The RNA workbench: best practices for RNA and high-throughput sequencing bioinformatics in Galaxy

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Received March 02, 2017; Revised April 13, 2017; Editorial Decision April 28, 2017; Accepted May 31, 2017

ABSTRACT

RNA-based regulation has become a major research topic in molecular biology. The analysis of epigenetic and expression data is therefore incomplete if RNAbased regulation is not taken into account. Thus, it is increasingly important but not vet standard to combine RNA-centric data and analysis tools with other types of experimental data such as RNA-seq or ChIP-seq. Here, we present the RNA workbench, a comprehensive set of analysis tools and consolidated workflows that enable the researcher to combine these two worlds. Based on the Galaxy framework the workbench guarantees simple access, easy extension, flexible adaption to personal and security needs, and sophisticated analyses that are independent of command-line knowledge. Currently, it includes more than 50 bioinformatics tools that are dedicated to different research areas of RNA biology including RNA structure analysis, RNA alignment, RNA annotation, RNA-protein interaction, ribosome profiling, RNA-seq analysis and RNA target prediction. The workbench is developed and maintained by experts in RNA bioinformatics and the Galaxy framework. Together with the growing community evolving around this workbench, we are committed to keep the workbench up-to-date for future standards and needs, providing researchers with a reliable and robust framework for RNA data analysis. Availability: The RNA workbench is available at https://github.com/bgruening/galaxy-rna-workbench.

INTRODUCTION

Since recent advances in high-throughput sequencing (HTS) emphasized the importance and versatile role of (non-coding) RNAs, there is high demand for integrated computational analyses investigating RNA-mediated regulation. Previously existing workbenches (such as *miARma-Seq* (1) *RAP* (2) and the UEA Small RNA Workbench (3)) were focused on providing tools for the analysis of RNA deep sequencing data and do not contain RNA centric tools.

We addressed these needs by developing the RNA workbench. Based on the Galaxy framework (4) it combines a

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 $^{{\}hbox{$\mathbb{C}$ The Author(s) 2017. Published by Oxford University Press on behalf of Nucleic Acids Research.}}$

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comprehensive set of tools for the analysis of RNA structures, RNA alignments, RNA-RNA and RNA-protein interactions, RNA sequencing, ribosome profiling, genome annotation and many more. So far, we integrated more than 50 RNA-related tools, including suites like the ViennaRNA package, covering this broad variety of use-cases (a complete list of tools can be found on GitHub). Every available tool works as a single building-block that can be connected with other tools to create computational pipelines. Datasets can be incorporated in a similar manner, facilitating an intersection of diverse data sources such as DNA methylation with RNA-seq experiments. Input and output datasets can be defined by the user, and can be as diverse as the adapted set of tools. Established data types for sequence and/or structure information are accepted as input. Output data types follow the same principle, can be converted to different formats, or ultimately used to draw plots and create figures. The workbench provides tools for visualizations of RNA structure datasets, such as dot-bracket strings, and RNA 2D or 3D structures. The workbench also covers a broad range of RNA secondary structure prediction and analysis tools such as RNAfold (5) or LocARNA (6,7).

GOALS OF THE RNA WORKBENCH

The main driving force behind the development of the RNA workbench is the goal to establish a central, redistributable workbench for scientists and programmers working with RNA-related data, and build a sustainable community around it. This platform is unique in combining available tools, workflows and training material, as well as providing easy access for experimentalists. Simultaneously, it serves as a central hub for programmers, which can easily integrate and deploy their existing or novel tools and workflows. The RNA workbench is based on three pillars: (i) a comprehensive set of RNA-bioinformatics tools, (ii) easy and stable dissemination via Galaxy and Docker and (iii) a set of predefined workflows and associated descriptions/training material. The latter is needed for two reasons: first, it facilitates the use of the RNA workbench for researchers with limited bioinformatics experience, and second, it allows to integrate the workbench in the daily lab work by combining RNArelated analysis tasks with workflows for RNA-seq analysis.

