ \sigma_{t}^{2} &= & \mathbf{0.756} \ + & \mathbf{0.126} \ \varepsilon_{t-1}^{2} + \ \mathbf{0.826} \ \sigma_{t-1}^{2}. \\ & (0.151)^{*} & (0.017)^{*} \ & (0.021)^{*} \end{aligned}$$

Unconditional  $\sigma^2 = 0.756 / (1 - 0.126 - 0.826) = 15.4630$ Log likelihood: 4795.08

<u>Note</u>:  $\alpha_1 + \beta_1 = .952 < 1.$  (Very persistent.)



Above, we plot the time-varying variance. Certain events are clearly different, for example, the 1930 great depression, with a peak variance of 282 (18 times unconditional variance!). The covid-19 volatility similar to the 2008-2009 financial crisis recession. ¶

#### **GARCH:** Forecasting and Persistence

Consider the forecast in a GARCH(1,1) model:  $\sigma_{t+1}^2 = \omega + \alpha_1 \varepsilon_t^2 + \beta_1 \sigma_t^2 = \omega + \sigma_t^2 (\alpha_1 z_t^2 + \beta_1) \qquad (\varepsilon_t^2 = \sigma_t^2 z_t^2)$ 

Taking expectation at time t

 $E_t[\sigma_{t+1}^2] = \omega + \sigma_t^2(\alpha_1 1 + \beta_1)$ Then, by repeated substitutions:  $E_t[\sigma_{t+j}^2] = \omega * [\sum_{i=0}^{j-1} (\alpha_1 + \beta_1)^i] + \sigma_t^2 (\alpha_1 + \beta_1)^j$ 

Assuming  $(\alpha_1 + \beta_1) < 1$ , as  $j \to \infty$ , the forecast reverts to the unconditional variance:  $\sigma^2 = \omega/(1 - \alpha_1 - \beta_1)$ .

When  $\alpha_1 + \beta_1 = 1$ , today's volatility affect future forecasts forever:  $E_t[\sigma_{t+j}^2] = \sigma_t^2 + j\omega$ 

**Example 1**: We want to forecast next month (September 2020) variance for CHF/USD changes. Recall we estimated  $\sigma_t^2$ :

 $\sigma_t^2 = 0.00012 + 0.19003 \varepsilon_{t-1}^2 + 0.71007 \sigma_{t-1}^2.$ getting  $\sigma_{2020:9}^2 = 0.003672220$  (= $\sigma_{2020:9} = \text{sqrt}(0.00367) = 6.1\%$ ) We based the  $\sigma_{2020:10}^2$  forecast on:  $E_t[\sigma_{t+j}^2] = \omega * [\sum_{i=0}^{j-1} (\alpha_1 + \beta_1)^i] + \sigma_t^2 (\alpha_1 + \beta_1)^j$ 

Then,  $(\alpha_1 + \beta_1) = 0.190 + 0.710 = 0.900$  $E_{2020:9}[\sigma_{2020:10}^2] = 0.00012 + 0.00367 * (0.9) = 0.003423$ 

We also forecast  $\sigma^2_{2020:12}$ 

$$E_{2020:9}[\sigma_{2020:12}^2] = 0.00012 * \{1+(0.9)+(0.9)^2\} + 0.00367 * (0.9)^3 = 0.00300063$$

We forecast volatility for March 2021:

 $E_{2020:6}[\sigma_{2021:03}^2] = 0.00012 * \{1 + (0.9) + (0.9)^2 + \dots + (0.9)^5\} + 0.00367 * (0.9)^6$ = 0.002512659

<u>Remark</u>: We observe that as the forecast horizon increases  $(j \rightarrow \infty)$ , the forecast reverts to the unconditional variance:

 $\omega/(1 - \alpha_1 - \beta_1) = 0.00012/(1 - 0.9) = 0.0012$  $\Rightarrow \sigma = \text{sqrt}(0.0012) = 0.0346 \qquad (3.46\% \approx \text{close to sample SD} = 0.0346$ 

#### **3.36%**). ¶

**Example 2**: On August 2020, we forecast the December's variance for the S&P500 changes. Recall we estimated  $\sigma_t^2$ :

Recall we estimated  $\sigma_t^2$ :  $\sigma_t^2 = 0.756 + 0.125 \varepsilon_{t-1}^2 + 0.826 \sigma_{t-1}^2$ , getting  $\sigma_{2020:8}^2 = 43.037841$ 

We based the 
$$\sigma_{2020:12}^2$$
 forecast on:  

$$E_t[\sigma_{t+j}^2] = \omega * [\sum_{i=0}^{j-1} (\alpha_1 + \beta_1)^i] + \sigma_t^2 (\alpha_1 + \beta_1)^j$$

Then, since  $(\alpha_1 + \beta_1) = 0.952$  $E_{2020:8}[\sigma^2_{2020:12}] = 0.756 * \{1+(0.952)+(0.952)^2+(0.952)^3\} +$ 

 $+43.037841 * (0.952)^4 = 38.02797$ 

Lower variance forecasted for the end of the year, but still far from the unconditional variance of 15.4.  $\P$ 

#### **GARCH:** Forecasting – Application to VaR

**Example**: In September 2020, Swiss Cruises wants to construct a VaR-mean for the USD 1 M receivable in 30 days (October). Data Receivable: USD 1 M  $S_{t=2020:9} = 1.45$  CHF/USD  $e_{f,t=2020:9} = 0.01934126$ TE<sub>t=2020:9</sub> = USD 1M \* 1.45 CHF/USD = CHF 1.45M.

$$\begin{split} E_{2020:9}[\sigma_{2020:10}^2] &= \textbf{0.003423} \implies \operatorname{sqrt}(\textbf{0.003423}) = 0.05851 \ (5.85\%) \\ \operatorname{VaR-mean}(.99) &= \operatorname{CHF} \textbf{1.45M} * \{E_{2020:9}[e_{f,t=2020:10}] - 2.33 * \operatorname{sqrt}(E_{2020:9}[\sigma_{2020:10}^2])\} \end{split}$$

 $E_{2020:9}[e_{f,t=2020:10}] = -0.00211 + 0.026 * e_{f,t=2020:9} \\ = -0.00211 + 0.026 * 0.01934126 = -0.001607$ 

VaR-mean(.99) = CHF 1.45M \* (-0.001607 - 2.33 \* sqrt(0.003423)) = CHF -0.1999941 M

<u>Interpretation of VaR-mean</u>: Relative to today's valuation (or *expected valuation*, according to RWM), the maximum *expected loss* with a 99% "chance" is **CHF -0.20 M**.

