RUM 2 (RNA-Seq Unified Mapper)

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Agenda

- Intro
- 2 The RUM pipeline
- Tour of the new RUM
 - Installation
 - Command-line interface
 - Job state management
 - Work distribution
- Demo
 - Web resources
 - Run tiny job

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Who am I?

- Working at Penn (ITMAT) since January
- Software engineering background
- Experience in a variety of languages, (Perl, Java, Clojure (lisp), Python, Ruby)
- ... and applications

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What is RUM?

- "RUM is an alignment, junction calling, and feature quantification pipeline specifically designed for Illumina RNA-Seq data"
- Written by Gregory Grant (ggrant@grant.org)
- Runs on Linux / UNIX / Mac
- Distributed (optionally)

What is RUM?

Inputs

- RNA-Seq reads: FASTA or FASTQ, paired or single
- Organism-specific index

What is RUM?

Inputs

 RNA-Seq reads: FASTA or FASTQ, paired or single

Outputs

Unique and non-unique alignments

Organism-specific index

- Coverage plots
- Feature quantifications
- Junction calls
- List of novel inferred internal exons



RUM has three distinct phases:

- Preprocessing
- Processing
- Postprocessing

Phase 1: Preprocessing



- Perform some quality checks on reads
- Split reads into N chunks
- Allows reads to be processed by N nodes on a cluster

Phase 2: Processing

- Align all reads against genome using Bowtie
- Align all reads against transcriptome using Bowtie
- Merge genome and transcriptome alignments and identify unmapped reads
- Align unmapped reads against genome using BLAT
- "This leverages the advantages of both genome and transcriptome mapping as well as combining the speed of Bowtie with the sensitivity and flexibility of Blat."
- Merge Bowtie and Blat alignments

Phase 3: Postprocessing

- Merge alignments for all chunks together
- Produce coverage plots, junction files
- Find novel internal exons
- Generate some reports



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RUM 2 Goals

- More code reuse for maintainability
- Improve inter-process communication
- More robust job state management
- Automated tests

Enhancements in RUM 2

- Standard installation process
- New command-line interface
- Get status of running job
- Restart a job where it left off
- More reliable kill command
- Run a chunk or postprocessing by itself
- Relocatable indexes
- SAM file is closer to conforming to standard

Installing RUM

- Uses standard Perl Makefile.PL
- Should be familiar to system administrators
- Download tarball from https://github.com/PGFI/rum/downloads
- perl Makefile.PL
- make install (optional)
- Then install indexes...

Installing Indexes

- RUM needs an index for each organism you want to align against
- Index includes genome, gene annotations, and binary index files for Bowtie
- Index installation is now separate from code installation / upgrade
- Run rum_indexes; it will guide you through the process

Command-line interface

Usage is rum_runner ACTION [OPTIONS] where action is one of:

- align Run an alignment
- status Check the status of a job
- stop Stop a job (can be restarted later)
- kill Stop a job and clean it up (to restart from scratch)
- clean Remove output files for a job
- help Get help
- version Show version number

Running an alignment

Use "rum_runner align" to run an alignment:

```
rum_runner align \
  --output ~/sample123/results \
  --index ~/rum_indexes/hg19 \
  --name TestJob \
  --chunks 25 \
  ~/sample123/forward.fq ~/sample123/reverse.fq
```

Job status

- Use rum_runner status to check on the status of a running job.
 - \$ rum_runner status -o ~/sample123/results

Processing in 25 chunks

XX XX XXX

XX XX XXX

XX XX

XX XX XXX

XX XX XXX

XX XX XXX

XXX

X X

X X

X X

X X X X

X X

```
XX XX XXXX XXXXX
                    Run bowtie on transcriptome
     XX XX XXXX XXXXX Parse transcriptome Bowtie output
X XXX
     XX XX XXXX XXXXX Merge unique mappers together
X XXX
X XXX
     XX XX XXXX
               XXXXX Merge non-unique mappers together
X XXX
     XX XX XXXX XXXXX Make unmapped reads file for blat
X XXX
     XX XX XXXX XXXXX Run blat on unmapped reads
     XX XX XXXX XXXXX Run mdust on unmapped reads
X XXX
     XX XX XXXX XXXXX Parse blat output
X XXX
     XX XX XXXX
               XXXXX Merge bowtie and blat results
X XXX
X XX
     XX XX XXX
                XXXX Clean up RUM files
X XX
     XX XX
           XXX
                XXXX Produce RUM_Unique
X XX
     XX XX
          XXX
                XXXX Sort RUM Unique by location
```

XXXX Create SAM file

XXXX Generate quants

Sort RUM_NU

XXXX Create non-unique stats

XXXX

XXXX

XXXX Sort cleaned non-unique mappers by ID

Remove duplicates from NU

Job status

```
Postprocessing
-----
X Merge RUM NU files
X Make non-unique coverage
X Merge RUM_Unique files
X Compute mapping statistics
X Make unique coverage
X Finish mapping stats
X Merge SAM headers
X Concatenate SAM files
X Merge quants
 make_junctions
 Sort junctions (all, bed) by location
 Sort junctions (all, rum) by location
 Sort junctions (high-quality, bed) by location
 Get inferred internal exons
 Quantify novel exons
All the chunk error log files are empty. That's good.
Main error log file is empty. That's good.
RUM is running (job ids 815718, 815720).
```

Job state management

- Model the workflow as a state machine
- A completed step transitions the job from one state to another
- Stitch steps together into a workflow
- Determine the state of a job by looking at which output files exist
- Similar to GNU Make (but simpler)
- Basis for a lot of additional features

Recovering from errors



- In case of failure...
- RUM 2 allows easier recovery
- Running "rum_runner align" again, RUM will determine what state the job was in when it failed
- Just resumes at the next uncompleted step
- Can save a lot of time when recovering from infrastructure failure

Killing a job

To stop a job and remove all of its output:

```
rum_runner kill -o dir
```

 Useful if you've run a job with incorrect parameters and need to start over



- Automatic support for one multi-core machine (will run each chunk in a separate process by default)
- Built-in support for Sun Grid Engine, with --qsub option
- Easily extensible for other platforms

For other platforms, you can either extend a Perl class to provide support, or set up some scripts to run the parts of the job on different machines:

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Run preprocessing alone:

```
rum_runner -o <dir>> --preprocess
```

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Run preprocessing alone:

```
rum_runner -o <dir>> --preprocess
```

Run chunks one at a time:

```
rum_runner -o <dir> --process --chunk 8
```

Run postprocessing alone:

```
rum_runner -o <dir>> --postprocess
```

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Web resources

```
Main github page https://github.com/PGFI/rum

User guide https://github.com/PGFI/rum/wiki

Issues https://github.com/PGFI/rum/issues

Downloads https://github.com/PGFI/rum/downloads
```

To the command line...