

Steps toward reproducible research

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Slides: bit.ly/jsm2016



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

The results in Table 1 don't seem to correspond to those in Figure 2.

In what order do I run these scripts?

Where did we get this data file?

Why did I omit those samples?

How did I make that figure?

“Your script is now giving an error.”

“The attached is similar to the code we used.”

Reproducible

vs.

Replicable

Reproducible

vs.

Correct

Steps toward reproducible research

kbroman.org/steps2rr

1. Organize your data & code

File organization and naming
are powerful weapons against chaos.

– Jenny Bryan

1. Organize your data & code

Your closest collaborator is you six months ago,
but you don't reply to emails.

(paraphrasing [Mark Holder](#))

1. Organize your data & code

```
RawData/           Notes/  
DerivedData/      Refs/  
  
Python/           ReadMe.txt  
R/                ToDo.txt  
Ruby/             Makefile
```

Chaos

```
AimeeNullSims/      Deuterium/          Ping/
AimeeResults/       ExtractData4Gary/   Ping2/
AnnotationFiles/    FromAimee/          Ping3/
Brian/              GoldStandard/       Ping4/
Chr6_extrageno/     HumanGWAS/          Play/
Chr6_segdis/        Insulin/            Prdm9/
ChrisPlaisier/      Int2_for_Mark/      RBM_PlasmaUrine_2012-03-08/
Code4Aimee/         Islet_2011-05/     Slco1a6/
CompAnnot/          MappingProbes/      StudyLineupMethods/
CondScans/          MultiProbes/        kidney_chr6.R
D20_2012-02-14/    NewMap/             pck2_sucla2.R
D20_cellcycle/     Notes/              penalties.txt
D20corr/           NullSims/           transeQTL4Lude/
Data4Aimee/         NullSims_2009-09-10/
Data4Tram/          PepIns_2012-02-09/
```


2. Everything with a script

If you do something once,
you'll do it 1000 times.

3. Automate the process (GNU Make)

```
R/analysis.html: R/analysis.Rmd Data/cleandata.csv
  cd R;R -e "rmarkdown::render('analysis.Rmd')"
```

```
Data/cleandata.csv: R/prepData.R RawData/rawdata.csv
  cd R;R CMD BATCH prepData.R
```

```
RawData/rawdata.csv: Python/xls2csv.py RawData/rawdata.xls
  Python/xls2csv.py RawData/rawdata.xls > RawData/rawdata.csv
```

3. Automate the process (GNU Make)

```
R/analysis.html: R/analysis.Rmd Data/cleandata.csv
  cd R;R -e "rmarkdown::render('analysis.Rmd')"
```

```
Data/cleandata.csv: R/prepData.R RawData/rawdata.csv
  cd R;R CMD BATCH prepData.R
```

```
RawData/rawdata.csv: Python/xls2csv.py RawData/rawdata.xls
  Python/xls2csv.py RawData/rawdata.xls > RawData/rawdata.csv
```

3. Automate the process (GNU Make)

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R/analysis.html: R/analysis.Rmd Data/cleandata.csv
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Data/cleandata.csv: R/prepData.R RawData/rawdata.csv
  cd R;R CMD BATCH prepData.R
```

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RawData/rawdata.csv: Python/xls2csv.py RawData/rawdata.xls
  Python/xls2csv.py RawData/rawdata.xls > RawData/rawdata.csv
```

3. Automate the process (GNU Make)

```
R/analysis.html: R/analysis.Rmd Data/cleandata.csv  
    cd R;R -e "rmarkdown::render('analysis.Rmd')"  
  
Data/cleandata.csv: R/prepData.R RawData/rawdata.csv  
    cd R;R CMD BATCH prepData.R  
  
RawData/rawdata.csv: Python/xls2csv.py RawData/rawdata.xls  
    Python/xls2csv.py RawData/rawdata.xls > RawData/rawdata.csv
```

4. Turn scripts into reproducible reports

Gough project diagnostics

Karl Broman, 3 March 2014

Combine genotypes and phenotypes

I've combined the initial genotypes (using the re-clustered genotypes for plates 14-16) with the well-behaved portion of the re-run genotypes. I'm focusing on 36813 markers that are informative (though, as we'll see, there are still a lot of badly behaved and basically non-informative markers that need to be removed). I've combined data on replicate samples, to give one set of genotype calls for each sample.

