Probability and Statistics FRE6871 & FRE7241, Spring 2025

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Pseudo-Random Numbers

Pseudo-random numbers are deterministic sequences of numbers which have some of the properties of random numbers, but they are not truly random numbers.

Pseudo-random number generators depend on a *seed* value, and produce the same sequence of numbers for a given *seed* value.

The function set.seed() initializes the random number generator by specifying the *seed* value.

The choice of *seed* value isn't important, and a given value is just good as any other one.

The function runif() produces random numbers from the *uniform* distribution.

The function rnorm() produces random numbers from the *normal* distribution.

The function rt() produces random numbers from the *t-distribution* with *df* degrees of freedom.

- > # Initialize the random number generator
- > set.seed(1121, "Mersenne-Twister", sample.kind="Rejection")
- > runif(3) # three numbers from uniform distribution
- > runif(3) # Simulate another three numbers
- > # Initialize the random number generator
- > set.seed(1121, "Mersenne-Twister", sample.kind="Rejection")
- > runif(3) # Simulate another three numbers
- > # Simulate random number from standard normal distribution
 > rnorm(1)
- > # Simulate five standard normal random numbers > rnorm(5)
- > # Simulate five non-standard normal random numbers
- > rnorm(n=5, mean=1, sd=2) # Match arguments by name
- > # Simulate t-distribution with 2 degrees of freedom
- > rt(n=5, df=2)

The Logistic Map

> # Define logistic map function
> log_map <- function(x, r=4) r*x*(1-x)</pre>

> log_map(0.25, 4) > # Plot logistic map

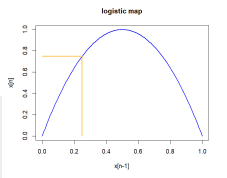
The *logistic map* is a recurrence relation which produces a deterministic sequence of numbers:

$$x_n = rx_{n-1}(1-x_{n-1})$$

If the seed value x_0 is in the interval (0,1) and if r=4, then the sequence x_n is also contained in the interval (0,1).

The function curve() plots a function defined by its name.

```
> x11(width=6, height=5)
> curve(expr=log_map, type="1", xlim=c(0, 1),
+ xlab="x[n-1]", ylab="x[n]", lwd=2, col="blue",
+ main="logistic map")
> lines(x=c(0, 0.25), y=c(0.75, 0.75), lwd=2, col="orange")
> lines(x=c(0.25, 0.25), v=c(0.0.75), lwd=2, col="orange")
```



Generating Pseudo-Random Numbers Using Logistic Map

The logistic map can be used to calculate sequences of pseudo-random numbers.

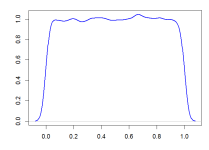
For most seed values x_0 and r = 4, the logistic map produces a pseudo-random sequence, but it's not uniformly distributed.

The inverse cosine function acos() transforms a logistic map sequence into a uniformly distributed sequence,

$$u_n = \arccos(1 - 2x_n)/\pi$$

```
> # Calculate uniformly distributed pseudo-random sequence
> # using logistic map function.
> unifun <- function(seedv, n=10) {
    # Pre-allocate vector instead of "growing" it
    datay <- numeric(n)
   # initialize
   datav[1] <- seedv
   # Perform loop
   for (i in 2:n) {
     datav[i] <- 4*datav[i-1]*(1-datav[i-1])
   } # end for
    acos(1-2*datav)/pi
```

uniform pseudo-random number density



- > unifun(seedv=0.1, n=15)
- > plot(
- density(unifun(seedy=runif(1), n=1e5)),
- xlab="", vlab="", lwd=2, col="blue",
- main="uniform pseudo-random number density")

end unifun

Generating Binomial Random Numbers

A binomial trial is a coin flip, that results in either a success or failure.

The binomial distribution specifies the probability of obtaining a certain number of successes in a sequence of independent binomial trials.

Let p be the probability of obtaining a success in a binomial trial, and let (1-p) be the probability of failure

p = 0.5 corresponds to flipping an unbiased coin.

The probability of obtaining k successes in nindependent binomial trials is equal to:

$$\binom{n}{k}p^k(1-p)^{(n-k)}$$

The function rbinom() produces random numbers from the *binomial* distribution

- > # Initialize the random number generator
- > set.seed(1121, "Mersenne-Twister", sample.kind="Rejection")
- > # Flip unbiased coin once, 20 times
- > # Number of heads after flipping twice, 20 times
- > rbinom(n=20, size=1, 0.5) > rbinom(n=20, size=2, 0.5)
- > # Number of heads after flipping thrice, 20 times
- > rbinom(n=20, size=3, 0.5)
- > # Number of heads after flipping biased coin thrice, 20 times > rbinom(n=20, size=3, 0.8)
- > # Number of heads after flipping biased coin thrice, 20 times
- > rbinom(n=20, size=3, 0.2) > # Flip unbiased coin once, 20 times
- > sample(x=0:1, size=20, replace=TRUE) # Fast
- > as.numeric(runif(20) < 0.5) # Slower

Generating Random Samples and Permutations

A sample is a subset of elements taken from a set of data elements.

The function sample() selects a random sample from a vector of data elements.

By default the *size* of the sample (the size argument) is equal to the number of elements in the data vector.

So the call sample(da_ta) produces a random permutation of all the elements of da_ta.

The function sample() with replace=TRUE selects samples with replacement (the default is replace=FALSE).

Monte Carlo simulation consists of generating random samples from a given probability distribution.

The Monte Carlo data samples can then be used to calculate different parameters of the probability distribution (moments, quantiles, etc.), and its functionals.

- > # Permutation of five numbers
- > sample(x=5)
 > # Permutation of four strings
- > sample(x=c("apple", "grape", "orange", "peach"))
- > # Sample of size three
 > sample(x=5, size=3)
- > sample(x=5, Size=3) > # Sample with replacement
- > # Sample with replacement > sample(x=5, replace=TRUE)
- > sample(# Sample of strings
- + x=c("apple", "grape", "orange", "peach"),
- + size=12,
- + replace=TRUE)
 > # Binomial sample: flip coin once, 20 times
- > sample(x=0:1, size=20, replace=TRUE)
- > # Flip unbiased coin once, 20 times
- > as.numeric(runif(20) > 0.5) # Slower

[1] 1.01

Statistical Estimators

A data sample is a set of observations $\{x_1, \ldots, x_n\}$ of a random variable x.

Let x follow a probability distribution with population $\it mean$ equal to μ and population $\it standard$ $\it deviation$ equal to $\sigma.$

A statistic is a function of the data sample: $f(x_1, ..., x_n)$, so it is itself a random variable.

A statistical *estimator* is a *statistic* that provides an estimate of a distribution *parameter*.

For example:

$$\bar{x} = \frac{1}{n} \sum_{i=1}^{n} x_i$$

Is an estimator of the population mean of the distribution.

```
> # Initialize the random number generator
> set.seed(1121, "Mersenne-Twister", sample.kind="Rejection")
> # Sample from Standard Normal Distribution
> datav <- rnorm(1000)
> mean(datav) # Sample mean
[1] -0.0354
> median(datav) # Sample median
[1] -0.0153
> sd(datav) # Sample standard deviation
```

Estimators of Higher Moments

The estimators of the moments of a probability distribution, based on a *sample* of data x_i , are given by:

Sample mean:
$$\bar{x} = \frac{1}{n} \sum_{i=1}^{n} x_i$$

Sample variance:
$$\hat{\sigma}^2 = \frac{1}{n-1} \sum_{i=1}^n (x_i - \bar{x})^2$$

Their expected values are equal to the population mean and standard deviation:

$$\begin{array}{ll} \mathbb{E}[\bar{\mathbf{x}}] = \mu \quad \text{and} \quad \mathbb{E}[\hat{\sigma}] = \sigma \mathbb{E}[\hat{\sigma}] = \sigma$$

The third and fourth moments are equal to:

$$\mu 3 = \frac{n}{(n-1)(n-2)} \sum_{i=1}^{n} (x_i - \bar{x})^3$$

$$\mu 4 = \frac{n}{(n-1)^2} \sum_{i=1}^{n} (x_i - \bar{x})^4$$

The skewness and kurtosis are equal to the moments scaled by the standard deviation:

$$\varsigma = \frac{\mu 3}{\sigma^3}, \quad \kappa = \frac{\mu 4}{\sigma^4}$$

- > # VTI returns
- > retp <- na.omit(rutils::etfenv\$returns\$VTI)
- > # Number of observations
- > nrows <- NROW(retp)
- > # Mean of VTI returns
- > meanv <- mean(retp)
 > # Standard deviation of VTI returns
- > stdev <- sd(retp)
- > # Standardize returns
- > retp <- (retp meanv)/stdev > # Skewness and kurtosis of VTI returns
- > nrows/((nrows-1)*(nrows-2))*sum(retp^3)
- [1] -0.385
- > nrows/(nrows-1)^2*sum(retp^4)
- [1] 13.8
- > # Random normal returns
- > retp <- rnorm(nrows)
- > # Mean and standard deviation of random normal returns
 > mean(retp); sd(retp)
- > mean(retp
 - [1] 0.976
 - > # Skewness and kurtosis of random normal returns
- > nrows/((nrows-1)*(nrows-2))*sum(retp^3)
- [1] -0.0464
- > nrows/(nrows-1)^2*sum(retp^4)
 [1] 2.74
- [1] 2.7

The normal distribution has skewness equal to zero $\varsigma=0$, and kurtosis equal to three $\kappa=3$.

Estimators of Quantiles

The *quantile* corresponding to a given *probability p*, is the value of the *random variable x*, such that the probability of obtaining values less than *x* is equal to the *probability p*.

The *quantile* of a data sample can be calculated by first sorting the sample, and then finding the value corresponding closest to the given *probability p*.

The function quantile() calculates the sample quantiles. It uses interpolation to improve the accuracy. Information about the different interpolation methods can be found by typing ?quantile.

The function sort() returns a vector sorted into ascending order.

The function pnorm() calculates the cumulative normal distribution, i.e. the cumulative probability for a given quantile value.

The function qnorm() calculates the inverse cumulative *normal* distribution, i.e. the quantile for a given probability value.

The function dnorm() calculates the normal probability density.

```
> # Calculate cumulative standard normal distribution
> c(pnorm(-2), pnorm(2))
> # Calculate inverse cumulative standard normal distribution
> c(qnorm(0.75), qnorm(0.25))
> # Initialize the random number generator
> set.seed(1121, "Mersenne-Twister", sample.kind="Rejection")
> # Sample from Standard Normal Distribution
> nrows <- 1000
> datay <- rnorm(nrows)
> # Sample mean - MC estimate
> mean(datav)
> # Sample standard deviation - MC estimate
> sd(datav)
> # Monte Carlo estimate of cumulative probability
> c(pnorm(1), sum(datav < 1)/nrows)
> # Monte Carlo estimate of quantile
> confl <- 0.99
> qnorm(confl)
> cutoff <- confl*nrows
> datay <- sort(datay)
> c(datay[cutoff], quantile(datay, probs=confl))
> # Read the source code of quantile()
> stats:::quantile.default
> # microbenchmark quantile
> library(microbenchmark)
> summary(microbenchmark(
    monte carlo=datav[cutoff].
```

times=100))[, c(1, 4, 5)] # end microbenchmark summary

quanty=quantile(datay, probs=confl).

