Control Statements, Operators and Debugging FRE6871 & FRE7241, Fall 2024

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September 20, 2024



Logical Operators

R has the following logical operators:

- "<" less than.</p>
- "<=" less than or equal to,</p>
- ">" greater than,
- ">=" greater than or equal to,
- "==" exactly equal to,
- "!=" not equal to,
- "!x" Not x.
- "x & y" x AND y,
- "x | y" x OR y,

These operators are applied to vectors element-wise.

```
> TRUE | FALSE
> TRUE | NA
> vec1 <- c(2, 4, 6)
> vec1 < 5 # Element-wise comparison
> (vec1 < 5) & (vec1 > 3)
> vec1[(vec1 < 5) & (vec1 > 3)]
> vec2 <- c(-10, 0, 10)
> vec1 < vec2
> c(FALSE, TRUE, FALSE) & c(TRUE, TRUE, FALSE)
> c(FALSE, TRUE, FALSE) | c(TRUE, TRUE, FALSE)
```

Long Form Logical Operators

R also has two long form logical operators:

- "x && y" x AND y,
- "x || y" x OR y,

These operators differ from the short form operators in two ways:

- They only evaluate single Boolean values, not Boolean vectors,
- They short-circuit (stop evaluation as soon as the expression is determined),

Rule of Thumb

Use "&&" and "||" in if-clauses,

```
> FALSE && TRUE

> FALSE || TRUE

> echo_true <- function() {cat("echo_true\t"); TRUE}

> echo_true <- function() {cat("echo_false\t"); FALSE}

> echo_true() || echo_false()

> echo_true() || echo_false()

> echo_true() || echo_false()

> echo_true() || echo_false()

> teno_true() || echo_false()

> teno_true() || echo_false()

> #Works (does nothing) using '&&'

> if (is.matrix(vecv) & (vecv[2, 3] > 0)) {

+ vecv[2, 3] <- 1

+ }

> #No short-circuit so fails (produces an error)

> if (is.matrix(vecv) & (vecv[2, 3] > 0)) {

+ vecv[2, 3] <- 1
```

+ }

Arithmetic Operators

Arithmetic *operators* perform arithmetic operations on numeric or complex vectors,

- "+" performs addition,
- "-" performs subtraction,
- "*" performs multiplication,
- "/" performs division,
- "^" and "**" perform exponentiation,

> ?Arithmetic

> 4.7 * 0.5 # Multiplication > 4.7 / 0.5 # Division

> 4.7 / 0.5 # Division

> # Exponentiation > 2**3

> 2**3

Comparing Objects With identical() and all.equal()

The function identical() tests if two objects are exactly the same, and always returns a single logical TRUE or FALSE (never NA or logical vectors).

For atomic arguments identical() often gives the same result as the "==" operator, but it's not synonymous with it in general.

The "==" operator applies the *recycling rule* to vector arguments and returns logical vectors, but identical() doesn't and returns a single logical value.

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

The variable .Machine contains information about the numerical characteristics of the computer R is running on, such as the largest double and integer numbers, and the *machine precision*.

```
> numv <- 2
> numv==2
> identical(numv, 2)
> identical(numv. NULL)
> # This doesn't work:
> # numv==NUI.I.
> is.null(numv)
> vecv <- c(2, 4, 6)
> vecv==2
> identical(vecv. 2)
> # numv is equal to "1.0" within machine precision
> numv <- 1.0 + 2*sqrt(.Machine$double.eps)
> all.equal(numv, 1.0)
> # Info machine precision of computer R is running on
> # ?.Machine
> # Machine precision
> .Machine$double.eps
```

Lookup and Matching Using which() and match()

The function which() returns the indices of the TRUE elements of a Boolean vector or array.

If the argument is an array and arr.ind=TRUE, then which() returns a matrix with rows containing the indices of the TRUE elements.

The functions which.max() and which.min() return the index of the minimum or maximum of a numeric or Boolean vector.

match() returns the index of the vector element that exactly matches its first argument.

If it doesn't find an exact match then it returns NA.

The expressions match(x, vecv) and min(which(vecv == x)) produce the same result, but match() can be faster for large vectors.

```
> vecv <- sample(1e3, 1e3)
> maty <- matrix(vecv. ncol=4)
> which(vecv == 5)
> match(5, vecv)
> # Equivalent but slower than above
> (1:NROW(vecv))[vecv == 5]
> which(vecv < 5)
> # Find indices of TRUE elements of Roolean matrix
> which((matv == 5)|(matv == 6), arr.ind=TRUE)
> # Equivalent but slower than above
> arravInd(which((matv == 5)|(matv == 6)).
     dim(matv), dimnames(matv))
> # Find index of largest element
> which.max(vecv)
> which(vecv == max(vecv))
> # Find index of smallest element
> which.min(vecv)
> # Benchmark match() versus which()
> all.equal(match(5, vecv), min(which(vecv == 5)))
> library(microbenchmark)
> summary(microbenchmark(
    match=match(5, vecv).
```

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

which=min(which(vecv == 5)),

Lookup and Matching Using %in% and any()

The binary operator %in% returns a Boolean vector with TRUE values corresponding to elements that have matches.

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

%in% never returns NA, so it's preferred in if() statements.

any() returns TRUE if at least one element of a Boolean vector is TRUE, and FALSE otherwise.

The function pmatch() performs partial matching of strings.

```
> # Does 5 belong in vecv?
> 5 %in% vecv
> match(5, vecv, nomatch=0) > 0
> # Does (-5) belong in vecv?
> (-5) %in% vecv
> c(5, -5) %in% vecv
> match(-5, vecv)
> # Equivalent to "5 %in% vecv"
> any(vecv == 5)
> # Equivalent to "(-5) %in% vecv"
> any(vecv == (-5))
> # Any negative values in vecv?
> any(vecv < 0)
> # Example of use in if() statement
> if (any(vecv < 2))
+ cat("vector contains small values\n")
> # Partial matching of strings
> pmatch("med", c("mean", "median", "mode"))
```

Finding Closest Match Using findInterval()

The function match() returns the index of the vector element that *exactly* matches its first argument.

If $\mathtt{match}(\tt)$ doesn't find an exact match then it returns \mathtt{NA}

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

If there's an exact match, then findInterval() returns the same index as function match().

If there's no exact match, then findInterval() finds the element of "vec" that is closest to, but not greater than, the element of "x".

- > # Display the formal arguments of findInterval
 > args(findInterval)
- > # Get index of the element of "vec" that matches 5
- > findInterval(x=5, vec=c(3, 5, 7))
- > match(5, c(3, 5, 7))
- > # No exact match
- > findInterval(x=6, vec=c(3, 5, 7))
- > match(6, c(3, 5, 7))
- > # Indices of "vec" that match elements of "x"
- > findInterval(x=1:8, vec=c(3, 5, 7))
- > # Return only indices of inside intervals
- > findInterval(x=1:8, vec=c(3, 5, 7), all.inside=TRUE)
- > # Make rightmost interval inclusive
- > findInterval(x=1:8, vec=c(3, 5, 7), rightmost.closed=TRUE)

Assignment Operators

The standard assignment operator in R is "<-".

