# FRE6871 R in Finance

Lecture#2, Fall 2023

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#### Benchmarking the Speed of R Code

The function system.time() calculates the execution time (in seconds) used to evaluate a given expression.

system.time() returns the "user time" (execution time of user instructions), the "system time" (execution time of operating system calls), and "elapsed time" (total execution time, including system latency waiting).

The function microbenchmark() from package microbenchmark calculates and compares the execution time of R expressions (in milliseconds), and is more accurate than system.time().

The time it takes to execute an expression is not always the same, since it depends on the state of the processor, caching, etc.

microbenchmark() executes the expression many times, and returns the distribution of total execution times in a *data frame*.

```
> library(microbenchmark)
> vecv < runif(1e6)
> # sqrt() and "^0.5" are the same
> all.equal(sqrt(vecv), vecv^0.5)
> # sqrt() is much faster than "^0.5"
> system.time(vecv^0.5)
> microbenchmark(
```

+ power = vecv^0.5,
+ sqrt = sqrt(vecv),

times=10)

The "times" parameter is the number of times the expression is evaluated.

The choice of the "times" parameter is a tradeoff between the time it takes to run microbenchmark(), and the desired accuracy,

#### Writing Fast R Code Using Compiled C++ Functions

Compiled C++ functions directly call compiled C++ or Fortran code, which performs the calculations and returns the result back to R.

This makes compiled C++ functions much faster than interpreted functions, which have to be parsed by R.

 $\operatorname{sum}()$  is much faster than  $\operatorname{mean}()$ , because  $\operatorname{sum}()$  is a compiled function, while  $\operatorname{mean}()$  is an interpreted function.

Given a single argument, any() is equivalent to %in%, but is much faster because it's a *compiled* function.

%in% is a wrapper for match() defined as follows:
"%in%" <- function(x, table) match(x, table,
nomatch=0) > 0.

The function all.equal() tests the equality of two objects to within the square root of the *machine* precision.

```
> # sum() is a compiled primitive function
> s11m
> # mean() is a generic function
> mean
> vecv <- runif(1e6)
> # sum() is much faster than mean()
> all.equal(mean(vecv), sum(vecv)/NROW(vecv))
> library(microbenchmark)
> summary(microbenchmark(
    mean = mean(vecv).
    sum = sum(vecv)/NROW(vecv).
    times=10))[, c(1, 4, 5)]
> # any() is a compiled primitive function
> anv
> # any() is much faster than %in% wrapper for match()
> all.equal(1 %in% vecv. anv(vecv == 1))
> summary(microbenchmark(
    inop = {1 %in% vecv}.
    anvfun = anv(vecv == 1).
```

times=10))[, c(1, 4, 5)]

## Increasing Speed of Loops by Pre-allocating Memory

 $\ensuremath{\mathbb{R}}$  performs automatic memory management as users assign values to objects.

R doesn't require allocating the full memory for vectors or lists, and allows appending new data to existing objects ("growing" them).

For example, R allows assigning a value to a vector element that doesn't exist yet.

This forces R to allocate additional memory for that element, which carries a small speed penalty.

But when data is appended to an object using the functions c(), append(), cbind(), or rbind(), then R allocates memory for the whole new object and copies all the existing values, which is very memory intensive and slow.

It is therefore preferable to pre-allocate memory for large objects before performing loops.

The function numeric(k) returns a numeric vector of zeros of length k, while numeric(0) returns an empty (zero length) numeric vector (not to be confused with a NULL object).

```
> vecv <- rnorm(5000)
> summary(microbenchmark(
+ # Compiled C++ function
    cpp = cumsum(vecv), # end for
+ # Allocate full memory for cumulative sum
   forloop = {cumsumv <- numeric(NROW(vecv))
     cumsumv[1] <- vecv[1]
     for (i in 2:NROW(vecv)) {
        cumsumv[i] <- cumsumv[i-1] + vecv[i]
     }}. # end for
+ # Allocate zero memory for cumulative sum
   growvec = {cumsumv <- numeric(0)
      cumsumv[1] <- vecv[1]
      for (i in 2:NROW(vecv)) {
+ # Add new element to "cumsumv" ("grow" it)
        cumsumv[i] <- cumsumv[i-1] + vecv[i]
     }}. # end for
+ # Allocate zero memory for cumulative sum
    combine = {cumsumv <- numeric(0)
     cumsumv[1] <- vecv[1]
      for (i in 2:NROW(vecv)) {
 # Add new element to "cumsumv" ("grow" it)
        cumsumv <- c(cumsumv, vecv[i])
     }}, # end for
    times=10))[, c(1, 4, 5)]
```

#### Benchmarking the Speed of R Code

The function system.time() calculates the execution time (in seconds) used to evaluate a given expression.

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The function microbenchmark() from package microbenchmark calculates and compares the execution time of R expressions (in milliseconds), and is more accurate than system.time().

The time it takes to execute an expression is not always the same, since it depends on the state of the processor, caching, etc.

microbenchmark() executes the expression many times, and returns the distribution of total execution times.

- > library(microbenchmark)
  > vecv <- runif(1e6)
- > # sqrt() and "^0.5" are the same
  > all.equal(sqrt(vecv), vecv^0.5)
  > # sqrt() is much faster than "^0.5"
- > system.time(vecv^0.5)
- > microbenchmark(
  + power = vecv^0.5.
- + sqrt = sqrt(vecv),
- + times=10)

The "times" parameter is the number of times the expression is evaluated.

The choice of the "times" parameter is a tradeoff between the time it takes to run microbenchmark(), and the desired accuracy,

#### Vectorized Functions for Vector Computations

Vectorized functions accept vectors as their arguments, and return a vector of the same length as their value.

Many vectorized functions are also compiled (they pass their data to compiled C++ code), which makes them very fast.

The following *vectorized compiled* functions calculate cumulative values over large vectors:

- cummax()
- cummin()
- cumsum()
- cumprod()

Standard arithmetic operations ("+", "-", etc.) can be applied to vectors, and are implemented as vectorized compiled functions.

ifelse() and which() are vectorized compiled functions for logical operations.

But many *vectorized* functions perform their calculations in R code, and are therefore slow, but convenient to use.

```
> vector1 <- rnorm(1000000)
> vector2 <- rnorm(1000000)
> big_vector <- numeric(1000000)
> # Sum two vectors in two different ways
> summary(microbenchmark(
    # Sum vectors using "for" loop
    rloop = (for (i in 1:NROW(vector1)) {
      big_vector[i] <- vector1[i] + vector2[i]
   F).
    # Sum vectors using vectorized "+"
    vecvized = (vector1 + vector2).
    times=10))[, c(1, 4, 5)] # end microbenchmark summary
> # Allocate memory for cumulative sum
> cumsumv <- numeric(NROW(big_vector))
> cumsumv[1] <- big vector[1]
> # Calculate cumulative sum in two different ways
> summary(microbenchmark(
+ # Cumulative sum using "for" loop
    rloop = (for (i in 2:NROW(big vector)) {
      cumsumv[i] <- cumsumv[i-1] + big vector[i]
    1).
+ # Cumulative sum using "cumsum"
    vecvized = cumsum(big vector).
    times=10))[, c(1, 4, 5)] # end microbenchmark summary
```

#### **Vectorized Functions for Matrix Computations**

apply() loops are very inefficient for calculating statistics over rows and columns of very large matrices.

