

# HiC-Bench Manual

# Stephen M. Kelly <sup>1,2</sup>, Charalampos Lazaris <sup>3,4,5</sup>, Aristotelis Tsirigos <sup>1,2,3,4,5</sup>

- <sup>1</sup> Applied Bioinformatics Center, Office of Collaborative Science, NYU School of Medicine, NY 10016, USA
- <sup>2</sup> Genome Technology Center, Office of Collaborative Science, NYU School of Medicine, NY 10016, USA
  - <sup>3</sup> Department of Pathology, NYU School of Medicine, New York, New York 10016, USA

# April 6, 2016

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<sup>&</sup>lt;sup>4</sup> NYU Cancer Institute and Helen L. and Martin S. Kimmel Center for Stem Cell Biology, NYU School of Medicine, New York, New York 10016, USA
<sup>5</sup> Center for Health Informatics & Bioinformatics, NYU School of Medicine, NY 10016, USA

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## 1 Introduction

#### 1.1 Installation

The analysis pipeline can be installed by cloning its git repository from GitHub, located here: https://github.com/NYU-BFX/hic-bench. In the Terminal (OS X, Linux), run a command such as the following:

```
git clone https://github.com/NYU-BFX/hic-bench.git
```

Once a clone of the pipeline repository has been made, it will be used as a blank template to start future analysis; analysis is not performed directly in the pipeline repository.

### 1.2 Compile Binaries

Source code for needed binaries has been included in the repository, and must be compiled. Navigate to the code/src directory from within the Terminal, and run the command make to automatically run the compilation scripts needed. The program bedGraphToBigWig is also required, and available as a binary file from UCSC at their page here: http://hgdownload.cse.ucsc.edu/admin/exe/. To directly install a version compatible with the Linux operating system, navigate to the code/bin directory and run the command wgethttp://hgdownload.cse.ucsc.edu/admin/exe/linux.x86\_64/bedGraphToBigWig.

### 1.3 Setting up a new analysis

Assuming your pipeline repository clone exists at ~/hic-bench, use the following terminal command to create a new analysis:

```
~/hic-bench/code/code.main/pipeline-new-analysis hicseq-standard <project_name>
```

This will create a new directory at the given location, and copy into it all the basic files and sub-directories needed for analysis from the pipeline repository.

# 1.4 Setting input files

Manual setup for the pipeline input files requires the creation of the directories ct\_name\pipeline\input\fastq or ct\_name\pipeline\input\bam, corresponding to the type of input files to be used. Sub-directories within these should be created with the name of each sample to be included in the analysis. A naming scheme similar to the following is suggested:

Importantly, the '-' should be used as a delimiter, since this is recognized by the sample sheet creation script. Within each sub-directory, place all fastq / fastq.gz or bam files for the sample. Symlinks can be used if the files are not contained in the same location as the project analysis directory, and are preferable in order to save storage space. Since this part of the pipeline setup is custom for each analysis, it must be completed manually. A script used to automatically create the correct directories and symlinks might look like this:

```
#!/bin/bash
Fastq_dir="/data/sequence/results/smithlab/2016-01-28/fastq"
Inputs_dir="/home/$(whoami)/projects/SmithLab_HiC_2016-02-09/inputs/fastq"

# make inputs dir
mkdir -p "$Inputs_dir"

for i in $Fastq_dir/*.fastq.gz; do
echo "$i"

TMP_NAME=$(echo "$(basename "$i")" |sed -nr 's/^([[:digit:]]) [^[:alnum:]]([[:alpha:]]+) [^[:alnum:]].*$/THP1
-\2-\1/p')
echo "$TMP_NAME"
mkdir -p "$Inputs_dir/$TMP_NAME"
In -s "$i" "$Inputs_dir/$TMP_NAME"
done
```

# 1.5 Create project sample sheet

A sample sheet must be created for the analysis project. After the inputs directory has been set up, the follow command can be used to automatically create a sample sheet template:

```
inputs$ ./code/create-sample-sheet.tcsh <genome> <fragment-size>
```

