

The Bioconductor Project

INTRODUCTION TO BIOCONDUCTOR IN R



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Bioconductor



¹ Bioconductor (www.bioconductor.org)

What do we measure and why?

- **Structure:** elements, regions, size, order, relationships



- **Function:** expression, levels, regulation, phenotypes



How to install Bioconductor packages?

- Bioconductor has its own repository, way to install packages, and each release is designed to work with a specific version of R.
 - For this course, you'll be using Bioconductor version 3.6.
- Bioconductor version 3.7 or earlier uses **BiocLite**:

```
source("https://bioconductor.org/biocLite.R")
biocLite("packageName")
```

- Bioconductor version 3.8 and later uses **BiocManager**:

```
if (!requireNamespace("BiocManager"))
  install.packages("BiocManager")
BiocManager::install()
```

Bioconductor version and package version

- BiocInstaller works for Bioconductor version 3.7 or earlier

```
# Check Bioconductor version (For versions <= 3.7)
BiocInstaller::biocVersion()

# or

biocVersion()
# Load a package
library(packageName)
# Check versions for reproducibility
sessionInfo()
# or
packageVersion("packageName")
# Check package updates (Bioconductor version <= 3.7)
BiocInstaller::biocValid()

# or

biocValid()
```

Let's practice!

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The Role of S4 in Bioconductor

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S3

Positive

- CRAN, simple but powerful
- Flexible and interactive
- Uses a generic function
- Functionality depends on the first argument
- Example: `plot()` and `methods(plot)`

Negative

- Bad at validating types and naming conventions (dot not dot?)
- Inheritance works, but depends on the input

S4

Positive

- Formal definition of classes
- Bioconductor reusability
- Has validation of types
- Naming conventions

Example: `mydescriptor <- new("GenomeDescription")`

Negative

- Complex structure compared to S3

Is it S4 or not?

Ask if an object is S4

```
isS4(mydescriptor)
```

```
TRUE
```

`str` of S4 objects start with `Formal class`

```
str(mydescriptor)
```

```
Formal class 'GenomeDescription' [package "GenomeInfoDb"] with 7 slots
```

```
...
```

S4 class definition

A class describes a representation

- **name**
- **slots** (methods/fields)
- **contains** (inheritance definition)

```
MyEpicProject <- setClass(# Define class name with UpperCamelCase  
                         "MyEpicProject",  
                         # Define slots, helpful for validation  
                         slots = c(ini = "Date",  
                                   end = "Date",  
                                   milestone = "character"),  
                         # Define inheritance  
                         contains = "MyProject")
```

```
.S4methods(class = "GenomeDescription")
```

```
[1] commonName      organism       provider      providerVersion   releaseDate    releaseName     seqinfo  
[8] seqnames       show        toString      bsgenomeName
```

```
showMethods(classes = "GenomeDescription", where = search())
```

Object summary

```
show(myDescriptor)
```

```
| organism: ()  
| provider:  
| provider version:  
| release date:  
| release name:  
| ---  
| seqlengths:
```