Standard Errors of Estimators

Statistical estimators are functions of samples (which are random variables), and therefore are themselves random variables

The standard error (SE) of an estimator is defined as its standard deviation (not to be confused with the population standard deviation of the underlying random variable).

For example, the *standard error* of the estimator of the mean is equal to:

$$\sigma_{\mu} = \frac{\sigma}{\sqrt{n}}$$

Where σ is the population standard deviation (which is usually unkown).

The estimator of this standard error is equal to:

$$SE_{\mu} = \frac{\hat{\sigma}}{\sqrt{n}}$$

where: $\hat{\sigma}^2 = \frac{1}{n-1} \sum_{i=1}^n (x_i - \bar{x})^2$ is the sample standard deviation (the estimator of the population standard deviation).

- > # Initialize the random number generator
- > set.seed(1121, "Mersenne-Twister", sample.kind="Rejection")
- > # Sample from Standard Normal Distribution
- > nrows <- 1000 > datay <- rnorm(nrows)
- > datav <- rnorm(nrows)
- > # Sample mean > mean(datay)
- > # Sample standard deviation
- > sd(datav)
- > # Standard error of sample mean
- > sd(datav)/sqrt(nrows)

The Characteristic Function

The characteristic function $\hat{f}(t)$ is equal to the Fourier transform of the probability density function f(x):

$$\hat{f}(t) = \mathbb{E}[e^{itx}] = \int_{-\infty}^{\infty} f(x) e^{itx} dx$$

The normal probability density function:

$$\phi(x) = \frac{e^{-(x-\mu)^2/2\sigma^2}}{\sigma\sqrt{2\pi}}$$

Has the *characteristic function* $\hat{\phi}(t)$ equal to:

$$\hat{\phi}(t) = e^{i\mu t} e^{-(\sigma t)^2/2}$$

The probability function f(x) is equal to the inverse Fourier transform of the characteristic function $\hat{f}(t)$:

$$f(x) = \frac{1}{2\pi} \int_{-\infty}^{\infty} \hat{f}(t) e^{-itx} dt$$

The characteristic function of the first derivative of f(x) is equal to $(-it)\hat{f}(t)$, the characteristic function multiplied by (-it):

$$\frac{df(x)}{dx} = \frac{1}{2\pi} \int_{-\infty}^{\infty} -it \, \hat{f}(t) \, e^{-itx} \, dt$$

The characteristic function of the n-th derivative of f(x) is equal to $(-it)^n \hat{f}(t)$, the characteristic function multiplied by $(-it)^n$:

$$\frac{d^n f(x)}{dx^n} = \frac{1}{2\pi} \int_{-\infty}^{\infty} (-it)^n \hat{f}(t) e^{-itx} dt$$

The Moment Generating Function

The moment generating function $M_X(t)$ of a random variable x with the probability density function f(x) is equal to:

$$M_X(t) = \mathbb{E}[e^{tx}] = \int_{-\infty}^{\infty} f(x) e^{tx} dx$$

The *n*-th derivative of $M_X(t)$ with respect to t, at t=0 is equal to the *n*-th moment μ_n :

$$\mu_n = \frac{d^n M_X(t)}{dt^n}|_{t=0} = \frac{d^n \mathbb{E}[e^{tx}]}{dt^n}|_{t=0}$$
$$= \mathbb{E}[x^n e^{tx}]|_{t=0} = \mathbb{E}[x^n]$$

The moments μ_n are related to the central moments $\mathbb{E}[(x-\mu)^n]$ but they are not equal to them.

The *moment generating function* can be expressed as a series of its *moments*:

$$M_X(t) = \sum_{n=0}^{\infty} \frac{\mu_n t^n}{n!}$$

The moment generating function for the *normal* distribution is equal to:

$$M_X(t) = \exp(\mu t + \frac{1}{2}\sigma^2 t^2)$$

The characteristic function $\hat{f}(t)$ is equal to the moment generating function with a purely imaginary argument:

$$\hat{f}(t) = \mathbb{E}[e^{itx}] = M_X(it)$$

Cumulants of Probability Distributions

The cumulant generating function $K_X(t)$ is equal to the logarithm of the moment generating function:

$$K_X(t) = \log M_X(t)$$

The *n*-th derivative of $K_X(t)$ with respect to t, at t=0 is equal to the *n*-th cumulant κ_n :

$$\kappa_n = \frac{d^n K_X(t)}{dt^n}|_{t=0}$$

The *cumulants* are related to the *moments* of a distribution: with the first three cumulants being equal to the *central moments* (mean, variance, and skewness), while the higher order *cumulants* are polynomials of the *moments*.

The cumulant generating function $K_X(t)$ can be expanded into a power series of the cumulants:

$$K_X(t) = \sum_{n=1}^{\infty} \frac{\kappa_n t^n}{n!} = \mu t + \sigma^2 \frac{t^2}{2} + \sigma^3 s^3 \frac{t^3}{6} + \dots$$

The cumulant generating function for the *normal* distribution is equal to:

$$K_X(t) = \mu t + \frac{1}{2}\sigma^2 t^2$$

So that its first two *cumulants* are equal to the *mean* μ and the *variance* σ^2 , and the *cumulants* of order 3 and higher are all equal to zero.

The advantage of *cumulants* over the *moments* is that the *cumulants* of the sum of independent random variables are equal to the sum of their *cumulants*:

$$\begin{aligned} K_{(X+Y)}(t) &= \log \mathbb{E}[e^{t(X+Y)}] = \log (\mathbb{E}[e^{tX}]\mathbb{E}[e^{tY}]) \\ &= \log \mathbb{E}[e^{tX}] \log \mathbb{E}[e^{tY}] = K_X(t) + K_Y(t) \end{aligned}$$

The Hermite Polynomials

The n-th derivative of the standard normal distribution $\phi(x)$ is given by Rodrigues' formula:

$$\frac{d^n\phi(x)}{dx^n} = \frac{d^n}{dx^n} \frac{e^{-x^2/2}}{\sqrt{2\pi}} = (-1)^n H_n(x)\phi(x)$$

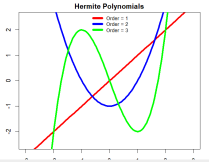
Where H_n are the Hermite polynomials.

The first four Hermite polynomials are equal to:

$$H_0(x) = 1$$
; $H_1(x) = x$
 $H_2(x) = x^2 - 1$; $H_3(x) = x^3 - 3x$

The even order polynomials are symmetric $H_{2n}(-x) = H_{2n}(x)$ while the odd order are antisymmetric $H_{2n-1}(-x) = -H_{2n-1}(x)$.

```
> # Define the Hermite polynomials
> hermitep <- function(x, n) {
      switch(n+1, 1, x, (x^2 - 1), (x^3 - 3*x), 0)
    # end hermitep
```



```
> colory <- c("red", "blue", "green")
> for (indeks in 1:3) { # Plot three curves
   curve(expr=hermitep(x, indeks).
   xlim=c(-3, 3), vlim=c(-2,5, 2,5),
   xlab="", ylab="", lwd=4, col=colorv[indeks],
   add=as.logical(indeks-1))
+ } # end for
```

Error in hermitep(x, indeks): could not find function "hermitep"

> # Add title and legend

> title(main="Hermite Polynomials", line=0.5)

Error in title(main = "Hermite Polynomials", line = 0.5): plot.new has not been called yet

> labelv <- paste("Order", 1:3, sep=" = ")

The Hermite Functions

The Hermite functions $\psi_n(x)$ are equal to:

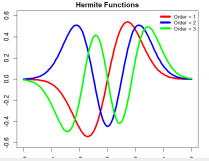
$$\psi_n(x) = \frac{1}{\sqrt{n!\sqrt{2\pi}}} e^{-x^2/4} H_n(x)$$

The Hermite functions form an orthonormal set:

$$\int_{-\infty}^{\infty} \psi_n(x) \, \psi_m(x) \, \mathrm{d}x = \begin{cases} 1 & \text{if } n = m \\ 0 & \text{if } n \neq m \end{cases}$$

The Hermite functions of increasing order oscillate more frequently, with the function of order n crossing zero n times

- > # Define the Hermite functions
- > hermitef <- function(x, n)
- $exp(-x^2/4)*hermitep(x, n)/(2*pi)^(0.25)/sqrt(factorial(n))$
- > # Integrate the Hermite functions > integrate(function(x, n, m)
- hermitef(x, n)*hermitef(x, m).
- lower=(-Inf), upper=Inf, n=2, m=3)



- > colorv <- c("red", "blue", "green")
- > for (indeks in 1:3) { # Plot three curves
- curve(expr=hermitef(x, indeks),
- xlim=c(-6, 6), ylim=c(-0.6, 0.6),xlab="", ylab="", lwd=4, col=colorv[indeks],
- add=as.logical(indeks-1))
- + } # end for
- > # Add title and legend
- > title(main="Hermite Functions", line=0.5)
- > labelv <- paste("Order", 1:3, sep=" = ")
- > legend("topright", inset=0.0, bty="n",
- + title=NULL, labely, cex=0.8, lwd=6, lty=1,
- - col=colorv)

draft: The Hermite Series

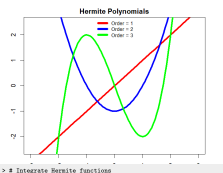
To-do: expand a probability distribution into a series of Hermite functions

The Hermite functions $\psi_i(x)$ form an orthonormal basis that can be used to expand a given probability distribution f(x) into a series:

$$f(x) = \sum_{i=0}^{n} f_i \, \psi_i(x)$$

The coefficients f_i are equal to:

$$f_i = \int_{-\infty}^{\infty} f(x) \, \psi_i(x) \, \mathrm{d}x$$



- > integrate(hermitep, lower=(-Inf), upper=Inf, n=2)
- > integrate(function(x, n, m) hermitep(x, n)*hermitep(x, m),
- lower=(-Inf), upper=Inf, n=2, m=3)
- > integrate(function(x, n, m) hermitep(x, n)*hermitep(x, m). lower=(-Inf), upper=Inf, n=2, m=2)
- > colorv <- c("red", "blue", "green")
- > for (indeks in 1:3) { # Plot three curves
- curve(expr=hermitep(x, indeks),
- xlim=c(-4, 4), ylim=c(-0.6, 0.6),
- xlab="", vlab="", lwd=3, col=colorv, add=as.logical(indeks-1))
- + } # end for
- > # Add title and legend
- > title(main="Hermite Functions", line=0.5)
- > labely <- paste("Order", 1:3, sep=" = ") Probability and Statistics

The Bell polynomials

The Bell polynomials B_n are the coefficients in the expansion of the exponent of a series:

$$\exp(\sum_{n=1}^{\infty} \kappa_n \frac{t^n}{n!}) = \sum_{n=0}^{\infty} B_n(\kappa 1, \dots, \kappa_n) \frac{t^n}{n!}$$

The first four Bell polynomials are equal to:

$$B_0 = 1$$

$$B1(\kappa 1) = \kappa 1$$

$$B2(\kappa 1, \kappa 2) = \kappa 1^2 + \kappa 2$$

$$B3(\kappa 1, \kappa 2, \kappa 3) = \kappa 1^3 + 3\kappa 1\kappa 2 + \kappa 3$$

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The Gram-Charlier Series

The Gram-Charlier series expresses the density function f(x) in terms of its cumulants and a basis function $\phi(x)$, with characteristic $\hat{\phi}(t)$.