R has very fast vectorized compiled functions for calculating sums and means of rows and columns:

- rowSums()
- colSums()
- rowMeans()
- colMeans()

These vectorized functions are also compiled functions. so they're very fast because they pass their data to compiled C++ code, which performs the loop calculations

- > # Calculate matrix of random data with 5.000 rows > matv <- matrix(rnorm(10000), ncol=2)
- > # Calculate row sums two different ways
- > all.equal(rowSums(matv), apply(matv, 1, sum))
- > summary(microbenchmark( rowsumv = rowSums(matv),
- apply = apply(matv, 1, sum),
- times=10))[, c(1, 4, 5)] # end microbenchmark summary

#### Package matrixStats for Fast Matrix Computations

The package *matrixStats* contains functions for calculating aggregations over matrix columns and rows, and other matrix computations, such as:

- estimating location and scale: rowRanges(), colRanges(), and rowMaxs(), rowMins(), etc.,
- testing and counting values: colAnyMissings(), colAnys(), etc.,
- cumulative functions: colCumsums(), colCummins(), etc.,
- binning and differencing: binCounts(), colDiffs(), etc.,

A summary of matrixStats functions can be found under:

https://cran.r-project.org/web/packages/matrixStats/vignettes/matrixStats-methods.html

The matrixStats functions are very fast because they are *compiled* functions (compiled from C++ code).

- > install.packages("matrixStats") # Install package matrixStats
  > library(matrixStats) # Load package matrixStats
  > # Calculate row minimum values two different ways
  > all.equal(matrixStats::rowMins(matv), do.call(pmin.int, lapply(1:
- + function(indeks) matv[, indeks])))
  > # Calculate row minimmum values three different ways
  > summarv(microbenchmark(
- + rowmins = matrixStats::rowMins(matv), + pmin = do.call(pmin.int. lapply(1:NCOL(matv).
- + function(indeks) matv[, indeks])),
- + as\_dframe = do.call(pmin.int, as.data.frame.matrix(matv)),
  + times=10))[, c(1, 4, 5)] # end microbenchmark summarv

> install.packages("Rfast") # Install package Rfast

#### Package Rfast for Fast Matrix and Numerical Computations

The package *Rfast* contains functions for fast matrix and numerical computations, such as:

- colMedians() and rowMedians() for matrix column and row medians,
- colCumSums(), colCumMins() for cumulative sums and min/max,
- eigen.sym() for performing eigenvalue matrix decomposition,

The Rfast functions are very fast because they are compiled functions (compiled from C++ code).

```
> library(Rfast) # Load package Rfast
> # Benchmark speed of calculating ranks
> vecv < 1e3
> all.equal(rank(vecv), Rfast::Rank(vecv))
> library(microbenchmark)
```

- > summary(microbenchmark( + rcode = rank(vecv).
- + Rfast = Rfast::Rank(vecv),
- + times=10))[, c(1, 4, 5)] # end microbenchmark summary
  > # Benchmark speed of calculating column medians
- > matv <- matrix(1e4, nc=10)
- > all.equal(matrixStats::colMedians(matv), Rfast::colMedians(matv))
  > summary(microbenchmark(
- + matrixStats = matrixStats::colMedians(matv),
- + Rfast = Rfast::colMedians(matv),
  - times=10))[, c(1, 4, 5)] # end microbenchmark summary

## Writing Fast R Code Using Vectorized Operations

R-style code is code that relies on *vectorized compiled* functions, instead of for() loops.

for() loops in R are slow because they call functions multiple times, and individual function calls are compute-intensive and slow.

The brackets "[]" operator is a *vectorized compiled* function, and is therefore very fast.

Vectorized assignments using brackets "[]" and Boolean or integer vectors to subset vectors or matrices are therefore preferable to for() loops.

R code that uses *vectorized compiled* functions can be as fast as C++ code.

R-style code is also very *expressive*, i.e. it allows performing complex operations with very few lines of code.

```
> summary(microbenchmark( # Assign values to vector three differen + # Fast vectorized assignment loop performed in C using brackets |
```

- brackets = {vecv <- numeric(10); vecv[] <- 2},
- + # Slow because loop is performed in R
- forloop = {vecv <- numeric(10)
- for (indeks in seq\_along(vecv))
- vecv[indeks] <- 2},
- + times=10))[, c(1, 4, 5)] # end microbenchmark summary
  > summary(microbenchmark( # Assign values to vector two different
- + # Fast vectorized assignment loop performed in C using brackets
- + brackets = {vecv <- numeric(10); vecv[4:7] <- rnorm(4)},
- + # Slow because loop is performed in R + forloop = {vecv <- numeric(10)
- + for (indeks in 4:7)
- + vecv[indeks] <- rnorm(1)},</pre>
- + times=10))[, c(1, 4, 5)] # end microbenchmark summary

#### **Vectorized Functions**

Functions which use vectorized operations and functions are automatically *vectorized* themselves.

Functions which only call other compiled C++ vectorized functions, are also very fast.

But not all functions are vectorized, or they're not vectorized with respect to their *parameters*.

Some vectorized functions perform their calculations in R code, and are therefore slow, but convenient to use.

```
> # Define function vectorized automatically
```

- > myfun <- function(input, param) {
- + param\*input
  + } # end myfun
- > # "input" is vectorized
- > myfun(input=1:3, param=2)
  > # "param" is vectorized
- > myfun(input=10, param=2:4)
- > # Define vectors of parameters of rnorm()
- > stdevs <- structure(1:3, names=paste0("sd=", 1:3))
  > means <- structure(-1:1, names=paste0("mean=", -1:1))</pre>
- > # "sd" argument of rnorm() isn't vectorized
- > rnorm(1, sd=stdevs)
- > # "mean" argument of rnorm() isn't vectorized
- > rnorm(1, mean=means)

## Performing sapply() Loops Over Function Parameters

Many functions aren't vectorized with respect to their parameters.

Performing sapply() loops over a function's parameters produces vector output.

```
> # Loop over stdevs produces vector output
> set.seed(1121)
> sapply(stdevs, function(stdev) rnorm(n=2, sd=stdev))
> set.seed(1121)
> sapply(stdevs, rnorm, n=2, mean=0)
> # Loop over means
> set.seed(1121)
> sapply(means, function(meanv) rnorm(n=2, mean=meanv))
> # Same
> set.seed(1121)
> sapply(means, rnorm, n=2)
```

#### Vectorized if-else Statements Using Function ifelse()

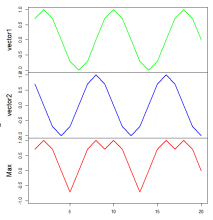
The function ifelse() performs vectorized if-else statements on vectors.

ifelse() is much faster than performing an element-wise loop in R.

> # Create two numeric vectors

```
> vector1 <- sin(0.25*pi*1:20)
> vector2 <- cos(0.25*pi*1:20)
> # Create third vector using 'ifelse'
> vector3 <- ifelse(vector1 > vector2, vector1, vector2)
> # cbind all three together
> vector3 <- cbind(vector1, vector2, vector3)
> colnames(vector3)[3] <- "Max"
> # Set plotting parameters
> xi1(vidth=6, height=7)
> par(ona=c(0, 1, 1, 1), mar=c(0, 2, 2, 1),
+ mgp=c(2, 1, 0), cex.lab=0.5, cex.axis=1.0, cex.main=1.8, cex.i
> # Plot matrix
> zoo::plot.zoo(vector3, lwd=2, ylim=c(-1, 1),
+ xlab="", col=c("green", "blue", "red"),
+ main=""ifelse() Calculates The Max of Two Data Sets")
```

#### ifelse() Calculates The Max of Two Data Sets



# Parallel Computing in R

#### Parallel Computing in R

Parallel computing means splitting a computing task into separate sub-tasks, and then simultaneously computing the sub-tasks on several computers or CPU cores.

