Putting it all together





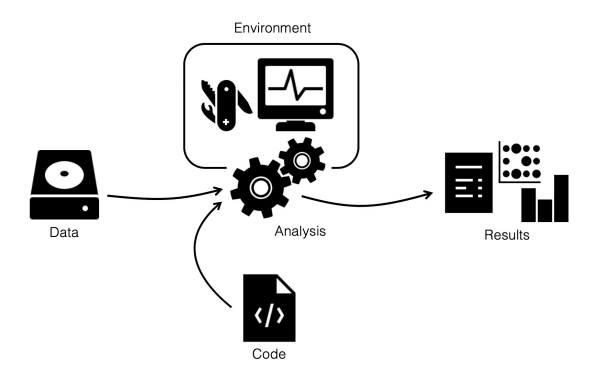
Putting it all together

- 1. Lecture: Summary of the week & how to put all the tools and procedures together
- 2. Q&A: How to implement these procedures on a day-to-day basis
- 3. Project: Time for you to apply the tools on one of your own research projects





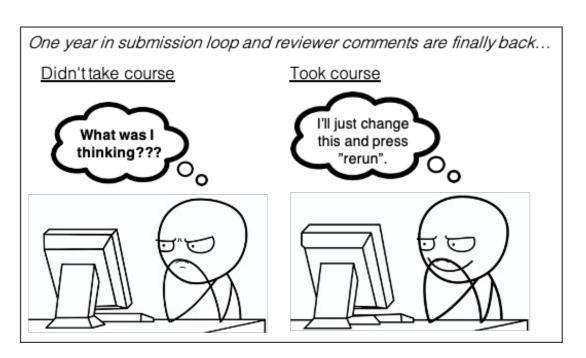
Take control of your research project by making its different components reproducible







By moving towards a reproducible way of working you will quickly realize that you at the same time make your own life a lot easier!







What have we learned?



- How to use the version control system Git to track changes to code
- How to use the package and environment manager Conda
- How to use the workflow manager
 Snakemake
- How to use R Markdown to generate automated reports
- How to use Jupyter notebooks to document your analysis
- How to use Docker and Singularity to distribute containerized computational environments





Divide your work into distinct projects

- Keep all files needed to go from raw data to final results in a dedicated directory
- Use relevant subdirectories
- Many software support the "project way of working", e.g. Rstudio and the text editors
 Sublime Text and Atom
- Use Git to create structured and version controlled project repositories





Everything can be a project

Project directory templates, e.g. NBIS project template:

```
project
I- doc/
                       documentation for the study
|- data/
                       raw and primary data, essentially all input files, never edit!
   |- raw_external/
  |- raw_internal/
  |- meta/
                       all code needed to go from input files to final results
|- code/
notebooks/
|- intermediate/
                       output files from different analysis steps, can be deleted
                       temporary files that can be safely deleted or lost
|- scratch/
|- logs/
                       logs from the different analysis steps
- results/
                       output from workflows and analyses
  |- figures/
  |- tables/
  |- reports/
                       sets which parts of the repository that should be git tracked
gitignore
|- Snakefile
                       project workflow, carries out analysis contained in code/
                       configuration of the project workflow
|- config.yml
|- environment.yml
                       software dependencies list, used to create a project environment
|- Dockerfile
                       recipe to create a project container
```

- https://github.com/NBISweden/project_template
- https://github.com/snakemake-workflows/cookiecutter-snakemake-workflow





Treasure your data

- Keep your input data read-only consider it static
- Don't create different versions of the input data write a script, R Markdown document, Jupyter notebook or a Snakemake workflow if you need to preprocess your input data so that the steps can be recreated
- Backup! Keep redundant copies in different physical locations
- Upload your raw data as soon as possible to a public data repository





Organize your coding

- Avoid generating files interactively or doing things by hand there is no way to track how they were made
- Write scripts, R Markdown documents, Jupyter notebooks or Snakemake workflows for reproducible results to connect raw data to final results
- Keep the parameters separate (e.g. at top of file or in a separate configuration file)





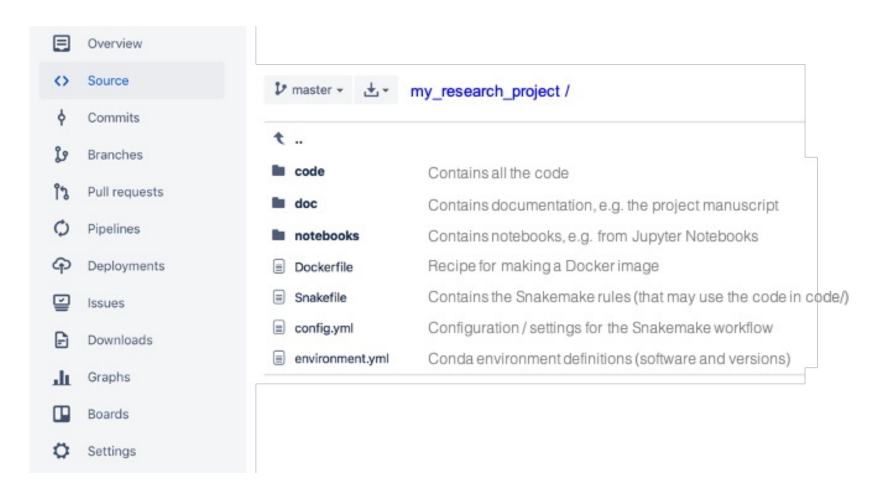
Discuss in your groups

Which of these advices are most relevant to your work?

