

Software carpentry and reproducible data analysis

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What do we mean by a reproducibility crisis?

Open access, freely available online

Essay

Why Most Published Research Findings Are False

John P. A. Ioannidis

Summary

There is increasing concern that most current published research findings are false. The probability that a research claim is true may depend on study power and bias, the number of other studies on the same question, and, importantly, the ratio of true to no relationships among the relationships probed in each scientific field. In this framework, a research finding is less likely to be true when the studies

factors that influence this problem and some corollaries thereof.

Modeling the Framework for False Positive Findings

Several methodologists have pointed out [9–11] that the high rate of nonreplication (lack of confirmation) of research discoveries is a consequence of the convenient, yet ill-founded strategy of claiming conclusive research findings solely on

is characteristic of the field and can vary a lot depending on whether the field targets highly likely relationships or searches for only one or a few true relationships among thousands and millions of hypotheses that may be postulated. Let us also consider, for computational simplicity, circumscribed fields where either there is only one true relationship (among many that can be hypothesized) or the power is similar to find any of the

Ioannidis, et al. PLoS Medicine 2005, 2(8):e124

Replicability of a given scientific finding across different labs?

or

Reproducibility of your methods and results?

Questions researchers should ask themselves

- If this computer disappears, can I continue my project unimpeded?
- Will I be able to easily re-run this analysis in a year?
- Can I easily run this analysis on 1,000 more datasets?
- Will someone else be able to easily run and understand this analysis without talking to me?

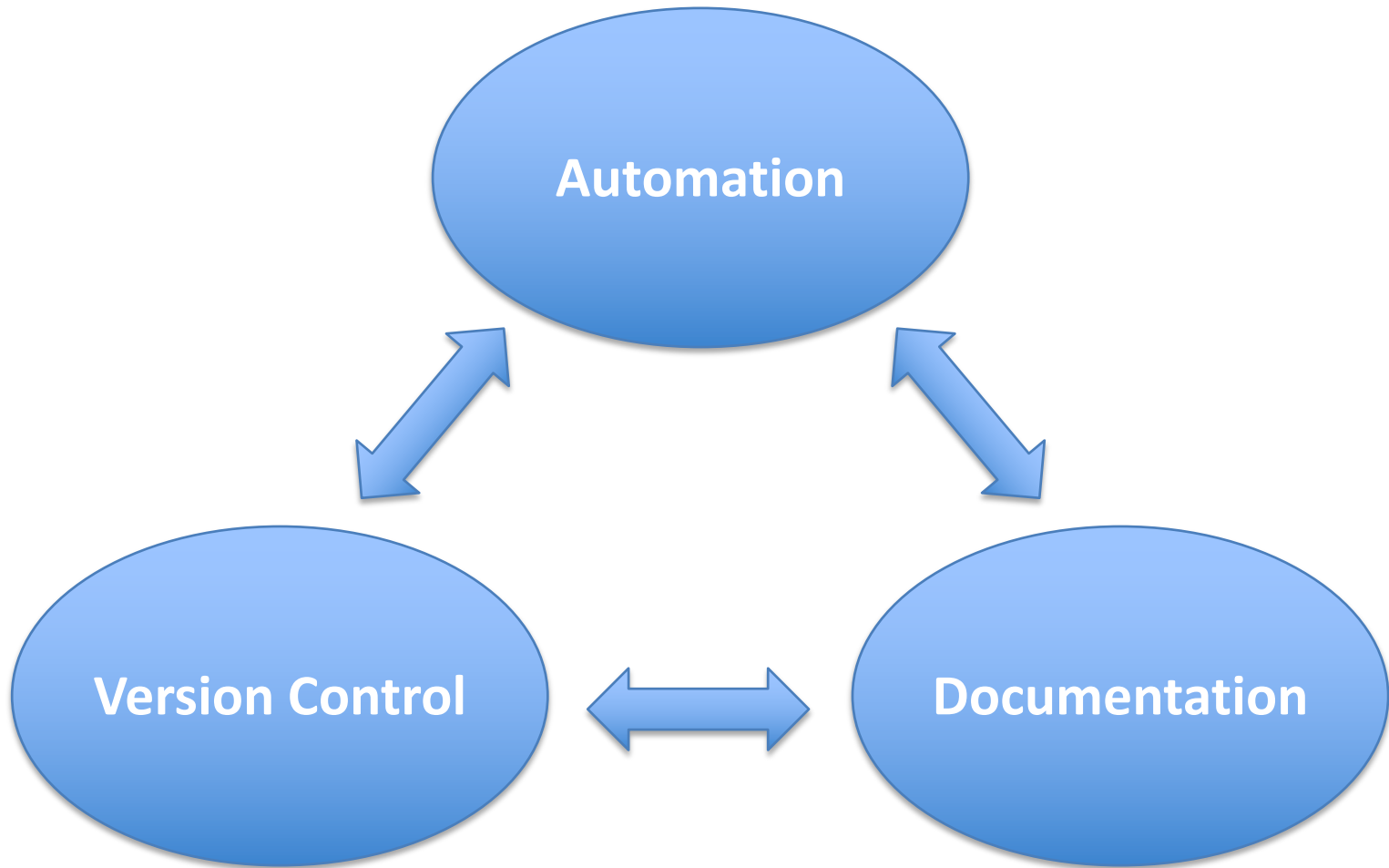
What makes research reproducible?