Both "<-" and "=" are valid assignment operators in R.

The "<-" operator may cause an error if R confuses it with the "<" logical operator.

But they differ in *scope* and *precedence* ("<-" has higher precedence than "=").

The "=" operator is used for named arguments in function calls

When variables are assigned within an argument list using the "=" operator, their *scope* is limited to the function.

Rule of Thumb:

Use "<-" in R scripts and inside functions, Use "=" only in function calls.

```
> numv1 <- 3 # "<-" and "=" are valid assignment operators
> numv1
> numv1 = 3
> numv1
> 2<-3 # "<" operator confused with "<-"
> 2 <-3 # "<" operator confused with "<-"
> 2 <-3 # Add space or brackets to avoid confusion
> # "=" assignment within argument list
> median(x=1:10)
> x # x doesn't exist outside the function
> # "<-" assignment within argument list
> median(x <- 1:10)
> x # x exists outside the function
> x # x exists outside the function
> x # x exists outside the function
```

The assign() Function

The assign() function assigns a value to an object in a specified *environment*, by referencing it using a character string (name).

assign() can be used to either assign values to existing variables, or to create new variables.

assign() looks for the object name in the specified *environment*, and assigns a value to it.

If assign() can't find the object name, then it creates it.

assign() expects a character string as its argument.

If a object name is passed to assign(), then it evaluates that object to get the string it contains.

If the object doesn't contain a string, then assign() produces an error.

```
> mvvar <- 1 # Create new object
> assign(x="myvar", value=2) # Assign value to existing object
> myvar
> rm(myvar) # Remove myvar
> assign(x="myvar", value=3) # Create new object from name
> myvar
> # Create new object in new environment
> envy <- new.env() # Create new environment
> assign("myvar", 3, envir=envv) # Assign value to name
> ls(envv) # List objects in "envv"
> envv$myvar
> rm(list=ls()) # Delete all objects in workspace
> symboln <- "myvar" # Define symbol containing string "myvar"
> assign(symboln, 1) # Assign value to "myvar"
> 1s()
> myvar
> assign("symboln", "new_var")
> assign(symboln, 1) # Assign value to "new_var"
> 1s()
> symboln <- 10
```

> assign(symboln, 1) # Can't assign to non-string

Applying assign() to Lists of Names

assign() allows creating new objects from listv or vectors of names (character strings), such as column names.

```
> rm(list=ls()) # Delete all objects in workspace
> # Create individual vectors from column names of EuStockMarkets >
for (colname in colnames(EuStockMarkets)) {
    # Assign column values to column names
    assign(colname, EuStockMarkets[, colname])
    } # end for
> ls()
> head(DAX)
> head(EuStockMarkets[, "DAX"])
> identical(DAX, EuStockMarkets[, "DAX"])
```

Retrieving Objects Using get()

The function get() accepts a character string and returns the value of the corresponding object in a specified environment.

get() retrieves objects that are referenced using character strings, instead of their names.

The functions get() and assign() allow retrieving and assigning values to objects that are referenced using character strings.

The function mget() accepts a vector of strings and returns a list of the corresponding objects.

- > # Create new environment
- > envv <- new.env()
- > # Pass string as name to create new object
- > assign("myvar1", 2, envir=envv) > # Create new object using \$ string referencing
- > envv\$myvar2 <- 1
- > # List objects in new environment > ls(envv)
- > # Reference an object by name
- > envv\$myvar1
- > # Reference an object by string name using get
- > get("myvar1", envir=envv) > # Retrieve and assign value to object
- > assign("myvar1",
- 2*get("myvar1", envir=envv),
- envir=envv)
- > get("myvar1", envir=envv)
- > # Return all objects in an environment
- > mget(ls(envv), envir=envv) > # Delete environment
- > rm(envv)

> newv

Metaprogramming in R

A powerful feature of R is non-standard evaluation (aka metaprogramming or programming on the language).

Unevaluated expressions are objects that represent R

formulas and commands.

Metaprogramming allows creating and manipulating

Metaprogramming allows creating and manipulating unevaluated R expressions and then executing them as needed.

The book $Advanced\ R$ by Hadley Wickham provides a good explanation of metaprogramming in R and R expressions.

R interprets character strings that are not in quotes "" as symbols or expressions.

The function as.symbol() converts a character string into a *symbol* object.

The function parse() converts a character string into an unevaluated *expression* object.

The function eval() evaluates a *symbol* or *expression* in a specified *environment*.

```
> rm(list=ls()) # Delete all objects in workspace
> # Convert string to symbol
> as.symbol("some_string")
> # The "name" class is synonymous with a symbol
> class(as.symbol("some string"))
> # Symbols are created during assignments
> symboln <- 2
> # Evaluate symbol (same as typing it)
> eval(symboln)
> # Convert string into a symbol and evaluate it
> eval(as.svmbol("svmboln"))
> # Convert string into unevaluated expression
> expv <- parse(text="newv <- symboln")
> expv
> class(expv)
> 1s()
> eval(expv) # Evaluate expression
> ls() # Expression evaluation created new object
```

Manipulating Symbols and Expressions

The function quote() accepts symbols and expressions, and returns an expression object without evaluating it.

The function quote() creates unevaluated *expression* objects which later can be evaluated by functions.

The function substitute() replaces objects in unevaluated expressions with their corresponding values, and returns an *expression*.

substitute() looks up the object names in either named listv (symbol-value pairs) or in environments, and evaluates the objects in them.

substitute() is often used inside functions to substitute formal arguments with the names of the actual arguments they are bound to in a function call.

```
> # Create the expression "1+1"
> quote(1+1)
> # Evaluate the expression "1+1"
> eval(quote(1+1))
> # Create an expression containing several commands
> expv <- quote({x <- 1; y <- 2; x+y})
> expv
> # Evaluate all the commands in the expression
> eval(expv)
> 1s()
> # Return an expression without evaluating it
> newv <- 2*symboln
> expv <- quote(symboln + newv)
> expv
> eval(expv) # Evaluate expression
> # Substitute objects in an expression
> expv <- substitute(symboln + newv,
         env=list(symbol=1, newv=2))
> expv
> eval(expv) # Evaluate expression
> # Get_input() substitutes its formal argument with the actual arg
> get_input <- function(inputv) {
    substitute(inputy)
+ } # end get_input
> mvvar <- 2
> get input(mvvar)
> eval(get_input(myvar))
```

Converting Symbols and Expressions Into Strings

The function deparse() is the opposite of parse(), and it converts *symbols* and *expressions* into character strings.

The combination of functions deparse(substitute()) returns a character string representing the actual argument passed into a function.