We also derive this value, using the sample mean and sample SD: sample mean = -0.00259sample SD = 0.033357 $\Rightarrow$  VaR-mean(.99) = CHF 1.45M \* \* (-0.00259 - 2.33 \* 0.033357) = = CHF - 0.1164491

<u>Remark</u>: The GARCH forecast reflects the higher than average uncertainty in 2020:9 (Covid-19, presidential elections).  $\P$ 

# **GARCH: Rugarch Package**

GARCH estimation requires numerical optimization, which is dependent on initial values. The R package does a good job at estimating ARMA-GARCH models, allowing for different models and performing a lot of specification tests.

You need to specify the model ("*specs*") first, for example, you want to estimate an AR(1)-GARCH(1,1) with a constant in the mean. Then, you estimate the model with the *ugarchfit* command.

Conditional Variance Dynamics

\_\_\_\_\_

GARCH Model : sGARCH(1,1) Mean Model : ARFIMA(1,0,0) Distribution : norm Optimal Parameters

			-	
	Estimate	Std. Er	ror t valu	e Pr(> t )
mu	0.004695	0.001052	4.4651	8e-06
ar1	0.277567	0.025120	11.0496	0e+00
omega	0.000075	0.000015	4.8968	1e-06
alpha1	0.126715	0.017529	7.2289	0e+00
beta1	0.826194	0.020600	40.1061	0e+00

ח 1 ע	G( 1 1 F			
Robust	Standard Erro	rs:	$  = \mathbf{D}_{\mathbf{u}}(\mathbf{x}   \mathbf{z}   \mathbf{z} $	
	Estimate Std.	Error t val	100  Pr( t )	
mu	0.004695	0.001145	4.1018 0.000041	
arl	0.2//56/	0.022948	12.0957 0.000000	
omega	0.000075	0.000021	3.6307 0.000283	
alphal	0.126715	0.026943	<b>4.7031</b> 0.000003	
betal	0.826194	0.028409	<b>29.0821</b> 0.000000	
LogLik	elihood: 3472			
Inform	ation Criteria			
Akaike	-3.8591			
Bayes	-3.8438			
Shibata	a -3.8591			
Hannar	n-Quinn -3.853	4		
		_		
Weight	ted Ljung-Box	Test on Sta	andardized Residual	S
		statistic p	-value	
Lag[1]		0.3178 0.5	57294	
Lag[2*	(p+q)+(p+q)-1	][2] 2.5	5441 0.08393	
Lag[4*	(p+a)+(p+a)-1	1151 <b>6.9</b>	210 0.02072	$\Rightarrow$ Need to add more lags in mean.
d.o.f=1		][-]		,
H0:N	o serial correla	tion		
Weight	ted Liung-Box	Test on Sta	andardized Squared	Residuals
811				

statis	tic p-value	
Lag[1] 0.191	5 0.6617	
Lag[2*(p+q)+(p+q)-1][5]	1.1353 0.8284	
Lag[4*(p+q)+(p+q)-1][9]	1.6161 0.9455	$\Rightarrow$ No evidence of extra ARCH lags

# IGARCH

Recall the technical detail: The standard GARCH model:  $\sigma_t^2 = \omega + \alpha(L)\varepsilon^2 + \beta(L)\sigma^2$ is covariance stationary if  $\alpha(1) + \beta(1) < 1$ .

But strict stationarity does not require such a stringent restriction

In the GARCH(1,1) model, if  $\alpha_1 + \beta_1 = 1$ , we have the Integrated GARCH (IGARCH) model.

In the IGARCH model, the autoregressive polynomial in the ARMA representation has a unit root: a shock to the conditional variance is "*persistent*."

Variance forecasts are generated with:  $E_t[\sigma_{t+j}^2] = \sigma_t^2 + j\omega$ 

 $\Rightarrow$  today's variance remains important for all future forecasts. This is persistence!

Variance forecasts are generated with:  $E_t[\sigma_{t+j}^2] = \sigma_t^2 + j\omega$ 

That is, today's variance remains important for future forecasts of all horizons.

In practice (see previous Example 2 for the S&P 500 data), it is often found that  $\alpha_1 + \beta_1$  are close to 1.

#### **GARCH: Variations – GARCH-in-mean**

The time-varying variance affects mean returns: Mean equation:  $y_t = X_t \gamma + \delta \sigma_t^2 + \varepsilon_t$ ,  $\varepsilon_t \sim N(0, \sigma_t^2)$ Variance equation:  $\sigma_t^2 = \omega + \alpha_1 \varepsilon_{t-1}^2 + \beta_1 \sigma_{t-1}^2$ 

We have a dynamic mean-variance relations. It describes a specific form of the risk-return tradeoff.

Finance intuition says that  $\delta$  has to be positive and significant. However, in empirical work, it does not work well:  $\delta$  is not significant or negative.

