Bio 723: Statistical Computing for Biologists

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Modern biological data is...

- ► Heterogeneous sequence data, species distributions, protein concentrations, . . .
- ► High-dimensional genome-wide, GIS, ...
- ► Copious time series, population surveys, global monitoring stations, . . .

To be an effective analyst of biological data you must have. . .

- ▶ Biological knowledge and intuition What does the data mean? What are the key questions? What are interesting patterns or findings in the data?
- Computational skills sort, filter, transform, and transform data and work with in an efficient and reproducible manner
- Statistical skills build and interpret quantitative statements (models)
 about patterns in your data, distinguish interesting "signal" from "noise" of
 natural biological variability and experimental design

Course Goals

- 1. Learn to visualize and explore complex biological data using R, a programming language and statistical computing environment
- Introduce multivariate statistics from a geometric perspective, emphasizing the geometry of vector spaces.
- Illustrate how to carry out common scientific computing tasks like simulation and building bioinformatics pipelines
- Provide the tools and knowledge to conduct reproducible computational and statistical research.

Syllabus

- 1. R Basics, data munging, and visualization
- 2. Introduction to multivariate statistics from a geometric perspective
- 3. A survey of common machine learning methods clustering, classification, dimensionality reduction
- 4. Building bioinformatics pipelines

Course mechanics

- ► Instructor and TA contact information
- ▶ Texts
- Grading
- ► Expectations and Policies

Class structure

1. Lectures

- ► Typically 60-75 minutes
 - Emphasize the mathematical basis of the methods/approaches from both a geometric and algebraic basis
 - Discuss algorithms underlying the methods
- 2. Hands-on
- ► Walk through some examples
 - Apply the techniques and concepts to real data
 - Highlight available R libraries

Goals for today's class session

- ► Make sure you have a working R/RStudio environment
- Make sure you know how to evaluate commands in the R terminal
- Make sure you know how to create and knit an RMarkdown document
- Review R data types numerics (double, integer, complex), logicals, character
- ▶ Review R data structures vectors, lists, data frames



In-class demonstration of RStudio interface.

RMarkdown
In-class demonstration of creating and knitting an RMarkdown document.
in-class demonstration of creating and knitting an rimarkdown document.

Data types

- ▶ Refers to the types of values that can be represented in a computer program
- ▶ Determine the representation of values in memory
- ▶ Constrains the operations you can perform on those values

In R data types are usually inferred by the interpretter rather than specified by the user, though occassionally you may find it necessary to specify a data type.

Numeric data types in R

► Floating point values ('doubles') – represent real numbers (continuous values)

```
> x <- 10.0
> typeof(x)
[1] "double"
```

▶ Integers – represent whole numbers. The default numeric type is double, so you must explicitly ask for integers.

```
> x <- as.integer(10)
> typeof(x)
[1] "integer"
```

Complex numbers – numbers with a real and "imaginary" part. We won't be explicitly using complex numbers in this course, but they sometimes appear unexpectedly in some calculations.

```
> x <- 1+1i
> typeof(x)
[1] "complex"
```

Arithmetic operations on numerical data types

```
> 10 + 2 \# addition
[1] 12
> 10 - 2 # subtraction
Γ17 8
> 10 * 2 # multiplication
Γ1 20
> 10 / 2 # division
Γ17 5
> 10 ^ 2 # exponentiation
Γ1] 100
> 10 ** 2 # alternate exponentiation
Γ17 100
> 10 %% 4 # modulus (remainder after division)
[1] 2
> 4-5 / 2 # operator precedence matters!
[1] 1.5
> (4-5)/2 # parentheses help you specify/disambiguate precedence.
[1] -0.5
```

Basic mathematical functions

```
> sqrt(10) # square root
[1] 3.162278
> 10 ^ 0.5 # same as square root
[1] 3.162278
> sqrt(-1) # NaN means "Not a Number"
[1] NaN
> sqrt(-1 + 0i) # But works if we use complex type
[1] 0+1i
> exp(1) # exponential function
[1] 2.718282
> log(100) # log base e
[1] 4.60517
> log10(100) # log base 10
Γ17 2
> log2(8) # log base 2
Γ17 3
> factorial(5) # factorial function: 5 * 4 * 3 * 3 * 1
Γ1] 120
> pi # R knows some useful constants
[1] 3.141593
> cos(2*pi) # cosine, also sin, tan, ...
Γ1 1
```

```
Logical (Boolean) type

> x <- TRUE
> typeof(x)

[1] "logical"
> y <- FALSE
> typeof(FALSE)

[1] "logical"
> !TRUE # logical negation of TRUE

[1] FALSE
> !y # logical negation (NOT) of value in y

[1] TRUE
```

Numerical comparison operators return logical types:

```
> 4 < 5  # less than
[1] TRUE
> 10 >= 9  # greater than or equal to
[1] TRUE
```

Comparison operations can be assigned to variables:

```
> x <- 1 > 2
> y <- 2 >= 2
> x & y # Logical AND
[1] FALSE
> x | y # Logical OR
[1] TRUE
> isTRUE(x) # returns true if x is logical and true
[1] FALSE
```

Character (string) type

```
> x <- "Hello" # enclosed in double quotes
> y <- 'World' # or single quotes
> z <- 'You said "Hello World"  # allows nesting</pre>
```

Simple character functions:

```
> paste(x, y) # concatenate strings
[1] "Hello World"
> paste(x, y, sep = "") # concatenate with no space
[1] "HelloWorld"
> strsplit("Hello world!", split=" ") # split on space
[[1]]
[1] "Hello" "world!"
```

The stringr package contains many useful string manipulation functions.

