TRUFA: User's manual

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1 What is this web server for ?

"RNA-seq, also called whole-transcriptome shotgun sequencing, refers to the use of high-throughput sequencing technologies for characterizing the RNA content and composition of a given sample" (Wolf, 2013). Until now, analyzing RNA-seq data remains a Bioinformatic challenge. This web server is designed to make analysis of RNA-seq data in a fast and user-friendly manner, by using cluster computing and reducing the amount of Bioinformatic knowledge necessary.

1.1 Quick background on RNA-seq analysis

Recently and especially with the development of new massively parallel sequencing method, the cost of sequencing whole transcriptomes fell drastically. With a thousand dollars, you can get Gigas worth of transcriptome sequences.

In the case of Illumina sequencing, you will receive as output 1 (single-end) or 2 (paired-end) files per samples in fastq format (.fastq, .fq or compressed fastq such as fq.tar.gz, fq.gz ...).

In order to get high throughput, the original RNA sequences have been shredded before sequencing and so your fastq files are filled with millions to billions of sequences, between 50 and 250 bp long each depending on the Illumina sequencing technology. These sequences are called "reads".

The first step in the RNA-seq analysis will be to clean the reads. Various cleanings can be performed, such as removing duplicates, adapters, poor quality bases/reads, putative contaminants etc...

The second step of the analysis is to reassemble the shredded pieces, the reads, into contigs (named transcripts in the case of RNA-seq) in order to reconstruct the original RNAs. To do so, two main methods are available:

- With a reference genome available, you will use "mapping" methods.
- Without a reference genome: you will use "de novo assembly" methods.

So far, TRUFA can only perform *de novo* assembly using the Trinity software. But more is to come ...

Once you obtained the assembly, the next step is to identify those contigs. This is particularly interesting if you are looking for specific genes. For example, if you are interested by venom toxins you could download a venom protein sequences database and then blast your assembly against it. You'll get hits, representing mRNA which could represent precursors of venom proteins. Another way to identify sequences is to scan your assembly for protein profiles (using HMMER for example).

Recently, a specific vocabulary called "Gene Ontology" has been developed in order to help in the identification process. The idea is to link biologically meaningful keywords to nucleotide sequences to identify their biological role, molecular function and location in cellular components. This can be performed using Blast2GO.

Finally, the last main step of an RNA-seq analysis will be to quantify the expression. One of the interest of having high throughput data is that the number of reads sequenced per contigs should reflect the expression levels of the corresponding gene. To quantify the expression, the first step will be to align all reads back to the transcripts (using BOWTIE for example). Then, programs such as eXpress can provide you with expression values (called RPKM in the case of single-end reads and FPKM in the case of paired-end reads).

For more basics on RNA-seq

Wolf (2013) Martin and Wang (2011)

1.2 What can be done with TRUFA?

So far, TRUFA is allowing you to perform the following steps (programs used are specified in the parentheses):

- Reads cleaning:
 - Quality control (FastQC)
 - Quality trimming and duplicates removal (Prinseq)
 - Trimming adapters (Cutadapt)
 - Filtering out potential contaminants (Blat)
- De novo assembly of your reads (Trinity)
- Reads mapping (Bowtie and Bowtie2)
- Contigs (i.e transcripts) identification:
 - based on sequence alignment (Blat, Blast)
 - based on protein profiles (HMMER)
 - Annotation with GO terms (Blast against nr and Blast2GO)
- Expression quantification:
 - providing RPKMs, FPKMs (eXpress, RSEM)

All the steps of the pipeline can be run as a whole or independently, depending on your needs. In the case of making a complete pipeline, your reads files will be cleaned, then assembled into transcripts and finally the transcripts will be identified and quantified.

2 Input:

The input files can be of multiple sources, depending on the kind of analysis you want to perform and the data you have available. So far the accepted inputs are:

• 1 or 2 Illumina reads files (i.e. single-end or paired-end reads) in fastq or compressed (extensions .tar.gz or .gz) fastq format

• 1 file with already assembled reads, i.e contigs in fasta format



Mark Important!

On top of the necessary inputs, you will need to specify some important information related to the library construction:

- Size of the insert (for Trinity and Bowtie)
- Adapters sequences (for Cutadapt)
- Strand specific reads? (for Trinity and Bowtie)

3 **Output:**

So far, TRUFA is providing the classical outputs of a RNA-seq analysis. For a complete analysis, you will get (type of file precised between parenthesis):

- Prinseq report of reads quality (html)
- FastQC report of reads quality (zip and html)
- Reads files after complete cleaning (fastq)
- Assembly file (fasta)
- File with alignment of the reads against the transcripts (bam)
- Results of the Blast2GO annotation (dat)
- Transcript expression quantification (txt)

Other useful programs: 4

Part of Trufa's output can be visualized with programs such as:

- Blast2GO http://www.blast2go.com/b2ghome
- IGV http://www.broadinstitute.org/igv/

Quick Start:

- Tablet http://bioinf.scri.ac.uk/tablet/
- RNAseq Viewer http://bioinfo.au.tsinghua.edu.cn/software/RNAseqViewer/

Other programs for RNA-seq analysis can be found at:

- http://omictools.com/
- Wikipedia: RNA-seq tools

5 Quick Start:

5.1 Launch a Job

This is a quick demonstration on how the web server can be used.