Knowing **what has been done** and **how has it been done**.

- Enumerate the “set pieces”: data, code, parameters, software versions.
- Explain the rationale behind making certain choices
- Ensure consistency in space and time: from computer to computer, or this year to next.

Decide to make your research reproducible

- **We do it because it helps us.**
- It shouldn't be an extra burden.
- We get better at those things that we practice
- If we do it all the time, no extra effort is needed.



Overview of today's session

Time	Subject
9:00 – 10:15	Intro & Shell scripting basics
10:15 – 10:30	BREAK
10:30 – 11:45	Shell scripting (continued) & Markdown
11:45 – 12:05	Best Practices in Scientific Research
12:05 – 1:15	LUNCH
1:15 – 3:00	Git basics
3:00 – 3:20	BREAK
3:20 – 5:00	Github for team projects

Required items

- Software carpentry website in browser
 - <https://software-carpentry.org/lessons>
- Command-line interface
- Text editor
- Git
- Account on Github

SHELL SCRIPTING

<http://swcarpentry.github.io/shell-novice>

Why use a command-line interface?

- Run analyses **iteratively**.
- Run analyses **reproducibly**.
- Build powerful pipelines (combining tools).
- Use specialized analysis software.
- Access high-performance computing systems.

Basic shell commands

- **pwd** : display working directory
- **ls** : list files
- **cd** : change directory
- **cp** : copy file
- **mv** : move file
- **rm** : remove file
- **mkdir** : make directory
- **cat** : display contents of input
- **head** : display the first few lines of its input
- **tail** : display the last few lines of its input
- **sort** : sort its inputs
- **wc** : count lines, words, and characters in its inputs
- ***** : matches zero or more characters in a filename
- **>** : redirects a command's output to a file
- **|** : pipeline output of the first command as input to the second

Loops

- ```
for filename in *.dat; do
 echo $filename;
done
```
- ```
for i in $(seq 1 0.5 10); do
    echo $i;
done
```
- ```
while read -r line; do
 echo "$line"
done < "$filename"
```
- ```
i="0"
while [ $i -lt 4 ]; do
    echo $i;
    i=$[$i+1]
done
```

Numeric comparison

- -eq is equal to
 if ["\$a" -eq "\$b"]
- -ne is not equal to
- -gt is greater than
- -ge is greater than or equal to
- -lt is less than
- -le is less than or equal to
- < is less than (within double parentheses)
 (("\$a" < "\$b"))
- <= is less than or equal to (within double parentheses)
- > is greater than (within double parentheses)
- >= is greater than or equal to (within double parentheses)

String comparison

- = is equal to
if ["\$a" = "\$b"]
- == is equal to
- != is not equal to
- < is less than, in ASCII alphabetical order
- > is greater than, in ASCII alphabetical order
- -z string is null, that is, has zero length

Named argument passing example

```
#!/bin/bash

usage() { echo "Usage: $0 [-s <45|90>] [-p <string>]"; exit 0;}

while getopts ":s:p:" o; do
    case "${o}" in
        s)
            s=${OPTARG}
            ((s == 45 || s == 90)) || usage
            ;;
        p)
            p=${OPTARG}
            ;;
        *)
            usage
            ;;
    esac
done
shift "$((OPTIND-1))"

if [ -z "${s}" ] || [ -z "${p}" ]; then
    usage
fi
echo "s = ${s}"
echo "p = ${p}"
```