Data structures

- Represent different ways collections of data are stored in memory or accessed by the user
- ► Homogenous or heterogeneous with respect to data types
- Different data structures are more efficient for particular modes of access or to represent different types operations

Vectors

Homogenous ordered list of items

```
> x <- c(2, 4, 6, 8)
> y <- c("hello", "world", "how", "are", "you?")</pre>
```

Vectors always have a length (possibly zero) and a type

```
> length(x)
[1] 4
> typeof(x)
[1] "double"
> length(y)
[1] 5
> typeof(y)
[1] "character"
```

Indexing vectors

Accessing the objects in a vector is accomplished by "indexing".

The elements of the vector are assigned indices $1 \dots n$ where n is the length of the vector

```
> x <- c(2,4,6,8)
> length(x)
[1] 4
> x[1]
[1] 2
> x[2]
[1] 4
> x[4]
[1] 8
> x[4]
[1] 8
> x[length(x)] # why might this be preferred way to get last value?
[1] 8
```

Indexing past the end of a vector returns NA (Not Available) values:

```
> x[5]
[1] NA
```

Single objects of core data types in $\ensuremath{\mathsf{R}}$ are themselves vectors

```
> x <- 1
> length(x)
[1] 1
> x[1] # can be indexed like any other vectors
[1] 1
> is.vector(x) # function to test whether something is vector
[1] TRUE
```

Elements of a vector will be coerced to be the same type

```
> x <- c(1+1i, 2+1i, 'Fred', 10)
> x
[1] "1+1i" "2+1i" "Fred" "10"
> typeof(x)
[1] "character"
```

Arithmetic operators and most math functions work on numerical vectors

Arithmetic operators and comparison work element-by-element on vectors.

```
> x <- c(2, 4, 6, 8)
> y <- c(0, 1, 2, 10)
> x + y
[1] 2 5 8 18
> x * y
[1] 0 4 12 80
> x<sup>2</sup>
[1] 4 16 36 64
> sqrt(x)
[1] 1.414214 2.000000 2.449490 2.828427
> x < y
[1] FALSE FALSE TRUE</pre>
```

Vectors can be indexed by other vectors, including logical vectors

```
> x <- c(2, 4, 6, 8)
> x[c(2,4)] # get 2nd and 4th elements of x
[1] 4 8

> x <- c(2, 4, 6, 8)
> y <- c(0, 1, 2, 10)
> x[x > y] # get elements of x where x > y
[1] 2 4 6
```

Lists

Lists in R are like vectors but the elements of a list are arbitrary objects (even other lists). Lists are "heteregeneous".

```
> x <- list('Bob',27, 10, c(720,710))
> typeof(x)
[1] "list"
> x
[[1]]
[1] "Bob"
[[2]]
Γ17 27
[[3]]
[1] 10
[[4]]
[1] 720 710
```

Indexing Lists

Items in lists are accessed in a different manner than vectors.

- ► Typically you use double brackets ([[]])to return the element at index i
- ▶ Single brackets always return a list containing the element at index i

```
> x <- list('Bob', 27, 10, c(720,710))
> x[1]
[[1]]
[1] "Bob"
> typeof(x[1])
[1] "list"
> x[[1]]
[1] "Bob"
> typeof(x[[1]])
[1] "character"
```

List elements can have names

```
> x <- list(name='Bob',age=27, years.in.school=10)
> x
$name
[1] "Bob"

$age
[1] 27

$years.in.school
[1] 10
```

Named list objects can be accessed via the \$ operator

```
> x$years.in.school
[1] 10
```

The names of list elements can be accessed with the names() function

```
> names(x)
[1] "name" "age" "years.in.school"
```

Data frames

Data frames represent data tables (data sets)

- ▶ Each column in the table has the same number of rows
- Every item in a given column has to be of the same type.
- ► Columns in a data frame must have names

Data frames: shape and column indexing

Shape

```
> nrow(example.df) # number of rows
[1] 3
> ncol(example.df) # number of columns
[1] 3
```

Indexing columns by name or position

Indexing with a vector to subset columns

```
> example.df[c("name","grade")]
   name grade
1 Paul C
2 Mary A
3 Peter B
```

Data frames: indexing rows

To index one or more rows of a data frame, specifying the value(s) to index on followed by a comma:

You can simultaneously index both rows and columns:

```
> example.df[c(1,3), c("name", "age")]
    name age
1  Paul 40
3  Peter 28
```

Matrices

In R matrices are two-dimensional collections of elements all of which have the same mode or type (homogenous).

Matrices don't by default have explicit row or column names, but they can be assigned/retrieved using the rownames() and colnames() functions respectively.

We will typically prefer data frames to matrices, because data frames are more flexible. But we'll return to matrices when we get into linear algebra later in the semester.