- > # Define symbol > myvar <- 10
- > # Convert symbol value into string
- > deparse(myvar)
- > # Convert symbol into string without evaluating it > deparse(quote(myvar))
- > # Substitute object with value from named list
- > symboln <- 2
 > deparse(substitute(symboln + myvar, env=list(myvar=2)))
- > # Create string with name of input argument
- > get_name <- function(inputv) {
- + names(inputv) <- deparse(substitute(inputv))
- + inputv
- + } # end get_name
- > get_name(myvar)

The Parenthesis "()" and Curly Braces "{}" Operators

The parenthesis "()" and curly braces " $\{\}$ " operators are used to enclose and to group (combine) expressions.

The parenthesis "()" and curly braces " $\{\}$ " operators are functions, and they return values.

An expression enclosed by the parenthesis "()" operator is evaluated separatately from other expressions, and its result is returned.

Enclosing expressions in parenthesis makes them less ambiguous.

The curly braces "{}" operator can group several expressions, that can be written either on separate lines, or be separated by the semicolon ";" operator.

The curly braces "{}" operator returns the last expression it encloses.

Both the parenthesis "()" and curly braces "{}" operators are functions, and executing them requires a little additional processing time.

The square braces (brackets) " []" operator subsets (references) the elements of vectors, matrices, and listv.

```
> # Expressions enclosed in parenthesis are less ambiguous
> -2:5
> (-2):5
> -(2:5)
> # Expressions enclosed in parenthesis are less ambiguous
> -2*3+5
> -2*(3+5)
> # Expressions can be separated by semicolons or by lines
> {1+2; 2*3; 1:5}
> # or
> {1+2
+ 2*3
+ 1:5}
> matv <- matrix(nr=3, nc=4)
> matv <- 0
> # Subset whole matrix
> mat.v[] <- 0
> # Parenthesis and braces require a little additional processing t
> library(microbenchmark)
> summary(microbenchmark(
    basep=sqrt(rnorm(10000)^2),
   parven=sqrt(((((rnorm(10000)^2))))),
    bra ce=sqrt({{{ffrnorm(10000)^2}}}}),
    times=10))[, c(1, 4, 5)] # end microbenchmark summary
```

The "if () else" Control Statement

R has the familiar "if () $\{...\}$ else $\{...\}$ " statement to control execution flow depending on logical conditions.

The logical conditions must be either a Boolean or numeric type, otherwise an error is produced.

The "else" statement can also be omitted.

"if" statements can be nested using multiple "else if" statements.

```
> numv1 <- 1
>
    if (numv1) { # Numeric zero is FALSE, all other numbers are TRUE
+ numv2 <- 4
+ } else if (numv1 == 0) { # 'else if' together on same line
+ numv2 <- 0
+ } else { # 'else' together with curly braces
+ numv2 <- -4
+ } # end if
> numv2
```

> centerf(myvar, "mean")

> centerf(mvvar, "median")

> centerf(mvvar, "mean narm")

The switch() Control Statement

The function switch() matches its first argument "EXPR" with one of the symbols in the following arguments, evaluates the corresponding expression, and returns it.

The arguments that follow the first argument "EXPR" should be given as <code>symbol=value</code> pairs.

If "EXPR" is a character string, then the expression bound to that symbol is returned by switch().

If "EXPR" is an integer, then switch() returns the expression from that position.

If switch() can't match "EXPR" to any symbol, then it returns NULL invisibly.

Using switch() is a convenient alternative to a cascade of "if () else" statements.

The function match.arg() matches a string to one of the possible values, and returns the matched value, or produces an error if it can't match it.

```
> switch("a", a="aaahh", b="bee", c="see", d=2,
         "else this")
> switch("c", a="aaahh", b="bee", c="see", d=2,
         "else this")
> switch(3, a="aaahh", b="bee", c="see", d=2,
         "else this")
> switch("cc", a="aaahh", b="bee", c="see", d=2,
         "else this")
> # Measure of central tendency
> centerf <- function(inputy, method=c("mean", "mean narm", "median
+ # validate "method" argument
    method <- match.arg(method)
    switch(method.
+ mean=mean(inputy).
  mean_narm=mean(inputv, na.rm=TRUE),
+ median=median(inputy))
+ } # end centerf
> mvvar <- rnorm(100, mean=2)
```

Iteration Using for() and while() Loops

```
The for() loop statement:
                                                             > colorl <- list("red", "white", "blue")
                                                             > # Loop over list
> for (indeks in vecv) {expvs}
                                                             > for (colory in color)) {
iterates the dummy variable indeks over the elements
                                                                 print(colory)
                                                             + } # end for
of the vector or list color1, and evaluates in a loop the
                                                             > # Loop over vector
ex_pressions contained in the body of the for() loop.
                                                             > for (indeks in 1:3) {
                                                                 print(color1[[indeks]])
Upon loop exit the dummy variable indeks is left equal
                                                             + } # end for
to the last element of the vector colorl.
                                                             > # While loops require initialization
while() loops start by testing their logical condition,
                                                             > indeks <- 1
and they repeat executing the loop body until that
                                                             > # While loop
                                                             > while (indeks < 4) {
condition is FALSE
                                                                 print(color1[[indeks]])
But while() loops risk producing infinite loops if not
                                                                 indeks <- indeks + 1
                                                             + } # end while
written properly, so Use Them With Care!
```

Performing Loops Using for() and apply()

The for() loop doesn't return a value, so values calculated in the for() loop body must be assigned to variables in the parent environment, or otherwise they are lost.

The expressions in the for() loop body have access to variables in the parent environment in which the for() loop is executed, and they can modify those variables.

So even though for() loops don't return a value, they can be used to perform calculations on variables in the parent environment, but this is discouraged since it can produce errors that are hard to debug.

Rule of Thumb:

- for() loops are preferred for producing side effects, like plotting or reading and writing data to files,
- apply() loops are preferred for performing calculations which produce vectors or matrices of values.

```
> vecv <- integer(7)
> # Loop over a vector and overwrite it
> for (i in seq_along(vecv)) {
   cat("Changing element:", i, "\n")
   vecv[i] <- i^2
+ } # end for
> # Modifying vecy inside sapply() has no effect
> vecv <- integer(7)
> vecv
> sapply(seg along(yecy), function(i) {
+ vecv[i] <- i^2
+ }) # end sapply
> vecv
> # Super-assignment operator "<<-" allows modifying vecy
> sapply(seq_along(vecv), function(i) {
+ vecv[i] <<- i^2 # "<<-" !!!
+ }) # end sapply
> vecv
> # sapply() loop returns vector of values
> vecv <- sapply(seg along(vecv), function(i) (i^2))
```

> fib sea

Fibonacci Sequence Using for() Loop

The *Fibonacci* sequence of integers is defined by the recurrence relation:

$$F_n = F_{n-1} + F_{n-2},$$

 $F_1 = 0, F_2 = 1,$
 $F_n = 0, 1, 1, 2, 3, 5, 8, 13, \dots$

The Fibonacci sequence was invented by the Indian mathematician Virahanka in the 8th century AD, and later described by the Italian mathematician Fibonacci in his famous treatise Liber Abaci.