Building on the shoulders of giants

In order to achieve long-term sustainability, we provide the essentials of our work on *BioConda* (https://bioconda. github.io) and BioContainers (8) (http://biocontainers.pro) for reproducible deployments of tools into Galaxy. Using easy-to-distribute packages for all tool dependencies also enables automatic continuous integration tests for all developed tools and the workbench. After a tool passes the tests and gets accepted it will be made available via an automatic deployment into the Galaxy ToolShed (https://toolshed.g2. bx.psu.edu) (9). From the ToolShed, Galaxy administrators can easily install desired tools and workflows.

Easily accessible and reproducible analysis platform

For the fast dissemination of the RNA workbench, as well as for an easy integration with other HTS analysis tasks, we implemented the RNA workbench within the Galaxy framework. A major advantage of relying on Galaxy as the core framework is that it is possible to leverage its scalability, which enables the RNA workbench to run on single CPU installations as well as on large multi-node high performance computing environments. Furthermore, Galaxy provides researchers with means to reproduce their own workflow analyses, enabling them to rerun entire pipelines, or publish and share them with others. The RNA workbench is containerized, i.e., administrators can deploy it via *Docker*. That makes it possible to have all tool installation dependencies already resolved, while still keeping maintenance tasks to a minimum. The provided layer of virtualization also allows the handling of user-defined input data in a secure and compartmentalized way, a key requirement for researchers working on sensitive data (e.g. patient data in clinics). Running the containerized RNA workbench simply requires installing *Docker* and starting the Galaxy RNA workbench image. Furthermore, containerizing Galaxy enables a customized Galaxy instance with a selected subset of tools dedicated to specific data analysis tasks, while keeping deployment and installation simple.

RNA-BIOINFORMATICS TOOLS

In its current state, the RNA workbench includes more than 50 tools covering all aspects of RNA research. In a community effort, these tools will be kept up-to-date and adapted to future needs. New tools and new ways to visualize data provided to the user will also be integrated. A current overview of tools available in the RNA workbench can be found at http://bgruening.github.io/galaxyrna-workbench/.

In the following, we will highlight a few of the integrated tools.

The *ViennaRNA* package (5) consists of a suite of tools centered around the prediction of secondary structures of RNAs based on the thermodynamic Turner energy model. Thus, it covers prediction of optimal and suboptimal structures from single sequences as well as alignments, prediction of ensemble base pair probabilities, accessibility of sequences, and RNA-RNA interaction prediction. Importantly, predictions can be flexibly controlled by hard and soft structure constraints; the latter enables the inclusion of structure probing data.

AREsite2 (10) is a resource for the investigation of AU, GU and U-rich elements (ARE, GRE, URE) in human and model organisms. It provides information on genomic location, genomic context, RNA secondary structure context and conservation of annotated motifs in the whole gene body including introns. It is integrated into the RNA workbench via its REST interface, which provides search results directly in Galaxy for further analysis.

LocARNA (6,7) provides a comparative analysis of multiple (unaligned) RNAs by simultaneous folding and alignment, implementing a fast variant of the Sankoff algorithm. Beyond pairwise and multiple alignments, it computes reliabilities of alignment columns and provides very fast analysis by simultaneous folding and matching. Finally, *LocARNA* supports anchor and structure constraints, which improve its applicability in practice.

doRiNA (11) is a database of RNA interactions in post-transcriptional regulation. The combined action of RNA-binding proteins (RBPs) and microRNAs (miRNAs) is believed to form the backbone of post-transcriptional regulation. doRiNA is implemented as data source tool inside the RNA workbench. This means that the Galaxy user is redirected to the post-transcriptional interaction database and can make selections using the optimized doRiNA interface. Once the selection is done, the data is streamed directly to Galaxy and can be freely analyzed with other tools.

The *Infernal* (12) tool suite can construct probabilistic models, also called covariance models (CM), that represent the sequence and structure of an RNA family from a multiple sequence alignment with consensus secondary structure. The covariance model can be used to find more members of this RNA family via homology search.