5.1.1 Upload demo files



- In the Upload area, **Browse** your computer for the reads file "reads_left.fq.tar.gz"
- Before clicking **Send**, select **compressed reads file** in the drop-down menu (instead of **undefined**)
- Once successfully uploaded, do the same for the reads file "reads_right.fq.tar.gz"
- Once both files have been uploaded, you are ready to "start a job".

Specifying the file type in the dropdown menu is mandatory and will help you by filtering the correct inputs for each steps of the analysis.

5.1.2 Start the job



- Go to Start a Job
- Select Paired-end reads (2 fastq files)
- Specify the 2 input files: "reads_left.fq.tar.gz" for **left reads** and "reads_right.fq.tar.gz" for **right reads**
- Go down to **RNA-seq steps** section and to the **Reads cleaning** tab.
- Check the boxes **Duplicated reads** and **Quality trimming** in the Prinseq section
- Check the box FastQC in the Post-cleaning quality control section
- Then go to the Reads Assembly and Mapping tab
- Check the boxes Assemble with Trinity and Align reads against contigs with Bowtie2
- Then go to the Contigs identification tab
- Go down to Blast2GO searches section
- Check the box Blast2GO
- Then go to the Expression quantification tab
- Check **eXpress**
- Then go down to **Launching Analysis** and hit the **Start** button.

A message confirming that the Job has been sent should appear. You can go back to **Home** in the main menu and here click on the corresponding Job in the Job list. You should be able to see the current state of your job there.

Quick Start:

5.2 Check output

Once your job has been completed, here are some ways to check the generated outputs.



- Go to File Manager in the main menu
- Put your username and password
- Browse the **Jobs** folder for the corresponding Job name

5.2.1 Check reads quality:



- Go to **STAT** and then **prinseq_report**

Here you should have 4 html files (2 for each reads file, one report after the removal of the duplicates and one report after the trimming process) that you can download and then open locally with your internet browser. Information about Prinseq statistics can be found in the "quality control" section of Prinseq manual:

http://prinseq.sourceforge.net/manual.html #QC



- Go to STAT and then fastqc_report

Here you should have 2 zip files (1 for each reads file). Download them, extract them and open locally the html files with your internet browser. Information about the FastQC output can be found here:

http://www.bioinformatics.babraham.ac.uk/projects/fastqc/

5.2.2 Check the assembly:



- Go to ASSEMBLY_MAPPING and then trinity

Here you will find the assembly file **Trinity.fasta**

Quick Start:



- Go to STAT and then assembly_qc

Here you will find text files displaying statistics related to your assembly which will help you to determine the quality of your assembly.

5.2.3 Check reads mapping against transcripts



- Go to ASSEMBLY_MAPPING and then bowtie2
- Download the file aligned_reads.bam and aligned_reads.bam.bai

You can use these two files to visualize the alignments using programs such as Tablet http://bioinf.scri.ac.uk/tablet/.

5.2.4 Check the Blast2GO results:



- Go to IDENTIFICATION and then b2go
- Download the file out_b2go.dat
- Open it by starting Blast2GO program locally and going to File, Load B2GO-Project (.dat)

Blast2GO can be obtained from http://www.blast2go.com/b2ghome

5.2.5 Check the eXpress results:



- Go to EXPRESSION and then express

Here you will find two text files ("params.xprs" and "results.xprs"). The "results.xprs" file is giving the expression quantification for all isoforms produced by Trinity. More information about eXpress outputs can be found at http://bio.math.berkeley.edu/eXpress/manual.html

6 Running an analysis:

This explains in more details each parts which can be found in the "Start a Job" page of the web server. This manual is far from pretending to replace the manual of each program which are implemented in TRUFA. Consequently, the manuals of each of these programs should be studied by the user in order to tune them and understand their output properly.

6.1 Type of input:

Essentially, all the steps of the RNA-seq pipeline on the web server can be realized with only reads files (1 or 2) as input. For the purpose of testing and parameters tuning, later steps can be as well directly performed with an assembly (fasta file with already assembled reads).