Very often variables are initialized to NULL before the start of iteration.

A more efficient way to perform iteration is by pre-allocating the vector.

The function numeric() returns an zero length numeric vector.

The function numeric(k) returns a numeric vector of zeros of length k.

```
> # fib_seq <- numeric()  # zero length numeric vector
> # Pre-allocate vector instead of "growing" it
> fib_seq <- numeric(10)
> fib_seq[1] <- 0  # Initialize
> for (i in 3:10) { # Perform recurrence loop
+ fib_seq[i] <- fib_seq[i-1] + fib_seq[i-2]
+ } # end for</pre>
```

Allocating Memory to Vectors and Matrices

 ${\tt R}$ automatically allocates memory to new objects as needed during runtime, but at the cost of slowing down calculations.

Allocating memory of the correct mode speeds up calculations by avoiding automatic memory allocation by R.

The functions character(), integer(), and numeric() return zero-length vectors of the specified *mode*.

Zero length vectors are not the same as NULL objects.

The function character(k) returns a character vector of empty strings of length k.

The function integer(k) returns a integer vector of zeros of length k.

The function numeric(k) returns a numeric vector of zeros of length k.

The function vector() by default returns a Boolean vector, unless the *mode* is specified.

The function matrix() by default returns a Boolean matrix containing NA values, unless the *mode* is specified.

```
> # Allocate character vector
> character()
> character(5)
> is.character(character(5))
> # Allocate integer vector
> integer()
> integer(5)
> is.integer(integer(5))
> is.numeric(integer(5))
> # Allocate numeric vector
> numeric()
> numeric(5)
> is.integer(numeric(5))
> is.numeric(numeric(5))
> # Allocate Boolean vector
> vector()
> vector(length=5)
> # Allocate numeric vector
> vector(length=5, mode="numeric")
> is.null(vector())
> # Allocate Boolean matrix
> matrix()
> is.null(matrix())
> # Allocate integer matrix
> matrix(NA_integer_, nrow=3, ncol=2)
> is.integer(matrix(NA integer , nrow=3, ncol=2))
```

> # Allocate numeric matrix

> matrix(NA real , nrow=3, ncol=2)

> is.numeric(matrix(NA real , nrow=3, ncol=2))

Logical Operators Applied to Vectors and Matrices

When logical operators are applied to vectors and matrices, they are applied element-wise, producing Boolean vectors and matrices.

Coercing Vectors Into Matrices

Vectors can be coerced into matrices by adding a dimension attribute.

The dimnames attribute can be assigned a named list to convert it into a named matrix.

The function structure() adds attributes (specified as symbol=value pairs) to an object, and returns it.

```
> matv <- 1:6  # Create a vector
> class(matv)  # Get its class
> # Is it vector or matrix?
> c(is.vector(matv), is.matrix(matv))
> structure(matv, dim=c(2, 3))  # Matrix object
> # Adding dimension attribute coerces into matrix
> dim(matv) <- c(2, 3)
> class(matv)  # Get its class
> # Is it vector or matrix?
> c(is.vector(matv), is.matrix(matv))
> # Assign dimnames attribute
```

> dimnames(matv) <- list(rows=c("row1", "row2"),
+ columns=c("col1", "col2", "col3"))</pre>

> matv

Coercing Matrices Into Other Types

Matrices can be explicitly coerced using the "as.*" coercion functions.

But coercion functions strip the attributes from an object.

```
> matv <- matrix(1:10, 2, 5) # Create matrix
> matv
```

- > # as.numeric strips dim attribute from matrix
- > as.numeric(matv)
- > # Explicitly coerce to "character" > matv <- as.character(matv)
- > c(typeof(matv), mode(matv), class(matv)) > # Coercion converted matrix to vector
- > c(is.matrix(matv), is.vector(matv))

Binding Vectors and Matrices Together

Vectors can be bound into matrices using the functions cbind() and rbind().

The *recycling rule* allows operations on vectors of different lengths:

- Vectors are scanned from left to right,
- Shorter vectors are extended in length by recycling their values until they match the length of longer vectors,

```
> vec1 <- 1:3 # Define vector

> vec2 <- 6:4 # Define vector

> # Bind vectors into columns

> cbind(vec1, vec2)

> # Bind vectors into rows

> rbind(vec1, vec2)

> # Extend to four elements

> vec2 <- c(vec2, 7)

> # Recycling rule applied

> cbind(vec1, vec2)

> # Another example of recycling rule

> 1:6 + c(10, 20)
```

Replicating Objects Using rep()

The function rep() replicates vectors and listv a given number of times.

rep() accepts a vector or list "x", and an integer specifying the type and number of replications.

Argument "times" replicates the whole vector a given number of times.

Argument "each" replicates each vector element a given number of times.

Argument "length.out" replicates the whole vector a certain number of times, so that the output vector length is equal to "length.out".

- > # Replicate a single element
- > rep("a", 5)
- > # Replicate the whole vector several times
- > rep(c("a", "b"), 5)
- > rep(c("a", "b"), times=5)
- > # Replicate the first element, then the second, etc. > rep(c("a", "b"), each=5)
- > # Replicate to specified length
- > rep(c("a", "b"), length.out=5)
- rep(c(a, b), length.out-b)

Multiplying Vectors and Matrices

The multiplication "*" operator performs element-wise (element-by-element) multiplication of vectors and matrices.

By default the matrix elements are multiplied column-wise by the vector elements: the first matrix element in the first column is multiplied by the first vector element, then the second matrix column is multiplied by the remaining vector elements, etc.

The *recycling rule* is applied to the vector elements as needed

The transpose function t() can be applied if we want to perform row-wise multiplication.

But the transpose function t() is very slow for large matrices.

A better choice is to use functions lapply() and do.call().

```
> # Define vector and matrix
> vec1 <- c(2, 4, 3)
> matv <- matrix(sample(1:12), ncol=3)
> # Multiply columns of matrix by vector
> vec1*matv
> # Or
> matv*vec1
> # Multiply rows of matrix by vector
> t(vec1*t(matv))
> # Multiply rows of matrix by vector - transpose is very slow
> matrixp <- lapply(1:NCOL(matv),
    function(x) vec1[x]*matv[, x])
> do.call(cbind, matrixp)
> library(microbenchmark)
> summary(microbenchmark(
    trans=t(vec1*t(matv)),
    lapp={
      matrixp <- lapply(1:NCOL(matv), function(x) vec1[x]*matv[, x]
      do.call(cbind, matrixp)
    times=10))[, c(1, 4, 5)] # end microbenchmark summary
```

Matrix Inner Multiplication

The %*% operator performs inner (scalar) multiplication of vectors and matrices

Inner multiplication multiplies the rows of one matrix with the columns of another matrix, so that each pair produces a single number:

$$C_{i,j} = \sum_{k=1}^n A_{i,k} B_{k,j}$$

Inner multiplication produces a vector or matrix with a reduced dimension

Inner multiplication requires the dimensions of the matrices to be conformable (number of columns in the first matrix must be equal to the number of rows in the second).