The *characteristic functions* $\hat{f}(t)$ and $\hat{\phi}(t)$ can be expressed in terms of their *cumulants* as:

$$\hat{f}(t) = e^{K_X(it)} = \exp(\sum_{n=1}^{\infty} \kappa_n \frac{(it)^n}{n!})$$

$$\hat{\phi}(t) = \exp(\sum_{n=1}^{\infty} \phi_n \frac{(it)^n}{n!})$$

Then $\hat{f}(t)$ can be expressed in terms of $\hat{\phi}(t)$ as:

$$\hat{f}(t) = \exp\left[\sum_{n=1}^{\infty} (\kappa_n - \phi_n) \frac{(it)^n}{n!}\right] \hat{\phi}(t)$$

The basis function $\phi(x)$ can be chosen to be a normal distribution, with mean and standard deviation equal to that of f(x), and with all its normal cumulants ϕ_n of order 3 and higher equal to zero.

Then we get a series starting at n=3:

$$\hat{f}(t) = \exp[\sum_{n=3}^{\infty} \kappa_n \frac{(it)^n}{n!}] \, \hat{\phi}(t)$$

We can expand the exponent and collect terms with the same power of t using the *Bell polynomials* B_n :

$$\hat{f}(t) = \sum_{n=0}^{\infty} B_n(0, 0, \kappa 3, \dots, \kappa_n) \frac{(it)^n}{n!} \hat{\phi}(t)$$

The inverse Fourier transform of the above equation gives the probability function f(x):

$$f(x) = \sum_{n=0}^{\infty} B_n(0, 0, \kappa 3, \dots, \kappa_n) \frac{(-1)^n}{n!} \frac{d^n \phi(x)}{dx^n}$$

The derivatives of the *normal* distribution $\phi(x)$ can be expressed using the *Hermite polynomials* H_n so that the *Gram-Charlier series* becomes:

$$f(x) = \phi(x) \sum_{n=0}^{\infty} \frac{B_n(0, 0, \kappa 3, \dots, \kappa_n)}{n! \sigma^n} H_n(\frac{x - \mu}{\sigma})$$

draft: The Edgeworth Expansion

The Edgeworth expansion expresses the probability density function f(x) as a series of its cumulants and a basis function $\phi(x)$ (with characteristic function $\hat{\phi}(t)$).

The characteristic functions $\hat{f}(t)$ and $\hat{\phi}(t)$ can be expressed in terms of their corresponding cumulants as:

$$\hat{f}(t) = e^{K_X(it)} = \exp(\sum_{n=1}^{\infty} \kappa_n \frac{(it)^n}{n!})$$

$$\hat{\phi}(t) = \exp(\sum_{n=1}^{\infty} \phi_n \frac{(it)^n}{n!})$$

Then $\hat{f}(t)$ can be expressed in terms of $\hat{\phi}(t)$ as:

$$\hat{f}(t) = \exp[\sum_{n=1}^{\infty} (\kappa_n - \phi_n) \frac{(it)^n}{n!}] \hat{\phi}(t)$$

If the basis function $\phi(x)$ is chosen to be the normal distribution, with mean and standard deviation equal to that of f(x), and since the normal cumulants of order 3 and higher are all equal to zero, then we get:

$$\hat{f}(t) = \exp\left[\sum_{n=3}^{\infty} \kappa_n (-1)^n \frac{(it)^n}{n!}\right] \hat{\phi}(t)$$

The *inverse Fourier transform* of the above equation gives the *probability function* f(x):

$$f(x) = \exp\left[\sum_{n=3}^{\infty} \kappa_n (-1)^n \frac{d^n}{dx^n}\right] \phi(x)$$

Now expand the exponent in a series and collect terms with the same order of the derivative to obtain:

$$f(x) = \sum_{n=0}^{\infty} B_n(0, 0, \kappa 3, \dots, \kappa_n) (-1)^n \frac{d^n \phi(x)}{dx^n}$$

draft: The Cornish-Fisher Expansion

The Cornish-Fisher expansion expresses the quantiles of a distribution as a series of its moments.

The Edgeworth Expansion The cumulant generating function $K_X(t)$ is equal to the logarithm of the moment generating function:

$$K_X(t) = \log M_X(t)$$

The *n*-th derivative of $K_X(t)$ with respect to t, at t=0 is equal to the *n*-th cumulant κ_n :

$$\kappa_n = \frac{d^n K_X(t)}{dt^n}|_{t=0}$$

The *cumulants* are related to the *moments* of the distribution: the first three cumulants are equal to the *central moments* (mean, variance, and skewness), while the higher order *cumulants* can be expressed as polynomials of the *central moments*.

The cumulant generating function $K_X(t)$ can be expanded into a power series of the cumulants:

$$\mathcal{K}_{X}(t) = \sum_{n=1}^{n} rac{\kappa_{n}t^{n}}{n!} = \mu t + \sigma^{2}t^{2}/2 +$$

The *n*-th moment μ_n is not equal to the central moment $\mathbb{E}[(x-\mu)^n]$.

The moment generating function $M_X(t)$ of a random variable x with the probability function f(x) is equal to:

> set.seed(1121, "Mersenne-Twister", sample.kind="Rejection")

> # Sample from Standard Normal Distribution

> nrows <- 1000

> datav <- rnorm(nrows)

> # Sample mean
> mean(datav)

> # Sample standard deviation

> sd(datav)

Hypothesis Testing

Hypothesis Tests are designed to test the validity of *null hypotheses*, and they consist of:

- A null hypothesis,
- A test statistic derived from the data sample,
- A p-value: the conditional probability of observing the test statistic value, assuming the null hypothesis is TRUE,
- A significance level α corresponding to a critical value

The *p*-value is compared to the *significance level* and if the *p*-value is less than the *significance level* α , then the *null hypothesis* is rejected.

It's possible for the *null hypothesis* to be TRUE, but to obtain a very small *p*-value purely by chance.

The *p*-value is the probability of erroneously rejecting a TRUE *null hypothesis*, due to the randomness of the data sample.

```
> ### Perform two-tailed test that sample is
> ### from Standard Normal Distribution (mean=0, SD=1)
> # generate vector of samples and store in data frame
> test_frame <- data.frame(samples=rnorm(1e4))
> # get p-values for all the samples
> test_frame$p_values <- sapply(test_frame$samples,
          function(x) 2*pnorm(-abs(x)))
> # Significance level, two-tailed test, critical value=2*SD
> signif_confl <- 2*(1-pnorm(2))
> # Compare p_values to significance level
> test_frame$result <-
    test_frame$p_values > signif_level
> # Number of null rejections
> sum(!test frame$result) / NROW(test frame)
> # Show null rejections
> head(test_frame[!test_frame$result, ])
```

The p-value is a conditional probability, and is not equal to the un-conditional probability of the hypothesis being TRUE.

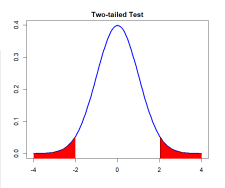
In statistics we cannot *prove* that a hypothesis is TRUE or not, but we can attempt to invalidate it, and conclude that it's unlikely to be TRUE, given the test statistic value and its *p*-value.

Two-tailed Hypothesis Tests

In two-tailed hypothesis tests, both tails of the probability distribution contribute to the *p*-value.

Two-tailed hypothesis tests are applied for testing if the absolute value of a sample exceeds the critical value.

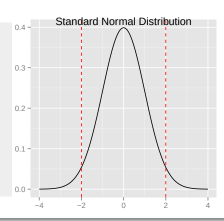
```
> # Plot the Normal probability distribution
> curve(expr=dnorm(x, sd=1), type="1", xlim=c(-4, 4),
+ xlab="", ylab="", lwd=3, col="blue")
> title(main="Two-tailed Test", line=0.5)
> # Plot tails of the distribution using polygons
> x1 <- 2: x2 <- 4
> # Plot right tail using polygon
> xvar <- seq(x1, x2, length=100)
> vxvar <- dnorm(xvar, sd=1)
> vxvar[1] <- (-1)
> vxvar[NROW(vxvar)] <- (-1)
> polygon(x=xvar, y=yxvar, col="red")
> # Plot left tail using polygon
> vxvar <- dnorm(-xvar, sd=1)
> vxvar[1] <- (-1)
> vxvar[NROW(vxvar)] <- (-1)
> polygon(x=(-xvar), y=yxvar, col="red")
```



Visualizing Hypothesis Testing Using Package ggplot2

In two-tailed hypothesis tests, both tails of the probability distribution contribute to the p-value.

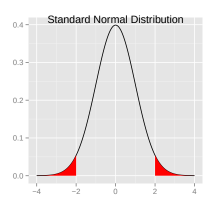
```
> library(ggplot2) # Load ggplot2
> qplot( # Simple ggplot2
     main="Standard Normal Distribution",
     c(-4, 4),
     stat="function",
     fun=dnorm.
     geom="line".
     xlab=NULL, vlab=NULL
     ) + # end aplot
 theme( # Modify plot theme
     plot.title=element_text(vjust=-1.0),
     plot.background=element blank()
     ) + # end theme
 geom vline( # Add vertical line
   aes(xintercept=c(-2.0, 2.0)).
   colour="red".
   linetype="dashed"
   ) # end geom vline
```



Visualizing Hypothesis Testing Using ggplot2 (cont.)

In two-tailed hypothesis tests, both tails of the probability distribution contribute to the p-value.

```
> ### Create ggplot2 with shaded area
> xvar <- -400:400/100
> norm frame <- data.frame(xvar=xvar.
                  d.norm=dnorm(xvar))
> norm_frame$shade <- ifelse(
             abs(norm_frame$xvar) >= 2,
             norm_frame$d.norm, NA)
> ggplot( # Main function
   data=norm_frame,
   mapping=aes(x=xvar, y=d.norm)
 ) + # end ggplot
+ # Plot line
   geom_line() +
+ # Plot shaded area
   geom_ribbon(aes(ymin=0, ymax=shade), fill="red") +
+ # No axis labels
   xlab("") + ylab("") +
+ # Add title
   ggtitle("Standard Normal Distribution") +
+ # Modify plot theme
   theme(
 plot.title=element_text(vjust=-1.0),
 plot.background=element_blank()
   ) # end theme
```



Student's t-test for the Distribution Mean

Student's t-test is designed to test the *null hypothesis* that a sample: $\{x_1, \ldots, x_n\}$ was obtained from a normal distribution with a *mean* equal to μ .

The test statistic is equal to the t-ratio:

$$t = \frac{\bar{x} - \mu}{\hat{\sigma} / \sqrt{n}}$$

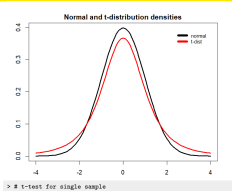
Where $\bar{x} = \frac{1}{n} \sum_{i=1}^{n} x_i$ is the sample mean and $\hat{\sigma}^2 = \frac{1}{n-1} \sum_{i=1}^{n} (x_i - \bar{x})^2$ is the sample variance.

Under the *null hypothesis* the *t-ratio* follows the *t-distribution* with *n* degrees of freedom, with the probability density function:

$$f(t) = \frac{\Gamma((n+1)/2)}{\sqrt{\pi n} \Gamma(n/2)} (1 + t^2/n)^{-(n+1)/2}$$

Student's t-test can also be used to test if two different normally distributed samples have equal population means.