PARalyzer (13) generates a high resolution map of interaction sites between RNA-binding proteins and their targets. The algorithm utilizes the deep sequencing reads generated by the PAR-CLIP (Photoactivatable-Ribonucleoside-Enhanced Crosslinking and Immunoprecipitation) protocol. The use of photoactivatable nucleotides in the PAR-CLIP protocol results in more efficient crosslinking between the RNA-binding protein and its target relative to other CLIP methods; in addition a nucleotide substitution occurs at the site of crosslinking, providing for single-nucleotide resolution binding information. PARalyzer utilizes this nucleotide substitution in a kernel density estimate classifier to generate the high resolution set of protein-RNA interaction sites.

FuMa (14) can generate an integration report on predicted fusion genes from most RNA-seq fusion gene detection software. It automatically orders the result based on the frequencies of the fusion genes such that frequently predicted fusion genes can be extracted.

WORKFLOWS

One of the core concepts of the RNA workbench is the definition of standard workflows as a minimal set of building blocks around which a researcher can compose and tailor specific pipelines. For example, a researcher wants to analyze the effects of an RNA-binding protein (RBP) in regard to expression levels in wild-type compared to knockout or knockdown of the RBP of interest. In this case, one needs to combine the detection of differentially expressed genes in the two conditions with the information of publicly available CLIP-data, as provided for example by the doRiNA (11) database, to differentiate between direct and indirect targets. Workflows for the analysis of differentially expressed genes are part of the RNA workbench, as well as an interface to doRiNA, such that it becomes an easy task to design a new workflow combining these analysis steps.

In Galaxy, workflows are typically created in two different ways: (i) from an existing history, which stores all tools applied in a previous analysis together with all pertinent parameters, or (ii) from scratch, using a graphical editor via drag-and-drop of tools from the tool panel into

the workflow editor. Within workflows, tools can be freely combined to ensure a maximum of flexibility in their usage and connectivity between different analysis steps, e.g. RNA structure analysis tools and RNA-seq data analysis. Various format converters embedded in Galaxy allow combining diverse analysis outputs. Easy sharing of workflows with other Galaxy users guarantees highly reproducible and transparent research. In other words, the workflows ensure that all analysis steps, tools and parameters of an experiment are documented and visible to researchers, readers and reviewers. Workflows can also be submitted to the Galaxy ToolShed or *myexperiment.org* (15) for further distribution. The RNA workbench currently includes publicly available standard workflows for RNA data analysis, e.g. for RNAseq. These workflows contain all required steps such as quality control, mapping, differential expression analysis, and visualization of results. Provided workflows can easily be extended or modified, e.g. to use other read mappers available in Galaxy.

In the following, we will describe two sample workflows, one closely related to the detection of ncRNAs, which is a common task in RNA-related research. The other workflow is related to the analysis of RNA-seq data and is often needed as a subworkflow for more complex analysis tasks. These workflows are well annotated and described in the RNA workbench and extended by interactive Galaxy *tours*.

Analysis of (unaligned) non-coding RNAs

An important task is to test for the existence of a functional structure in a non-coding RNA. However, the secondary structure of structured non-coding RNAs is not significantly more stable compared to random sequences (16). Thus, putative functional structures can only be detected using information about conservation. Our workflow for non-coding RNAs performs the typical analysis steps required to detect conserved secondary structures, given a set of unaligned RNA sequences. It computes a sequence and a structure-based alignment by MAFFT (17) and LocARNA, respectively, and analyzes them with RNAcode (18) and RNAz (19) with appropriate parameter settings. RNAz and RNAcode both work on a given alignment. RNAz tests whether a consensus secondary structure is significantly conserved, whereas RNAcode differentiates coding from non-coding RNAs. Together these tools provide information, whether the RNAs are related and conserve a common secondary structure. In addition, a covariance model is built from the LocARNA alignment and subsequently used to search the given sequence database for RNAs with similar sequence- and structure-conservation. This workflow resembles the core of RNAlien (20), which is based on the same tools and is integrated into the RNA workbench. Going beyond the presented workflow, RNAlien automatically gathers sequences via homology search starting from a single sequence and constructs RNA family models in an iterative process.