The function drop() removes any extra dimensions of length one.

The functions rowSums() and colSums() calculate the sums of rows and columns, and they're very fast because they pass their data to compiled C++ code.

- > vec2 <- 6:4 # Define vector
- > # Multiply two vectors element-by-element
- > vec1 * vec2
- > # Calculate inner product
- > vec1 %*% vec2
- > # Calculate inner product and drop dimensions > drop(vec1 %*% vec2)
- > # Multiply columns of matrix by vector
- > matv %*% vec1 # Single column matrix
- > drop(matv %*% vec1) # vector > rowSums(t(vec1 * t(matv)))
- > # using rowSums() and t() is 10 times slower than %*%
- > library(microbenchmark)
- > summary(microbenchmark(
- inner=drop(matv %*% vec1),
- transp=rowSums(t(vec1 * t(matv))),
- times=10))[, c(1, 4, 5)] # end microbenchmark summary

Matrix Transpose

The function t() returns the transpose of a matrix.

The function crossprod() also performs inner (scalar) multiplication, exactly the same as the %*% operator, but is slightly faster.

```
> # Multiply matrix by vector fails because dimensions aren't confo
> vec1 %*% matv
```

- > # Works after transpose
- > drop(vec1 %*% t(matv))
- > # Calculate inner product
- > crossprod(vec1, vec2)
- > # Create matrix and vector
- > matv <- matrix(1:3000, ncol=3)
- > tmatv <- t(matv) > vecv <- 1:3
- > # crossprod() is slightly faster than "%*%" operator
- > summary(microbenchmark(
- cross_prod=crossprod(tmatv, vecv),
- inner_prod=matv %*% vecv,
- times=10))[, c(1, 4, 5)] # end microbenchmark summary

Matrix Outer Multiplication

An *outer* product consists of all possible products of pairs of elements of two objects:

$$C_{i,j} = A_i \cdot B_j$$

An *outer* product of a function consists of applying it to all possible pairs of elements of two objects:

$$C_{i,j} = f(A_i, B_j)$$

Outer multiplication produces an object with dimension equal to the sum of the factors' dimensions, and with the number of elements equal to the product of the factors' elements.

The function outer() calculates the *outer* product of two matrices, and by default multiplies the elements of its arguments.

outer() can also calculate the values of a vectorized function of two variables passed to the "FUN" argument.

```
> # Define named vectors

> vec1 <- sample(1:4)

> vec1

> vec2 <- sample(1:3)

> names(vec2) <- paste0("col", 1:3, "=", vec2)

> vec2

> # Calculate outer product of two vectors

> matv <- outer(vec1, vec2)

> # Calculate vectorized function spanned over two vectors

> matv <- outer(vec1, vec2,

+ FUN=function(x1, x2) x2*sin(x1))

> matv
```

Flattening a List of Vectors to a Matrix Using do.call()

A list of vectors can be flattened into a matrix using the functions do.call() and either rbind() or cbind().

If the list contains vectors of different lengths, then R applies the recycling rule.

If the list contains a NULL element, that element is skipped.

```
> # Create list of vectors
> listv <- lapply(1:3, function(x) sample(6))
> # Bind list elements into matrix - doesn't work
> rbind(listv)
> # Bind list elements into matrix - tedious
> rbind(listv[[1]], listv[[2]], listv[[3]])
> # Bind list elements into matrix - works!
> do.call(rbind, listv)
> # Create numeric list
> listv <- list(1, 2, 3, 4)
> do.call(rbind, listv) # Returns single column matrix
> do.call(cbind, listv) # Returns single row matrix
> # Recycling rule applied
> do.call(cbind, list(1:2, 3:5))
> # NULL element is skipped
> do.call(cbind, list(1, NULL, 3, 4))
> # NA element isn't skipped
```

> do.call(cbind, list(1, NA, 3, 4))

Efficient Binding of Lists Into Matrices

A list of vectors can be flattened into a matrix using the functions do.call() and either rbind() or cbind().

But for large vectors this procedure can be very slow, and often causes an out of memory error.

The function do_call_rbind() efficiently combines a list of vectors into a matrix.

do_call_rbind() produces the same result as
do.call(rbind, list_var), but using recursion.

do_call_rbind() calls lapply in a loop, each time binding neighboring list elements and dividing the length of the list by half.

do_call_rbind() is the same function as
do.call.rbind() from package qmao:
 https://r-forge.r-project.org/R/?group.id=1113

```
> listv <- lapply(1:5, rnorm, n=10)
> maty <- do.call(rbind, listy)
> dim(mat.v)
> do call rbind <- function(listy) {
    while (NROW(listv) > 1) {
+ # Index of odd list elements
      odd index <- seg(from=1, to=NROW(listv), bv=2)
+ # Bind odd and even elements, and divide listy by half
      listy <- lapply(odd index, function(indeks) {
+ if (indeks==NROW(listy)) return(listy[[indeks]])
+ rbind(listv[[indeks]], listv[[indeks+1]])
      }) # end lapply
    } # end while
+ # listv has only one element - return it
    listv[[1]]
+ } # end do call rbind
> all.equal(matv. do call rbind(listv))
```

Filtering Data Frames Using subset()

Filtering means extracting rows from a data frame that satisfy a logical condition.

Data frames can be filtered using Boolean vectors and brackets "[]" operators.

The function subset() filters data frames, by applying logical conditions to its columns, using the column names.

subset() provides a succinct notation and discards NA
values, but it's slightly slower than using Boolean
vectors and brackets "[]" operators.

```
> airquality[(airquality$Solar.R > 320 & !is.na(airquality$Solar.R)
> subset(x=airquality, subset=(Solar.R > 320))
> summary(microbenchmark(
+ subset=subset(x=airquality, subset=(Solar.R > 320)),
+ brackets=airquality([airquality$Solar.R > 320 &
```

+ !is.na(airquality\$Solar.R)),], + times=10))[, c(1, 4, 5)] # end microbenchmark summary

> dim(splitiris\$setosa)
> head(splitiris\$setosa, 2)
> all.equal(setosa, splitiris\$setosa)

Splitting Data Frames Using factor Categorical Variables

The function split() divides an object into a list of objects, according to a factor (categorical variable).

The list's names attribute is equal to the factor levels.

```
> unique(iris$$pecies) # Species has three distinct values
> # $plit into separate data frames by hand
> setosa <- iris[iris$pecies=="setosa", ]
> versi <- iris[iris$pecies=="versicolor", ]
> virgin <- iris[iris$pecies=="virginica", ]
> dim(setosa)
> head(setosa, 2)
> # $plit iris into list based on Species
> splitiris <- split(iris, iris$Species)
> str(splitiris, max.confl=1)
> names(splitiris)
```

The split-apply-combine Procedure

The split-apply-combine procedure consists of:

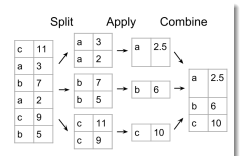
- dividing an object into a list, according to a factor (attribute).
- applying a function to each list element.
- o combining the results.