Using the aci-b cluster

#Logging in

```
ssh -X <username>@aci-b.aci.ics.psu.edu
```

#Submitting jobs to the open queue

```
qsub -A open -l nodes=1:ppn=1 -l walltime=0:05:00 script.sh
```

#Checking job status

```
qstat
```

#Deleting a job

```
qdel <job id>
```

#Available software modules

```
module avail
```

#Load software modules

```
module load r/3.3
```

Example cluster script header

```
#PBS -l walltime=1:00:00
```

```
#PBS -l nodes=1:ppn=1
```

```
#PBS -j oe
```

```
#PBS -A open
```

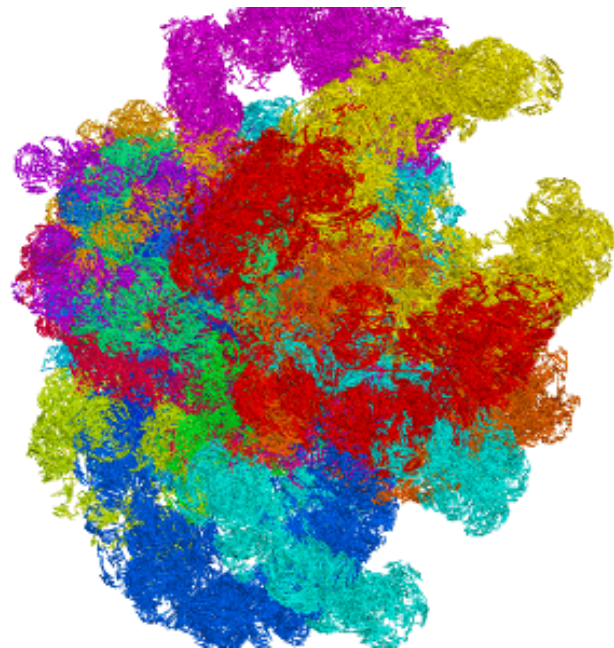
```
#PBS -l mem=1gb
```

```
module load r/3.3
```

```
module load python/2.7.8
```

Reproducibility in my lab: miniMDS

miniMDS: 3D structural inference from high-resolution Hi-C data
Rieber & Mahony, **Bioinformatics**, 2017



<http://github.com/seqcode/miniMDS>

The screenshot shows three views of the GitHub repository `seqcode/miniMDS`:

- Code View:** Shows the main repository page with 176 commits. The sidebar lists files like `scripts`, `README.md`, and `array_tools.py`.
- Documentation View:** Shows the `README.md` file content. The `minimds` section describes the tool for inferring 3D structures from normalized H approximation to multidimensional scaling (MDS). The `Installation` section lists prerequisites: python 2.7, numpy, scikit-learn, pymp, mayavi (optional), ImageMagick (optional), scipy (optional), and matplotlib (optional).
- Figure regeneration View:** Shows the `scripts` repository with a list of commits. The commits include:
 - Lila fixed method call
 - updated readme
 - updated figure scripts
 - updated figure scripts
 - fixed method call
 - updated figure scripts
 - added out path
 - fixed bug
 - fixed method call
 - updated figure scripts
 - fixed time command
 - changed parameters to defaults

Code

Documentation

Figure regeneration

Many bioinformatics tools are not executable!

Name	Type	Results from 1-Kbp resolution data
miniMDS	Optimization	Success
BACH	Modeling	Bug
ShRec3D	Optimization	Success
MOGEN (Trieu and Cheng, 2014)	Optimization	Exceeds maximum input size
3D-GNOME (Szałaj et al., 2016)	Optimization	Success (graphical output only)
ChromSDE (Zhang et al., 2013)	Optimization	Failed to produce output in reasonable time
tRex (Park and Lin, 2016)	Modeling	Requires fragment length
MCMC5 (Rousseau et al., 2011)	Modeling	Requires standard deviation
PASTIS (Varoquaux et al., 2014)	Modeling	Failed to install IPOPT
TAD-bit (Baù and Marti-Renom, 2012)	Optimization	Failed to produce output in reasonable time
HSA (Zou et al., 2016)	Modeling	Memory error
Chromosome3D (Adhikari et al., 2016)	Optimization	Failed to produce output in reasonable time
InfMod3DGen (Wang et al., 2015)	Modeling	Bug

But we're not perfect...