Where genome is hg19, hg38, etc.. The fragment-size entry is optional and should be a numeric argument such as 300, representing the library size of the sequencing sample. After creation of the sample sheet (sample-sheet.tsv), a manual review process is required to match the correct control or input samples with experimental samples, verify proper grouping names, files, and other entries. If not entered prior, fragment-size should be filled in for each sample. This process can be completed within Microsoft Excel, but saving the file in Excel should be avoided due to the introduction of invisible formatting errors by Microsoft Office products. It is advisable to instead copy the finalized sheet from Excel and paste directly into a terminal text editor such as vi or nano for saving under the file name sample-sheet.tsv.

# 1.6 Running the Pipeline

After navigating to the parent directory of the analysis project, run the pipeline with:

```
./code.main/pipeline-execute PROJECT-NAME E-MAIL
```

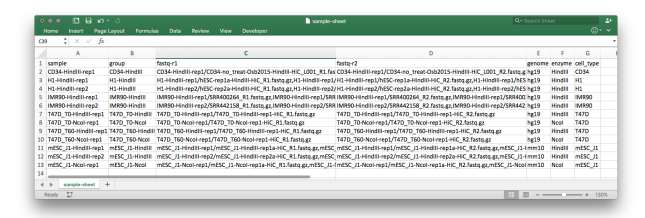


Figure 1: Example sample sheet

# 1.7 Dependencies

This pipeline was developed for use in a High Performance Computing environment, running CentOS 6. Additionally, tcsh and bash shells are required, along with R version 3.2.0. The following includes software used in the HiC-Seq pipeline:

```
OGS/Grid Engine 2011.11
Linux 2.6.32-573.3.1.el6.x86_64 #1 SMP Thu Aug 13 22:55:16 UTC 2015 x86_64 GNU/Linux
tcsh 6.17.00 (Astron) 2009-07-10 (x86_64-unknown-linux)
GNU bash, version 4.1.2(1)-release (x86_64-redhat-linux-gnu)
armatus/2014-05-19
bedtools/2.22.0
bowtie2/2.2.6
caltads/0.1.0
ghmm/0.9
java/1.7
matlab/R2013a
picard-tools
python/2.7.3
r/3.2.0
r/3.2.3
samtools/1.2.1
```

The following R packages are used in the pipeline:

```
plyr 1.8.1

vennDiagram 1.6.16
flsa 1.05
genlasso 1.3
ggplot2 1.0.1
optparse 1.3.0
pastecs 1.3-18
plotrix 3.5-11
reshape2 1.4.1
zoo 1.7-12
preprocessCore 1.24.0
MASS 7.3-35
gplots 2.17.0
reshape 0.8.5
corrplot 0.73
RColorBrewer 1.1-2
lattice 0.20-33
grid
stringr 1.0.0
```

For software information specific to the creation of this document, see Section 5.4