The *split-apply-combine* procedure is also called the *map-reduce* procedure, or simply *data pivoting*, and it's similar to *pivot tables* in *Excel*.

Data pivoting can be performed data frames, by aggregating its columns based on categorical data stored in one of its columns.

You can read more about the *split-apply-combine* procedure in Hadley Wickham's paper:

http://www.jstatsoft.org/v40/i01/paper



Data Pivoting Example

Data pivoting can be performed through successive applications of functions split(), apply(), and unlist().

A data frame can be pivoted either by first splitting it into a list of data frames and then aggregating, or by splitting just a single column and aggregating it.

The function split() divides an object into a list of objects, according to a factor (categorical variable).

The list's names attribute is equal to the factor levels.

The functional aggregate() pivots the columns of a data frame

aggregate() can accept a "formula" argument with the column names, or it can accept "x" and "by" arguments with the columns.

aggregate() returns a data frame containing the names of the groups (factor confls).

- > unique(mtcars\$cyl) # cyl has three unique values > # Split mpg column based on number of cylinders
- > split(mtcars\$mpg, mtcars\$cyl)
- > # Split mtcars data frame based on number of cylinders
- > split_cars <- split(mtcars, mtcars\$cyl) > str(split_cars, max.confl=1)
- > names(split cars)
- > # Aggregate the mean mpg over split mtcars data frame
- > sapply(split cars, function(x) mean(x\$mpg)) > # Or: split mpg column and aggregate the mean
- > sapply(split(mtcars\$mpg, mtcars\$cyl), mean) > # Same but using with()
- > with(mtcars, sapply(split(mpg, cyl), mean))
- > # Or: aggregate() using formula syntax > aggregate(x=(mpg ~ cyl), data=mtcars, FUN=mean)
- > # Or: aggregate() using data frame syntax
- > aggregate(x=mtcars\$mpg, by=list(cyl=mtcars\$cyl), FUN=mean)
- > # Or: using name for mpg
- > aggregate(x=list(mpg=mtcars\$mpg), bv=list(cvl=mtcars\$cvl), FUN=me > # Aggregate() all columns
- > aggregate(x=mtcars, by=list(cyl=mtcars\$cyl), FUN=mean)
- > # Aggregate multiple columns using formula syntax
- > aggregate(x=(cbind(mpg, hp) ~ cvl), data=mtcars, FUN=mean)

The tapply() Functional

The functional tapply() is a specialized version of the apply() functional, that applies a function to elements of a *jagged array*.

A *jagged array* is a list consisting of vectors or matrices of different lengths.

tapply() accepts a vector of values "X", a factor "INDEX", and a function "FUN".

tapply() first groups the elements of "X" according to the factor "INDEX", transforming it into a jagged array, and then applies "FUN" to each element of the jagged array.

tapply() applies a function to sub-vectors aggregated using a factor, and performs *data pivoting* in a single function call

The by() function is a wrapper for tapply().

The with() function evaluates an expression in an environment constructed from the data.

Data Pivoting Returning a Matrix

Sometimes data pivoting returns a list of vectors.

A list of vectors can be flattened into a matrix using the functions do.call() and either rbind() or cbind().

The function do.call() executes a function call using a function name and a list of arguments.

do.call() passes the list elements individually, instead of passing the whole list as one argument:

do.call(fun, list)= fun(list[[1]], list[[2]],
...)

- +) # end sapply > cardata # sapply() produces a matrix
- > # Now same using lapply
 > cardata <- lapply(split_cars, function(x) {</pre>
- + c(mean=mean(x\mpg), max=max(x\mpg), min=min(x\mpg)) +} # end anonymous function
- +) # end anonymous
- > is.list(cardata) # lapply produces a list
- > # do.call flattens list into a matrix
- > do.call(cbind, cardata)

Data Pivoting of Panel Data

The data frame panel_data contains fundamental financial data for S&P500 stocks.

The Industry column has 22 unique elements, while the Sector column has 10 unique elements. Each Industry belongs to a single Sector, but each Sector may have several Industries that belong to it.

The functional aggregate() allows aggregating over the Industry column, by performing *data pivoting*.

```
> # Download CRSPpanel.txt from the NYU share drive
> # Read the file using read.table() with header and sep arguments
> paneld <- read.table(file="/Users/jerzy/Develop/lecture_slides/da
                     header=TRUE, sep="\t")
> paneld[1:5, 1:5]
> attach(paneld)
> # Split paneld based on Industry column
> panelds <- split(paneld, Industry)
> # Number of companies in each Industry
> sapply(panelds, NROW)
> # Number of Sectors that each Industry belongs to
> sapply(panelds, function(x) {
    NROW(unique(x$Sector))
+ }) # end sapply
> # Nr
> aggregate(x=(Sector ~ Industry),
    data=paneld, FUN=function(x) NROW(unique(x)))
> # Industries and the Sector to which they belong
> aggregate(x=(Sector ~ Industry), data=paneld, FUN=unique)
> # Or
> aggregate(x=Sector, by=list(Industry), FUN=unique)
> sapply(unique(Industry), function(x) {
    Sector[match(x, Industry)]
+ }) # end sapply
```

> # Or use tapply() (returns an array)

> # Coerce secind2 into a jagged array

> secind2 <- drop(as.matrix(secind2))
> all.equal(secind2, secind)

Data Pivoting Returning a Jagged Array

A $jagged\ array$ is a list consisting of vectors or matrices of different lengths.

The functional aggregate() returns a data frame, so it's output must be coerced if the data pivoting attempts to return a jagged array.

The functional tapply() returns an array, so it's output must be coerced if the data pivoting attempts to return a jagged array.

tapply() accepts a vector of values "X", a factor "INDEX", and a function "FUN".

tapply() first groups the elements of "X" according to the factor "INDEX", transforming it into a jagged array, and then applies "FUN" to each element of the jagged array.

tapply() applies a function to sub-vectors aggregated using a factor, and performs *data pivoting* in a single function call

> # Split paneld based on Sector column > panelds <- split(paneld, Sector) > # Number of companies in each Sector > sapply(panelds, NROW) > # Industries belonging to each Sector (jagged array) > secind <- sapply(panelds, function(x) unique(x\$Industry)) > # Or use aggregate() (returns a data frame) > secind2 <- aggregate(x=(Industry ~ Sector), data=paneld, FUN=function(x) unique(x)) > # Or use aggregate() with "by" argument > secind2 <- aggregate(x=Industry, by=list(Sector), FUN=function(x) as.vector(unique(x))) > # Coerce secind2 into a jagged array > namev <- secind2[, 1] > secind2 <- secind2[, 2] > names(secind2) <- namev > all.equal(secind2, secind)

> secind2 <- tapply(X=Industry, INDEX=Sector, FUN=unique)

Data Pivoting Over Multiple Columns

Data pivoting over multiple columns can be performed by splitting the data frame and then performing an sapply() loop using an anonymous function.