First, you should specify and upload the necessary input files:

- For Reads cleaning: You will need one or two fastq files (i.e. single or paired-end reads files)
- For Assembly: You will need one or two fastq files (i.e. single or pairedend reads files)
- For Identification, you will need either: a fasta file with the reads already assembled OR 1 or 2 fastq reads files and generate an assembly.
- For Reads mapping: 1 fasta file with the reads already assembled and the corresponding reads in 1 or 2 fastq files OR 1 or 2 fastq reads files and input or generate an assembly.
- For Expression quantification: 1 fasta file with the reads already assembled and the corresponding reads in 1 or 2 fastq files OR 1 or 2 fastq reads files and input or generate an assembly (see How to generate an assembly).

Then, by specifying in the **Run a Job** part the type of input and the input files, different possibilities will be available to you, depending on the type and number of input files you specified.

6.2 Cleaning step:

In this well, you will find programs to perform reads cleaning prior to the assembly.

- FastQC: Will give an idea on how to clean your reads. Are your reads of poor quality? Got hexamer priming biais? Got contaminants like adapters sequences in my reads?

 More info on FASTQC homepage
- Cutadapt: Allows you to remove the adapter sequences you might have found during the FASTQC quality control. Adapter sequences must be specified further down in the "Cutadapt options".

 More on Cutadapt
- Prinseq: Performs duplicate removal and reads trimming. Lot's of options can be tuned further down the "Start a Job" page in "Duplication options" and "Trimming options". At this stage, removing duplicates is mainly to gain computation time during the assembly process. For expression quantification purposes, it seems more recommended to mark duplicates using programs such as Picard.

 More info on Prinseq homepage
- Blat: performs homology searches but much faster than Blast and can be therefore used to search homologies between large set of reads and a database, in order to remove potential contaminants before assembly. Custom databases of potential contaminants can be added by Uploading them as "nucleotide seqs database for blasting" in fasta format. Then the corresponding database can be checked in the "Nucleotide db" button in the "Blast against potential contaminants" area. Blat parameters cannot be tuned on TRUFA yet and is used with default parameters. Custom scripts are automatically removing for the next steps of the analysis the reads with at least one blat hit for any of the selected database

 More info on Blat

6.3 Assembly and Mapping step:

In this well, the programs will perform assembly and/or mapping of the reads. If Cutadapt, Prinseq or Blat actions have been checked in the "Cleaning

step" well, then the output of this cleaning step will be used as input for the assembly and mapping steps. If no cleaning options have been selected, raw reads will be used for the assembly/mapping.

- Trinity: performs de novo assembly of the reads. Several options are important to tune, such as the minimum contig length and the strand specificity if required. This can be tuned in the "Trinity options". More info on Trinity
- Assembly quality checks: provide various statistics helping to judge the quality of the assembly. This includes the CEGMA program, blast searches and Trinity scripts and custom scripts for statistics and plots.
- Bowtie2: align the reads back to the transcripts. on top of being necessary for the expression quantification, this mapping and the amount of reads which align back are an indicator of the quality of the assembly. It's important to verify that the insert size of your library is comprise between the minimum and the maximum insert size in "Bowtie2 options".

More info on Bowtie2

6.4 Identification step:

This part of the pipeline aims at annotating the transcripts newly assembled and link nucleotide sequences to a particular biological function.

• Blat: Again, Blat is here to perform fast homology searches against classical databases such as NCBI nr or Uniref90. Blat parameters can not be tuned yet and are set to default. Custom databases can be added by Uploading them as "nucleotide (or protein) seqs database for blasting" in fasta format. Then the corresponding database can be checked in the respective button ("nucleotide" or "protein" db) in the "Custom Blat searches" area.

More info on Blat

• HMMER: performs as well homology searches but is able to detect more remote homologs with HMM protein profiles. The HMMER search can be performed on the whole PFAMA database or against one/multiple custom hmm profiles previously generated (TRUFA does not generate HMM profile files) and **Uploaded** as "**protein hmm profile for HMMER** in .hmm format.

More info on HMMER

• Blast+ against nr and B2GO: To get the GO terms annotations for each transcripts, B2GO first needs the results of a blast search against the NCBI nr database. The Blast+ job is the most computation intensive from TRUFA and can in certain cases stop before the end of the search is reached. If you notice that B2GO is not producing any output, please contact us, this could be the issue. More info on Blast+ More info on Blast2GO

6.5 Expression quantification step:

This step allows the user to produce classical expression quantification outputs with for example RPKM or FPKM values useful to further test for differential expression (not yet available on TRUFA).

6.6 Visualizing the output:

6.6.1 Cleaning outputs:

6.6.2 Assembly-Mapping outputs:

Assembly and reads alignment can be visualized using IGV by importing the assembly file (Trinity.fas) as a genome and loading the corresponding bam file(s).

6.6.3 Identification outputs:

6.6.4 Expression outputs:

7 Acknowledgments

References

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