#### **GARCH: Variations – Asymmetric GJR**

GJR-GARCH model – Glosten, Jagannathan & Runkle (JF, 1993):

$$\sigma_t^2 = \omega + \sum_{i=1}^{q} \alpha_i \varepsilon_{t-i}^2 + \sum_{i=1}^{q} \gamma_i \varepsilon_{t-i}^2 * I_{t-i} + \sum_{j=1}^{p} \beta_j \sigma_{t-j}^2$$

where  $I_{t-1} = 1$  if  $\varepsilon_{t-1} < 0$ ; = 0 otherwise.

Using the indicator variable  $I_{t-i}$ , this model captures sign (asymmetric) effects in volatility: Negative news ( $\varepsilon_{t-1} < 0$ ) increase the conditional volatility (*leverage effect*).

where  $I_{t-1} = 1$  if  $\varepsilon_{t-1} < 0$ ; = 0 otherwise.

When

$$\begin{array}{ll} \varepsilon_{t-1} < 0 & \Rightarrow & \sigma_t^2 = \omega + (\alpha_1 + \gamma_1) \varepsilon_{t-1}^2 + \beta_1 \sigma_{t-1}^2 \\ \varepsilon_{t-1} > 0 & \Rightarrow & \sigma_t^2 = \omega + \alpha_1 \varepsilon_{t-1}^2 + \beta_1 \sigma_{t-1}^2 \end{array}$$

This is a very popular variation of the GARCH models. The leverage effect is significant.

There is another variation, the Exponential GARCH, or EGARCH, that also captures the asymmetric effect of negative news on the conditional variance.

# **GARCH: Variations – NARCH**

Non-linear ARCH model NARCH – Higgins and Bera (1992) and Hentschel (1995).

These models apply the Box-Cox-type transformation to the conditional variance:

$$\sigma_t^{\gamma} = \omega + \sum_{i=1}^{q} \alpha_i |\varepsilon_{t-i} - \kappa|^{\gamma} + \sum_{j=1}^{\nu} \beta_j \sigma_{t-j}^{\gamma}$$

Special case:  $\gamma = 2$  (standard GARCH model).

<u>Note</u>: The variance depends on both the size and the sign of the variance which helps to capture leverage type (asymmetric) effects.

# **ARCH Estimation: MLE – Regularity Conditions**

<u>Technical Note</u>: The appeal of MLE is the optimal properties of the resulting estimators under ideal conditions. However, these ideal conditions, which are called "*regularity conditions*," are difficult to verify for ARCH models

• Block-diagonality

In many applications of ARCH models, the parameters can be partitioned into mean parameters,  $\theta_1$ , and variance parameters,  $\theta_2$ . Thus, the Information matrix ( $\approx$ Hessian) is *block-diagonal*.

Not a bad result:

- Regression can be consistently done with OLS.

- Asymptotically efficient estimates for the ARCH parameters can be obtained on the basis of the OLS residuals.

But:

- Conventional OLS standard errors could be terrible.

– When testing for autocorrelation, in the presence of ARCH, the conventional Bartlett s.e.  $-T^{1/2}$  – could seriously underestimate the true standard errors.

# **ARCH Estimation: Non-Normality**

The basic GARCH model allows a certain amount of leptokurtosis. It is often insufficient to explain real world data.

<u>Solution</u>: Assume a distribution, other than the normal, that can produce fatter tails in the distribution.

• *t* Distribution - Bollerslev (1987)

The t distribution has a degrees of freedom parameter which allows greater kurtosis. The likelihood function for observation *t* is:

 $l_t = \ln(\Gamma(0.5(\nu+1))\Gamma(0.5\nu)^{-1}(\nu-2)^{-1/2}(1+z_t(\nu-2)^{-1})^{-(\nu+1)/2}) - 0.5\ln(\sigma^2_t)$ 

where  $\Gamma$  is the gamma function and v is the degrees of freedom. As  $\upsilon \to \infty$ , this tends to the normal distribution.

#### **ARCH: Testing**

Standard BP test, where we have an ARCH(q) as the alternative H<sub>1</sub>. Then, we test H<sub>0</sub>:  $\alpha_1 = \alpha_2 = ... = \alpha_q = 0$ .

Steps:

- Step 1. (Same as BP's Step 1). Run OLS on DGP:  $y = X \beta + \varepsilon.$  Keep residuals,  $e_t$ - Step 2. (Auxiliary Regression). Regress  $e_t^2$  on  $e_{t-1}^2$ , ....,  $e_{t-q}^2$   $e_t^2 = \alpha_0 + \alpha_1 e_{t-1}^2 + .... + \alpha_q e_{t-q}^2 + v_t$  Keep R<sup>2</sup>, say  $R_{e2}^2$ . - Step 3. Compute the statistic:  $LM = (T-q) R_{e2}^2 \xrightarrow{d} \chi_q^2$ .

**Example:** We do an ARCH Test with 4 lags, for the AR(1) residuals of log changes in the CHF/USD (T = 593):

```
yyy <- z;
T <- length(yyy)
xx_1 <- z[-T]
yy <- z[-1]
fit2 <- lm(yy ~ xx_1 -1)
res d <- fit2$residuals
```

# Step 1: extract residuals

```
p lag <-4
e2 lag <- matrix(0, T-p lag, p lag)
                                                       # matrix to put lag e^2
resid r2 <- res d^2
a <-1
while (a \le p \text{ lag})
 e2 lag[,a] \leq resid r2[a:(T-p lag+a-1)]
a <- a+1
}
fit lm2 \le lm(resid r2[(p lag+1):T] \sim e2 lag)
                                                               # Step 2: Auxiliary Regression
r2 e1 <- summary(fit lm2)$r.squared
                                                               # extract R^2
\lim t <- (T-p lag)*r2 e1
                                                               # LM test: Sample size * R^2
> lm t
                        \Rightarrow Reject H<sub>0</sub> (No ARCH) with a p-value of 0.001.
[1] 17.08195
```

#### **ARCH: Testing – Ignoring ARCH**

In ARCH Models, testing as usual: LR, Wald, and LM tests. Suppose ARCH is detected, but ARCH is ignored. What are the consequences of ignoring ARCH?