There are many different packages that allow parallel computing in R, most importantly package *parallel*, and packages foreach, doParallel, and related packages:

```
http://cran.r-project.org/web/views/HighPerformanceComputing.html \\ http://blog.revolutionanalytics.com/high-performance-computing/http://gforge.se/2015/02/how-to-go-parallel-in-r-basics-tips/
```

#### R Base Package parallel

The package parallel provides functions for parallel computing using multiple cores of CPUs,

The package parallel is part of the standard R distribution, so it doesn't need to be installed.

```
http://adv\text{-}r.had.co.nz/Profiling.html\#parallelise
```

https://github.com/tobigithub/R-parallel/wiki/R-parallel-package-overview

#### Packages foreach, doParallel, and Related Packages

http://blog.revolutionanalytics.com/2015/10/updates-to-the-foreach-package-and-its-friends.html

## Parallel Computing Using Package parallel

The package *parallel* provides functions for parallel computing using multiple cores of CPUs.

The package *parallel* is part of the standard R distribution, so it doesn't need to be installed.

Different functions from package *parallel* need to be called depending on the operating system (*Windows, Mac-OSX*, or *Linux*).

Parallel computing requires additional resources and time for distributing the computing tasks and collecting the output, which produces a computing overhead.

Therefore parallel computing can actually be slower for small computations, or for computations that can't be naturally separated into sub-tasks.

- > library(parallel) # Load package parallel
  > # Get short description
- > packageDescription("parallel")
- > # Load help page
  > help(package="parallel")
- > help(package="parallel")
- > # List all objects in "parallel"
- > ls("package:parallel")

# Performing Parallel Loops Using Package parallel

Some computing tasks naturally lend themselves to parallel computing, like for example performing loops.

Different functions from package parallel need to be called depending on the operating system (Windows. Mac-OSX. or Linux).

The function mclapply() performs loops (similar to lapply()) using parallel computing on several CPU cores under Mac-OSX or Linux.

Under Windows, a cluster of R processes (one per each CPU core) need to be started first, by calling the function makeCluster().

Mac-OSX and Linux don't require calling the function makeCluster()

The function parLapply() is similar to lapply(), and performs loops under Windows using parallel computing on several CPU cores.

```
> # Define function that pauses execution
> paws <- function(x, sleep_time=0.01) {
+ Sys.sleep(sleep_time)
+ } # end paws
> library(parallel) # Load package parallel
> # Calculate number of available cores
> ncores <- detectCores() - 1
> # Initialize compute cluster under Windows
> cluster <- makeCluster(ncores)
> # Perform parallel loop under Windows
> outv <- parLapply(cluster, 1:10, paws)
> # Perform parallel loop under Mac-OSX or Linux
> outv <- mclapply(1:10, paws, mc.cores=ncores)
> library(microbenchmark) # Load package microbenchmark
> # Compare speed of lapply versus parallel computing
> summary(microbenchmark(
```

# parallel = parLapply(cluster, 1:10, paws), + parallel = mclapply(1:10, paws, mc.cores=ncores),

standard = lapply(1:10, paws),

times=10)

+ )[, c(1, 4, 5)]

#### Computing Advantage of Parallel Computing

Parallel computing provides an increasing advantage for larger number of loop iterations.

The function stopCluster() stops the R processes running on several CPU cores.

The function plot() by default plots a scatterplot, but can also plot lines using the argument type="1".

The function lines() adds lines to a plot.

The function legend() adds a legend to a plot.

```
> # Compare speed of lapply with parallel computing
> runv <- 3:10
> timev <- sapply(runv, function(nruns) {
      summary(microbenchmark(
+ standard = lapply(1:nruns, paws),
+ # parallel = parLapply(cluster, 1:nruns, paws),
+ parallel = mclapply(1:nruns, paws, mc.cores=ncores),
+ times=10))[, 4]
      }) # end sapply
> timev <- t(timev)
> colnames(timev) <- c("standard", "parallel")
> rownames(timev) <- runv
> # Stop R processes over cluster under Windows
> stopCluster(cluster)
```

#### Compute times 8 standard parallel 8 8 4 20 10 number of iterations in loop

```
> x11(width=6, height=5)
> plot(x=rownames(timev),
       y=timev[, "standard"],
       type="1", lwd=2, col="blue",
       main="Compute times",
       xlab="Number of iterations in loop", ylab="",
       vlim=c(0, max(timev[, "standard"])))
> lines(x=rownames(timev).
+ v=timev[, "parallel"], lwd=2, col="green")
> legend(x="topleft", legend=colnames(timev),
+ inset=0.1, cex=1.0, btv="n", bg="white",
+ v.intersp=0.3, lwd=2, ltv=1, col=c("blue", "green"))
```

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# Parallel Computing Over Matrices

Very often we need to perform time consuming calculations over columns of matrices.

The function parCapply() performs an apply loop over columns of matrices using parallel computing on several CPU cores

- > # Calculate matrix of random data
  > matv <- matrix(rnorm(1e5), ncol=100)</pre>
- > # Define aggregation function over column of matrix
  > aggfun <- function(column) {</pre>
- + output <- 0
- + for (indeks in 1:NROW(column))
  + output <- output + column[indeks]</pre>
- output
- + output
- + } # end aggfun
- > # Perform parallel aggregations over columns of matrix
- > aggs <- parCapply(cluster, matv, aggfun)
- > # Compare speed of apply with parallel computing
- > summary(microbenchmark(
- + apply=apply(matv, MARGIN=2, aggfun),
- + parapply=parCapply(cluster, matv, aggfun),
  + times=10)
- + )[, c(1, 4, 5)]
- > # Stop R processes over cluster under Windows
- > stopCluster(cluster)

#### Initializing Parallel Clusters Under Windows

Under *Windows* the child processes in the parallel compute cluster don't inherit data and objects from their parent process.

Therefore the required data must be either passed into parLapply() via the dots "..." argument, or by calling the function clusterExport().

Objects from packages must be either referenced using the double-colon operator "::", or the packages must be loaded in the child processes.

```
> basep <- 2
> # Fails because child processes don't know basep:
> parLapply(cluster, 2:4, function(exponent) basep^exponent)
> # basep passed to child via dots ... argument:
> parLapply(cluster, 2:4,
      function(exponent, basep) basep^exponent,
      basep=basep)
> # basep passed to child via clusterExport:
> clusterExport(cluster, "basep")
> parLapply(cluster, 2:4,
      function(exponent) basep^exponent)
> # Fails because child processes don't know zoo::index():
> parSapply(cluster, c("VTI", "IEF", "DBC"),
      function(symbol)
        NROW(zoo::index(get(symbol, envir=rutils::etfenv))))
> # zoo function referenced using "::" in child process:
> parSapply(cluster, c("VTI", "IEF", "DBC"),
      function(symbol)
        NROW(zoo::index(get(symbol, envir=rutils::etfenv))))
> # Package zoo loaded in child process:
> parSapply(cluster, c("VTI", "IEF", "DBC"),
      function(symbol) {
        stopifnot("package:zoo" %in% search() || require("zoo", qui
        NROW(zoo::index(get(symbol, envir=rutils::etfenv)))
      }) # end parSapply
> # Stop R processes over cluster under Windows
> stopCluster(cluster)
```

#### Reproducible Parallel Simulations Under Windows

Simulations use pseudo-random number generators, and in order to perform reproducible results, they must set the *seed* value, so that the number generators produce the same sequence of pseudo-random numbers.