To give an other example, in the context of $\mu ORFs$ detection, RNA-seq analysis, the identification of non-coding RNAs with *RNAcode* and *RNAz* and the detection of transcription start sites can be used to determine new, short transcripts that are expressed and do not exhibit secondary

structure conservation (i.e. are likely not functional ncR-NAs). Subsequent analysis of Ribo-seq data can then provide additional evidence for a new transcript that may code for a small protein. For all these tasks, partial workflows and required tools are already integrated in our RNA workbench, which implies that it is easy to set up a new workflow for a more complex task.

RNA-seg analysis: trimming, mapping and read count

As mentioned before, the analysis of RNA-centric data like CLIP-seq requires the combination with other type of data, and very often RNA-seq. For that reason, we provide a standard RNA-seq workflow that can easily be combined with other workflows. The RNA-seq workflow (as shown in Figure 1) takes a list of RNA-seq datasets as input and successively executes a series of analysis steps - adapter & quality trimming, mapping to a reference genome and read count per annotated gene. The input allows two conditions, e.g. treatment versus control and it also accepts single-end and paired-end reads for each condition. At the trimming step, the workflow employs Trim Galore! (21,22) to perform adapter trimming. Then, *TopHat2* (23) is used to map the trimmed reads against the reference sequences, which should be provided by the user. As last step, the workflow executes HTSeq-count (24) to generate read counts per annotated gene for each condition and for each sequencing type. A reference annotation in Gene Transfer Format (GTF), e.g. provided by Ensembl (25) is required at this step. The final read counts can be used for the downstream assessment of differential expression using tools like DESeq2 (26). The current workflow can serve as a template that can be modified by the user according to different needs, for instance, replacement of tools or modification of the wrapping strategy.

IMPLEMENTATION

The workbench is implemented as portable virtualized container based on Galaxy. The Galaxy framework allows for reproducible and transparent scientific research which makes it easy to access, deploy and scale—conceptualized as a web service. The foundation of the workbench container is a generic Galaxy Docker instance (http:// bgruening.github.io/docker-galaxy-stable/). On-top of this, pre-configured Galaxy tools can be automatically installed from the Galaxy ToolShed using the Galaxy API BioBlend (27). In Galaxy, tool dependencies are automatically resolved via BioConda, which is the bioinformatics channel for the Conda package manager. Bio Conda facilitates software packaging and enables installation at a user level, keeping track of different versions of the same software in virtual environments. These features are in line with the scope of Galaxy; maintaining large numbers of dependencies in a reproducible way. Therefore, all available tools within the RNA workbench are also distributed as BioConda packages and BioContainers, which are persistent, frozen, containerized versions of *Conda* packages. The RNA workbench ships with a variety of tools, tours, documentation, workflows and data that have been added as additional layers on top of the generic Docker instance. During development, the software has been tested extensively in a continuous integration setup (CI) at different levels: Galaxy itself, tool integration in Galaxy (IUC, galaxytools channels), dependencies (BioConda) and at the workbench level. Together with a strict version management on all levels, this contributes to a high degree of error-control and reproducibility. The RNA workbench started in January 2015 - with constant development over 2 years, and extensive testing in local and public Galaxy instances, such as the Freiburg Galaxy instance, the MDC instance in Berlin and Erasmus MC's Galaxian. More than 500 users accessed the RNA tools during the last two years and the virtualized *Docker* instance was already downloaded >500 times. Moreover, due to an open and transparent development process, there is a growing community that contributes to our workbench, which guarantees the sustainability of the RNA workbench project and maintenance of the underlying Docker/rkt images.

USING THE RNA WORKBENCH

Installation: The RNA workbench can be installed under OSX and Windows using the graphical tool Kitematic (https://kitematic.com), or with the following Linux command:

docker run -d -p 8080:80 bgruening/galaxy-rnaworkbench

This installation is production-ready and can be configured to use external computer clusters or cloud environments. Due to the very modular system, it is also possible to install all or only a few tools of the RNA workbench on available Galaxy servers. Just get in contact with your local Galaxy administrator. When using the RNA workbench Docker image, the user has full administration rights, which enables customization independent of potential user restrictions.