• Ignoring ARCH

- Suppose yt has an AR structure:

 $y_t = \gamma_0 + \gamma_1 y_{t-1} + \varepsilon_t$ ,  $\varepsilon_t | I_{t-1} \sim N(0, \sigma^2_t)$ . with ARCH structure in the error term, but ARCH is ignored. Then, we fit the AR(1) model using OLS.

- Simulations find that OLS t-test with no correction for ARCH spuriously reject H<sub>0</sub>:  $\phi_1 = 0$  with arbitrarily high probability for sufficiently large *T*.

- If White's (1980) SE are used, the results are better. NW SE help less.

**Figure.** From Hamilton (2008). Fraction of samples in which OLS *t-test* leads to rejection of H<sub>0</sub>:  $\phi_1 = 0$  as a function of *T* for regression with Normal errors (solid blue line) and Student's *t* errors (dashed green line).

<u>Note</u>:  $H_0$  is actually true & the *t*-test is evaluated at the 5% level.



#### **ARCH: Which Model to Use**

Questions 1) Lots of ARCH models. Which one to use?

2) Choice of *p* and *q*. How many lags to use?

Hansen and Lunde (2004) compared lots of ARCH models:

- It turns out that the GARCH(1, 1) is a great starting model.
- Add a leverage effect for financial series and it's even better.
- A *t*-distribution is also a good addition.

# **RV Models: Intuition**

The idea of realized volatility is to estimate the latent (unobserved) variance using the realized data, without any modeling. Recall the definition of sample variance:

$$s^{2} = \frac{1}{(T-1)} \sum_{i=1}^{T} (x_{i} - \overline{x})^{2}$$

Suppose we want to calculate the daily variance for stock returns. We know how to compute it: we use daily information, for T days, and apply the above definition.

Alternatively, we use hourly data for the whole day (with *k* hours). Since hourly returns are very small, ignoring  $\overline{x}$  seems OK. We use  $r_{t,i}^2$  as the *i*<sup>th</sup> hourly variance on day *t*. Then, we add  $r_{t,i}^2$  over the day:

$$Variance_t = \sum_{i=1}^k r_{t,i}^2$$

In more general terms, we use higher frequency data to estimate a lower frequency variance:

$$V_t = \sum_{i=1}^k r_{t,i}^2$$

where  $r_{t,i}$  is the realized returns in (higher frequency) interval *i* of the (lower frequency) period *t*.

We estimate the *t-frequency* variance, using *k i-intervals*. If we have daily returns and we want to estimate the monthly variance, then, *k* is equal to the number of days in a month.

It can be shown that as the interval *i* becomes smaller  $(i \rightarrow 0)$ ,

$$RV_t \rightarrow \text{Return Variation } [t-1, t].$$

That is, with an increasing number of observations we get an accurate measure of the latent variance.

# **RV Models: High Frequency**

Note that RV is a model-free measure of variation –i.e., no need for ARCH-family specifications. The measure is called *realized variance* (RV). The square root of the realized variance is the *realized volatility* (RVol, RealVol):

$$RVol_t = sqrt(RV_t)$$

Given the previous theoretical result, RV is commonly used with intra-daily data, called *high frequency* (HF) data.

It lead to a revolution in the field of volatility, creating new models and new ways of thinking about volatility and how to model it.

We usually associate realized volatility with an observable proxy of the unobserved volatility.

# **RV Models: High Frequency – Tick Data**

As mentioned above, the theory behind realized variation measures dictates that the sampling frequency, or k in the RV<sub>t</sub> formula above, goes to  $\infty$ . Then, use the highest frequency available, say millisecond to millisecond returns.

Intra-daily data applications are the most common. But, when using intra-daily data, RV calculations are affected by microstructure effects: bid-ask bounce, infrequent trading, calendar effects, etc.  $r_{t,i}$  does not look uncorrelated.

Example: The bid-ask bounce induces serial correlation in intra-day returns, which biases RVt.

The usual solutions:

(1) Filter data using an ARMA model to get rid of the autocorrelations and/or dummy variables to get rid of calendar effects.

Then, used the filtered data to compute RV<sub>t</sub>.

(2) Sample at frequencies where the impact of microstructure effects is minimized and/or eliminated.

We will follow solution (2).

# **RV Models: High Frequency – Practice**

In intra-daily RV estimation, it is common to use 10' intervals. They have good properties. However, there are estimations with 1' intervals.

Some studies suggest using an *optimal* frequency, where optimal frequency is the one that minimizes the MSE.

Hansen and Lunde (2006) find that for very liquid assets, such as the S&P 500 index, a 5' sampling frequency provides a reasonable choice. Thus, to calculate daily RV, we need to add 78 five-minute intervals.

**Example:** Based on TAQ (*Trade and Quote*) NYSE data, we use 5' realized returns to calculate 30' variances –i.e., we use six 5' intervals. Then, the 30' variance, or RV<sub>t=30-min</sub>, is:

$$RV_{t=30-min} = \sum_{j=1}^{k=6} r_{t,j}^2$$
,  $t = 1, 2, \dots, T=15$ 

 $r_{t,j}$  is the 5' return during the j<sup>th</sup> interval on the half hour t. Then, we calculate 30' variances for the whole day –i.e., we calculate 13 variances, since the trading day goes from 9:30 AM to 4:00 PM.

The Realized Volatility, RVol, is:

$$RVol_{t=30-min} = \sqrt{RV_{t=30-min}}$$

**Example:** Below, we show the first transaction of the **SPY TAQ** (*Trade and Quote*) data (tickby-tick *trade* data) on **January 2, 2014**.