Student's t-test is not valid for random variables that do not follow the normal distribution.



```
> t.test(rnorm(100))
> # t-test for two samples
> t.test(rnorm(100),
+ rnorm(100, mean=1))
```

> # Plot the normal and t-distribution densities
> x11(width=6, height=5)

> par(mar=c(3, 3, 3, 1), oma=c(0, 0, 0, 0))
> curve(expr=dnorm, xlim=c(-4, 4),

+ xlab="", ylab="", lwd=3)
> curve(expr=dt(x, df=3),

+ xlab="", ylab="", lwd=3, + col="red", add=TRUE)

> # Add title > title(main="Normal and t-distribution densities", line=0.5) > # Add legend

draft: The Analysis of Variance (ANOVA)

The Analysis of Variance (ANOVA) to test if the sub-samples of the data have the same mean.

ANOVA provides a statistical test of whether two or more population means are equal, and therefore generalizes the t-test beyond two means. Student's t-test is designed to test the null hypothesis that a sample: $\{x_1, \ldots, x_n\}$ was obtained from a normal distribution with a mean equal to μ .

For example, ANOVA is widely used to study the effect of medical treatments, with the null hypothesis being that the treatments have no effect and that differences are due to random chance

The test statistic is equal to the t-ratio:

Jerzy Pawlowski (NYU Tandon)

$$t = \frac{\bar{x} - \mu}{\hat{\sigma}/\sqrt{n}}$$

Where $\bar{x} = \frac{1}{n} \sum_{i=1}^{n} x_i$ is the sample mean and $\hat{\sigma}^2 = \frac{1}{n-1} \sum_{i=1}^n (x_i - \bar{x})^2$ is the sample variance.

Under the null hypothesis the t-ratio follows the t-distribution with n degrees of freedom, with the probability density function:

$$f(x) = \frac{\Gamma((n+1)/2)}{\sqrt{\pi n} \Gamma(n/2)} (1 + x^2/n)^{-(n+1)/2}$$

Student's t-test can also be used to test if two different normally distributed samples have equal population

Normal and t-distribution densities 4.0 0.0 -2 0 > # t-test for single sample

> t.test(rnorm(100)) > # t-test for two samples > t.test(rnorm(100), rnorm(100, mean=1)) > # Plot the normal and t-distribution densities > x11(width=6, height=5) > par(mar=c(3, 3, 3, 1), oma=c(0, 0, 0, 0)) > curve(expr=dnorm, xlim=c(-4, 4), xlab="", ylab="", lwd=3) > curve(expr=dt(x, df=3),

xlab="", ylab="", lwd=3,

> # Add legend

col="red", add=TRUE) > # Add title > title(main="Normal and t-distribution densities", line=0.5)

The Shapiro-Wilk Test of Normality

The Shapiro-Wilk test is designed to test the null hypothesis that a sample: $\{x_1, \ldots, x_n\}$ is from a normally distributed population.

The test statistic is equal to:

$$W = \frac{\left(\sum_{i=1}^{n} a_{i} x_{(i)}\right)^{2}}{\sum_{i=1}^{n} (x_{i} - \bar{x})^{2}}$$

Where the: $\{a_1, \ldots, a_n\}$ are proportional to the *order* statistics of random variables from the normal distribution.

 $x_{(k)}$ is the *k*-th *order statistic*, and is equal to the *k*-th smallest value in the sample: $\{x_1, \ldots, x_n\}$.

The *Shapiro-Wilk* statistic follows its own distribution, and is less than or equal to 1.

The *Shapiro-Wilk* statistic is close to 1 for samples from normal distributions.

The *p*-value for *VTI* returns is extremely small, and we conclude that the *null hypothesis* is FALSE, and the *VTI* returns are not from a normally distributed population.

The *Shapiro-Wilk* test is not reliable for large sample sizes, so it's limited to less than 5000 sample size.

- > # Calculate VTI percentage returns
- > library(rutils)
- > retp <- as.numeric(na.omit(rutils::etfenv\$returns\$VTI))[1:499]
- > # Reduce number of output digits
- > ndigits <- options(digits=5)
- > # Shapiro-Wilk test for normal distribution
- > nrows <- NROW(retp)
- > shapiro.test(rnorm(nrows))

Shapiro-Wilk normality test

data: rnorm(nrows)

W = 0.997, p-value = 0.45 > # Shapiro-Wilk test for VTI returns

> shapiro.test(retp)

Shapiro-Wilk normality test

data: retp

W = 0.991, p-value = 0.0029

> # Shapiro-Wilk test for uniform distribution

> shapiro.test(runif(nrows))

Shapiro-Wilk normality test

data: runif(nrows)

- W = 0.953, p-value = 1.5e-11
- > # Restore output digits
- > options(digits=ndigits\$digits)

The Jarque-Bera Test of Normality

The Jarque-Bera test is designed to test the null hypothesis that a sample: $\{x_1, \ldots, x_n\}$ is from a normally distributed population.

The test statistic is equal to:

$$JB = \frac{n}{6}(\varsigma^2 + \frac{1}{4}(\kappa - 3)^2)$$

Where the skewness and kurtosis are defined as:

$$\varsigma = \frac{1}{n} \sum_{i=1}^{n} \left(\frac{x_i - \bar{x}}{\hat{\sigma}} \right)^3 \qquad \kappa = \frac{1}{n} \sum_{i=1}^{n} \left(\frac{x_i - \bar{x}}{\hat{\sigma}} \right)^4$$

The Jarque-Bera statistic asymptotically follows the chi-squared distribution with two degrees of freedom.

The Jarque-Bera statistic is small for samples from normal distributions

The p-value for VTI returns is extremely small, and we conclude that the null hypothesis is FALSE, and the VTI returns are not from a normally distributed population.

- > library(tseries) # Load package tseries
- > # Jarque-Bera test for normal distribution > jarque.bera.test(rnorm(nrows))

Jarque Bera Test

data: rnorm(nrows)

- X-squared = 2, df = 2, p-value = 0.3> # Jarque-Bera test for VTI returns
- > iarque.bera.test(retp)

Jarque Bera Test

data: retp

X-squared = 22, df = 2, p-value = 2e-05 > # Jarque-Bera test for uniform distribution

> jarque.bera.test(runif(nrows))

Jarque Bera Test

data: runif(nrows)

X-squared = 29, df = 2, p-value = 5e-07

The Kolmogorov-Smirnov Test for Probability Distributions

The Kolmogorov-Smirnov test null hypothesis is that two samples: $\{x_1, \ldots, x_n\}$ and $\{y_1, \ldots, y_n\}$ were obtained from the same probability distribution.

The *Kolmogorov-Smirnov* statistic depends on the maximum difference between two empirical cumulative distribution functions (cumulative frequencies):

$$D = \sup_{i} |P(x_i) - P(y_i)|$$

The function ks.test() performs the Kolmogorov-Smirnov test and returns the statistic and its p-value invisibly.

The second argument to ks.test() can be either a numeric vector of cumulative frequencies, or a name of a cumulative distribution function.

The Kolmogorov-Smirnov test can be used as a goodness of fit test, to test if a set of observations fits a probability distribution.

The K-S test is more sensitive to the tails of the distribution because it depends on the maximum difference between the two empirical cumulative frequencies.

- > # KS test for normal distribution
- > kstest <- ks.test(rnorm(100), pnorm)
- > kstest\$p.value
- > # KS test for uniform distribution
- > ks.test(runif(100), pnorm)
- > # KS test for two shifted normal distributions
- > ks.test(rnorm(100), rnorm(100, mean=0.1))
 > ks.test(rnorm(100), rnorm(100, mean=1.0))
- > # KS test for two different normal distributions
- > ks.test(rnorm(100), rnorm(100, sd=2.0))
- > KS.test(rnorm(100), rnorm(100, sd=2.0))
 > # KS test for VTI returns vs normal distribution
- > retp <- as.numeric(na.omit(rutils::etfenv\$returns\$VTI))
- > retp <- as.numeric(na.omit(ntils..etrenv@returns@vii > retp <- (retp - mean(retp))/sd(retp)
- > ks.test(retp, pnorm)
- > ks.test(retp, phori

Chi-squared Distribution

Let z_1, \ldots, z_k be independent standard *Normal* random variables.

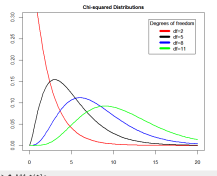
Then the random variable $X = \sum_{i=1}^k z_i^2$ is distributed according to the *Chi-squared* distribution with k degrees of freedom: $X \sim \chi_k^2$, and its probability density function is given by:

$$f(x) = \frac{x^{k/2-1} e^{-x/2}}{2^{k/2} \Gamma(k/2)}$$

The *Chi-squared* distribution with k degrees of freedom has mean equal to k and variance equal to 2k.

> # Degrees of freedom

+ } # end for



> # Add title
> title(main="Chi-squared Distributions", line=0.5)
> # Add legend
> labelv <- paste("df", degf, sep="=")
> legend("topright", inset=0.05, bty="n",
+ title="Degrees of freedom", labelv,

cex=0.8, lwd=6, ltv=1, col=colorv)

The Chi-squared Test for the Goodness of Fit

Goodness of Fit tests are designed to test if a set of observations fits an assumed theoretical probability distribution.

The Chi-squared test tests if a frequency of counts fits the specified distribution.

The Chi-squared statistic is the sum of squared differences between the observed frequencies o; and the theoretical frequencies p_i :

$$\chi^2 = N \sum_{i=1}^n \frac{(o_i - p_i)^2}{p_i}$$

Where N is the total number of observations.

The null hypothesis is that the observed frequencies are consistent with the theoretical distribution

The function chisq.test() performs the Chi-squared test and returns the statistic and its p-value invisibly.

The parameter breaks in the function hist() should be chosen large enough to capture the shape of the frequency distribution.

- > # Observed frequencies from random normal data
- > histp <- hist(rnorm(1e3, mean=0), breaks=100, plot=FALSE)
- > countsn <- histp\$counts > # Theoretical frequencies
- > countst <- rutils::diffit(pnorm(histp\$breaks))
- > # Perform Chi-squared test for normal data
- > chisq.test(x=countsn, p=countst, rescale.p=TRUE, simulate.p.value
- > # Return p-value > chisqtest <- chisq.test(x=countsn, p=countst, rescale.p=TRUE, sim
- > chisqtest\$p.value
- > # Observed frequencies from shifted normal data
- > histp <- hist(rnorm(1e3, mean=2), breaks=100, plot=FALSE)
- > countsn <- histp\$counts/sum(histp\$counts) > # Theoretical frequencies
- > countst <- rutils::diffit(pnorm(histp\$breaks))
- > # Perform Chi-squared test for shifted normal data
- > chisq.test(x=countsn, p=countst, rescale.p=TRUE, simulate.p.value
- > # Calculate histogram of VTI returns
- > histp <- hist(retp, breaks=100, plot=FALSE)
- > countsn <- histp\$counts
- > # Calculate cumulative probabilities and then difference them
- > countst <- pt((histp\$breaks-loc)/scalev, df=2)
- > countst <- rutils::diffit(countst)
- > # Perform Chi-squared test for VTI returns
- > chisq.test(x=countsn, p=countst, rescale.p=TRUE, simulate.p.value

Sorting and Ranking Data

The function sort() returns a vector sorted into ascending order, from smallest to largest.

A permutation is a re-ordering of the elements of a vector.

The permutation index specifies how the elements are

re-ordered in a permutation.

The function order() calculates the permutation index

to sort a given vector into ascending order.