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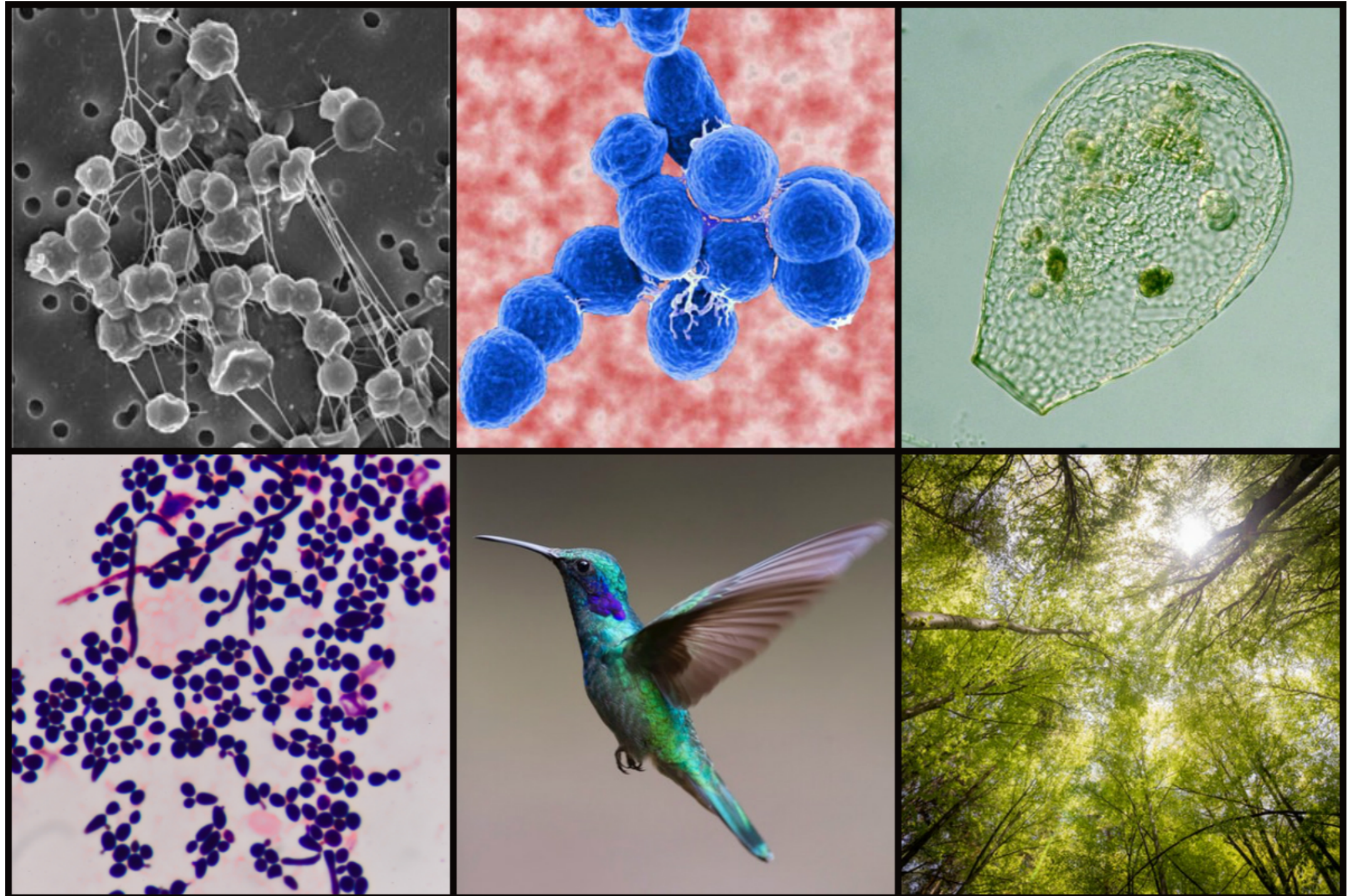
Introducing biology of genomic datasets