SYMBOL	DATE	TIME	PRICE	SIZE
SPY	20140102	9:30:00	183.98	500

SPY	20140102	9:30:00	183.98	500
SPY	20140102	9:30:00	183.98	200
SPY	20140102	9:30:00	183.98	500
SPY	20140102	9:30:00	183.98	1000
SPY	20140102	9:0:00	183.98	1000
SPY	20140102	9:30:00	183.98	800
SPY	20140102	9:30:00	183.98	100
SPY	20140102	9:30:00	183.98	100
SPY	20140102	9:30:00	183.97	200
SPY	20140102	9:30:00	183.98	100
SPY	20140102	9:30:00	183.97	200
SPY	20140102	9:30:00	183.98	1000
SPY	20140102	9:30:00	183.97	100
SPY	20140102	9:30:00	183.98	1000
SPY	20140102	9:30:00	183.98	2600
SPY	20140102	9:30:00	183.98	1000
SPY	20140102	9:30:00	183.97	400

**Example:** Below, we show the first transaction of the **AAPL TAQ** (*Trade and Quote*) data (tick-by-tick *quote* data) on January 2, 2014: 4 AM

SYMBOL	DATE	TIME	BID	OFR	BIDSIZ	OFRSIZ	MODE	ΕX
AAPL	20140102	4:00:00	455.39	0	1	0	12	Т
AAPL	20140102	4:00:00	553.5	558	2	2	12	Р
AAPL	20140102	4:00:01	455.39	561.02	1	2	12	Т
AAPL	20140102	4:00:45	552.1	558	1	2	12	Р
AAPL	20140102	4:00:51	552.1	558.4	1	2	12	Р
AAPL	20140102	4:00:51	552.1	558.8	1	2	12	Р
AAPL	20140102	4:00:51	552.1	559	1	1	12	Р
AAPL	20140102	4:01:14	553	559	1	1	12	Р
AAPL	20140102	4:01:30	553.01	561.02	1	2	12	Т
AAPL	20140102	4:01:43	553.01	559	1	1	12	Т
AAPL	20140102	4:01:44	553.05	559	1	1	12	Р

AAPL	20140102 4:01:49	455.39	559	1	1	12	Т
AAPL	20140102 4:01:49	553.61	559	1	1	12	Т
AAPL	201401024:02:02	553.05	559	1	2	12	Р
AAPL	20140102 4:02:04	455.39	559	1	1	12	Т
AAPL	201401024:02:04	548.28	559	1	1	12	Т
AAPL	201401024:02:33	553.05	558.83	1	2	12	Р
AAPL	201401024:02:33	555.17	558.83	2	2	12	Р
AAPL	20140102 4:03:50	555.2	558.83	5	2	12	Р

#### **RV Models: High Frequency – Working with Tick Data**

**Example:** We read **SPY trade data** for 2014:Jan. > HF da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/SPY 2014.csv", head=TRUE, sep=",") > summary(HF da) G127 **SYMBOL** DATE TIME PRICE SIZE SPY:6800865 Min. :20140102 9:30:00 : 21436 Min. :176.6 Min. : 1 Min. :0 1st Qu.:20140110 16:00:00: 11352 1st Qu.:178.9 1st Qu.: 100 1st Ou.:0 Median :20140121 9:30:01 : 5922 Median :182.6 Median : 100 Median :0 Mean :20140119 15:59:59: 4090 Mean :181.4 Mean : 337 Mean :0 3rd Qu.:20140128 15:59:55: 3198 3rd Qu.:183.5 3rd Qu.: 300 3rd Qu.:0 Max. :20140131 15:50:00: 2916 Max. :189.2 Max. Max. :0 :4715350 (Other) :6751951 COND CORR ΕX Min. :0.0e+00 :3351783 Т :1649158 (a)1st Ou.:0.0e+00 Р :2888182 :1335135 F Median :0.0e+00: 524409 Ζ :1182126 Mean :1.9e-04 0 : 18057 D :1062382 3rd Qu.:0.0e+00 4 : 9098 Κ : 437900 Max. :1.2e+01 : 8142 : 356539 J 6 (Other): 1194 (Other): 777625

• Now, we calculate using 5'-returns a daily realized volatility for the first 4 days in 2014 (2014:01:02 - 2014:01:07). Originally, we have T = 1,048,570.

pt <- as.POSIXct(paste(HF\_da\$DATE, HF\_da\$TIME), format="%Y%m%d %H:%M:%S")
library(xts)
hf\_1 <- xts(x=HF\_da, order.by = pt) # Define a specific time series data set</pre>



spy\_ret <- log(spy\_p[-1]/spy\_p[-T])
plot(spy\_ret, type="l", ylab="Return", main="Tick by Tick Return (2014:01:02 - 2014:01:07)")
mean(spy\_ret)
sd(spy\_ret)</pre>





Very noisy data, with lots of "jumps": Mean tick by tick return: -3.7365e-09 Tick-by-tick SD: 6.3163e-05

• For the whole month of January 2020:



> mean(spy\_ret)
[1] -4.796933e-09
> sd(spy\_ret)
[1] 7.804991e-05

• We plot the autocorrelogram for the TAQ SPY data:



TAQ SPY Data: January 2014



1.000 -0.469 -0.013 -0.010 0.014 -0.008 0.000 -0.002 -0.001 0.000 0.000

• We aggregate the tick-by-tick data in **5' intervals** using the function *aggregateTrades* in the R package *highfrequency*. It needs as an input an xts object (hf\_1, for us).