Applying the function and an () twice: and an () and an () twice: and an () and an () twice: and an () and an ()

Applying the function order() twice: order(order()), calculates the permutation index to sort the vector from ascending order into its original unsorted order.

The permutation index produced by: order(order()) is the reverse of the permutation index produced by: order().

The function rank() calculates the ranks of the elements, according to their magnitude, from smallest to largest.

The ranks of the elements are equal to the reverse permutation index.

```
> # Sort a vector into ascending order
> datav <- round(runif(7), 3)
> sorty <- sort(datay)
> datav # original data
[1] 0.391 0.375 0.717 0.114 0.895 0.853 0.977
> sorty # sorted data
[1] 0.114 0.375 0.391 0.717 0.853 0.895 0.977
> # Calculate index to sort into ascending order
> indeks <- order(datay)
> indeks # permutation index to sort
[1] 4 2 1 3 6 5 7
> all.equal(sortv, datav[indeks])
[1] TRUE
> # Sort the ordered vector back to its original unsorted order
> indeks <- order(order(datay))
> indeks # permutation index to unsort
[1] 3 2 4 1 6 5 7
> all.equal(datav, sortv[indeks])
[1] TRUE
> # Calculate ranks of the vector elements
> rank(datay)
[1] 3 2 4 1 6 5 7
> all.equal(rank(datav), indeks)
```

[1] TRUE

The Mean and Median Estimators of Location

The *mean* and the *median* are both estimators of the *location* (centrality) of a distribution.

For *normally* distributed data, the *mean* has a smaller standard error than the *median* (it is more *efficient*).

But for distributions with very large *kurtosis* (fat tails), the *median* may be more *efficient* than the *mean*, because it's less sensitive to data outliers.

In addition, the *median* is often defined even for distributions for which the *mean* and *variance* are infinite.

For *symmetric* distributions, the expected values of the sample *mean* and the *median* are equal to each other, and equal to the population mean.

But for *skewed* distributions (for example asset returns), the *median* estimator is *biased* (its expected value is not equal to the population mean).

```
> # VTT returns
> retp <- as.numeric(na.omit(rutils::etfenv$returns[, "VTI"]))
> nrows <- NROW(retp)
> retp <- 100*(retp-mean(retp))/sd(retp)
> # Simulate normal random data
> ndata <- rnorm(nrows, sd=100)
> # Bootstrap the mean and median estimators
> bootd <- sapply(1:1e3, function(x) {
    # Simulate data
    ndata <- rnorm(nrows, sd=100)
    retp <- retp[sample.int(nrows, replace=TRUE)]
    c(n mean=mean(ndata).
     n median=median(ndata).
     vti mean=mean(retp).
      vti_median=median(retp))
+ }) # end sapply
> bootd <- t(bootd)
> # Analyze bootstrapped data
> head(bootd)
> sum(is.na(bootd))
> # Means and medians from bootstrap
> apply(bootd, MARGIN=2, function(x)
    c(mean=mean(x), stderror=sd(x)))
> # Parallel bootstrap under Windows
> library(parallel) # Load package parallel
> ncores <- detectCores() - 1 # Number of cores
> compclust <- makeCluster(ncores) # initialize compute cluster
> bootd <- parLapply(compclust, 1:1e4,
   function(x, nrows, retp) {
      # Simulate data
      ndata <- rnorm(nrows, sd=100)
      retp <- retp[sample.int(nrows, replace=TRUE)]
      c(n_mean=mean(ndata),
+ n median=median(ndata).
+ vti mean=mean(retp).
+ vti_median=median(retp))
+ }, nrows, retp) # end parLapply
```

The Bias-Variance Tradeoff

The tradeoff between *unbiased* estimators but with higher variance, and efficient estimators with lower variance but with some bias is called the bias-variance tradeoff.

Let $\hat{\theta}$ be an estimator of the parameter θ , with expected value $\mathbb{E}[\hat{\theta}] = \bar{\theta}$ (which may not necessarily be equal to θ).

The accuracy of the estimator $\hat{\theta}$ can be measured by its mean squared error (MSE), equal to the expected value of the squared difference $(\hat{\theta} - \theta)^2$:

$$\begin{split} \mathsf{MSE} &= \mathbb{E}[(\hat{\theta} - \theta)^2] = \mathbb{E}[(\hat{\theta} - \bar{\theta} + \bar{\theta} - \theta)^2] = \\ \mathbb{E}[(\hat{\theta} - \bar{\theta})^2 + 2(\hat{\theta} - \bar{\theta})(\bar{\theta} - \theta) + (\bar{\theta} - \theta)^2] = \\ \mathbb{E}[(\hat{\theta} - \bar{\theta})^2] + (\bar{\theta} - \theta)^2 = \mathsf{var}(\bar{\theta}) + \mathsf{bias}(\bar{\theta})^2 \end{split}$$

Since
$$\mathbb{E}[(\hat{\theta} - \bar{\theta})(\bar{\theta} - \theta)] = (\bar{\theta} - \theta)\mathbb{E}[(\hat{\theta} - \bar{\theta})] = 0$$

The above formula shows that the *MSE* is equal to the sum of the estimator *variance* plus the square of the estimator *bias*.

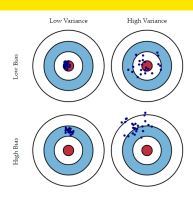


Fig. 1 Graphical illustration of bias and variance.

- > # Bias and variance from bootstrap
 > bias_var <- apply(bootd, MARGIN=2,</pre>
- + function(x) c(bias=mean(x), variance=var(x)))
- > # MSE of mean
- > bias_var[1, 3]^2 + bias_var[2, 3]
- > # MSE of median
- > bias_var[1, 4]^2 + bias_var[2, 4]

draft: The Hodges-Lehmann Estimator of Location

For distributions which are both *leptokurtic* (fat tailed) and *skewed*, the *median* estimator is more *efficient* but it's *biased*, while the *mean* is *unbiased* but it's less *efficient*

The *Hodges-Lehmann* estimator of location is a compromise between the *mean* and the *median* estimators.

The Hodges-Lehmann estimator is the median of the means of all the possible one and two-element subsets.

For a dataset with n measurements, the set of all possible one or two-element subsets of it has n(n+1)/2 elements. For each such subset, the mean is computed. Finally, the median of these n(n+1)/2 averages is defined to be the Hodges-Lehmann estimator of location

A subset is obtained by removing a few elements, and the subset mean is the mean of this subset.

Nonparametric Estimators distribution free methods, which do not rely on assumptions that the data are drawn from a given parametric family of probability distributions. As such it is the opposite of parametric statistics. accuracy

The Median Absolute Deviation (MAD) is a robust measure of dispersion (variability), defined using the median instead of the mean:

```
> retp <- as.numeric(na.omit(rutils::etfenv$returns[, "VTI"]))
> nrows <- NROW(retp)
> retp <- 100*(retp-mean(retp))/sd(retp)
> # Simulate normal random data
> ndata <- rnorm(nrows. sd=100)
> # Hodges-Lehmann estimator
> # Bootstrap the mean and median estimators
> bootd <- sapply(1:1e3, function(x) {
    # Simulate data
    ndata <- rnorm(nrows, sd=100)
    retp <- retp[sample.int(nrows, replace=TRUE)]
    c(n mean=mean(ndata),
      n_median=median(ndata),
      vti_mean=mean(retp),
      vti median=median(retp))
+ }) # end sapply
> bootd <- t(bootd)
> # Analyze bootstrapped data
> head(bootd)
> sum(is.na(bootd))
> # Means and medians from bootstrap
> apply(bootd, MARGIN=2, function(x)
    c(mean=mean(x), stderror=sd(x)))
```

draft: Nonparametric Estimators of Location

Robust Order statistics, which are based on the ranks of observations, is one example of such statistics.

The Median Absolute Deviation (MAD) is a robust measure of dispersion (variability), defined using the median instead of the mean:

$$MAD = median(abs(x_i - median(x)))$$

The advantage of *MAD* is that it's always well defined, even for data that has infinite variance.

Explain breakdown point of estimators breakdown point of estimators

For normally distributed data, the MAD has a larger standard error than the standard deviation.

But for distributions with fat tails (like asset returns), the standard deviation has a larger standard error than the MAD.

The MAD for normally distributed data is equal to $\Phi^{-1}(0.75) \cdot \hat{\sigma} = 0.6745 \cdot \hat{\sigma}$.

The function mad() calculates the MAD and divides it by $\Phi^{-1}(0.75)$ to make it comparable to the standard deviation.

```
> nrous <- 1e3
> datay <- rnorm(nrows)
> sd(datav)
> mad(datav)
> median(abs(datav - median(datav)))
> median(abs(datav - median(datav)))/qnorm(0.75)
> # Bootstrap of sd and mad estimators
> bootd <- sapply(1:1e4, function(x) {
   samplev <- datay[sample.int(nrows, replace=TRUE)]
   c(sd=sd(samplev), mad=mad(samplev))
+ }) # end sapply
> bootd <- t(bootd)
> # Analyze bootstrapped variance
> head(bootd)
> sum(is.na(bootd))
> # Means and standard errors from bootstrap
> apply(bootd, MARGIN=2, function(x)
+ c(mean=mean(x), stderror=sd(x)))
> # Parallel bootstrap under Windows
> library(parallel) # Load package parallel
> ncores <- detectCores() - 1 # Number of cores
> compclust <- makeCluster(ncores) # initialize compute cluster
> bootd <- parLapply(compclust, 1:1e4,
  function(x, datav) {
     samplev <- datav[sample.int(nrows, replace=TRUE)]
      c(sd=sd(samplev), mad=mad(samplev))
   }, datav=datav) # end parLapply
> stopCluster(compclust) # Stop R processes over cluster
> # Parallel bootstrap under Mac-OSX or Linux
> bootd <- mclapply(1:1e4, function(x) {
      samplev <- datav[sample.int(nrows, replace=TRUE)]
     c(sd=sd(samplev), mad=mad(samplev))
    }, mc.cores=ncores) # end mclapply
> # Means and standard errors from bootstrap
> bootd <- rutils::do call(rbind, bootd)
> apply(bootd, MARGIN=2, function(x)
+ c(mean=mean(x), stderror=sd(x)))
```

draft: Robust Estimators and Influence Functions

The influence function measures the sensitivity of an estimator to changes in the values of individual data points.

But for distributions with very large kurtosis (fat tails), the *median* may have a smaller standard error than the mean, because it's less sensitive to outliers.

Statistical estimators are functions of samples (which are random variables), and therefore are themselves random variables

The standard error (SE) of an estimator is defined as its standard deviation (not to be confused with the population standard deviation of the underlying random variable).

For example, the *standard error* of the estimator of the mean is equal to:

$$\sigma_{\mu} = \frac{\sigma}{\sqrt{n}}$$

Where σ is the population standard deviation (which is usually unknown).

The estimator of this standard error is equal to:

$$SE_{\mu} = \frac{\hat{\sigma}}{\sqrt{n}}$$

where: $\hat{\sigma}^2 = \frac{1}{n-1} \sum_{i=1}^n (x_i - \bar{x})^2$ is the sample standard deviation (the estimator of the population

- > set.seed(1121, "Mersenne-Twister", sample.kind="Rejection")
- > # Sample from Standard Normal Distribution
- > nrows <- 1e3
- > datav <- rnorm(nrows)
- > # Sample mean > mean(datay)
- > # Sample standard deviation
- > sd(datav)

The Wilcoxon Signed Rank Test for Distribution Symmetry

The null hypothesis of the Wilcoxon Signed Rank test is that the data sample is symmetric about a given location.

