Extending MetAMOS - new methods and new integrations

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Biodiversity analysis of metagenomic and metatranscriptomic data acquired from next-generation sequencing (NGS) requires following multiple analytic steps, often independent from each other with exception of passing output files of previous step as input for the following. If parameterization of steps following one after another is independent from one another, they may be pipelined. There are three most popular pipelines used for NGS analyses: QIIME, mothur and MetAMOS. In this work we describe our extensions to the latter. One is supplementing MetAMOS' default modes with taxonomic and metabolic biodiversity using metagenomics and metatranscriptomics data and the other provides a web-based interface to run predefined analyses that is easy to integrate with laboratory information management systems.
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Abstract

Biodiversity analysis of metagenomic and metatranscriptomic data acquired from next-generation sequencing (NGS) requires following multiple analytic steps, often independent from each other with exception of passing output files of previous step as input for the following. If parameterization of steps following one after another is independent from one another, they may be pipelined. There are three most popular pipelines used for NGS analyses: QIIME, mothur and MetAMOS. In this work we describe our extensions to the last package. One extension is supplementing MetAMOS’ default modes with taxonomic and metabolic analyses on metagenomics and metatranscriptomics data and the other extension provides a web-based interface to run predefined analyses that is easy to integrate with laboratory information management systems.

Keywords: metagenomics, metatranscriptomics, NGS, pipeline, web service

Introduction

There is a number of standardized protocols available for NGS data dubbed Standard Operating Procedures (SOPs). However, it is not uncommon for researchers to perform own sets of analyses, depending on specific research topic needs. The most popular pipelines for NGS analyses are QIIME (Caporaso et al. 2010), mothur (Schloss et al. 2009), and MetAMOS (Schloss et al. 2009; Treangen et al. 2013). The first two are focused on biodiversity assessment while the latter is aiming at assembly. Proper choice, deployment and pipelining of tools used in such analyses is a task non-trivial even for an experienced bioinformatician, and may pose a big problem for non-specialists.

The most easy to extended is definitely QIIME, as is it already a set of Python scripts. However its installation is so challenging, that the installation method recommended by QIIME authors is through VirtualBox or to use cloud computing instances with QIIME pre-installed. Mothur, being extremely portable due implementation in binary files, is hard to extend. In our case,
requiring work with multiple instances of the software in a dynamic hardware environment, we set out to work with MetAMOS. MetAMOS focus - namely creation of automated, reproducible, assembly & analysis pipeline was in full alignment with our project. Moreover, it enabled fast and efficient implementation of extension of functionalities, developed by our team. This work presents an example of the solution to an effective cooperation between computational and experimental biologists, merging complex analysis pipeline with user-friendly interface through combination of MetAMOS with custom pipeline and web-interface.

**Custom pipeline**

Synthesis of research needs and technical capabilities of the research infrastructure required development of in-house tool called bipype (Fig 1). “bipype” stands for bioinformatics-python-pipe.
Fig. 1. Chart with bipype functionalities

Bipype accepts three types of inputs: amplicons, WGS (whole genome sequences) and metatranscriptomic data. Bipype may work with paired-end and single read sequences. For amplicon (prokaryotic or fungal) data, if needed, paired-end reads are merged and sequence reconstruction is performed. Reconstructed 16S or ITS sequences are searched in proper reference databases and hits to related taxonomic units are counted.

For WGS reads three paths are available. In one they will either be used for reconstruction of contigs, which will be further used for reference database search (outside of the pipeline). In the
second they will be used directly in taxonomic diversity search, in third they will be compared to
sequences related to metabolic pathways.