# 2 Default Pipeline Components

## 2.1 Parent Directory Overview

A default pipeline will have the following basic structure within its parent directory:

```
\verb|hicseq.analysis-for-hicbench||
              at570
                     14 Feb 14 19:28
                                       _01a-align -> pipeline/align
Irwxrwxrwx
                                       02a-filter -> pipeline/filter
              at570
                     15 Feb 14 19:28
Irwxrwxrwx
                                       at570
                     21 Feb 14 19:28
Irwxrwxrwx
                     15 Feb 14 19:28
                                       _03a—tracks —> pipelinė/tracks
              at570
Irwxrwxrwx
                                       __04a-matrix-filtered -> pipeline/matrix-filtered
              at570
Irwxrwxrwx
                     24 Feb 14 19:28
              at570
                     20 Feb 14 19:28
Irwxrwxrwx
                                       _05a-matrix-prep -> pipeline/matrix-prep
              at570
                     18 Feb 14 19:28
Irwxrwxrwx
                                       _06a-matrix-ic -> pipeline/matrix-ic
Irwxrwxrwx
              at570
                     23 Feb 14 19:28
                                       _07a-matrix-hicnorm -> pipeline/matrix-hicnorm
Irwxrwxrwx
              at570
                     21 Feb 14 19:28
                                       _08a-matrix-stats -> pipeline/matrix-stats
              at570
                     25 Feb 14 19:28
Irwxrwxrwx
                                       _09a-compare-matrices -> pipeline/compare-matrices
              at570
at570
Irwxrwxrwx
                     31 Feb 14 19:28
                                       _09b-compare-matrices-stats -> pipeline/compare-matrices-stats
                                       _10a-boundary-scores -> pipeline/boundary-scores
                     24 Feb 14 19:28
Irwxrwxrwx
                                     __10b-boundary-scores-pca -> pipeline/boundary-scores-pca
Irwxrwxrwx
              at570
                     28 Feb 14 19:28
Irwxrwxrwx
              at570
                     16 Feb 14 19:28
                                       _11a-domains -> pipeline/domains
                                     __12a-compare-boundaries -> pipeline/compare-boundaries
Irwxrwxrwx
              at570
                     27 Feb 14 19:28
Irwxrwxrwx
              at570
                     33 Feb 14 19:28
                                      _12b-compare-boundaries-stats -> pipeline/compare-boundaries-stats
                                     __13a-hicplotter -> pipeline/hicplotter
Irwxrwxrwx\\
              at570
                     19 Feb 14 19:28
Irwxrwxrwx
              at570
                     21 Feb 14 19:28
                                      __14a-interactions -> pipeline/interactions
Irwxrwxrwx
              at570
                     20 Feb 14 19:28
                                     __15a-annotations -> pipeline/annotations
Irwxrwxrwx
              at570
                     26 Feb 14 19:28
                                      _15b-annotations-stats -> pipeline/annotations-stats
              at570
                     30 Feb 18 11:16 code -> code.repo/code.hicseq-standard
Irwxrwxrwx
              at570
                     14 Nov 12 11:55 code.main -> code/code.main
Irwxrwxrwx
drwxr-xr-x 10 at570 238 Feb 15 19:53 code.repo
Irwxrwxrwx
              at570
                    36 Mar 10 16:30 data -> /ifs/home/at570/pipeline-master/data
drwxr-xr-x
            5 at 570 230 Jan 5 09:24 inputs
drwxr-xr-x 25 at570 834 Feb 14 19:28 pipeline
           1 at570 981 Jan 5 19:40 run
-rwxr-xr-x
            1 at570 554 Dec 18 17:02 run.dry
```

The following components can be seen here:

- \_\_01a-align ... \_\_15b-annotations-stats: Symlinks to each step in the pipeline, in alphanumeric order of execution.
- code: Symlink to the directory containing scripts and code specific to the current analysis type e.g. ChIP-Seq.
- code.main: Symlink to the directory containing scripts and code used for all pipelines.
- code.repo: Directory containing all code for the project, copied from the main pipeline repository.
- data: Symlink to a directory containing reference genome data; set this in your original repository clone.
- inputs: Directory containing information on the files used as inputs.
- pipeline: Directory containing the files needed for each step in the pipeline.
- project\_notes: A bare directory in which you can place miscellaneous notes and documents concerning the analysis.
- run: File containing code for running the pipeline.
- run.dry: File containing code for testing the pipeline without execution of pipeline steps.

#### 2.2 Code Directories

The code needed for the execution of the analysis pipeline is divided among several sub-directories, based on usage. Within an analysis pipeline, the directory code.repo contains all of these sub-directories.

- bin: A directory containing symlinks to binary files for programs used by the pipeline.
- code.chipseq-standard, code.hicseq-standard: Directories containing scripts specific to the execution of each step in the the given type of pipeline analysis.
- code.main: A directory containing code and scripts used for all analysis pipelines.