The null hypothesis is rejected if the data sample is not centered on the given location or if it's skewed.

For a single sample of data, the Wilcoxon Signed Rank test tests if the distribution is symmetric about its location

For symmetric distributions, the Wilcoxon test only requires that the distributions have equal medians, but they can have different standard deviations, and shapes (skewness)

But for skewed distributions, the Wilcoxon test requires that the distributions be the same

The Wilcoxon Signed Rank test is a nonparametric generalization of the Student's t-test.

The null hypothesis of the Wilcoxon Signed Rank test is that the two data samples, x_i and y_i , were obtained from similar probability distributions.

The function wilcox.test() with parameter paired=TRUE calculates the Wilcoxon Signed Rank test statistic and its p-value.

If a single argument is passed into wilcox.test() then it tests if the data has zero median

```
> nrous <- 1e3
> # Wilcoxon test for normal distribution
> normv <- rnorm(nrows)
> wilcox.test(normy)
> wilcox.test(normv+1)
> # Skewed distribution with median=0
> lognormy <- exp(normy)
> # lognormv <- rlnorm(nrows, sdlog=1)
> mean(lognormy); median(lognormy)
> # Skewed distribution with median!=0
> wilcox.test(lognormy)
> # Skewed distribution with median=0
> wilcox.test(lognormy-median(lognormy))
> # Skewed distributions with median!=0
> wilcox.test(lognormv, normv, paired=TRUE)
> # Two distributions with median=0
> wilcox.test(lognormv-median(lognormv), normv-median(normv),
        paired=TRUE)
> # Skewed distribution with median=0
> wilcox.test(lognormv-median(lognormv))
> # Skewed distribution with mean=0
> mean(lognormv); median(lognormv)
> wilcox.test(lognormv-median(lognormv))
> # Same as
> wilcox.test(lognormv-median(lognormv), rep(0, nrows), paired=TRUE
> # Skewed distributions with median!=0
```

> wilcox.test(lognormv-median(lognormv), normv-median(normv),

> # Normal samples with different standard deviations

> wilcox.test(lognormv, normv, paired=TRUE) > # Two distributions with median=0

paired=TRUE)

draft: The W Statistic of the Wilcoxon Signed Rank Test

Let $\{x_1, \ldots, x_n\}$ and $\{y_1, \ldots, y_n\}$ be two samples of data, which form pairs of observations:

$$\{x_1, y_1\}, \ldots, \{x_n, y_n\}.$$

The *W* statistic of the *Wilcoxon Signed Rank* test is equal to the sum of the ranks of the absolute differences $r_i = \text{rank}(|x_i - y_i|)$, weighted by their signs:

$$W = \sum_{i=1}^n \operatorname{sgn}(x_i - y_i) r_i$$

The function wilcox.test() returns the V statistic, not the the W statistic:

$$V = \sum_{i=1}^n \mathsf{H}(x_i - y_i) r_i$$

Where H(x) = 1 if x > 0, and 0 otherwise.

- > # Wilcoxon test for random data around 0 > unifv <- (runif(nrows) - 0.5) > wilcoxt <- wilcox.test(unifv) > # Calculate V statistic of Wilcoxon test > wilcoxt\$statistic > sum(rank(abs(unifv))[unifv>0]) > # Calculate W statistic of Wilcoxon test > sum(sigm(unifv)+rank(abs(unifv))) > # Two sets of normal data
- > sample1 <- rnorm(nrows)
- > sample2 <- rnorm(nrows, mean=0.1)
 > # Wilcoxon test
- > wilcoxt <- wilcox.test(sample1, sample2, paired=TRUE)
 > wilcoxt\$statistic
- > # Calculate V statistic of Wilcoxon test
 > datav <- (sample1 sample2)</pre>
- > sum(rank(abs(datav))[datav>0])
- > # Calculate W statistic of Wilcoxon test
- > sum(sign(datav)*rank(abs(datav)))

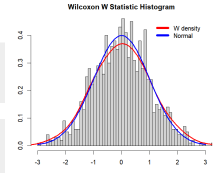
draft: The Distribution of the W Statistic

The W statistic follows a distribution without a simple formula, which converges to the normal distribution for large sample size n, with an expected value equal to 0 and a variance equal to $\frac{n(n+1)(2n+1)}{6}$.

> lines(density(wilcoxw, bw=0.4), lwd=3, col="red")
> curve(expr=dnorm, add=TRUE, lwd=3, col="blue")

> legend("topright", inset=0.05, bty="n",
+ leg=c("W density", "Normal"),
+ lwd=6, lty=1, col=c("red", "blue"))

> # Add legend



draft: The Wilcoxon Signed Rank Two-Sample Test

The *null hypothesis* of the *Wilcoxon Signed Rank* test is that the two data samples, x_i and y_i , were obtained from *similar* probability distributions.

Fix?

For symmetric distributions, the Wilcoxon test only requires that the distributions have equal medians, but they can have different standard deviations. and shapes (skewness)

Fix?

But for *skewed* distributions, the *Wilcoxon* test requires that the distributions be the same.

The Wilcoxon Signed Rank test is a nonparametric generalization of the Student's t-test.

The function wilcox.test() with parameter paired=TRUE calculates the *Wilcoxon Signed Rank* test statistic and its *p*-value.

If a single argument is passed into wilcox.test() then it tests if the data has zero *median*.

```
> nrous <- 1e3
> # Wilcoxon test for normal distribution
> normv <- rnorm(nrows)
> wilcox.test(normy)
> wilcox.test(normv+1)
> # Skewed distribution with median=0
> lognormy <- rlnorm(nrows, sdlog=1)
> mean(lognormv); median(lognormv)
> # Skewed distribution with median!=0
> wilcox.test(lognormy)
> # Skewed distribution with median=0
> wilcox.test(lognormv-median(lognormv))
> # Skewed distributions with median!=0
> wilcox.test(lognormv, normv, paired=TRUE)
> # Two distributions with median=0
> wilcox.test(lognormv-median(lognormv), normv-median(normv),
        paired=TRUE)
> # Skewed distribution with median=0
> wilcox.test(lognormv-median(lognormv))
> # Skewed distribution with mean=0
> mean(lognormv); median(lognormv)
> wilcox.test(lognormv-median(lognormv))
> # Same as
> wilcox.test(lognormv-median(lognormv), rep(0, nrows), paired=TRUE
> # Skewed distributions with median!=0
> wilcox.test(lognormv, normv, paired=TRUE)
> # Two distributions with median=0
> wilcox.test(lognormv-median(lognormv), normv-median(normv),
        paired=TRUE)
> # Normal samples with different standard deviations
> sample1 <- rnorm(nrows, sd=1)
```

draft: The W Statistic of the Wilcoxon Signed Rank Two-Sample Test

Let $\{x_1,\ldots,x_n\}$ and $\{y_1,\ldots,y_n\}$ be two samples of data, which form pairs of observations:

$$\{x_1, y_1\}, \ldots, \{x_n, y_n\}.$$

The *W* statistic of the *Wilcoxon Signed Rank* test is equal to the sum of the ranks of the absolute differences $r_i = \text{rank}(|x_i - y_i|)$, weighted by their signs:

$$W = \sum_{i=1}^n \operatorname{sgn}(x_i - y_i) r_i$$

The function wilcox.test() returns the V statistic, not the the W statistic:

$$V = \sum_{i=1}^n \mathsf{H}(x_i - y_i) r_i$$

Where H(x) = 1 if x > 0, and 0 otherwise.

- > # Wilcoxon test for random data around 0 > unifv <- (runif(arovs) - 0.5) > wilcoxt <- wilcox.test(unifv) > # Calculate V statistic of Wilcoxon test > wilcoxt§statistic > sum(rank(abs(unifv))[unifv>0] > # Calculate W statistic of Wilcoxon test > sum(sign(unifv)*rank(abs(unifv))) + Two sets of normal data
- > sample1 <- rnorm(nrows)
 > sample2 <- rnorm(nrows, mean=0.1)</pre>
- > # Wilcoxon test
 > wilcoxt <- wilcox.test(sample1, sample2, paired=TRUE)</pre>
- > wilcoxt\$statistic > # Calculate V statistic of Wilcoxon test
- > # Calculate V statistic of Wilcoxon temples > datav <- (samples samples)
- > sum(rank(abs(datav))[datav>0])
- > # Calculate W statistic of Wilcoxon test
- > sum(sign(datav)*rank(abs(datav)))

draft: The Distribution of the W Statistic

The W statistic follows a distribution without a simple formula, which converges to the normal distribution for large sample size n, with an expected value equal to 0 and a variance equal to $\frac{n(n+1)(2n+1)}{6}$.

```
> # Calculate distribution of Wilcoxon W statistic
> wilcoxu <- sapply(1:1e3, function(x) {
+ datav <- (runif(nrows) - 0.5)
+ sum(sign(datav)*rank(abs(datav)))
+ }) # end sapply
> wilcoxw <- wilcoxw/sqrt(nrows*(nrows*1)*(2*nrows*1)/6)
> var(wilcoxw)
```

```
> hist(wilcoxw, col="lightgrey",

* xlab="returns", breaks=50, xlim=c(-3, 3),

* ylab="frequency", freq=F&LSE,

* main="Wilcoxon W Statistic Histogram")

> lines(density(wilcoxw, bb=0.4), lwd=3, col="red")

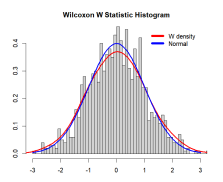
> curve(exp="ednorm, add=TRUE, lwd=3, col="blue")

* Add legend

> legend("topright", inset=0.05, bty="n",

+ leg=c("W density", "Normal"),

+ lwd=6, lty=1, col=c("red", "blue"))
```



draft: The Wilcoxon Signed Rank Test Versus Student's t-test

The *Wilcoxon* test can be considered to be a *nonparametric* analogue of the *Student's t-test*, but it tests for the *medians*, rather than the *means*.

The Wilcoxon test is nonparametric because it doesn't assume any type of sample distribution, unlike the Student's t-test which assumes that the sample is taken from the normal distribution.

The *Wilcoxon* test is more *robust* with respect to data outliers because it only depends on the ranks of the sample differences $(x_i - y_i)$, not the differences themselves.

Therefore the *Wilcoxon* test reports fewer *false positive* cases when there are outliers.

For many distributions, the *Wilcoxon* test has greater sensitivity than the *Student's t-test*.

The *sensitivity* of a statistical test is the ability to correctly identify *true positive* cases (when the null hypothesis is FALSE).

```
> # Wilcoxon test for two normal distributions
> sample1 <- rnorm(1e2)</pre>
```

- > sample2 <- rnorm(1e2, mean=0.1)
- > wilcox.test(sample1, sample2, paired=TRUE)\$p.value
- > t.test(sample1, sample2)\$p.value
- > # Wilcoxon test with data outliers
 > sample2 <- rnorm(1e2)</pre>
- > sample2 (1807m(162) > sample2[1:3] <- sample2[1:3] + 1e3
- > wilcox.test(sample1, sample2, paired=TRUE)\$p.value
- > t.test(sample1, sample2)\$p.value

The *sensitivity* of a statistical test is equal to the *true* positive rate, i.e the fraction of FALSE null hypothesis cases that are correctly classified as FALSE.

The *specificity* of a statistical test is the ability to correctly identify *true negative* cases (when the null hypothesis is TRUE).