Training

For self-empowering the user, documentation and training of the RNA workbench are important. We included an extensive set of documentation in traditional formats, e.g. tool descriptions and 'README' files.

We also provide training sessions around HTS data analyses and RNA-seq data analysis. The training materials ranging from the introduction to Galaxy, to usage and maintenance of Galaxy and the RNA workbench are freely accessible for self-paced studies at the Galaxyproject Github repository (http://galaxyproject.github.io/trainingmaterial). This training material is constantly improved and extended in an international community effort, including ELIXIR and EMBL. For HTS data analyses we provide training as a specific introduction to the topic with selfexplanatory presentation slides, a hands-on training documentation describing the analysis workflow, all necessary input files ready-to-use via Zenodo, a Galaxy Interactive Tour, and a tailor-made Galaxy Docker image for the corresponding data analysis.

To provide an even more intense training experience within the RNA workbench, we also included interactive training such as the Galaxy Interactive Tours. Such tours guide users through an entire analysis in an interactive

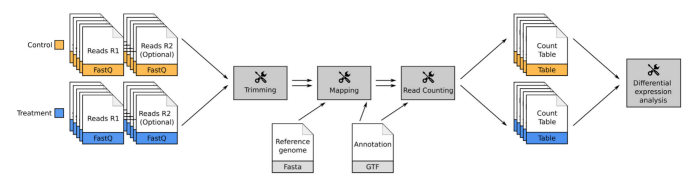


Figure 1. The workflow for analyzing RNA-seq data. The workflow tolerates single-end and paired-end reads derived from different conditions. It employs *TopHat2* for mapping and *HTSeq-count* to create the read counts. The final outputs contain read count per annotated gene for each condition and for each sequencing type.

and explorative way. It combines advantages from training videos and detailed protocols. Production of training videos is very time-consuming and tend to become outdated very soon, due to tool version changes or renewed workflows. In contrast to conventional screencasts, a Galaxy Interactive Tour can be easily updated and improved to guide the Galaxy user step-by-step, e.g. through a whole HTS analysis starting from uploading the data to using complex analysis tools. Exemplary, the RNA workbench currently integrates two Galaxy Interactive Tours. The first one introduces a new user to the Galaxy interface and its usage with an RNAseg example dataset. The second one illustrates secondary structure prediction of RNA molecules using parts of the ViennaRNA package. To show how Galaxy Interactive Tours can interactively guide users through the necessary steps of HTS analyses, the tours are also provided as online screencasts.

Visualization

Following data reduction as a key element of explorative research, there is a need for meaningful figures and visualizations that summarize results. The RNA workbench includes standard interactive plotting tools to draw bar charts and scatter plots from all kinds of tabular data and allows for connections to *Integrated Genome Browser* (29) and *UCSC* (30) like any other Galaxy instance. On top of this, we included three visualizations specific to RNA research. An interactive DotPlot visualization for secondary structures in EPS format (Figure 2b), a 2D visualization for the common dot-bracket format (Figure 2a) and a 3D visualization capable of visualizing PDB, SDF and MOL files containing three-dimensional coordinates (Figure 2c).

COMMUNITY

The RNA workbench project is an open source project that strives to create a community interested in accessible and reproducible RNA-related research. Knowing that real sustainability can only come true with a strong community we are aiming at more open participation, reward, and inclusion. We are working together with Galaxy, *BioConda*, *BioContainers* and *BioJS* and coordinating efforts to not reinvent the wheel but joining forces to create the new generation of bioinformatics infrastructure together. In the RNA

workbench community, we practice the organizations on GitHub, IRC, and Gitter and welcome everyone to contribute on every level to improve the entire stack from documentation to tools and scientific workflows. Support will be provided through the same channels.