- From a different manuscript submitted from my lab (not from Lila) last year:
 - Reviewer 2:

...

“It looks easy to use - it's a java command line program. But I got the "You don't have permission to access on this server." when trying to download the executable jar file. So I can't really assess this.”

...

BEST PRACTICES FOR SCIENTIFIC COMPUTING

Wilson, *et al.* PLoS Biology, 2014 12(1):e1001745

Rule 1: Write Programs for People, not Computers

1.1: Keep it simple.

1.2: Make names consistent, distinctive, and meaningful.

1.3: Make code style and formatting consistent.

Rule 2: Let the Computer Do the Work

- 2.1: Make the computer repeat tasks.**

- 2.2: Save recent commands in a file for re-use.**

- 2.3: Use a build tool to automate workflows.**

Rule 3: Make Incremental Changes

3.1: Small steps with frequent feedback

3.2: Use a version control system.

3.3: Version control EVERYTHING
(...except big data)

Rule 4: Don't Repeat Yourself (or Others)

- 4.1: Every piece of data must have a single authoritative representation
- 4.2: Modularize code rather than copying and pasting.
- 4.3: Re-use code instead of rewriting it.

Rule 5: Plan for Mistakes

5.1: Add assertions to programs to check their operation.

5.2: Use an off-the-shelf unit testing library.

5.3: Turn bugs into test cases.

5.4: Use a symbolic debugger.

Rule 6: Optimize Software Only After It Works Correctly

- 6.1: Use a profiler to identify bottlenecks.**

- 6.2: Write code in the highest-level language possible.**

Rule 7: Document Design and Purpose, Not Mechanics

7.1: Document interfaces and reasons not implementations.

7.2: Refactor code in preference to explaining how it works.

7.3: Embed the documentation for a piece of software in that software.

Rule 8: Collaborate

8.1: Use code reviews.

8.2: Use pair programming

8.3: Use an issue tracking tool.

VERSION CONTROL WITH GIT

<http://swcarpentry.github.io/git-novice>

THIS IS GIT. IT TRACKS COLLABORATIVE WORK
ON PROJECTS THROUGH A BEAUTIFUL
DISTRIBUTED GRAPH THEORY TREE MODEL.

COOL. HOW DO WE USE IT?

NO IDEA. JUST MEMORIZIZE THESE SHELL
COMMANDS AND TYPE THEM TO SYNC UP.
IF YOU GET ERRORS, SAVE YOUR WORK
ELSEWHERE, DELETE THE PROJECT,
AND DOWNLOAD A FRESH COPY.



Git commands

\$ git init [project-name]

Creates a new local repository with the specified name

\$ git clone [url]

Downloads a project and its entire version history

\$ git status

Lists all new or modified files to be committed

\$ git add [file]

Snapshots the file in preparation for versioning

\$ git reset [file]

Unstages the file, but preserve its contents

\$ git diff

Shows file differences not yet staged

\$ git diff --staged

Shows file differences between staging and the last file version

\$ git commit -m "[message]"

Records file snapshots permanently in version history

\$ git push [alias] [branch]

Uploads all local branch commits to GitHub

\$ git pull

Downloads bookmark history and incorporates changes

	COMMENT	DATE
O	CREATED MAIN LOOP & TIMING CONTROL	14 HOURS AGO
O	ENABLED CONFIG FILE PARSING	9 HOURS AGO
O	MISC BUGFIXES	5 HOURS AGO
O	CODE ADDITIONS/EDITS	4 HOURS AGO
O	MORE CODE	4 HOURS AGO
O	HERE HAVE CODE	4 HOURS AGO
O	AAAAAAA	3 HOURS AGO
O	ADKFJSLKDFJSOKLFJ	3 HOURS AGO
O	MY HANDS ARE TYPING WORDS	2 HOURS AGO
O	HAAAAAAAAANDS	2 HOURS AGO

AS A PROJECT DRAGS ON, MY GIT COMMIT
MESSAGES GET LESS AND LESS INFORMATIVE.

DISCUSSION & WRAP-UP

Discussion points

- What did we cover today that was new to you?
- How can we put all the pieces together?
- How will you put all of this into practice?