The *specificity* of a statistical test is equal to the *true negative* rate, i.e the fraction of TRUE null hypothesis cases that are correctly classified as TRUE.

draft: The Mann-Whitney Test for Distribution Similarity

The Mann-Whitney test null hypothesis is that the two samples, x_i and y_i , were obtained from probability distributions with the same median (location).

The function wilcox.test() with parameter paired=FALSE (the default) calculates the Mann-Whitney test statistic and its p-value.

Let $\{x_1, \ldots, x_{n1}\}$ and $\{y_1, \ldots, y_{n2}\}$ be two samples of data, and let $\{z_1, \ldots, z_n\}$ be the combined data sample, with n = n1 + n2.

In contrast to the Wilcoxon Signed Rank test, the two samples don't have to be of equal length $(n1 \neq n2)$.

The Mann-Whitney test is also known as the Wilcoxon Rank Sum test, or the Mann-Whitney-Wilcoxon test.

```
> # Data samples
> datay <- sort(rnorm(38))
> indeks <- c(1:9, 20:29)
> # Or
> datay <- sort(rnorm(398))
> indeks <- c(1:99, 200:299)
> sample1 <- datav[indeks]
> sample2 <- datav[-indeks]
> # Or
> indeks <- sample(1:nrows, size=nrows/2)
> sample1 <- retp[indeks]
> sample2 <- (-retp[-indeks])
> sample1 <- rpois(1e2, lambda=4)
> sample1 <- rnorm(1e2)
> sample1 <- (sample1- median(sample1))
> sample2 <- runif(1e2)
> sample2 <- (sample2- median(sample2))
> moments::moment(sample1, order=3)
> moments::moment(sample2, order=3)
> # Mann-Whitney test for normal distribution
> wilcox.test(sample1, sample2, paired=FALSE)
> wilcox.test(sample1, sample2, paired=TRUE)
> blue <- rgb(0, 0, 1, alpha=0.5)
> red <- rgb(1, 0, 0, alpha=0.5)
> barplot(sample2, col=red)
> barplot(sample1, col=blue, add=TRUE)
> hist(sample1)
> # Mann-Whitney test for normal distribution
> datay <- rnorm(nrows, sd=100)
> wilcox.test(datav.paired=FALSE)
```

draft: The U Statistic of the Mann-Whitney Test

Let z_i be the combined data of two sub-samples: x_i and v_i , with n = n1 + n2.

Let $r_i = \text{rank}(z_i)$ be the ranks of the sample, and r_i^1 and r_i^2 be the ranks of the sub-samples.

The sum of all the ranks is equal to:

$$\sum_{i=1}^{n} r_i = \sum_{i=1}^{n} r_i^1 + \sum_{i=1}^{n^2} r_i^2 = \frac{n(n+1)}{2}$$

The Mann-Whitney test statistics U1 and U2 are equal to the sum of the ranks minus a term accounting for the size of the samples:

$$U1 = \sum_{i=1}^{n1} r_i^1 - \frac{n1(n1+1)}{2}$$

$$U2 = \sum_{i=1}^{n2} r_i^2 - \frac{n2(n2+1)}{2}$$

So that U1 + U2 = n1n2.

The U statistic follows a distribution without a simple formula, which converges to the normal distribution for large sample size n.

In the extreme case when the ranks of the first sample are all greater than the second sample $(r_i^1 > r_j^2)$, then U2 = 0 and U1 = n1n2.

```
> # Data samples
> sample1 <- rnorm(200)
> sample2 <- rnorm(100, mean=0.1)
> # Mann-Whitney-Wilcoxon rank sum test
> wilcoxt <- wilcox.test(sample1, sample2,
+ paired=FALSE)
> wilcoxt$statistic
> # Calculate U statistics of Mann-Whitney-Wilcoxon test
> datav <- c(sample1, sample2)
> ranks <- rank(datav)
> ranks (2 rank(datav)
> sum(ranks[1:200]) - 100*201
> sum(ranks[201:3001) - 50*101
```

draft: The U Statistic of the Wilcoxon Rank Sum Test

The Wilcoxon Rank Sum test statistic U is equal to the sum of the ranks $r_i = \text{rank}(|x_i - y_i|)$ of the absolute differences weighted by their signs:

$$U = \sum_{i=1}^{n} \operatorname{sgn}(x_i - y_i) r_i$$

The statistic U follows a distribution without a simple formula, which converges to the normal distribution for large sample size n, with an expected value equal to 0 and a variance equal to $\frac{n(n+1)(2n+1)}{6}$.

The Wilcoxon Rank Sum test is nonparametric because it doesn't assume any type of sample distribution, unlike the Student's t-test which assumes that the sample is taken from the normal distribution.

The Wilcoxon Rank Sum test is more robust with respect to data outliers because it only depends on the ranks of the sample differences $(x_i - y_i)$, not the differences themselves.

- > # Wilcoxon test for random data around 0 > nrows <- 1e3
- > datav <- (runif(nrows) 0.5)
- > wilcoxt <- wilcox.test(datav)
 > # Calculate V statistic of Wilcoxon test
- > wilcoxt\$statistic
- > sum(rank(abs(datav))[datav>0])
- > # Calculate W statistic of Wilcoxon test
- > sum(sign(datav)*rank(abs(datav)))
- > # Calculate distributon of Wilcoxon W statistic
- > wilcoxw <- sapply(1:1e3, function(x) {
- + datav <- (runif(nrows) 0.5)
 + sum(sign(datav)*rank(abs(datav)))</pre>
- + }) # end sapply
- > wilcoxw <- wilcoxw/sqrt(nrows*(nrows+1)*(2*nrows+1)/6)
 > var(wilcoxw)
- > hist(wilcoxw)

The function wilcox.test() returns the V statistic, not the the W statistic:

$$V = \sum_{i=1}^n \mathsf{H}(x_i - y_i) r_i$$

Where H(x) = 1 if x > 0, and 0 otherwise.

The Kruskal-Wallis Test for the Distribution Similarity

The *Kruskal-Wallis* test is designed to test the *null hypothesis* that sub-samples of the data corresponding to different categories follow similar distributions.

The Kruskal-Wallis test can be used as a type of nonparametric ANOVA test, to test if the sub-samples of the data have the same mean.

For example, given the heights of several different species of trees, the *Kruskal-Wallis* test can test if all the species have the same height.

The Kruskal-Wallis test can also test if samples have different skewness, even if they have the same means.

The function kruskal.test() accepts a vector of sample data and a factor specifying the categories, and calculates the *Kruskal-Wallis* statistic and its *p*-value.

The function kruskal.test() can also accept the data as a formula combined with a matrix or data frame.

```
> # iris data frame
> aggregate(Sepal.Length ~ Species, data=iris,
    FUN=function(x) c(mean=mean(x), sd=sd(x)))
> # Kruskal-Wallis test for iris data
> ktest <- kruskal.test(Sepal.Length ~ Species, data=iris)
> str(ktest)
> ktest$statistic
> # Kruskal-Wallis test for independent normal distributions
> sample1 <- rnorm(1e3)
> sample2 <- rnorm(1e3)
> groupv <- c(rep(TRUE, 1e3), rep(FALSE, 1e3))
> kruskal.test(x=c(sample1, sample2), g=groupv)
> # Kruskal-Wallis test for shifted normal distributions
> kruskal.test(x=c(sample1+1, sample2), g=groupv)
> # Kruskal-Wallis test for beta distributions
> sample1 <- rbeta(1e3, 2, 8) + 0.3
> sample2 <- rbeta(1e3, 8, 2) - 0.3
> mean(sample1); mean(sample2)
> kruskal.test(x=c(sample1, sample2), g=groupv)
> # Plot the beta distributions
> x11()
> plot(density(sample1), col="blue", lwd=3,
       xlim=range(c(sample1, sample2)), xlab="samples",
       main="Two samples from beta distributions with equal means")
> lines(density(sample2), col="red", lwd=3)
```

The Kruskal-Wallis Test Statistic

Given a data sample x_i with n elements, and a factor of k categories, the sample can be divided into k sub-samples x_i^j .

Let $r_i = \text{rank}(x_i)$ be the ranks of the sample, and r_i^I be the ranks of the sub-samples.

The Kruskal-Wallis test statistic H is proportional to the sum of squared differences between the average rank of the sample $\bar{r}=\frac{n+1}{2}$, minus the average ranks of the sub-samples \bar{r}_i :

$$H = \frac{12}{n(n+1)} \sum_{j=1}^{k} (\frac{n+1}{2} - \bar{r}_j)^2 n_j$$

Where the sum is over all the k categories, and n_j is the number of elements in sub-sample j.

The H statistic follows a distribution without a simple formula, which is approximately equal to the ${\it chi-squared}$ distribution with k-1 degrees of freedom.

- > # Kruskal-Wallis test for iris data
- > ktest <- kruskal.test(Sepal.Length ~ Species, data=iris)
- > # Calculate Kruskal-Wallis test Statistic
- > nrows <- NROW(iris)
- > irisdf <- data.frame(ranks=rank(iris\$Sepal.Length),
- + species=iris\$Species)
 > kruskal_stat <- (12/nrows/(nrows+1))*sum(
- + aggregate(ranks ~ species, data=irisdf,
- + FUN=function(x) {NROW(x)*((nrows+1)/2 mean(x))^2})[, 2])
- > c(ktest=unname(ktest\$statistic), k_stat=kruskal_stat)

The Kruskal-Wallis Test with Data Outliers

The Kruskal-Wallis test is nonparametric because it doesn't assume any type of sample distribution.

The Kruskal-Wallis test is also robust with respect to data outliers, since it only depends on the ranks of the sample.

When a few data outliers are added to the data, Student's t-test rejects the null hypothesis that the means are equal, but the Kruskal-Wallis test still accepts the null hypothesis that the distributions are similar.

- > # Kruskal-Wallis test with data outliers
- > sample1 <- rnorm(1e3) > sample2 <- rnorm(1e3)
- > sample2 (- Inform(1e3) > sample2[1:11] <- sample2[1:11] + 50
- > groupv <- c(rep(TRUE, 1e3), rep(FALSE, 1e3))
- > kruskal.test(x=c(sample1, sample2), g=groupv)\$p.value
- > t.test(sample1, sample2)\$p.value

> nrous <- 1e3 > datay <- rnorm(nrows)

draft: Robust Estimators of Dispersion

The Median Absolute Deviation (MAD) is a robust measure of dispersion (variability), defined using the median instead of the mean:

$$MAD = median(abs(x_i - median(x)))$$

The advantage of MAD is that it's always well defined. even for data that has infinite variance.