Splitting the *data frame* allows aggregations over multiple columns.

An anonymous function allows applying different aggregations on the same column.

- > # Average ROE in each Industry
 > sapply(split(ROE, Industry), mean)
- > # Average, min, and max ROE in each Industry
- > t(sapply(split(ROE, Industry), FUN=function(x)
- + c(mean=mean(x), max=max(x), min=min(x))))
- > # Split paneld based on Industry column
- > panelds <- split(paneld, Industry)
- > # Average ROE and EPS in each Industry
- > t(sapply(panelds, FUN=function(x)
 - c(mean_roe=mean(x\$ROE),
 - mean_eps=mean(x\$EPS.EXCLUDE.EI))))
- > # Or: split paneld based on Industry column
- > panelds <- split(paneld[, c("ROE", "EPS.EXCLUDE.EI")],
- + paneld\$Industry)
- > # Average ROE and EPS in each Industry
- > t(sapply(panelds, FUN=function(x) sapply(x, mean)))
- > # Average ROE and EPS using aggregate()
- > aggregate(x=paneld[, c("ROE", "EPS.EXCLUDE.EI")],
- + by=list(paneld\$Industry), FUN=mean)

draft: Compare for() Loops With split-apply-combine Procedure

from: Wickham Split Apply Combine for Data Analysis

The split-apply-combine procedure allows for analyzing data frames, by spliting an analysis into smaller steps, and then combining them together.

in data preparation, for performing group-wise ranking. standardization, or normalization.

creating summaries

(aggregations) of data by groups. During modeling, for fitting

separate models to individual panels of data.

The split-apply-combine procedure is similar to the map-reduce procedure for processing large data.

also similar to pivot tables in Excel

The split(), apply() and combine Procedure

The lapply() and sapply() functions are specialized versions

```
> library(plyr)
> one <- ozone[1, 1, ]
> month <- ordered(rep(1:12, length72))
> model <- rlm(one ~ month - 1)
> deseas <- resid(model)</p>
> deseasf <- function(value) rlm(value ~ month - 1)
> # For loops
> models <- as.list(rep(NA, 24 * 24))
> dim(models) <- c(24, 24)</pre>
> deseas <- array(NA, c(24, 24, 72))
> dimnames(deseas) <- dimnames(ozone)</p>
> for (i in seq_len(24)) {
+ for (j in seq_len(24)) {
+ mod <- deseasf(ozone[i, j, ])
+ models[[i, j]] <- mod
+ deseas[i, j, ] <- resid(mod)
+ }
+ }
> # Apply functions
> models <- apply(ozone, 1:2, deseasf)
> resids list <- lapply(models, resid)
> resids <- unlist(resids list)
> dim(resids) <- c(72, 24, 24)
> deseas <- aperm(resids, c(2, 3, 1))</pre>
> dimnames(deseas) <- dimnames(ozone)
```

draft: Examples of split-apply-combine Procedure

 $from: \\ http://4dpiecharts.com/2011/12/$

quick-primer-on-split-apply-combine-problems/

we have a data frame with one column containing the values to calculate a statistic for and another column containing the group to which that value belongs

```
> # InsectSprays dataset
> head(InsectSprays)
> # Split the count column by the spray column.
> count_by_spray <- with(InsectSprays, split(count, spray))
>
> # Next apply the statistic to each element of the list. Lets use the mean here.
> mean_by_spray <- lapply(count_by_spray, mean)
> # Finally combine the list as a vector
> unlist(mean_by_spray)
> # or in one line
> sapply(count_by_spray, mean)
> # Stan also use the functions tapply(), aggregate() and by():
> with(InsectSprays, tapply(count, spray, mean))
> with(InsectSprays, by(count, spray, mean))
> aggregate(count * spray, InsectSprays, mean)
```

Exception Conditions: Errors and Warnings

Exception conditions are R objects containing information about *errors* or *warnings* produced while evaluating expressions.

The function warning() produces a *warning* condition, but doesn't halt function execution, and returns its message to the warning handler.

The function stop() produces an *error* condition, halts function execution, and returns its message to the error handler.

The handling of warning conditions depends on the value of options("warn"):

- negative then warnings are ignored,
- zero then warnings are stored and printed after the top-confl function has completed,
- one warnings are printed as they occur,
- two or larger warnings are turned into errors,

The function suppressWarnings() evaluates its expressions and ignores all warnings.

```
> # ?options # Get info on global options
> getOption("warn") # Global option for "warn"
> options("warn") # Global option for "warn"
> getOption("error") # Global option for "error"
> calc_sqrt <- function(inputv) {
+ # Returns its argument
   if (inputv < 0) {
     warning("calc_sqrt: input is negative")
     NULL # Return NULL for negative argument
   } else {
      sqrt(inputv)
   } # end if
+ } # end calc_sqrt
> calc_sqrt(5)
> calc_sqrt(-1)
> options(warn=-1)
> calc_sqrt(-1)
> options(warn=0)
> calc_sqrt()
> options(warn=1)
> calc_sqrt()
> options(warn=3)
> calc sgrt()
```

Validating Function Arguments

Argument validation consists of first determining if any arguments are *missing*, and then determining if the arguments are of the correct *type*.

An argument is *missing* when the formal argument is not bound to an actual value in the function call.

The function missing() returns TRUE if an argument is missing, and FALSE otherwise.

Missing arguments can be detected by:

- assigning a NULL default value to formal arguments and then calling is.null() on them,
- calling the function missing() on the arguments.

The argument *type* can be validated using functions such as is.numeric(), is.character(), etc.

The function return() returns its argument and terminates futher function execution.

- > # Function valido validates its arguments
 > valido <- function(inputv=NULL) {</pre>
- + # Check if argument is valid and return double
 + if (is.null(inputy)) {
- return("valido: input is missing")
 - } else if (is.numeric(inputv)) {
- + 2*inputv
 + } else cat("valido: input not numeric")
- + } # end valido > valido(3)
- > valido("a")
- > valido()
- > # valido validates arguments using missing()
- > valido <- function(inputv) {
- + # Check if argument is valid and return double
 + if (missing(inputy)) {
- + return("valido: input is missing")
- } else if (is.numeric(inputv)) {
- + 2*inputv
- + } else cat("valido: input is not numeric")
 + } # end valido
- > valido(3)
- > valido("a")
- > valido()

Validating Assertions Inside Functions

If assertions about variables inside a function are FALSE, then stop() can be called to halt its execution.

Calling stop() is preferable to calling return(), or inserting cat() statements into the code.

Using stop() inside a function allows calling the function traceback(), if an error was produced.