 $\begin{array}{l} RVol_{t=2014:01:02}=0.0053344\\ RVol_{t=2014:01:03}=0.0043888\\ RVol_{t=2014:01:04}=0.0059836 \end{array}$ 

 $RVol_{t=2014:01:05} = 0.0052772$ 

We plot the autocorrelogram for the 5-minute TAQ SPY return data:



5-minute SPY Data: January 2014

<u>Note</u>: We have a negative  $1^{st}$ -order autocorrelation: -0.105, thought not significant. However, the autocorrelation of order 5 is significant.

• We plot the 10-minute TAQ SPY return data. Smoothing increases.



 $\begin{aligned} RVol_{t=2014:01:02} &= 0.005478294 \\ RVol_{t=2014:01:03} &= 0.004256046 \\ RVol_{t=2014:01:04} &= 0.006190508 \\ RVol_{t=2014:01:05} &= 0.005145601 \end{aligned}$ 

We plot the autocorrelogram for the 10' TAQ SPY return data:

#### 10-minute SPY Data: January 2014



Note: Now, none of the autocorrelations is significant. The **10-minute returns** look independent.  $\P$ 

# **RV Models: High Frequency – TAQ In Practice**

In practice, 10' returns are common. To form a daily measure for RV, we have 39 10-minute returns plus one overnite return (from 16:00 PM to next day 9:30 AM)

We have some technical issues working with tick data:

- Not all days the stock market is open from 9:30 AM to 16:00 PM, NYSE closes early on certain days (Christmas Eve, Thanksgiving).

- For many stocks, we do have lapses in trading. For these stocks, using 5' or 10' intervals may not work well.

- There are many suggested solutions to the problem of infrequent trading. Usual solution: interpolation from quote data.

- We have a lot of (discrete) jumps in the data.

**Example:** R script to compute *monthly* realized volatility for **MSCI USA daily returns** MSCI\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/MSCI\_daily.csv", head=TRUE, sep=",") x\_us <- MSCI\_da\$USAT <- length(x\_us) us r <- log(x us[-1]/x us[-T])

x <- us r	# US log returns from MSCI USA Index
$T \leq -length(x)$	
rvs=NULL	# create vector to fill with RV
i <- 1	
k <- 21	# k: observations per period (78 for 5' data)
while $(i < T - k)$ {	
$s_{2} < sum(x[i:(i+k)]^{2})$	# realized variance
$i \leq k + i$	
rvs <- rbind(rvs,s2)	
}	
rvol <- sqrt(rvs)	# realized volatility
mean(rvol)	# mean
sd(rvol)	# variance

**Example:** Using **daily MSCI USA** data we calculate 1-mo Realized Volatility (*k*=21 days) for log returns for the USA MSCI (1970: Jan – 2020: Oct).



Technical computing points:

We use k=21 days, which is an average of the trading days per month. Of course, not all months have the same amount of trading days. In 2019, February had the fewest (19) and October the most (23), but, in 2018, February and September (18) had the fewest and August the most (23). For us, k=21 days is an approximation.

To be precise, if we use daily data to calculate a monthly variance, we need to use an exact index of trading days, say,  $K=[k_1, k_2, k_3, ..., k_J]$  where  $k_i$  is the exact number of trading days in *month*-*year i*.

In addition, for daily data, we should not ignore the mean in the computation of RV.

```
Example: Below, the while loop in R is modified to incorporate the vector K (c1) of exact
trading days for each month.
MSCI cd <-
read.table("https://www.bauer.uh.edu/rsusmel/4397/MSCI d count days.txt",header=FALSE)
                                    # Vector with all exact days in a month
c1 \leq MSCI cd[,1]
n cl \leq- length(cl)
                                    # Total number of days in sample
                                     # Initialize empty vector to place RVs
rvs=NULL
                                    # index for the days for while loop
t <- 1
ti <- 1
                                     # index for the months for while loop
x_m = mean(x)
while (tj \le n c1) {
mj \le c1[tj]
                                    # reading exact number of days for month tj
                                            # daily returns (in deviation from mean) per month
xx <- x[t:(t+mj-1)] - x m
tj
s2 \leq sum(xx^2)
                     # RV for month tj
t \leq t + mj
```

#### Monthly RVOL for MSCI USA with exact day count



<u>Note</u>: The results (mean, SD and shape of RV) are very similar, but if used to compare to other monthly volatility estimates, these are the correct monthly RVol estimates.  $\P$ 

#### **RV Models: Log Approximation Rules**

The log approximations rules for the variance and SD are used to change frequencies for the RV and RVol. For example, suppose we are calculating RV based on frequency j,  $RV_{t=j}$ ; but we are interested in the J-period  $RV_{t=J}$ . Then, the J-period (with j intervals) realized variance and realized volatility can be calculated as

$$RV_{t=J} = J * RV_{t=j}$$
  

$$RVol_{t=J} = sqrt(J) * RVol_{t=j}$$

**Example:** We calculate using 5' data the daily realized variance,  $RV_{t=daily}$ . Then, the annual variance can be calculated as

 $RV_{t=annual} = 260 * RV_{t=daily}$ 

where 260 is the number of trading days in the year. The annualized RVOL is the squared root of  $RV_{annual}$ :

$$RVOL_{t=annual} = sqrt(260) * RVOL_{t=daily}$$

**Example:** Using daily data we calculate 3-mo Realized Volatility (k=66 days) for log returns for the MSCI (1970: March – 2020: Oct).

US MSCI Monthly Stock Return RVol



#### **RV Models: Properties**

Under some conditions (bounded kurtosis and autocorrelation of squared returns less than 1),  $RV_t$  is consistent.

Realized volatility is a measure. It has a distribution.

For returns, the distribution of RV is non-normal (as expected). It tends to be skewed right and leptokurtic.

Daily returns standardized by RVol measures are nearly Gaussian.

RV is highly persistent. (Check with a LB test.)