Are you going to change your way of working? If yes, how?



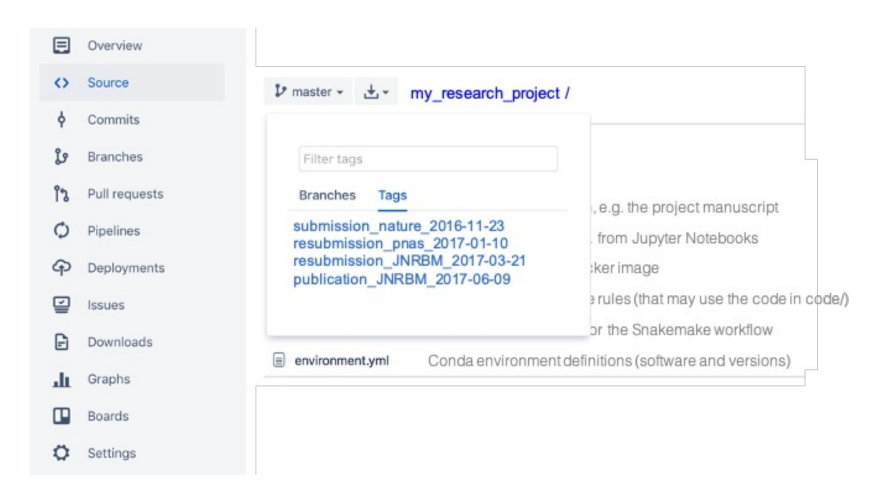




www.bitbucket.org













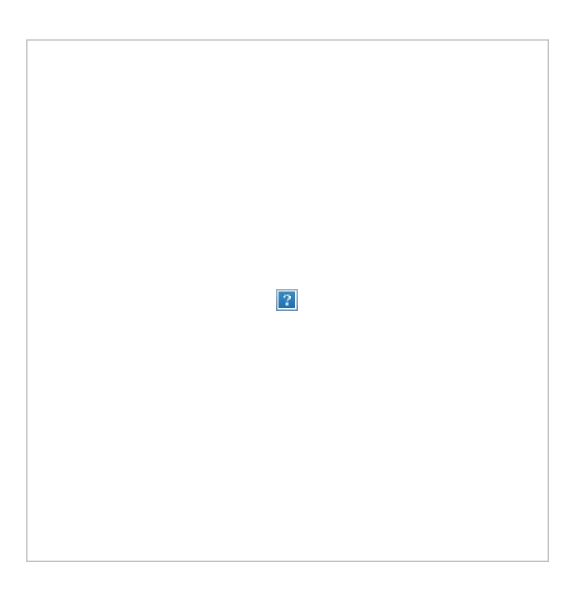
- Git clone and run code or workflow
- Git clone, create Conda env, and run code or workflow
- Git clone, Docker build, and run code or workflow in container
- Docker pull (from online repository) and run code or workflow in container





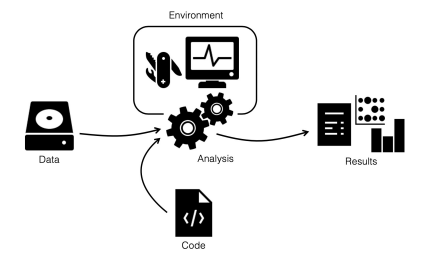
Example video:

Pulling the Docker image for the case study from an online repository and running the workflow in a container



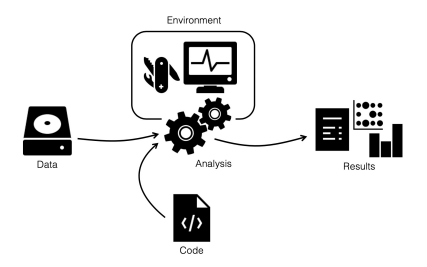












Minimal: write code in a reproducible way

Connect your results with the code:

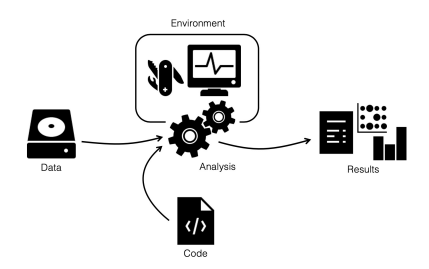
 Use R Markdown documents or Jupyter notebooks