There are 1497 genotyped mice and 1464 phenotyped mice. All of the mice in the phenotype data have genotypes, but there are 33 genotyped mice with no phenotypes, including 3 Gough mice and 30 F2 progeny.

4. Turn scripts into reproducible reports

Gough project diagnostics

Karl Broman, 3 March 2014

Comb

I've comb
the well-
informat
informat
give one

There are
data have
mice and

```
25 I've combined the initial genotypes (using the re-clustered genotypes
26 for plates 14-16) with the well-behaved portion of the re-run
27 genotypes. I'm focusing on `r totmar(g)` markers that are informative
28 (though, as we'll see, there are still a lot of badly behaved and
29 basically non-informative markers that need to be removed).
30 I've combined data on replicate samples, to give one set of genotype
31 calls for each sample.
32
33 There are `r nind(g)` genotyped mice and `r nrow(phe)` phenotyped
34 mice. All of the mice in the phenotype data have genotypes, but there
35 are `r sum(is.na(match(gid, pid)))` genotyped mice with no phenotypes,
36 including `r sum(g$pheno$gen[which(is.na(match(gid, pid)))]==0)`
37 Gough mice and `r sum(g$pheno$gen[which(is.na(match(gid, pid)))]==2)`
38 F2 progeny.
```

5. Turn repeated code into functions

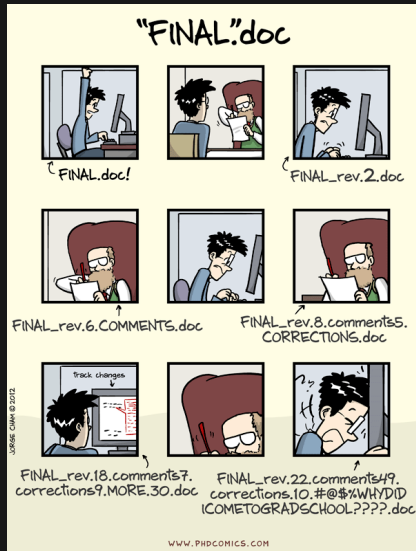
```
# Python
def read_genotypes (filename):
    "Read matrix of genotype data"
```

```
# R
plot_genotypes <-
function(genotypes , ...)
{
}
```


6. Create a package/module

Don't repeat yourself

7. Use version control (git/GitHub)



No “final” in file names

Deprecated/	hypo_prcomp.RData
ReadMe.txt	islet_int1_final.RData
adipose_int1_final.RData	islet_int2_final.RData
adipose_int2_final.RData	islet_mlratio_final.RData
adipose_mlratio_final.RData	islet_mlratio_nqrank_final.RData
adipose_mlratio_nqrank_final.RData	islet_prcomp.RData
adipose_prcomp.RData	kidney_int1_final.RData
aligned_genome_with_pmap.RData	kidney_int2_final.RData
batches_final.RData	kidney_mlratio_final.RData
batches_raw_final.RData	kidney_mlratio_nqrank_final.RData
cpl_final.RData	kidney_prcomp.RData
d2o_final.RData	lipomics_final_rev2.RData
gastroc_int1_final.RData	liverTG_final.RData
gastroc_int2_final.RData	liver_int1_final.RData
gastroc_mlratio_final.RData	liver_int2_final.RData
gastroc_mlratio_nqrank_final.RData	liver_mlratio_final.RData
gastroc_prcomp.RData	liver_mlratio_nqrank_final.RData
hypo_int1_final.RData	liver_prcomp.RData
hypo_int2_final.RData	mirna_final.RData
hypo_mlratio_final.RData	necropsy_final_rev2.RData
hypo_mlratio_final_old.RData	plasmaurine_final_rev.RData
hypo_mlratio_nqrank_final.RData	pmark.RData
hypo_mlratio_nqrank_final_old.RData	rbm_final.RData
hypo_omit.RData	

