## Writing reproducible reports

#### knitr with R Markdown

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► To estimate power?

- ► To estimate power?
- ► To estimate a p-value?

- ▶ To estimate power?
- ► To estimate a p-value?
- ► To estimate some other quantity?

# Data analysis reports

- ► Figures/tables + email
- ► Static Word document

 $\blacktriangleright \text{ LATEX} + \text{R} \rightarrow \text{PDF}$ 

ightharpoonup R Markdown = knitr + Markdown ightarrow Web page

What if the data change?

What if you used the wrong version of the data?

#### knitr in a knutshell

kbroman.org/knitr\_knutshell

#### knitr in a knutshell

kbroman.org/knitr\_knutshell

rmarkdown.rstudio.com

#### knitr code chunks

#### Input to knitr:

```
We see that this is an intercross with `r nind(sug)`
individuals. There are `r nphe(sug)` phenotypes, and genotype
data at `r totmar(sug)` markers across the `r nchr(sug)`
autosomes. The genotype data is quite complete.

```{r summary_plot, fig.height=8}
plot(sug)
```
```

#### Output from knitr:

```
We see that this is an intercross with 163 individuals. There are 6 phenotypes, and genotype data at 93 markers across the 19 autosomes. The genotype data is quite complete.

""r
plot(sug)
""![plot of chunk summary_plot](RmdFigs/summary_plot.png)
```

#### html

```
<!DOCTYPE html>
<html>
<head>
 <meta charset=utf-8"/>
 <title>Example html file</title>
</head>
<body>
<h1>Markdown example</h1>
Use a bit of <strong>bold</strong> or <em>italics</em>. Use
backticks to indicate <code > code > code > that will be rendered
in monospace.
<111>
This is part of a list
another item
</body>
</html>
```

### **CSS**

```
ul,ol {
  margin: 0 0 0 35px;
a {
  color: purple;
  text-decoration: none;
  background-color: transparent;
a:hover
  color: purple;
  background: #CAFFFF;
```

#### Markdown

```
# Markdown example
Use a bit of **bold** or italics . Use backticks to indicate
`code` that will be rendered in monospace.
- This is part of a list
- another item
Include blocks of code using three backticks:
. . .
x <- rnorm(100)
Or indent four spaces:
    mean(x)
    sd(x)
And it's easy to create links, like to
[Markdown] (http://daringfireball.net/projects/markdown/).
```

#### R Markdown

- ▶ R Markdown is a variant of Markdown, developed at RStudio.com
- ► Markdown + knitr + extras
- ► A few extra marks
- ► LATEX equations
- ► Bundle images into the final html file

#### YAML header

```
---
title: "knitr/R Markdown example"
author: "Karl Broman"
date: "28 January 2015"
output: html_document
---
```

```
title: "Another knitr/R Markdown example"
author: "[Karl Broman](https://kbroman.org)"
date: "`r Sys.Date()`"
output: word_document
---
```

?rmarkdown::html\_document

### ?rmarkdown::html\_document

- ► toc\_float
- ► toc\_depth
- ► code\_folding
- ► theme
- ► df\_print

## Code chunks, again

```
```{r knitr options, include=FALSE}
knitr::opts_chunk$set(fig.width=12, fig.height=4,
                      fig.path='Figs/', warning=FALSE,
                      message=FALSE)
set.seed(53079239)
### Preliminaries
Load the R/qtl package using the `library` function:
'``{r load gtl}
library(qtl)
To get help on the read.cross function in R, type the
following:
```{r help, eval=FALSE}
?read.cross
. . .
```

# Chunk options

echo=FALSE Don't include the code

results="hide" Don't include the output

include=FALSE Don't show code or output

eval=FALSE Don't evaluate the code at all

warning=FALSE Don't show R warnings

message=FALSE Don't show R messages

fig.width=# Width of figure

fig.height=# Height of figure

fig.path="Figs/" Path for figure files

There are lots of chunk options.

# Global chunk options

```
```{r knitr_options, include=FALSE}
knitr::opts_chunk$set(fig.width=12, fig.height=4,
                      fig.path='Figs/', warning=FALSE,
                      message=FALSE, include=FALSE,
                      echo=FALSE)
set.seed(53079239)
```{r make_plot, fig.width=8, include=TRUE}
x < - rnorm(100)
y < -2*x + rnorm(100)
plot(x, y)
```

- Use global chunk options rather than repeat the same options over and over.
- ► You can override the global values in specific chunks.

# Package options

```
'``{r package_options, include=FALSE}
knitr::opts_knit$set(progress = TRUE, verbose = TRUE)
```

- ▶ It's easy to confuse global chunk options with package options.
- ► I've not used package options.
- ► So focus on opts\_chunk\$set() not opts\_knit\$set().

#### In-line code

```
We see that this is an intercross with `r nind(sug)` individuals. There are `r nphe(sug)` phenotypes, and genotype data at `r totmar(sug)` markers across the `r nchr(sug)` autosomes. The genotype data is quite complete.
```

- ► Each bit of in-line code needs to be within one line; they can't span across lines.
- ► I'll often precede a paragraph with a code chunk with include=FALSE, defining various variables, to simplify the in-line code.
- ▶ Never hard-code a result or summary statistic again!