For normally distributed data, the MAD has a larger standard error than the standard deviation

But for distributions with fat tails (like asset returns), the standard deviation has a larger standard error than the MAD

The MAD for normally distributed data is equal to $\Phi^{-1}(0.75) \cdot \hat{\sigma} = 0.6745 \cdot \hat{\sigma}.$

The function mad() calculates the MAD and divides it by $\Phi^{-1}(0.75)$ to make it comparable to the standard deviation.

```
> sd(datav)
> mad(datav)
> median(abs(datav - median(datav)))
> median(abs(datav - median(datav)))/qnorm(0.75)
> # Bootstrap of sd and mad estimators
> bootd <- sapply(1:1e4, function(x) {
    samplev <- datav[sample.int(nrows, replace=TRUE)]
    c(sd=sd(samplev), mad=mad(samplev))
+ }) # end sapply
> bootd <- t(bootd)
> # Analyze bootstrapped variance
> head(bootd)
> sum(is.na(bootd))
> # Means and standard errors from bootstrap
> apply(bootd, MARGIN=2,
+ function(x) c(mean=mean(x), stderror=sd(x)))
> # Parallel bootstrap under Windows
> library(parallel) # Load package parallel
> ncores <- detectCores() - 1 # Number of cores
> compclust <- makeCluster(ncores) # initialize compute cluster
> bootd <- parLapply(compclust, 1:1e4,
   function(x, datav) {
      samplev <- datav[sample.int(nrows, replace=TRUE)]
      c(sd=sd(samplev), mad=mad(samplev))
    }, datav=datav) # end parLapply
> stopCluster(compclust) # Stop R processes over cluster
> # Parallel bootstrap under Mac-OSX or Linux
> bootd <- mclapply(1:1e4,
    function(x) {
      samplev <- datav[sample.int(nrows, replace=TRUE)]
      c(sd=sd(samplev), mad=mad(samplev))
    }, mc.cores=ncores) # end mclapply
> # Means and standard errors from bootstrap
> bootd <- rutils::do call(rbind, bootd)
> apply(bootd, MARGIN=2, function(x)
                                    May 12, 2025
```

draft: Robust Estimators of Skewness

The medcouple robust skewness estimator

The Median Absolute Deviation (MAD) is a robust measure of dispersion (variability), defined using the median instead of the mean:

$$\mathsf{MAD} = \mathsf{median}(\mathsf{abs}(x_i - \mathsf{median}(\mathbf{x})))$$

The advantage of MAD is that it's always well defined, even for data that has infinite variance

For normally distributed data, the MAD has a larger standard error than the standard deviation.

But for distributions with fat tails (like asset returns), the standard deviation has a larger standard error than the MAD.

The MAD for normally distributed data is equal to $\Phi^{-1}(0.75) \cdot \hat{\sigma} = 0.6745 \cdot \hat{\sigma}.$

The function mad() calculates the MAD and divides it by $\Phi^{-1}(0.75)$ to make it comparable to the standard deviation

```
> nrous <- 1e3
> datay <- rnorm(nrows)
> sd(datav)
> mad(datav)
> median(abs(datav - median(datav)))
> median(abs(datav - median(datav)))/qnorm(0.75)
> # Bootstrap of sd and mad estimators
> bootd <- sapply(1:1e4, function(x) {
   samplev <- datav[sample.int(nrows, replace=TRUE)]
   c(sd=sd(samplev), mad=mad(samplev))
+ }) # end sapply
> bootd <- t(bootd)
> # Analyze bootstrapped variance
> head(bootd)
> sum(is.na(bootd))
> # Means and standard errors from bootstrap
> apply(bootd, MARGIN=2, function(x)
+ c(mean=mean(x), stderror=sd(x)))
> # Parallel bootstrap under Windows
> library(parallel) # Load package parallel
> ncores <- detectCores() - 1 # Number of cores
> compclust <- makeCluster(ncores) # initialize compute cluster
> bootd <- parLapply(compclust, 1:1e4,
  function(x, datav) {
     samplev <- datav[sample.int(nrows, replace=TRUE)]
     c(sd=sd(samplev), mad=mad(samplev))
   }, datav=datav) # end parLapply
> stopCluster(compclust) # Stop R processes over cluster
> # Parallel bootstrap under Mac-OSX or Linux
> bootd <- mclapply(1:1e4, function(x) {
      samplev <- datav[sample.int(nrows, replace=TRUE)]
     c(sd=sd(samplev), mad=mad(samplev))
    }, mc.cores=ncores) # end mclapply
> # Means and standard errors from bootstrap
> bootd <- rutils::do call(rbind, bootd)
> apply(bootd, MARGIN=2, function(x)
+ c(mean=mean(x), stderror=sd(x)))
                                    May 12, 2025
```

Pearson Correlation

The covariance σ_{xy} between two sets of data, x_i and y_i , is defined as:

$$\sigma_{xy} = \frac{\sum_{i=1}^{n} (x_i - \bar{x})(y_i - \bar{y})}{n-1}$$

Where $\bar{x} = \frac{1}{n} \sum_{i=1}^{n} x_i$ and $\bar{y} = \frac{1}{n} \sum_{i=1}^{n} y_i$ are the *mean* values.

The function cov() calculates the covariance between two numeric vectors.

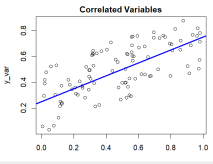
The Pearson correlation ρ_P is equal to the covariance divided by the standard deviations σ_x and σ_y :

$$\rho_P = \frac{\sigma_{xy}}{\sigma_x \sigma_y}$$

The function cor() calculates the correlation between two numeric vectors.

Depending on the argument "method", it calculates either the *Pearson* (default), *Spearman*, or *Kendall* correlations

Jerzy Pawlowski (NYU Tandon)



- > # Initialize the random number generator
- > set.seed(1121, "Mersenne-Twister", sample.kind="Rejection")
- > # Define variables and calculate correlation
- > nrows <- 100
 > xvar <- runif(nrows); yxvar <- runif(nrows)</pre>
- > cor(xvar, yxvar)
- > rho <- 0.5
- > yxvar <- rho*xvar + (1-rho)*yxvar
- > # Plot in x11 window
- > x11(width=5, height=4)
- > # Set plot parameters to reduce whitespace around plot

> # Correlate the variables and calculate correlation

- > par(mar=c(4, 4, 2, 1), oma=c(0.5, 0.5, 0, 0))
- > # Plot scatterplot and exact regression line
- > plot(xvar, yxvar, xlab="xvar", ylab="yxvar")
 > title(main="Correlated Variables", line=0.5)
- > abline(a=0.25, b=rho, lwd=3, col="blue")
 > # Calculate regression
- Probability and Statistics

draft: Standard Error of Pearson Correlation

```
See my comment to Jean Paul comment:
https://www.jstor.org/stable/2277400
```

https://www.tandfonline.com/doi/abs/10.1080/00220973.1956.11010555 https://stats.stackexchange.com/questions/73621/standard-error-from-

correlation-coefficient/262893

https://stats.stackexchange.com/questions/226380/derivation-of-thestandard-error-for-pearsons-correlation-coefficient

https://stats.stackexchange.com/questions/154362/confidence-interval-onpoint-biserial-correlation-coefficient

Let x be standard normally distributed with mean zero

$$E[x] = 0$$
 and standard deviation equal to one: $E[x^2] = 1$.

Let
$$y = \rho x + \varepsilon \sqrt{1 - \rho^2}$$
, where ε is standard normally distributed, with zero correlation to x : $E[x\varepsilon] = 0$.

Then v is also standard normally distributed, with correlation to x equal to ρ : $E[xy] = E[x(\rho x + \varepsilon \sqrt{1 - \rho^2})] = E[\rho x^2] = \rho.$

The variance of the correlation is equal to:

$$E[(xy - \rho)^{2}] = E[\rho^{2}x^{4} + 2x^{3}\varepsilon\rho\sqrt{1 - \rho^{2}} + x^{2}\varepsilon^{2}(1 - \rho^{2}) - 2\rho xy + \rho^{2}] = 1 - \rho^{2} + \rho^{2}E[x^{4}] = 3\rho^{2} + 1 - \rho^{2} - 2\rho^{2} + \rho^{2}.$$

$$E[\rho^{2}x^{4} + 2x^{3}\varepsilon\rho\sqrt{1 - \rho^{2}} + x^{2}\varepsilon^{2}(1 - \rho^{2})] = 1 - \rho^{2} + 3\rho^{2}$$

The standard error of

the correlation estimator is equal to: $\frac{1}{\sqrt{n-2}}$, and slowly decreases as the square root of n - the length of the data.

The function cor.test() performs a test of the

> # Simulation of sample correlation > nrows <- 1e4 > rho <- 0.99 > rho2 <- sqrt(1-rho^2) > datav <- sapply(1:1000, function(x) { xvar <- rnorm(nrows) vxvar <- (rho*xvar + rho2*rnorm(nrows)) cor(xvar, vxvar) + }) # end sapply > sd(datav) > # Correct formula > (1-rho^2)/sqrt(nrows-2) > # Incorrect formula > sart((1-rho^2)/(nrows-2)) > # Correlation > corv <- cor(xvar, yxvar) > # Standard error of correlation > stderror <- sqrt((1-corv^2)/(nrows-2)) > # t-value of correlation > corv/stderror > # 95% confidence intervals

> corv*c(1-qnorm(0.975)*stderror, 1+qnorm(0.975)*stderror)

> # Test statistical significance of correlation > cor.test(xvar, yxvar)

> rho2 <- sqrt(1-rho^2) > set.seed(1121, "Mersenne-Twister", sample.kind="Rejection")

> rho <- 0.9

> # Bootstrap of sample mean and median > bootd <- sapply(1:1000, function(x) { xvar <- rnorm(nrows)

yxvar <- (rho*xvar + rho2*rnorm(nrows)) c(rho=mean(yxvar*xvar), y_sd=sd(yxvar), cor=cor(xvar, yxvar))

Spearman Rank Correlation

The *Spearman* correlation ρ_S is equal to the *Pearson* correlation between the ranks rx_i and ry_i of the variables x_i and y_i :

$$\rho_{S} = \frac{\sum_{i=1}^{n} (rx_{i} - \bar{r}x)(ry_{i} - \bar{r}y)}{(n-1)\sigma_{rx}\sigma_{ry}}$$

If the ranks are all distinct integers, then the *Spearman* correlation ρ_S can be expressed as:

$$\rho_{S} = 1 - \frac{6\sum_{i=1}^{n} dr_{i}^{2}}{n(n^{2} - 1)}$$

Where $dr_i = rx_i - ry_i$ are the differences between the ranks.

The *Spearman* correlation is a *robust* measure of association because it depends on the ranks, so it's not sensitive to the extreme values of the variables x_i and y_i .

The *Spearman* correlation is considered a *nonparametric* estimator because it does not depend on the joint probability distribution of the variables x_i and y_i .

- > # Calculate correlations
- > cor(xvar, yxvar, method="pearson")
- > cor(xvar, yxvar, method="spearman")
 > # Test statistical significance of correlations
- > cor.test(xvar, yxvar, method="pearson")
- > cor.test(xvar, yxvar, method="spearman")

Kendall's au Correlation

The pair of observations $\{x_i, y_i\}$ is concordant with the pair $\{x_j, y_j\}$ if the signs of the differences $(rx_i - rx_j)$ and $(ry_i - ry_j)$ are the same, i.e. if their ranks follow the same order.

The Kendall correlation τ_K (Kendall's τ) is equal to the difference between the number of concordant pairs of observations, minus the number of discordant pairs:

$$\tau_K = \frac{2}{n(n-1)} \sum_{i < j}^n \operatorname{sgn}(rx_i - rx_j) \operatorname{sgn}(ry_i - ry_j)$$

The Kendall correlation τ_K is also a robust and nonparametric estimator of association, because it only depends on the ranks, so it's not sensitive to the extreme values of the variables x_i and y_i .

- > # Calculate correlations
- > cor(xvar, yxvar, method="pearson")
 > cor(xvar, yxvar, method="kendall")
- > # Test statistical significance of correlations
- > cor.test(xvar, yxvar, method="pearson")
- > cor.test(xvar, yxvar, method="kendall")