## 2.3 Data Directory

The reference genome information needed for analysis is contained in the data directory. This can be contained in an external location and symlinked to the project directory if it has not already been set in the cloned HiC-bench repository template. Our example data contains only the subdirectory genomes, which is configured as such:

```
data/genomes/hg19$
-rw-r-r- 1 at570 at570
                           36M Nov 23 12:43 HindIII.fragments.bed
-rw-r-r- 1 at570 at570 298M Mar 7 15:58 Mbol.fragments.bed
                           32M Nov 23 12:43 Ncol.fragments.bed
-rw-r-r- 1 at570 at570
Irwxrwxrwx 1 at570 at570
                            20 Nov 23 12:41 bowtie2.index -> genome/bowtie2.index
                          1.5K Nov 23 13:43 centrotelo.bed
-rw-r-r- 1 at570 at570
drwxr-xr-x 2 at570 at570 1.2K Mar 7 16:00 features-hicnorm
-rw-r-r- 1 at570 at570 2.2M Dec 30 22:58 gene-name.bed
-rw-r-r- 1 at570 at570 2.2M Nov 30 15:43 gene.bed
                            33 Nov 23 12:34 genome -> /ifs/home/at570/Data/Genomes/hg19
Irwxrwxrwx 1 at570 at570
-rw-r-r- 1 at570 at570 564 Nov 23 13:43 genome.bed
data/genomes/mm10$
                           36M Nov 23 12:47 HindIII.fragments.bed
-rw-r-r- 1 at570 at570
-rw-r-r- 1 at570 at570
                           37M Nov 23 12:47 Ncol.fragments.bed
                            20 Nov 23 12:47 bowtie2.index -> genome/bowtie2.index
Irwxrwxrwx 1 at570 at570
-rw-r-r--1 at570 at570 1.3K Nov 23 13:48 centrotelo.bed drwxr-xr-x 2 at570 at570 1.2K Mar 7 16:00 features-hicnorm
                          1.4M Dec 30 22:58 gene-name.bed
-rw-r-r- 1 at570 at570
-rw-r-r- 1 at570 at570
                          1.4M Nov 30 15:43 gene.bed
Irwxrwxrwx 1 at570 at570
                            33 Nov 23 12:47 genome -> /ifs/home/at570/Data/Genomes/mm10
-rw-r-r- 1 at570 at570
                           495 Nov 23 13:48 genome.bed
```

Also included are indexes for bowtie2, which can be obtained from bowtie-bio.sourceforge.net/bowtie2/manual.shtml or http://support.illumina.com/sequencing/sequencing\_software/igenome.html.

## 2.4 Inputs Directory

The inputs directory contains files needed to run the pipeline.

- README: File containing usage notes for the inputs directory.
- code: Symlink to code one level up in the parent directory.
- data: Symlink to data one level up in the parent directory.
- fastq: Directory containing sub-directories for each sample to be used in the analysis. This directory is not created automatically, it must be created and populated manually. Alternatively, the directory bam can be used in its place if .bam files are to be used.
- genomes: Symlink to the directory containing reference genome information, within the data directory.
- params: Directory containing the parameters files associated with the input files.
- sample-sheet.tsv: Sample sheet for pipeline execution.

### 2.4.1 FASTQ Directory

The contents of an example fastq directory can be seen here:

```
hicseq.analysis-for-hicbench/inputs/fastq$
                    97 Feb 12 15:43 CD34-HindIII-rep1
Irwxrwxrwx 1 at570
Irwxrwxrwx 1 at570
                    95 Feb 12 15:43 GM-HindIII-rep1
Irwxrwxrwx 1 at570
                    92 Feb 12 15:43 GM-Ncol-rep1
Irwxrwxrwx 1 at570 103 Feb 12 15:43 H1-HindIII-Ren2015 rep1
Irwxrwxrwx 1 at570 103 Feb 12 15:43 H1-HindIII-Ren2015 rep2
Irwxrwxrwx 1 at570
                    95 Feb 12 15:43 H1-HindIII-rep1
Irwxrwxrwx 1 at570
                    95 Feb 12 15:43 H1-HindIII-rep2
Irwxrwxrwx 1 at570
                    98 Feb 12 15:43 IMR90-HindIII-rep1
Irwxrwxrwx 1 at570
                    98 Feb 12 15:43 IMR90-HindIII-rep2
Irwxrwxrwx 1 at570 100 Feb 12 15:43 T47D_T0-HindIII-rep1
Irwxrwxrwx 1 at570
                   97 Feb 12 15:43 T47D_T0-Ncol-rep1
Irwxrwxrwx 1 at570 101 Feb 12 15:43 T47D_T60-HindIII-rep1
Irwxrwxrwx 1 at570
                   98 Feb 12 15:43 T47D_T60-Ncol-rep1
Irwxrwxrwx 1 at570 100 Feb 12 15:43 mESC_J1-HindIII-rep1
Irwxrwxrwx 1 at570 100 Feb 12 15:43 mESC_J1-HindIII-rep2
                    97 Feb 12 15:43 mESC J1-Ncol-rep1
Irwxrwxrwx 1 at570
```