# Python in R Markdown

You can have python code chunks in R Markdown. And information is remembered between chunks.

```
Time in the image is a second problem of the image is a secon
```

It seems like you can't use python in-line. But if load the package 'reticulate', you can get access to python objects with R code.

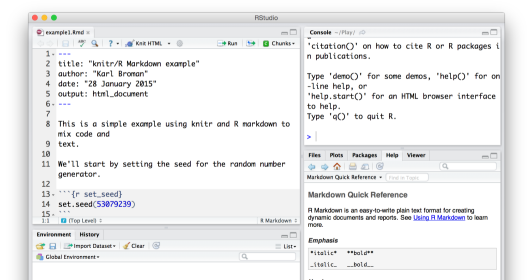
```
The first value in `x` is `r py$x[1]`, while the first value in `y` is `r py$y[1]`.
```

More at rstudio.github.io/reticulate/

## Rounding

- ightharpoonup cor(x,y) might produce 0.8992877, but I want 0.90.
- round(cor(x,y), 2), would give 0.9, but I want 0.90.
- You could use sprintf("%.2f", cor(x,y)), but sprintf("%.2f", -0.001) gives -0.00.
- ▶ Use the myround function in my R/broman package.
- ▶ myround(cor(x,y), 2) solves both issues.

### R Markdown → html, in RStudio



### R Markdown $\rightarrow$ html, in R

```
> library(rmarkdown)
> render("knitr_example.Rmd")
```

```
> rmarkdown::render("knitr_example.Rmd")
```

### R Markdown → html, GNU make

```
knitr_example.html: knitr_example.Rmd
R -e "rmarkdown::render('knitr_example.Rmd')"
```

### Need pandoc in your PATH

RStudio includes pandoc; you just need to add the relevant directory to your PATH.

#### Mac:

 $/{\tt Applications/RStudio.app/Contents/MacOS/pandoc}$ 

#### Windows:

"c:\Program Files\RStudio\bin\pandoc"

## Reproducible knitr documents

- ► Don't use absolute paths like ~/Data/blah.csv
- ► Keep all of the code and data in one directory (and its subdirectories)
- ► If you must use absolute paths, define the various directories with variables at the top of your document.
- ► Use R --vanilla or perhaps

```
R --no-save --no-restore --no-init-file --no-site-file
```

- ▶ Use GNU make to document the construction of the final product (tell future users what to do)
- ▶ Include a final chunk with getwd() and devtools::session\_info().
- ► For simulations, use set.seed in your first chunk.

# Controlling figures

```
```{r test_figure, dev.args=list(pointsize=18)}
x <- rnorm(100)
y <- 2*x + rnorm(100)
plot(x,y)
```</pre>
```

- ▶ The default is for knitr/R Markdown is to use the png() graphics device.
- ▶ Use another graphics device with the chunk option dev.
- ▶ Pass arguments to the graphics device via the chunk option dev.args.

#### **Tables**

```
'``{r kable}
x <- rnorm(100)
y <- 2*x + rnorm(100)
out <- lm(y ~ x)
coef_tab <- summary(out)$coef
library(knitr)
kable(coef_tab, digits=2)
'``</pre>
```

```
'``{r xtable, results="asis"}
library(xtable)
tab <- xtable(coef_tab, digits=c(0, 2, 2, 1, 3))
print(tab, type="html")
'``</pre>
```

```
library(gt)
gt( round(coef_tab, 2) )
```

## Important principles

Modify your desires to match the defaults.

Focus your compulsive behavior on things that matter.

What should a report contain?

## What should a report contain?

Karl -- this is very interesting, however you used an old version of the data (n=143 rather than n=226).

I'm really sorry you did all that work on the incomplete dataset.

Bruce

## What should a report contain?

- ► Explain your shared goals
- Describe the data
- Explain what you did
- ► Show your results
- ► Explain your conclusions
- ► When you're done, go back and write an *executive summary*

### Standard scientific article

- ► Abstract
- ► Introduction/background
- ► Materials and methods
- ► Results
- ► Conclusions/discussion

Why this format?

## Further suggestions

- ► Tailor the report to the audience
- Try not to be boring
- Limit equations and code; details in an appendix
- ► Break it up into sections; simple and clear language and structure
- Lots of figures, ideally interactive; explain the figures
- ▶ What do + and mean (regarding coefficients/effects)?

# Organizing projects

- ► RStudio Projects
- ▶ here package for R

## Other R Markdown-based things

- ▶ blogdown for websites
- bookdown for book-like objects
- xaringan for slides
- pagedown for paged documents (like resumes or letters)

# Interactive graphics tools

- ► plotly
- ▶ htmlwidgets
- ▶ leaflet
- ► networkD3
- ▶ DiagrammeR
- ▶ DT
- ▶ d3heatmap
- ► scatterD3