The function set.seed() initializes the random number generator by specifying the seed value, so that the number generator produces the same sequence of numbers for a given seed value.

But under *Windows* set.seed() doesn't initialize the random number generators of child processes, and they don't produce the same sequence of numbers.

The function clusterSetRNGStream() initializes the random number generators of child processes under Windows.

The function set.seed() does initialize the random number generators of child processes under *Mac-OSX* and *Linux*.

- > library(parallel) # Load package parallel
  > # Calculate number of available cores
- > ncores <- detectCores() 1
- > # Initialize compute cluster under Windows
- > cluster <- makeCluster(ncores)
- > # Set seed for cluster under Windows
- > # Doesn't work: set.seed(1121)
- > clusterSetRNGStream(cluster, 1121)
- > # Perform parallel loop under Windows
- > output <- parLapply(cluster, 1:70, rnorm, n=100)
  > sum(unlist(output))
- > # Stop R processes over cluster under Windows
- > stopCluster(cluster)
- > stopCluster(cluster)
- > # Perform parallel loop under Mac-OSX or Linux
- > output <- mclapply(1:10, rnorm, mc.cores=ncores, n=100)

#### Monte Carlo Simulation

 ${\it Monte \ Carlo} \ {\it simulation \ consists} \ {\it of \ generating \ random} \ {\it samples \ from \ a \ given \ probability \ distribution}.$ 

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

The *quantile* of a probability distribution is the value of the *random variable* x, such that the probability of values less than x is equal to the given *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 <code>?quantile</code>.

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

```
> set.seed(1121) # Reset random number generator
> # Sample from Standard Normal Distribution
> nsimu <- 1000
> datav <- rnorm(nsimu)
> # Sample mean - MC estimate
> mean(datay)
> # Sample standard deviation - MC estimate
> sd(datav)
> # Monte Carlo estimate of cumulative probability
> pnorm(-2)
> sum(datav < (-2))/nsimu
> # Monte Carlo estimate of quantile
> confl <- 0.02
> gnorm(confl) # Exact value
> cutoff <- confl*nsimu
> datay <- sort(datay)
> datav[cutoff] # Naive Monte Carlo value
> quantile(datay, probs=confl)
> # Analyze the source code of quantile()
> stats:::quantile.default
> # Microbenchmark quantile
> library(microbenchmark)
> summary(microbenchmark(
    monte_carlo = datav[cutoff],
    quanty = quantile(datay, probs=confl),
```

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

#### Standard Errors of Estimators Using Bootstrap Simulation

The bootstrap procedure uses Monte Carlo simulation to generate a distribution of estimator values.

The bootstrap procedure generates new data by randomly sampling with replacement from the observed (empirical) data set.

If the original data consists of simulated random numbers then we simply simulate another set of these random numbers

The bootstrapped datasets are used to recalculate the estimator many times, to provide a distribution of the estimator and its standard error

- > # Sample from Standard Normal Distribution
- > nsimu <- 1000; datav <- rnorm(nsimu)
- > # Sample mean and standard deviation > mean(datav); sd(datav)
- > # Bootstrap of sample mean and median
- > nboot <- 10000
- > bootd <- sapply(1:nboot, function(x) {
- # Sample from Standard Normal Distribution samplev <- rnorm(nsimu)
- c(mean=mean(samplev), median=median(samplev))
- + }) # end sapply
- > bootd[, 1:3] > bootd <- t(bootd)
- > # Standard error from formula
- > sd(datav)/sqrt(nsimu)
- > # Standard error of mean from bootstrap > sd(bootd[, "mean"])
- > # Standard error of median from bootstrap
- > sd(bootd[, "median"])

#### The Distribution of Estimators Using Bootstrap Simulation

The standard errors of estimators can be calculated using a bootstrap simulation.

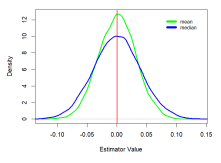
The *bootstrap* procedure generates new data by randomly sampling with replacement from the observed (empirical) data set.

The bootstrapped dataset is used to recalculate the estimator many times.

The *bootstrapped* estimator values are then used to calculate the probability distribution of the estimator and its standard error.

The function density() calculates a kernel estimate of the probability density for a sample of data.

#### Distribution of Bootstrapped Mean and Median



- > # Plot the densities of the bootstrap data
- > x11(width=6, height=5)
- > plot(density(bootd[, "mean"]), lwd=3, xlab="Estimator Value",
  + main="Distribution of Bootstrapped Mean and Median", col="gr
- > lines(density(bootd[, "median"]), lwd=3, col="blue")
- > abline(v=mean(bootd[, "mean"]), lwd=2, col="red")
- > legend("topright", inset=0.05, cex=0.8, title=NULL,
- + leg=c("mean", "median"), bty="n", y.intersp=0.4,
- + lwd=6, bg="white", col=c("green", "blue"))

#### **Bootstrapping Using Vectorized Operations**

Bootstrap simulations can be accelerated by using vectorized operations instead of R loops.

But using vectorized operations requires calculating a matrix of random data, instead of calculating random vectors in a loop.

This is another example of the tradeoff between speed and memory usage in simulations.

Faster code often requires more memory than slower code.

```
> set.seed(1121) # Reset random number generator
> nsimu <- 1000
> # Bootstrap of sample mean and median
> nhoot <- 100
> bootd <- sapply(1:nboot, function(x) median(rnorm(nsimu)))
> # Perform vectorized bootstrap
> set.seed(1121) # Reset random number generator
> # Calculate matrix of random data
> samplev <- matrix(rnorm(nboot*nsimu), ncol=nboot)
> booty <- matrixStats::colMedians(sampley)
> all.equal(bootd, booty)
> # Compare speed of loops with vectorized R code
> library(microbenchmark)
> summary(microbenchmark(
    loop = sapply(1:nboot, function(x) median(rnorm(nsimu))),
      samplev <- matrix(rnorm(nboot*nsimu), ncol=nboot)
      matrixStats::colMedians(samplev)
```

times=10))[, c(1, 4, 5)] # end microbenchmark summary

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#### Bootstrapping Standard Errors Using Parallel Computing

The *bootstrap* procedure performs a loop, which naturally lends itself to parallel computing.

Different functions from package *parallel* need to be called depending on the operating system (*Windows*, *Mac-OSX*, or *Linux*).

The function makeCluster() starts running R processes on several CPU cores under *Windows*.

The function parLapply() is similar to lapply(), and performs loops under *Windows* using parallel computing on several CPU cores.

The R processes started by makeCluster() don't inherit any data from the parent R process.

Therefore the required data must be either passed into parLapply() via the dots "..." argument, or by calling the function clusterExport().

The function mclapply() performs loops using parallel computing on several CPU cores under *Mac-OSX* or *Linux*.