The biodiversity data at taxonomical and functional levels are similar in their tree structure - we
provide display of results in a common form of html interactive comparative (multiple samples
presented in single file) tree (Fig. 3c.). Bipype is a portable solution with a few dependencies, but
to simplify analyses for the end-users, GUI presenting available analysis pathways and
(optional!!) parameters of each step was required. Below we present MetAMOS integration
process, meeting this requirement. Bipype source code is available at:

https://github.com/krassowski/bipype/

Integration with MetAMOS

As the number of customisable steps in MetAMOS is limited, we needed to overcome certain
inflexibility of this system by inserting the whole bipype run in the first possible step and
terminating the workflow afterwards. In order to do so, we added bipype to the list of tools used
by metAMOS as a fake assembler. We added a script that created empty files required by
metAMOS when it is checking if the assembly step finished successfully. Sample configuration
file for such fake assembler is shown in Fig. 2. Command line options for metAMOS allow
skipping most of the steps in any workflow, making it possible to finish immediately after
assembly.

```
[CONFIG]
name bipype_amplicons
input FASTQ
scaffoldOutput [RUNDIR]/Scaffolds.fasta
location python/bipype
output 16S_ITS.krona
threads --threads
unpaired [FIRST]
paired_interleaved [FIRST] [SECOND]
paired [FIRST] [SECOND]
commands mkdir [RUNDIR] && 
    bipype --out_dir [RUNDIR] --cutadapt use_paths both --mode run -ITS -16S -ot ITS
16S --input [INPUT] & & \n    bipype_cheat [RUNDIR]
```

Fig. 2. Sample configuration file
Web interface to MetAMOS

We have developed a web interface to analyses run on MetAMOS to let others not only access results but to run preconfigured analyses on their own. Fig. 3a shows the webpage where user can choose the type of analysis and library on which it would be run. Each type of analysis corresponds to one configuration file with appropriate bipype options. Status of analyses for each sample and workflow is stored in an SQL database. When user selects a workflow and a sample, web service checks if the corresponding job is finished and either shows the result (Fig. 3c), or its computation progress (Fig. 3b). If the job is not even started, the web service adds it to the database. A background service script periodically queries the database and runs queued jobs.

Additionally, we have added a simple results management, that is possibility to remove results in case re-computation is needed.

Fig. 3a. Screenshot of the interface: sample and workflow choice
Fig 3b. screenshot of the interface: job progress

Fig 3c. screenshot of the interface: job results

Web interface source code is available at:

https://bitbucket.org/Serpens/metamos_web_interface/src
Materials and Methods

bipype is written in Python v2.7.3 and links following tools: fastx-toolkit v0.0.13 (“FASTX-Toolkit” 2016), usearch v7.0.959_i86linux32 (Edgar 2010), FLASH v1.2.7 (Magoč and Salzberg 2011), bowtie2 v2.2.4 (Langmead et al. 2009), samtools v0.1.18 (Li et al. 2009), RAPsearch 2.12_64bits (Zhao, Tang, and Ye 2012), MetaVelvet v1.2.01 (Namiki et al. 2012), HUMAnN v0.99 (Abubucker et al. 2012), SARTools v1.2.0 (Varet et al. 2015), KRONA 2.0 (Ondov, Bergman, and Phillippy 2013). Presented workflow was performed on Illumina reads with varying insert lengths, either provided in form of parameter or read from filename.

Databases used as reference include: SILVA rRNA database (Griffith, Malachi, and Griffith 2004; Quast et al. 2013), Unified system for the DNA based fungal species linked to the classification (UNITE) (Kõljalg et al. 2013), The Reference Sequence (RefSeq) Database (Griffith, Malachi, and Griffith 2004), NCBI Taxonomy Database (Wheeler 2004), Kyoto Encyclopaedia of Genes and Genomes database (Kanehisa et al. 2014),

Web interface is build using MetAMOS v.1.5rc3, Python v2.7.3, Django v1.4.5, jQuery v1.11 and Bootstrap v3.1.1. It was tested with SQLite v3.7.13 and Apache v2.2.22.

Conclusions

We present a way to develop time and resource efficient customizable pipeline serving for metagenomic and metatranscriptomic analyses, by extending MetAMOS workflow engine. The proposed solution, besides abovementioned efficiencies presents following benefits:

- modularity allowing insertion of more custom tools and analyses (use of different assembly and display tools, search engines etc.),
- user-friendly web interface enabling easy access and steep learning curve for new team members, also enabling quick and repeated complete analyses by personnel not focused on software development,
- unified display methodology for both types of diversity data.

Our extensions are obviously available as open source under GNU GPLv2 license, allowing other researchers to build upon our work.
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References