Daily RV calculate with intra-daily data, it is found to be more robust than measures using daily data, like GARCH.

#### **RV Models: ACF and Persistence**

Like all volatility measures, RVOL is highly autocorrelated.

**Example:** We plot the ACF and PACF for the 1-mo Realized Volatility, based on daily data for the monthly USA MSCI data.


# **RV Models: Forecasting**

We can fit ARMA models to the RVOL series to generate forecasts.

**Example:** Based on the ACF and PACF, we fit an AR(2) model for the monthly RVOL, calculated from monthly data: > fit\_rvol\_ar2 <- arima(rvol, order=c(2,0,0)) > fit\_rvol\_ar2 Call: arima(x = rvol, order = c(2, 0, 0)) ar1 ar2 intercept 0.5631 0.0967 0.0433 s.e. 0.0396 0.0396 0.0023 sigma^2 estimated as 0.0004056: log likelihood = 1568.46, aic = -3128.92

> checkresiduals(fit\_rvol\_ar2)

Ljung-Box test

data: Residuals from ARIMA(2,0,0) with non-zero mean  $Q^* = 12.008$ , df = 7, p-value = 0.1003

Model df: 3. Total lags used: 10



• AR(2) model seems to pass diagnostic tests. Now, we forecast RVOL.

fcast\_rvol <- forecast(fit\_rvol\_ar2, h=12, level=.95) # h=number of step-ahead forecasts > fcast\_rvol

	Point Forecast	Lo 95	Hi 95	
632	0.05201688 0	0.01254198	11 0.091491	78
633	0.04937852 0	0.00407615	48 0.094680	88
634	0.04757422 -0	0.00058224	56 0.095730	69
635	0.04630317 -(	0.00317169	03 0.095778	04
636	0.04541302 -(	0.00469926	67 0.095525	32
637	0.04478891 -0	0.00563344	66 0.095211	26
638	0.04435142 -(	0.00622262	287 0.094925	46
639	0.04404473 -(	0.00660368	68 0.094693	15
640	0.04382975 -0	0.00685518	09 0.094514	67
641	0.04367904 -(	0.00702381	75 0.094381	90
642	0.04357339 -(	0.00713827	18 0.094285	06
643	0.04349934 -(	0.00721665	577 0.094215	33.¶

#### RVOL: Forecast 2020:Oct - 2021:Jun



<u>Note</u>: The VIX index ("*fear index*") is a forecast for the next 30-day volatility, derived from S&P 500 options. The VIX on Sep 30, 2020 was 26.37, that is, the volatility at the end of October is expected to be 26.37% annualized or 7.61% monthly, higher than 5.20%, but, well within the 95% C.I. (More on this later.)

## **RV Models: Forecasting – Using VIX**

Empirical work uses the VIX to calculate the implied volatility, IVt, for the S&P500. The VIX index is based on the S&P500 index options (on a panel of S&P 500 option prices), using the

"model-free" approach tailored to replicate the (annualized) risk-neutral volatility of a fixed 30day maturity.



**Example:** We use **VIX** to forecast monthly RV based on daily data (1990:May - 2020:Sep). We regress

 $RV_{t+1} = \alpha + \beta VIX_t + \varepsilon_t.$ 

Mid da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/Mid1 U B data.csv", head=TRUE, sep=",") v date <- Mid da\$Code VIX <- Mid da\$VIX # Extract VIX data T rv  $\leq$  length(rvol) # End of sample for RVol (2020:Oct) rvol 90 <- as.numeric(rvol ts[245:T rv])\*100 # RVol starting in 1990: May in % rvol 0 <- rvol 90[-1] # remove first observation ( $RV_{t+1}$ ) VIX  $m \leq VIX/sqrt(12)$ # VIX in monthly % lm rvol f <- lm(rvol  $0 \sim VIX m$ ) > summary(lm rvol f) Coefficients: Estimate Std. Error t value Pr(>|t|)-0.89301 0.28021 (Intercept) -3.187 0.00156 \*\* 0.04641 **20.469** < 2e-16 \*\*\* VIX m 0.94997 ---Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 1.967 on 363 degrees of freedom

Multiple R-squared: 0.5358, Adjusted R-squared: 0.5345 F-statistic: 419 on 1 and 363 DF, p-value: < 2.2e-16

Note: In sample, a strong positive predictive relation.





<u>Note</u>: There is good match between the two series. RVOL shocks (Financial crisis, Covid) are unexpected by IV.

• We also check the contemporaneous relation between RVOL and VIX.

lm\_rvol <- lm(rvol\_90[-length(rvol\_90)] ~ VIX\_m)
> summary(lm\_rvol)

Estimate Std. Error t value Pr(>|t|) (Intercept) -1.88538 0.20458 -9.216 <2e-16 \*\*\* VIX\_m 1.12543 0.03388 33.214 <2e-16 \*\*\* Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 1.436 on 363 degrees of freedom Multiple R-squared: 0.7524, Adjusted R-squared: 0.7517 F-statistic: 1103 on 1 and 363 DF, p-value: < 2.2e-16

Note: A strong contemporaneous relation. RVOL is highly correlated.

## **RV Models: Variance Risk Premium (VRP)**

The implied volatility of an option, calculated today, or  $IV_t$ , is a measure of the ("ex ante") expected variance over the remaining life of the option.

The Black-Scholes (BS) and similar models for option prices produce the same option prices as would be seen under modified probabilities in a world of investors who were indifferent to risk *(risk neutral)*.

IV & other parameters extracted from options market prices embed these modified "*risk neutral*" probabilities, that combine investors' objective predictions of the real world returns distribution with their risk preferences.

Under BS assumptions, IV and market volatility are the same. But, BS assumptions do not hold. The VRP uses this disparity.