The function stopCluster() stops the R processes running on several CPU cores.

```
> library(parallel) # Load package parallel
> ncores <- detectCores() - 1 # Number of cores
> cluster <- makeCluster(ncores) # Initialize compute cluster under
> set.seed(1121) # Reset random number generator
> # Sample from Standard Normal Distribution
> nsimu <- 1000
> # Bootstrap mean and median under Windows
> nboot <- 10000
> bootd <- parLapply(cluster, 1:nboot,
   function(x, datay, nsimu) {
   samplev <- rnorm(nsimu)
   c(mean=mean(samplev), median=median(samplev))
   }, datav=datav, nsimu*nsimu) # end parLapply
> # Bootstrap mean and median under Mac-OSX or Linux
> bootd <- mclapply(1:nboot,
   function(x) {
   samplev <- rnorm(nsimu)
   c(mean=mean(sampley), median=median(sampley))
   }, mc.cores=ncores) # end mclapply
> bootd <- rutils::do call(rbind, bootd)
> # Means and standard errors from bootstrap
> apply(bootd, MARGIN=2, function(x) c(mean=mean(x), stderror=sd(x)
> # Standard error from formula
```

> stopCluster(cluster) # Stop R processes over cluster under Windo

> sd(datav)/sqrt(nsimu)

#### Parallel Bootstrapping of the Median Absolute Deviation

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.

```
> nsimu <- 1000
> datav <- rnorm(nsimu)
> sd(datav): mad(datav)
> median(abs(datav - median(datav)))
> median(abs(datay - median(datay)))/gnorm(0.75)
> # Bootstrap of sd and mad estimators
> nboot <- 10000
> bootd <- sapply(1:nboot, function(x) {
   samplev <- rnorm(nsimu)
+ 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
> cluster <- makeCluster(ncores) # Initialize compute cluster
> bootd <- parLapply(cluster, 1:nboot,
+ function(x, datav) {
     samplev <- rnorm(nsimu)
     c(sd=sd(samplev), mad=mad(samplev))
   }, datav=datav) # end parLapply
> # Parallel bootstrap under Mac-OSX or Linux
> bootd <- mclapply(1:nboot, function(x) {
   samplev <- rnorm(nsimu)
   c(sd=sd(samplev), mad=mad(samplev))
+ }, mc.cores=ncores) # end mclapply
> stopCluster(cluster) # Stop R processes over cluster
> bootd <- rutils::do call(rbind, bootd)
> # Means and standard errors from bootstrap
> apply(bootd, MARGIN=2, function(x) c(mean=mean(x), stderror=sd(x))
```

#### Resampling From Empirical Datasets

Resampling is randomly selecting data from an existing dataset, to create a new dataset with similar properties to the existing dataset.

Resampling is usually performed with replacement, so that each draw is independent from the others.

Resampling is performed when it's not possible or convenient to obtain another set of empirical data, so we simulate a new data set by randomly sampling from the existing data.

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

The function sample.int() is a *method* that selects a random sample of *integers*.

The function sample.int() with argument replace=TRUE selects a sample with replacement (the integers can repeat).

The function sample.int() is a little faster than sample().

- > # Calculate time series of VII returns
  > library(rutils)
  > retp < rutils::etfenv\$returns\$VII
  > retp < na.omit(retp)
  > # Sample from VII returns
  > samplev <- retp[sample.int(nsimu, replace=TRUE)]
  > £ (sad=sGamplev), mad@mad(samplev))
  > # sample int() is a little faster than sample()
  > library(microbenchmark)
  > summary(microbenchmark)
- + sample = sample(1e3), + times=10))[, c(1, 4, 5)]

sample.int = sample.int(1e3),

> stderrors[2, ]/stderrors[1, ]

#### **Bootstrapping From Empirical Datasets**

Bootstrapping is usually performed by resampling from an observed (empirical) dataset.

Resampling consists of randomly selecting data from an existing dataset, with replacement.

Resampling produces a new *bootstrapped* dataset with similar properties to the existing dataset.

The bootstrapped dataset is used to recalculate the estimator many times.

The bootstrapped estimator values are then used to calculate the probability distribution of the estimator and its standard error

Bootstrapping shows that for asset returns, the *Median Absolute Deviation* (*MAD*) has a smaller relative standard error than the standard deviation.

Bootstrapping doesn't provide accurate estimates for estimators which are sensitive to the ordering and correlations in the data.

```
> # Sample from time series of VTI returns
> library(rutils)
> retp <- rutils::etfenv$returns$VTI
> retp <- na.omit(retp)
> nsimu <- NROW(retp)
> # Bootstrap sd and MAD under Windows
> library(parallel) # Load package parallel
> ncores <- detectCores() - 1 # Number of cores
> cluster <- makeCluster(ncores) # Initialize compute cluster under
> clusterSetRNGStream(cluster, 1121) # Reset random number generate
> nboot <- 10000
> bootd <- parLapply(cluster, 1:nboot,
   function(x, retp, nsimu) {
     samplev <- retp[sample.int(nsimu, replace=TRUE)]
     c(sd=sd(samplev), mad=mad(samplev))
   }, retp=retp, nsimu*nsimu) # end parLapply
> # Bootstrap sd and MAD under Mac-OSX or Linux
> bootd <- mclapply(1:nboot, function(x) {
      samplev <- retp[sample.int(nsimu, replace=TRUE)]
     c(sd=sd(samplev), mad=mad(samplev))
   }, mc.cores=ncores) # end mclapply
> stopCluster(cluster) # Stop R processes over cluster under Windo
> bootd <- rutils::do call(rbind, bootd)
> # Standard error of standard deviation assuming normal distributi
> sd(retp)/sqrt(nsimu)
> # Means and standard errors from bootstrap
> stderrors <- apply(bootd, MARGIN=2,
   function(x) c(mean=mean(x), stderror=sd(x)))
> stderrors
> # Relative standard errors
```

#### Standard Errors of Regression Coefficients Using Bootstrap

The standard errors of the regression coefficients can be calculated using a *bootstrap* simulation.

The bootstrap procedure creates new design matrices by randomly sampling with replacement from the regression design matrix.

Regressions are performed on the *bootstrapped* design matrices, and the regression coefficients are saved into a matrix of *bootstrapped* coefficients.

```
> # Initialize random number generator
> set.seed(1121)
> # Define predictor and response variables
> nsimu <- 100
> predm <- rnorm(nsimu, mean=2)
> noisev <- rnorm(nsimu)
> respv <- (-3 + 2*predm + noisev)
> desm <- cbind(respv, predm)
> # Calculate alpha and beta regression coefficients
> betav <- cov(desm[, 1], desm[, 2])/var(desm[, 2])
> alpha <- mean(desm[, 1]) - betav*mean(desm[, 2])
> x11(width=6, height=5)
> plot(respv ~ predm, data=desm)
> abline(a=alpha, b=betav, lwd=3, col="blue")
> # Bootstrap of beta regression coefficient
> nboot <- 100
> bootd <- sapply(1:nboot, function(x) {
    samplev <- sample.int(nsimu, replace=TRUE)
    desm <- desm[samplev, ]
+ cov(desm[, 1], desm[, 2])/var(desm[, 2])
+ }) # end sapply
```

## Distribution of Bootstrapped Regression Coefficients

The *bootstrapped* coefficient values can be used to calculate the probability distribution of the coefficients and their standard errors,

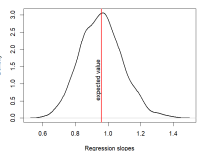
The function  ${\tt density}()$  calculates a kernel estimate of the probability density for a sample of data.

abline() plots a straight line on the existing plot.

The function text() draws text on a plot, and can be used to draw plot labels.

```
> # Mean and standard error of beta regression coefficient
> c(mean=mean(bootd), stderror=sd(bootd))
# Plot density of bootstrapped beta coefficients
> plot(density(bootd), lud=2, xlab="Regression slopes",
# main="Bootstrapped Regression Slopes")
> # Add line for expected value
> abline(v=mean(bootd), lud=2, col="red")
> text(x=mean(bootd)-0.01, v=1.0. labels="expected value".
```

#### **Bootstrapped Regression Slopes**



1wd=2, srt=90, pos=3)

#### Bootstrapping Regressions Using Parallel Computing

The *bootstrap* procedure performs a loop, which naturally lends itself to parallel computing.