The function traceback() prints the call stack, showing the function that produced the *error* condition. cat() statements inside the function body provide information about the state of its variables.

```
> # valido() validates its arguments and assertions

> valido < function(inputv) {

+ # Check if argument is valid and return double

+ if (missing(inputv)) {

+ stop("valido: input is missing")

+ } else if (!is.numeric(inputv)) {

+ cat("input =", inputv, "\n")

+ stop("valido: input is not numeric")

+ } else 2*inputv
```

- + } # end valido > valido(3)
- > valido("a") > valido()
- > # Print the call stack
- > traceback()

> valido("a")

Validating Assertions Using stopifnot()

R provides robust validation and debugging tools through *type* validation functions, and functions missing(), stop(), and stopifnot().

If the argument to function stopifnot() is FALSE, then it produces an *error* condition, and halts function execution.

stopifnot() is a convenience wrapper for stop(), and eliminates the need to use if () statements.

stopifnot() is often used to check the validity of function arguments.

stopifnot() can be inserted anywhere in the function body in order to check assertions about its variables.

```
> valido <- function(inputv) {
    # Check argument using long form '&&' operator
    + stopifnot(!missing(inputv) && is.numeric(inputv))
    + 2*inputv
    } # end valido
    > valido(3)
    > valido("a")
    > valido <- function(inputv) {
    # Check argument using logical '&' operator
    + stopifnot(!missing(inputv) & is.numeric(inputv))
    + 2*inputv
    } # end valido
    > valido
    > valido
    > valido)
```

Validating Function Arguments and Assertions

The functions stop() and stopifnot() halt function execution and produce *error* conditions if certain assertions are FALSE.

The type validation functions, such as is.numeric(), is.na(), etc., and missing(), allow for validation of arguments and variables inside functions.

cat() statements can provide information about the state of variables inside a function.

cat() statements don't return values, so they provide information even when a function produces an error.

```
> # sumtwo() returns the sum of its two arguments
> sumtwo <- function(input1, input2) { # Even more robust
+ # Check if at least one argument is not missing
    stopifnot(!missing(input1) && !missing(input2))
+ # Check if arguments are valid and return sum
    if (is.numeric(input1) && is.numeric(input2)) {
      input1 + input2 # Both valid
    } else if (is.numeric(input1)) {
      cat("input2 is not numeric\n")
      input1 # input1 is valid
   } else if (is.numeric(input2)) {
      cat("input1 is not numeric\n")
      input2 # input2 is valid
    } else {
      stop("none of the arguments are numeric")
     # end sumtwo
> sumtwo(1, 2)
> sumtwo(5, 'a')
> sumtwo('a', 5)
> sumtwo('a', 'b')
> sumtwo()
```

The R Debugger Facility

The function debug() flags a function for future debugging, but doesn't invoke the debugger.

After a function is flagged for debugging with the call "debug(myfun)", then the function call "myfun()" automatically invokes the debugger (browser).

When the debugger is first invoked, it prints the function code to the console, and produces a *browser* prompt: "Browse [2]>".

Once inside the debugger, the user can execute the function code one command at a time by pressing the *Enter* key.

The user can examine the function arguments and variables with standard R commands, and can also change the values of objects or create new ones.

The command "c" executes the remainder of the function code without pausing.

The command "Q" exits the debugger (browser).

The call "undebug(myfun)" at the R prompt unflags the function for debugging.

- > # Flag "valido" for debugging
- > debug(valido)
- > # Calling "valido" starts debugger > valido(3)
- > # unflag "valido" for debugging
- > undebug(valido)

Debugging Using browser()

As an alternative to flagging a function for debugging, the user can insert the function browser() into the function body.

browser() pauses the execution of a function and invokes the debugger (browser) at the point where browser() was called.

Once inside the debugger, the user can execute all the same browser commands as when using debug().

browser() is usually inserted just before the command that is suspected of producing an *error* condition.

Another alternative to flagging a function for debugging, or inserting browser() calls, is setting the "error" option equal to "recover".

Setting the "error" option equal to "recover" automatically invokes the debugger when an *error* condition is produced.

> options(error=NULL) # Set back to default "error" option

Using the Debugger in RStudio

RStudio has several built-in debugging facilities that complement those already installed in R:

- toggling breakpoints, instead of inserting browser() commands,
- stepping into functions,
- environment pane with environment stack, instead of calling 1s(),
- traceback pane, instead of calling traceback(),

RStudio provides an online debugging tutorial: https://support.rstudio.com/hc/en-us/articles/ 205612627-Debugging-with-RStudio



Handling Exception Conditions

The function tryCatch() executes functions and expressions, and handles any exception conditions produced when they are evaluated.

 ${\tt tryCatch()} \ first \ evaluates \ its \ "expression" \ argument.$

If no error or warning condition is produced then tryCatch() just returns the value of the expression.

If an exception condition is produced then
tryCatch() invokes error and warning handlers and
executes other expressions to provide information about
the exception condition.

If a handler is provided to tryCatch() then the error is captured by the handler, instead of being broadcast to the console.

At the end, tryCatch() evaluates the expression provided to the finally argument.

```
> str(tryCatch) # Get arguments of tryCatch()
> tryCatch( # Without error handler
  { # Evaluate expressions
     numv <- 101 # Assign
     stop("mv error") # Produce error
   ٦.
   finally=print(paste0("numv = ", numv))
    # end trvCatch
> trvCatch( # With error handler
      # Evaluate expressions
     numv <- 101 # Assign
     stop("my error") # Produce error
   # Error handler captures error condition
   error=function(msg) {
     print(paste0("Error handler: ", msg))
   }, # end error handler
   # Warning handler captures warning condition
   warning=function(msg) {
     print(paste0("Warning handler: ", msg))
   }, # end warning handler
   finally=print(paste0("numy = ", numy))
+ ) # end tryCatch
```

end apply

Error Conditions in Loops

If an *error* occurs in an apply() loop, then the loop exits without returning any result.

apply() collects the values returned by the function supplied to its FUN argument, and returns them only after the loop is finished.

If one of the function calls produces an error, then the loop is interrupted and apply() exits without returning any result.

The function tryCatch() captures errors, allowing loops to continue after the error condition.

```
> # Apply loop without tryCatch
> apply(matrix(1:5), 1, function(numv) { # Anonymous function
+ stopintot((numv == 3)) # Check for error
+ # Broadcast message to console
+ cat("(cat) numv = ", numv, "\n")
+ # Return a value
+ pasteO("(return) numv = ", numv)
+ } # and anonymous function
}
```

+) # end apply

Exception Handling in Loops

If the body of the function supplied to the FUN argument is wrapped in tryCatch(), then the loop can finish without interruption and return its results.

The messages produced by *errors* and *warnings* can be caught by *handlers* (functions) that are supplied to tryCatch().

The *error* and *warning* messages are bound (passed) to the formal arguments of the *handler* functions that are supplied to tryCatch().

tryCatch() always evaluates the expression provided to the finally argument, even after an error occurs.