Bwb tutorial: Agenda

1. Presentation
2. Demo
3. Q&A


Container: [https://hub.docker.com/r/biodepot/bwb](https://hub.docker.com/r/biodepot/bwb)

YouTube of tutorial: [https://www.youtube.com/watch?v=r_03_UG1mBg](https://www.youtube.com/watch?v=r_03_UG1mBg)

Building containerized workflows for RNA-seq data using the BioDepot-workflow-Builder (Bwb). Hung et al. [bioRxiv 099010](https://www.biorxiv.org/content/10.1101/099010)
Using BioDepot-workflow-Builder (Bwb) to create and execute reproducible bioinformatics workflows

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Motivation

Transitioning from laptop/desktop/cluster based analyses to cloud based analyses

How do we facilitate sharing new workflows reproducibly?

How do we facilitate executing workflows on the cloud?

How do we get adoption and input from biologists?
Reproducibility in biology

- **Irreproducible**
  - US$28.2B (50%)

- **Reproducible**
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Estimated US Annual Preclinical Research Spend

**Categories of Preclinical Irreproducibility**

- Biological Reagents and Reference Materials (36.1% of total)
- Study Design (27.6% of total)
- Data Analysis and Reporting (25.5% of total)
- Laboratory Protocols (10.8% of total)

The spectrum of reproducibility

Reproducibility exercise on a DREAM Challenge

Students in a 500-level bioinformatics class were asked to reproduce results from a crowd sourcing challenge in 2016-17. **Given:** time series gene expression data (time 0, 24 hours) across 4 different viruses (H1N1, H3N2, RSV, Rhinovirus) in 7 studies. **Goal:** build predictors to distinguish people who become contagious after exposure to flu and other respiratory viruses.
Question: How many do you think we can reproduce?

Challenges:

- Missing files: e.g. missing pre-processed data, need access to Google drive
- Environment setup: e.g. version conflict, memory, disk space
- Code issues: e.g. don’t have software, code simply doesn’t run, missing library
Challenges in reproducing bioinformatics analysis results

- Parameters not given or incorrectly given
- Environment assumed i.e. directory structure
- Configuration assumed
- Different OS/software versions
- Bioinformatics analyses usually involve a pipeline consisting of many tools.
- These tools often have graphical user interfaces.

Our solution: We share the computing environment, not just code with Bwb.
• Docker containers wrap up a piece of software in a complete file system that contains everything it needs to run: code, runtime, system tools, system libraries – anything you can install on a server.

• It will give the same results regardless of the host environment

• Docker Hub: repository of Docker containers

https://www.docker.com/what-docker
BioDepot-workflow-Builder (Bwb)

- **Drag-and-drop** form based interface
- **Modular**: Build modular bioinformatics workflows using widgets. Mix-and-match.
- **Reproducible results**: each module is encapsulated by a software container, computing environment and parameters are maintained
- **Extensible**: can add easily new widgets using a form-based builder.
- Support graphical output in modules
- Not affected by third party software upgrades.
- Easily deployed across different platforms.
Drag-and-drop GUI
Widgets

- Represent modular units in a workflow, such as an executable, script or a basic operation.
- Each widget executes a Docker container.
- Can be dragged to the canvas and connected to form workflows.
Workflows

When the pipeline is to be run, the Bwb/Orange engine follows the graph, executing each widget and then propagating the output to connected widgets which are then executed, until the entire graph is traversed.
Running Bwb

Bwb can be deployed on your local host or any cloud platform with Docker installed.

1. Download the Bwb Docker image
docker pull biodepot/bwb:latest

2. Start the Bwb container
docker run --rm -it -p 6080:6080 -v ${PWD}:/data \
   -v /var/run:/var/run -v /tmp/.X11-unix:/tmp/.X11-unix biodepot/bwb

To access the container open up a browser window and type in localhost (or the IP of the container) followed by the port number 6080 i.e. localhost:6080
Bwb: Implementation details

- Drag-and-drop interface based on OrangeML library (http://orange.biolab.si)
- OrangeML: widget is a manually written Python file
- Bwb widgets
  - stores the widget definition in 3 JSON (JavaScript Object Notation) files, 1 icon file and 1 auto-generated Python file in a directory.
  - parameters specified in widgets stored in JSON and xml files.
RNA-seq case study

Differential expression of genes:
(count the number of transcripts made by genes under different conditions)

- align short sequence reads to genome sequence to identify transcript (intensive C/C++ executables)
- statistical analyses to calculate which transcripts/genes are differentially expressed (R or Python libraries)
- visualization (R/Python/Jupyter

Example
Kallisto-sleuth RNA-seq workflow
Ex: RNA-seq workflow using Kallisto and Sleuth [Pachter Lab @ Caltech]

- **Kallisto** [Bray et al. 2016]
  - a program for fast RNA-Seq quantification based on pseudo-alignment.
  - Can quantify 30 million human reads in less than 3 minutes on a desktop computer using only the read sequences and a transcriptome index that itself takes less than 10 minutes to build.
  - Written in C++

- **Sleuth** [Pimentel et al. 2017]
  - Written in R. Shiny app with plots.
  - Dependences on other R packages
  - Recommended installation: biocLite or conda
  - Issues with the latest version of R
Ex: RNA-seq workflow using Kallisto and Sleuth [Pachter Lab]

- Sample data: GSE37704
- RNA-seq data in which lung fibroblasts in response to loss of the developmental transcription factor HOXA1 [Trapnell et al. 2013]
- 6 samples: 3 knockdown, 3 control

Download input fastq files and reference sequence → Kallisto index → Kallisto quant → Differential expression inference using sleuth
The Jupyter Notebook is an open-source web application that allows you to create and share documents that contain live code, equations, visualizations and explanatory text.
Demo
Take home message

• Reproducible results when containers are deployed across different hardware and software configurations.
• Dockerization makes it easy to swap out a component and not worry about dependencies. Each module can be run in a different environment.
• UI to create your own containerized widgets that run your customized code.
Future Work

• BwB
  – Scheduler
  – Support the Common Workflow Language and YAML

• Benchmarking bioinformatics workflows on the cloud and build predictive models

• Improve the scalability of workflows
Using Docker containers to address the reproducibility of bioinformatics analyses

GUIdock: Using Docker containers with a common graphics user interface to address the reproducibility of research. Ling-Hong Hung, Daniel Kristiyanto, Sung Bong Lee, Ka Yee Yeung. _PLOS One 2016, 11(4):e0152686_.

GUIdock-VNC: using a graphical desktop sharing system to provide a browser-based interface for containerized software. Varun Mittal, Ling-Hong Hung, Jayant Keswani, Daniel Kristiyanto, Sung Bong Lee, Ka Yee Yeung. _Gigascience 2017, 6(4): 1-6_.


Embedding containerized workflows inside data science notebooks enhances reproducibility. Jiaming Hu, Ling-Hong Hung, Ka Yee Yeung. _bioRxiv 309567_.

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For more information, visit https://github.com/BioDepot/
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