Different functions from package *parallel* need to be called depending on the operating system (*Windows*, *Mac-OSX*, or *Linux*).

The function makeCluster() starts running R processes on several CPU cores under *Windows*.

The function parLapply() is similar to lapply(), and performs loops under *Windows* using parallel computing on several CPU cores.

The R processes started by makeCluster() don't inherit any data from the parent R process.

Therefore the required data must be passed into parLapply() via the dots "..." argument.

The function mclapply() performs loops using parallel computing on several CPU cores under *Mac-OSX* or *Linux*.

The function stopCluster() stops the R processes running on several CPU cores.

```
> library(parallel) # Load package parallel
> ncores <- detectCores() - 1 # Number of cores
> cluster <- makeCluster(ncores) # Initialize compute cluster under
> # Bootstrap of regression under Windows
> bootd <- parLapply(cluster, 1:1000,
   function(x, desm) {
     samplev <- sample.int(nsimu, replace=TRUE)
     desm <- desm[samplev, ]
     cov(desm[, 1], desm[, 2])/var(desm[, 2])
    }, design=desm) # end parLapply
> # Bootstrap of regression under Mac-OSX or Linux
> bootd <- mclapply(1:1000.
   function(x) {
     samplev <- sample.int(nsimu, replace=TRUE)
     desm <- desm[samplev, ]
     cov(desm[, 1], desm[, 2])/var(desm[, 2])
   }, mc.cores=ncores) # end mclapply
> stopCluster(cluster) # Stop R processes over cluster under Windo
```

#### Analyzing the Bootstrap Data

The *bootstrap* loop produces a *list* which can be collapsed into a vector.

The function unlist() collapses a list with atomic elements into a vector (which can cause type coercion).

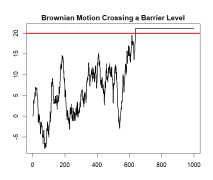
- > # Collapse the bootstrap list into a vector > class(bootd)
- > bootd <- unlist(bootd)
- $\gt$  # Mean and standard error of beta regression coefficient
- > c(mean=mean(bootd), stderror=sd(bootd))
- > # Plot density of bootstrapped beta coefficients
- > plot(density(bootd),
- + lwd=2, xlab="Regression slopes",
  + main="Bootstrapped Regression Slopes")
- > # Add line for expected value
- > abline(v=mean(bootd), lwd=2, col="red")
- > text(x=mean(bootd)-0.01, y=1.0, labels="expected value",
- + lwd=2, srt=90, pos=3)

#### Simulating Brownian Motion Using while() Loops

while() loops are often used in simulations, when the number of required loops is unknown in advance.

Below is an example of a simulation of the path of *Brownian Motion* crossing a barrier level.

```
> set.seed(1121) # Reset random number generator
> barl <- 20 # Barrier level
> nsimu <- 1000 # Number of simulation steps
> pathv <- numeric(nsimu) # Allocate path vector
> pathv[1] <- rnorm(1) # Initialize path
> it <- 2 # Initialize simulation index
> while ((it <= nsimu) && (pathv[it - 1] < barl)) {
+ # Simulate next step
+ pathv[it] <- pathv[it - 1] + rnorm(1)
+ it <- it + 1 # Advance index
+ } # end while
> # Fill remaining path after it crosses barl
> if (it <= nsimu)
   pathv[it:nsimu] <- pathv[it - 1]
> # Plot the Brownian motion
> x11(width=6, height=5)
> par(mar=c(3, 3, 2, 1), oma=c(1, 1, 1, 1))
> plot(pathv, type="1", col="black",
      lty="solid", lwd=2, xlab="", ylab="")
> abline(h=barl, lwd=3, col="red")
> title(main="Brownian Motion Crossing a Barrier Level", line=0.5)
```



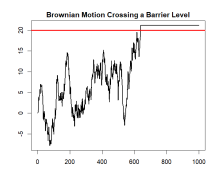
## Simulating Brownian Motion Using Vectorized Functions

Simulations in R can be accelerated by pre-computing a vector of random numbers, instead of generating them one at a time in a loop.

Vectors of random numbers allow using *vectorized* functions, instead of inefficient (slow) while() loops.

> set.seed(1121) # Reset random number generator

```
> barl <- 20 # Barrier level
> nsimu <- 1000 # Number of simulation steps
> # Simulate path of Brownian motion
> pathv <- cumsum(rnorm(nsimu))
> # Find index when path crosses barl
> crossp <- which(pathv > barl)
> # Fill remaining path after it crosses barl
> if (NROW(crossp)>0) {
   pathv[(crossp[1]+1):nsimu] <- pathv[crossp[1]]
+ } # end if
> # Plot the Brownian motion
> x11(width=6, height=5)
> par(mar=c(3, 3, 2, 1), oma=c(1, 1, 1, 1))
> plot(pathv, type="1", col="black",
      ltv="solid", lwd=2, xlab="", vlab="")
> abline(h=barl, lwd=3, col="red")
> title(main="Brownian Motion Crossing a Barrier Level", line=0.5)
```



The tradeoff between speed and memory usage: more memory may be used than necessary, since the simulation may stop before all the pre-computed random numbers are used up.

But the simulation is much faster because the path is simulated using  $\emph{vectorized}$  functions,

## Estimating the Statistics of Brownian Motion

The statistics of Brownian motion can be estimated by simulating multiple paths.

An example of a statistic is the expected value of Brownian motion at a fixed time horizon, which is the option payout for the strike price k:  $\mathbb{E}[(p_t - k)_+]$ .

Another statistic is the probability of Brownian motion crossing a boundary (barrier) b:  $\mathbb{E}[\mathbb{1}(p_t - b)]$ .

```
> # Define Brownian motion parameters
> sigmav <- 1.0 # Volatility
> drift <- 0.0 # Drift
> nsteps <- 1000 # Number of simulation steps
> nsimu <- 100 # Number of simulation paths
> # Simulate multiple paths of Brownian motion
> set.seed(1121)
> pathm <- rnorm(nsimu*nsteps, mean=drift, sd=sigmav)
> pathm <- matrix(pathm, nc=nsimu)
> pathm <- matrixStats::colCumsums(pathm)
> # Final distribution of paths
> mean(pathm[nsteps, ]); sd(pathm[nsteps, ])
> # Calculate option payout at maturity
> strikep <- 50 # Strike price
> payouts <- (pathm[nsteps, ] - strikep)
> sum(payouts[payouts > 0])/nsimu
> # Calculate probability of crossing the barrier at any point
> barl <- 50
> crossi <- (colSums(pathm > barl) > 0)
```

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Paths of Brownian Motion

- > # Plot in window
- > x11(width=6, height=5)
- > par(mar=c(4, 3, 2, 2), oma=c(0, 0, 0, 0), mgp=c(2.5, 1, 0))
- > # Select and plot full range of paths > ordern <- order(pathm[nsimu, ])
- > pathm[nsimu, ordern]
- > indeks <- ordern[seq(1, 100, 9)]
- > zoo::plot.zoo(pathm[, indeks], main="Paths of Brownian Motion",
- + xlab="time steps", ylab=NA, plot.type="single")
- > abline(h=strikep, col="red", lwd=3)
- > text(x=(nsimu-60), y=strikep, labels="strike price", pos=3, cex=1

> sum(crossi)/nsimu

#### Variance Reduction Using Antithetic Sampling

Variance reduction are techniques for increasing the precision of Monte Carlo simulations.