We define the variance risk premium (VRP) as the difference between the "ex-ante" *risk neutral* expectation at time t of the future return variation over the period [t, t+1] time interval and the ex-post realized return variation over the [t-1, t]:

$$\sqrt{\mathbf{RP}_t} = \mathbf{IV}_t - \mathbf{RV}_t.$$

It is an ad-hoc definition, we could have defined VRP<sub>t</sub> based on the expectation at time t for  $RV_{t+1}$ , in this case  $E_t[RV_{t+1}]$ . The one-step-ahead forecast can be obtained using an ARMA process for  $RV_t$ .

In practice, using  $E_t[RV_{t+1}]$  or  $RV_t$ , does not affect VRPt that much.

The are many ways to calculate IV: based on models, like the BS, or "*model free*," similar to how we calculated IV, in this case, using changes in option prices for different strike prices and computing an average.

**Example:** We plot IV<sub>t</sub>(=VIX), RV<sub>t</sub> & VRP<sub>t</sub> for the S&P500 Index (shaded blue area are U.S. recessions). Data: Monthly 1990-2008.



• Bollerslev et al. (2009) use 5' intervals to calculate RVt find that VRPt is a predictor of stock market excess returns at different horizons (t+h). That is, they regress:  $r_{t+h} - r_{f,t+h} = [log(P_t) - log(P_{t-1})] = \mu + \delta VRPt + \varepsilon_t$  They find that  $\hat{\delta}$  is positive and has a *t-stat*=1.76 for monthly data (*h*=1) and a *t-stat* = 2.86 for quarterly data (*h*=3). The R<sup>2</sup> is 1.07% for monthly data and 6.82% for quarterly data. For annual data the t-stat is not significant.

Monthly Return Horizon	1	3	6	9	12	15	18	24
Constant	-0.55	-2.08	1.12	3.63	4.62	4.84	5.61	6.48
	(-0.13)	(-0.56)	(0.33)	(1.15)	(1.50)	(1.59)	(1.81)	(2.07)
$IV_t - RV_t$	0.39	0.47	0.30	0.17	0.12	0.11	0.06	0.01
	(1.76)	(2.86)	(2.15)	(1.36)	(1.00)	(0.94)	(0.56)	(0.11)
Adj. R <sup>2</sup> (%)	1.07	6.82	5.42	2.30	1.23	1.00	0.05	-0.50

**Example:** We regress excess next-month returns, using the FF Mkt-RF factor as the dependent variable, on today's VRP:

FF\_da <- read.csv("https://www.bauer.uh.edu/rsusmel/4397/FF\_5\_factors.csv",header=TRUE)</td>x\_Mkt\_RF <- FF\_da\$Mkt\_RF</td>T\_FF <- length(x\_RF)</td>Mkt\_RF <- x\_Mkt\_RF[323:T\_FF]/100</td># Obs 332: 1990: Mayvrp <- VIX\_m^2 - rvol\_90[-length(rvol\_90)]^2</td># Variance risk premiumpred\_vrp <-lm(Mkt\_RF[-1] ~ vrp)</td>> summary(pred\_vrp)

Estimate Std. Error t value Pr(>|t|)(Intercept) 6.655e-03 2.335e-03 2.850 0.00462 \*\* vrp 1.815e-05 6.210e-05 0.292 0.77029  $\Rightarrow$  not significant ---Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 0.04346 on 363 degrees of freedom Multiple R-squared: 0.0002352, Adjusted R-squared: -0.002519. ¶

#### **Other Models: Parkinson's (1980) Estimator**

The Parkinson's (1980) estimator:  $s^2 t = \{ \sum_t [\ln(H_t) - \ln(L_t)]^2 / (4\ln(2)T) \},\$ where H<sub>t</sub> is the highest price and L<sub>t</sub> is the lowest price.

There is an RV counterpart, using HF data: Realized Range (RR):  $RR_t = \{ \sum_j [100 * (ln(H_{t,j}) - ln(L_{t,j}))]^2 / (4ln(2)) \},\$ where  $H_{t,j}$  and  $L_{t,j}$  are the highest and lowest price in the j<sup>th</sup> interval.

These "range" estimators are very good and very efficient.

These estimators can be applied to intra-daily data. The Realized Range works well with combined with other models.

## Stochastic volatility (SV/SVOL) models

Now, instead of a known volatility at time *t*, like ARCH models, we allow for a stochastic shock to  $\sigma_t$ ,  $\eta_t$  or  $\upsilon_t$ :

 $\sigma_t = \omega + \beta_1 \sigma_{t-1} + \eta_t, \qquad \upsilon_t \sim N(0, \sigma_\eta^2)$ 

Or using logs:

 $\log \sigma_t = \omega + \beta_1 \log \sigma_{t-1} + \upsilon_t, \qquad \upsilon_t \sim N(0, \sigma_v^2)$ 

The difference with ARCH models: The shocks that govern the volatility are not necessarily the shocks to the mean process,  $\epsilon_t$ 's.

Usually, the standard model centers log volatility around  $\omega$ :

 $\log \sigma_t = \omega + \beta_1 (\log \sigma_{t-1} - \omega) + v_t,$ 

Then,

$$\begin{split} E[\log(\sigma_t)] &= \omega \\ Var[\log(\sigma_t)] &= \kappa^2 = |\sigma_{\upsilon}^2/(1-\beta^2). \\ &\implies \text{Unconditional distribution: } \log(\sigma_t) \sim N(\omega, \kappa^2) \end{split}$$

Like ARCH models, SV models produce returns with kurtosis > 3 (and, also, positive autocorrelations between squared excess returns).

We have 3 SVOL parameters to estimate:  $\varphi = (\omega, \beta, \sigma_{\upsilon})$ .

Estimation: The modern approach uses Bayesian methods (MCMC), which are advanced for this class. Brooks discusses the estimation of SVOL.