DISCUSSION

In this work, we present the RNA workbench, maintained and developed by a constantly growing community. The presented workbench is unique as it allows to easily combine RNA-centric analysis with other types of experiments. It provides a set of tools, each one being available as *Bio-Conda* package as well as a Docker/rkt container (*BioContainers*). Based on the Galaxy *Docker* project, the proposed web server is more than the sum of its parts. It offers a comprehensive virtualized RNA workbench that can be deployed on every standard Linux, Windows and OSX computer, but can at the same time employ high-performance-or cloud-computing infrastructure.

Major advantages of our approach to deliver a dockerized workbench for RNA centric analysis are the ease of installation, the high number of pre-included tools, the flexibility in regard to extension with other tools and workflows and the high reproducibility and transparency of workflows. All tools that are available on the Galaxy *Toolshed* can be installed along with their automatically resolved dependencies with a single click in the Galaxy interface. Best practice pipelines for the analysis of RNA-seq data are provided with the *Docker* image and can easily be modified, extended or combined with other analysis pipelines via Galaxy's workflow editor GUI.

The RNA workbench was designed as a community project, and as such it is easy for users to contribute to the workbench with workflows, new tools and training material, keeping the workbench up-to-date and valuable for research. Moreover, all components such as tools, workflows, visualizations, interactive tours and training material can be easily integrated into any available Galaxy instance for teaching, learning or exploratory purposes.

The main difference to existing solutions such as miARma-Seq (1), RAP (2) and the UEA Small RNA Workbench (3) is that our RNA workbench combines the realm of RNA-centric analysis on sequence and structure level with modern high-throughput sequence analysis. In this re-

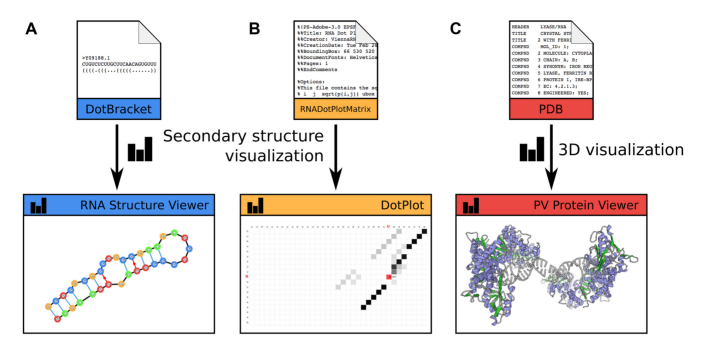


Figure 2. RNA structure visualization: The figure shows visualization for an IRE1 RNA sequence, retrieved from the Rfam database (28), via different backends integrated into the toolbox. (A) Secondary structure encoded in dot-bracket notation, can be displayed by the RNA structure viewer. (B) Base pairing probabilities are visualized as DotPlot. (C) Tertiary/Quaternary structure information encoded in protein-database format is rendered via Protein Viewer.

gard we provide well established tools for RNA structure prediction, analysis and visualization together with read mappers and expression analysis tools for HTS analysis.

ACKNOWLEDGEMENTS

We thank the de.NBI and ELIXIR projects for supporting bioinformatics infrastructure. Thanks also to the Galaxy community, especially to the Freiburg Galaxy Team, for developing, maintaining and supporting this great framework. We also like to acknowledge the BioConda and Bio-Containers community for setting new standards in reproducible software deployments. Thanks also to the BioJS community for great discussions about scientific visualizations and how we can make them more accessible. Moreover, the authors acknowledge the support of many upstream developers that helped us to integrate their tools into the RNA workbench and accepted patches.

FUNDING

Collaborative Research Center 992 Medical Epigenetics [DFG grant SFB 992/1 2012]; German Federal Ministry of Education and Research [BMBF grants 031 A538A/A538C RBC, 031L0101B/031L0101C de.NBI-epi, 031L0106 de.STAIR (de.NBI)]; Center for Translational Molecular Medicine (CTMM), TraIT project [05T-401 to Y.H.]. Funding for open access charge: German Govern-

Conflict of interest statement. None declared.

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