Naive Monte Carlo refers to Monte Carlo simulation without using variance reduction techniques.

Antithetic Sampling is a variance reduction technique in which a new random sample is computed from an existing sample, without generating new random numbers.

In the case of a *Normal* random sample  $\phi$ , the new antithetic sample is equal to minus the existing sample:  $\phi_{new} = -\phi$ .

In the case of a *Uniform* random sample  $\phi$ , the new *antithetic* sample is equal to 1 minus the existing sample:  $\phi_{\text{new}} = 1 - \phi$ .

Antithetic Sampling doubles the number of independent samples, so it reduces the standard error by  $\sqrt{2}$ .

Antithetic Sampling doesn't change any other parameters of the simulation.

```
> set.seed(1121) # Reset random number generator
> # Sample from Standard Normal Distribution
> nsimu <- 1000
> datav <- rnorm(nsimu)
> # Estimate the 95% quantile
> nboot <- 10000
> bootd <- sapply(1:nboot, function(x) {
    samplev <- datav[sample.int(nsimu, replace=TRUE)]
    quantile(sampley, 0.95)
+ }) # end sapply
> sd(bootd)
> # Estimate the 95% quantile using antithetic sampling
> bootd <- sapply(1:nboot, function(x) {
    samplev <- datav[sample.int(nsimu, replace=TRUE)]
    quantile(c(samplev, -samplev), 0.95)
+ }) # end sapply
> # Standard error of quantile from bootstrap
> sd(bootd)
```

> sart(2)\*sd(bootd)

## Simulating Rare Events Using Probability Tilting

Rare events can be simulated more accurately by *tilting* (deforming) their probability distribution, so that rare events occur more frequently.

A popular probability *tilting* method is exponential (Esscher) tilting:

$$p(x,\lambda) = \frac{\exp(\lambda x)p(x)}{\int_{-\infty}^{\infty} \exp(\lambda x)p(x)dx}$$

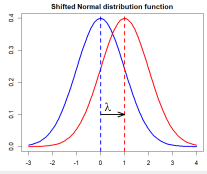
Where p(x) is the probability density,  $p(x, \lambda)$  is the tilted density, and  $\lambda$  is the tilt parameter.

For the Normal distribution  $\phi(x) = \frac{\exp(-x^2/2)}{\sqrt{2\pi}}$ , exponential tilting is equivalent to shifting the distribution by  $\lambda \colon x \to x + \lambda$ .

$$\phi(x,\lambda) = \frac{\exp(\lambda x) \exp(-x^2/2)}{\int_{-\infty}^{\infty} \exp(\lambda x) \exp(-x^2/2) dx} =$$

$$\frac{\exp(-(x-\lambda)^2/2)}{\sqrt{2\pi}} = \exp(x\lambda - \lambda^2/2) \cdot \phi(x,\lambda = 0)$$

Shifting the random variable  $x \to x + \lambda$  is equivalent to multiplying the distribution by the weight factor:  $\exp(x\lambda - \lambda^2/2)$ .



- > # Plot a Normal probability distribution
- > curve(expr=dnorm, xlim=c(-3, 4),
- + main="Shifted Normal distribution function", + xlab="", ylab="", lwd=3, col="blue")
- > # Add shifted Normal probability distribution
- > curve(expr=dnorm(x, mean=1), add=TRUE, lwd=3, col="red")
- > # Add vertical dashed lines
- > abline(v=0, lwd=3, col="blue", lty="dashed")
- > abline(v=1, lwd=3, col="red", lty="dashed")
  > arrows(x0=0, v0=0.1, x1=1, v1=0.1, lwd=3.
- + code=2, angle=20, length=grid::unit(0.2, "cm"))
- > text(x=0.3, 0.1, labels=bquote(lambda), pos=3, cex=2)

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#### Variance Reduction Using Importance Sampling

Importance sampling is a variance reduction technique for simulating rare events more accurately.

The *variance* of an estimate produced by simulation decreases with the number of events which contribute to the estimate:  $\sigma^2 \propto \frac{1}{a}$ .

Importance sampling simulates rare events more frequently by *tilting* the probability distribution, so that more events contribute to the estimate.

In standard Monte Carlo simulation, the simulated data points have equal probabilities.

But in *importance sampling*, the simulated data must be weighted (multiplied) to compensate for the tilting of the probability.

The tilt weights are equal to the ratio of the base probability distribution divided by the tilted distribution, which for the *Normal* distribution are equal to:

$$w_x = \frac{\phi(x, \lambda = 0)}{\phi(x, \lambda)} = \exp(-x\lambda + \lambda^2/2)$$

- > # Sample from Standard Normal Distribution > nsimu <- 1000 > datay <- rnorm(nsimu)
- > # Cumulative probability from formula > quantv <- (-2)
- > pnorm(quantv)
  > integrate(dnorm, lower=-Inf, upper=quantv)
- > # Cumulative probability from Naive Monte Carlo > sum(datav < quantv)/nsimu
- > # Generate importance sample > lambda <- (-1.5) # Tilt parameter
- > datat <- datav + lambda # Tilt the random numbers
- > # Cumulative probability from importance sample wrong! > sum(datat < quantv)/nsimu
- > # Cumulative probability from importance sample correct > weightv <- exp(-lambda\*datat + lambda^2/2)
- > sum((datat < quantv)\*weightv)/nsimu > # Bootstrap of standard errors of cumulative probability
- > nboot <- 1000
- > bootd <- sapply(1:nboot, function(x) {
  + datay <- rnorm(nsimu)</pre>
- + naivemc <- sum(datav < quantv)/nsimu
- + datav <- (datav + lambda) + weightv <- exp(-lambda\*datav + lambda^2/2)
- + isample <- sum((datav < quantv)\*weightv)/nsimu
- + isample <- sum((datav < quantv)\*weightv)/nsimu + c(naivemc=naivemc.impsample=isample)
- + c(naivemc=naivemc, impsample=isample)
- + }) # end sapply
- > apply(bootd, MARGIN=1, function(x) c(mean=mean(x), sd=sd(x)))

## Calculating Quantiles Using Importance Sampling

The quantiles can be calculated from the cumulative probabilities of the importance sample data.

The importance sample data points must be weighted to compensate for the tilting of the probability.

Importance sampling can be used to estimate the VaR (quantile) corresponding to a given confidence level.

The standard error of the VaR estimate using importance sampling can be several times smaller than that of naive Monte Carlo.

The reduction of standard error is greater for higher confidence levels

Naive Monte Carlo refers to Monte Carlo simulation without using variance reduction techniques.

The function findInterval() returns the indices of the intervals specified by "vec" that contain the elements of "x".

- > # Quantile from Naive Monte Carlo
- > confl <- 0.02 > qnorm(confl) # Exact value
- > datay <- sort(datay) # Must be sorted for importance sampling
- > cutoff <- nsimu\*confl
- > datav[cutoff] # Naive Monte Carlo value
- > # Importance sample weights
- > datat <- datay + lambda # Tilt the random numbers
- > weightv <- exp(-lambda\*datat + lambda^2/2) > # Cumulative probabilities using importance sample
- > cumprob <- cumsum(weightv)/nsimu
- > # Quantile from importance sample
- > datat[findInterval(confl, cumprob)]
- > # Bootstrap of standard errors of quantile
- > nboot <- 1000
- > bootd <- sapply(1:nboot, function(x) {
- datav <- sort(rnorm(nsimu))
- naivemc <- datav[cutoff]
- datat <- datay + lambda
- weightv <- exp(-lambda\*datat + lambda^2/2)
- cumprob <- cumsum(weightv)/nsimu
- isample <- datat[findInterval(confl. cumprob)] c(naivemc=naivemc, impsample=isample)
- + }) # end sapply
- > apply(bootd, MARGIN=1, function(x) c(mean=mean(x), sd=sd(x)))

## Calculating CVaR Using Importance Sampling

Importance sampling can be used to estimate the Conditional Value at Risk (*CVaR*) corresponding to a given *confidence level*.