No “final” in file names

```
Deprecated/  
ReadMe.txt  
adipose_int1_final.RData  
adipose_int2_final.RData  
adipose_mlratio_final.RData  
adipose_mlratio_nqrank_final.RData  
adipose_prcomp.RData  
aligned_genome_with_pmap.RData  
batches_final.RData  
batches_raw_final.RData  
cpl_final.RData  
d2o_final.RData  
gastroc_int1_final.RData  
gastroc_int2_final.RData  
gastroc_mlratio_final.RData  
gastroc_mlratio_nqrank_final.RData  
gastroc_prcomp.RData  
hypo_int1_final.RData  
hypo_int2_final.RData  
hypo_mlratio_final.RData  
hypo_mlratio_final_old.RData  
hypo_mlratio_nqrank_final.RData  
hypo_mlratio_nqrank_final_old.RData  
hypo_omit.RData  
hypo_prcomp.RData  
islet_int1_final.RData  
islet_int2_final.RData  
islet_mlratio_final.RData  
islet_mlratio_nqrank_final.RData  
islet_prcomp.RData  
kidney_int1_final.RData  
kidney_int2_final.RData  
kidney_mlratio_final.RData  
kidney_mlratio_nqrank_final.RData  
kidney_prcomp.RData  
lipomics_final_rev2.RData  
liverTG_final.RData  
liver_int1_final.RData  
liver_int2_final.RData  
liver_mlratio_final.RData  
liver_mlratio_nqrank_final.RData  
liver_prcomp.RData  
mirna_final.RData  
necropsy_final_rev2.RData  
plasmaurine_final_rev.RData  
pmark.RData  
rbm_final.RData
```

7. Use version control (git/GitHub)

PUBLIC kbroman / Talk_MAGIC Unwatch 1 Star 0 Fork 0

Fix two slight bugs in slides: [Browse code](#)

- 8-way RIL by selfing: map expansion = 1 at k=0
- Slight repair to definition of 3-pt coincidence

master

kbroman authored 4 months ago 1 parent e0e8608 commit 51d4aa9ceb104bbf26e8cbe185a5c7f8dc02a832

Showing 2 changed files with 5 additions and 3 deletions. [Show Diff Stats](#)

6 R/map_expansion_func.R [View file @ 51d4aa9](#)

```
... .. @@ -25,8 +25,10 @@ mesibA4 <- function(k)
25 25 #####
26 26 # Eight-way
27 27 #####
28 -mesif8 <- function(k)
29 - 4 - ((1)/(2))^(k-2)
+mesif8 <- function(k) {
29 + if(k==0) return(1)
30 + 4 - ((1)/(2))^(k-2)
31 +}
30 32
31 33 mesibX8 <- function(k)
32 34 ((14)/(3)) - ((30 + 14*sqrt(5))/(15)) * (((1+sqrt(5))/(4)))^k - ((30 - 14*sqrt(5))/(15)) * (((1-s
```

2 magic.tex [View file @ 51d4aa9](#)

```
... .. @@ -636,7 +636,7 @@
636 636
637 637 \hspace{20mm} {\color{myblue} = $\mathsf{Pr}(\text{rec'n in 23})$ |
638 638 \ \text{rec'n in 12})$ /
639 - Pr(\text{rec'n in 12})$}
639 + Pr(\text{rec'n in 23})$}
640 640
641 641 \item
642 642 No interference { \color{myblue} = 1 }
```

8. License your software

Pick a license, any license

– Jeff Atwood

The most important tool is the **mindset**,
when starting, that the end product
will be reproducible.

– Keith Baggerly

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