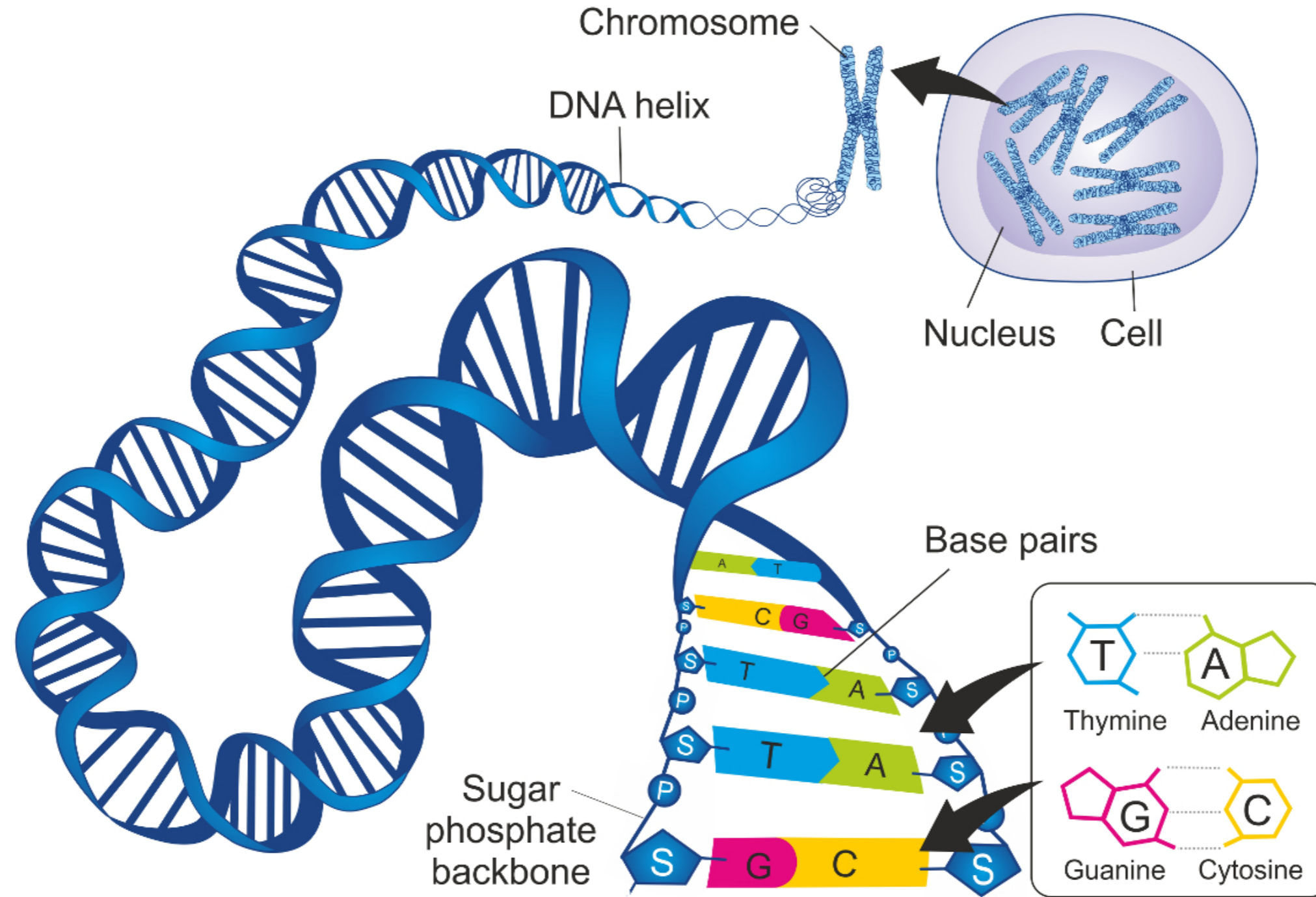
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Genome elements

- Genetic information DNA alphabet
- A set of chromosomes (highly variable number)
- Genes (carry heredity instructions)
 - coding and non-coding
- Proteins (responsible for specific functions)
 - DNA-to-RNA (transcription)
 - RNA-to-protein (translation)

Yeast

- A single cell microorganism
- The fungus that people love ❤
- Used for fermentation: beer, bread, kefir, kombucha, bioremediation, etc.
- Name: *Saccharomyces cerevisiae* or *S. cerevisiae*



BSgenome annotation package

```
# Load the package and store data into yeast
library(BSgenome.Scerevisiae.UCSC.sacCer3)
yeast <- BSgenome.Scerevisiae.UCSC.sacCer3
#interested in other genomes?
available.genomes()
```

Using accessors

```
# Chromosome number
length(yeast)
# Chromosome names
names(yeast)
# Sequence lengths
seqLengths(yeast)
```

Get sequences

S4 method for BSgenome

```
# S4 method getSeq() requires a BSgenome object
getSeq(yeast)
# Select chromosome sequence by name, one or many
getSeq(yeast, "chrM")
# Select start, end and/or width
# end = 10, selects first 10 base pairs of each chromosome
getSeq(yeast, end = 10)
```

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