First the VaR (quantile) is estimated, and then the expected value (CVaR) is estimated using it.

The standard error of the CVaR estimate using importance sampling can be several times smaller than that of naive Monte Carlo.

The reduction of standard error is greater for higher confidence levels

- > # VaR and CVaR from Naive Monte Carlo
  > varisk <- datay[cutoff]
- > sum((datav <= varisk)\*datav)/sum((datav <= varisk))
- > # CVaR from importance sample
- > varisk <- datat[findInterval(confl, cumprob)]
- > sum((datat <= varisk)\*datat\*weightv)/sum((datat <= varisk)\*weight
  > # CVaR from integration
- > integrate(function(x) x\*dnorm(x), low=-Inf, up=varisk)\$value/pnor > # Bootstrap of standard errors of CVaR
- > nboot <- 1000
- > bootd <- sapply(1:nboot, function(x) {
- + datav <- sort(rnorm(nsimu))
- + varisk <- datav[cutoff]
- + naivemc <- sum((datav <= varisk)\*datav)/sum((datav <= varisk))
- + datat <- datav + lambda
- + weightv <- exp(-lambda\*datat + lambda^2/2) + cumprob <- cumsum(weightv)/nsimu
- + cumprob <- cumsum(weightv)/nsimu
- + varisk <- datat[findInterval(confl, cumprob)]
- + isample <- sum((datat <= varisk)\*datat\*weightv)/sum((datat <= v. + c(naivemc=naivemc, impsample=isample)
  - + c(naivemc=naivemc, impsample=isample; + }) # end sapply
- + }) # end sapply
- > apply(bootd, MARGIN=1, function(x) c(mean=mean(x), sd=sd(x)))

## The Optimal Tilt Parameter for Importance Sampling

The tilt parameter  $\lambda$  should be chosen to minimize the standard error of the estimator.

The optimal tilt parameter depends on the estimator and on the required confidence level.

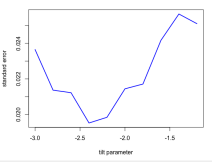
More tilting is needed at higher confidence levels, to provide enough significant data points.

When performing a loop over the tilt parameters, the same matrix of random data can be used for different tilt parameters.

The function Rfast::sort\_mat() sorts the columns of a matrix using very fast C++ code.

```
> # Calculate matrix of random data
> set.seed(1121) # Reset random number generator
> nsimu <- 1000; nboot <- 100
> datav <- matrix(rnorm(nboot*nsimu), ncol=nboot)
> datay <- Rfast::sort_mat(datay) # Sort the columns
> # Bootstrap function for VaR (quantile) for a single tilt parame' > lambdav <- seq(-3.0, -1.2, by=0.2)
> calc_vars <- function(lambda, confl=0.05) {
   datat <- datav + lambda # Tilt the random numbers
   weightv <- exp(-lambda*datat + lambda^2/2)
   # Calculate quantiles for columns
  sapply(1:nboot, function(it) {
     cumprob <- cumsum(weightv[, it])/nsimu
     datat[findInterval(confl, cumprob), it]
   }) # end sapply
   # end calc vars
> # Bootstrap vector of VaR for a single tilt parameter
> bootd <- calc_vars(-1.5)
```

#### Standard Frrors of Simulated VaR



- > # Define vector of tilt parameters
- > # Calculate vector of VaR for vector of tilt parameters
- > varisk <- sapply(lambdav, calc\_vars, confl=0.02) > # Calculate standard deviations of VaR for tilt parameters
- > stdevs <- apply(varisk, MARGIN=2, sd)
- > # Calculate the optimal tilt parameter
- > lambdav[which.min(stdevs)]
- > # Plot the standard deviations
- > x11(width=6, height=5)
- > plot(x=lambdav, y=stdevs,
- main="Standard Errors of Simulated VaR",
- xlab="tilt parameter", ylab="standard error",
- type="1", col="blue", lwd=2)

#### Importance Sampling of Brownian Motion

The statistics that depend on extreme paths of Brownian motion can be simulated more accurately using *importance sampling*.

The normally distributed variables  $x_i$  are shifted by the tilt parameter  $\lambda$  to obtain the importance sample variables  $x_i^{tilt}$ :  $x_i^{tilt} = x_i + \lambda$ .

The Brownian paths  $p_t$  are equal to the cumulative sums of the tilted variables  $x_t^{iilt}$ :  $p_t = \sum_{i=1}^t x_i^{tilt}$ .

Each tilted Brownian path has an associated weight factor equal to the product:  $\prod_{i=1}^{t} \exp(-x_i^{tilt}\lambda + \lambda^2/2)$ .

To compensate for the probability tilting, the statistics derived from the tilted Brownian paths must be multiplied by their weight factors.

```
> # Define Brownian motion parameters
> sigmav <- 1.0 # Volatility
> drift <- 0.0 # Drift
> nsteps <- 100 # Number of simulation steps
> nsimu <- 10000 # Number of simulation paths
> # Calculate matrix of normal variables
> set.seed(1121)
> datav <- rnorm(nsimu*nsteps, mean=drift, sd=sigmav)
> datav <- matrix(datav, nc=nsimu)
> # Simulate paths of Brownian motion
> pathm <- matrixStats::colCumsums(datav)
> # Tilt the datay
> lambda <- 0.04 # Tilt parameter
> datat <- datav + lambda # Tilt the random numbers
> patht <- matrixStats::colCumsums(datat)
> # Calculate path weights
> weightm <- exp(-lambda*datat + lambda^2/2)
> weightm <- matrixStats::colProds(weightm)
> # Nr
> weightm <- exp(-lambda*colSums(datat) + nsteps*lambda^2/2)
> # Calculate option payout using naive MC
> strikep <- 10 # Strike price
> payouts <- (pathm[nsteps, ] - strikep)
> sum(pavouts[pavouts > 0])/nsimu
> # Calculate option payout using importance sampling
> payouts <- (patht[nsteps, ] - strikep)
> sum((weightm*payouts)[payouts > 0])/nsimu
> # Calculate crossing probability using naive MC
> barl <- 10
> crossi <- (colSums(pathm > barl) > 0)
> sum(crossi)/nsimu
> # Calculate crossing probability using importance sampling
> crossi <- colSums(patht > barl) > 0
> sum(weightm*crossi)/nsimu
```

#### Homework Assignment

#### Required

- Study all the lecture slides in FRE6871\_Lecture2.pdf, and run all the code in FRE6871\_Lecture2.R,
- Study bootstrap simulation from the files bootstrap\_technique.pdf and doBootstrap\_primer.pdf,
- Study the Vasicek single factor model from Vasicek Portfolio Default Distribution.pdf,
- Study credit portfolio risk models from BOE Credit Risk Models.pdf and BIS Bank Capital Model.pdf,
- Study CDO models from Elizalde CDO Vasicek Credit Model.pdf,
- Study the CVAR credit portfolio risk measure from Danielsson CVAR Estimation Standard Error.pdf.

#### Recommended

Read about plotting from plot par cheatsheet.pdf and ggplot2 cheatsheet.pdf.
 You can download R Cheat Sheets here.