Take another step:

Convert your code into a Snakemake workflow







Minimal: write code in a reproducible way

Good: versioned and structured

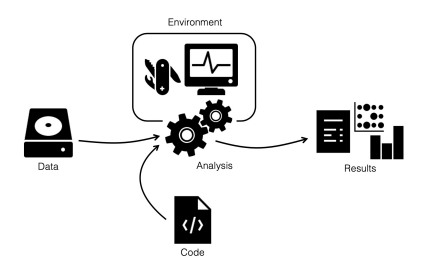
repository

Use Git for version controlling and collaboration:

- Create one Git repository per project
- Track your changes with Git
- Publish your code along with your results on GitHub or BitBucket







Minimal: write code in a reproducible way

Good: versioned and structured

repository

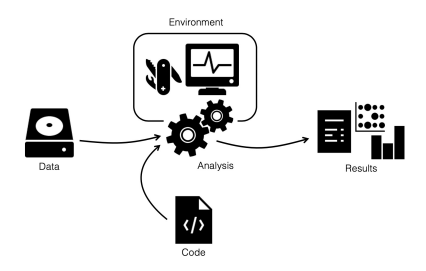
Better: organize software dependencies

Manage your depencencies:

 Use Conda to install software in environments that can be easily exported and installed on a different system







Minimal: write code in a reproducible way

Good: versioned and structured

repository

Better: organize software dependencies

Best: export everything!

Completely recreate the compute system:

 Consider packaging your project in a Docker or Singularity container





Discuss in your groups

Which of the suggested ambition levels seems most reasonable for your work?

Are you going to use one of the suggested combinations of tools in your research? Or are you planning to use a different combination?





Version control

- Git Widely used and a lot of tools available + GitHub/BitBucket.
- Mercurial Distributed model just like Git, close to sourceforge.
- Subversion Centralized model unlike git/mercurial; no local repository on your computer and somewhat easier to use.

Environment / package managers

- Conda General purpose environment and package manager. Community-hosted collections of tools at bioconda or conda-forge.
- Pip Package manager for Python, has a large repository at pypi.
- Apt/yum/brew Native package managers for different OS. Integrated in OS and might deal with e.g. update notifications better.
- Virtualenv Environment manager used to set up semi-isolated python environments.





Workflow managers

- Snakemake Based on Python, easily understandable format, relies on file names.
- Nextflow Based on Groovy, uses data pipes rather than file names to construct the workflow.
- Make Used in software development and has been around since the 70s. Flexible but notoriously obscure syntax.
- Galaxy attempts to make computational biology accessible to researchers without programming experience by using a GUI.





Literate programming

- Jupyter Create and share notebooks in a variety of languages and formats by using a web browser.
- R Markdown Developed by Rstudio, focuses on generating high-quality documents.
- Zeppelin Developed by Apache. Closely integrated with Spark for distributed computing and Big Data applications.
- Beaker Newcomer based on Ipython, just as Jupyter. Has a focus on integrating multiple languages in the same notebook.





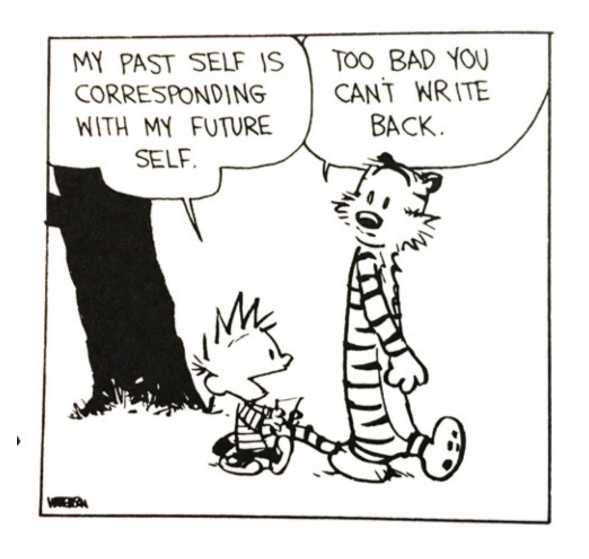
Containerization / virtualization

- Docker Used for packaging and isolating applications in containers. Dockerhub allows for convenient sharing. Requires root access.
- Singularity Simpler Docker alternative geared towards high performance computing. Does not require root.
- Shifter Similar ambition as Singularity, but less focus on mobility and more on resource management.
- VirtualBox/VMWare Virtualization rather than containerization. Less lightweight, but no reliance on host kernel.





"What's in it for me?"







NBIS Bioinformatics drop-in

Any questions related to reproducible research tools and concepts? Talk to an NBIS expert!

- Online (zoom)
- Every Tuesday, 14.00-15.00 (except public holidays)
- Check www.nbis.se/events for zoom link and more info





Q&A: How to implement these procedures on a dayto-day basis