Each directory name contains information about the sample, in the format <CellLine>-<treatment>-<SampleID>. This format can be modified to suit your purposes, though it is recommended to retain the "-" character as a delimiter since it is used downstream in the sample sheet generation steps. Each directory should contain all of the .fastq / .fastq.gz files associated with the sample; symlinks pointing to each file can be used as well, and are encouraged in order to save disk space. The same protocol should be followed if .bam files are to be used. As per standard Linux Terminal guidelines, spaces and special characters should be avoided in file names and directory names.

## 2.5 Pipeline Directory

The pipeline directory contains information for each step in the pipeline. An example pipeline directory will have the following structure:

```
hicseq.analysis-for-hicbench/pipeline$
drwxr-xr-x
            5 at 570 228 Feb 15 16:47 align
drwxr-xr-x
              at570 207 Jan 19 22:12 annotations
            5
              at570 213 Feb 16 17:24 annotations-stats
drwxr-xr-x
            5 at 570 211 Feb 6 17:46 boundary-scores
drwxr-xr-x
              at570 389 Mar 10 18:16 boundary-scores-pca
drwxr-xr-x
            1 at570
                     7 Dec 2 12:39 code -> ../code
Irwxrwxrwx
           1 at570
                    12 Dec 2 12:39 code.main -> ../code.main
Irwxrwxrwx
            5 at570 385 Jan 19 22:08 compare-boundaries
drwxr-xr-x
drwxr-xr-x 5 at570 437 Mar 10 18:16 compare-boundaries-stats
```

```
11 drwxr-xr-x 5 at570 212 Jan 19 16:04 compare-matrices
  drwxr-xr-x 5 at570 429 Mar 10 18:15 compare-matrices-stats
  drwxr-xr-x 4 at570 420 Jan 19 22:10 diff-domains
  drwxr-xr-x 5 at570 231 Jan 20 13:06 domains
  drwxr-xr-x 5 at570 229 Jan 19 16:19
  drwxr-xr-x 6 at570 256 Jan 19 15:53 filter-stats
  drwxr-xr-x 5 at570 206 Jan 19 22:11 hicplotter
   -rw-r-r- 1 at570 331 Feb 14 19:28 index txt
  Irwxrwxrwx 1 at570
                      9 Dec 2 12:39 inputs ->
              5 at 570 208 Jan 19 22:12 interactions
  drwxr-xr-x
  drwxr-xr-x 4 at570 362 Jan 19 16:01 matrix-estimated
              5 at570 211 Jan 19 16:08 matrix-filtered
  drwxr-xr-x
  drwxr-xr-x 5 at570 794 Feb
                              7 17:05 matrix-hicnorm
  drwxr-xr-x 5 at570 205 Jan 19 16:00 matrix-ic
  drwxr-xr-x
              5 at570 207 Jan 19 15:59 matrix-prep
  drwxr-xr-x
              5 at570 361 Jan 19 16:03 matrix-stats
  drwxr-xr-x 4 at570 158 Dec 22 17:44 template
  drwxr-xr-x 5 at570 229 Jan 19 15:55 tracks
```

- align ... qc: Directories containing the information for each pipeline step.
- code: Symlink to the directory containing code specific to current the analysis type.
- code.main: Symlink to the directory containing code used for all analyses.
- inputs: Symlink to the inputs directory containing the .fastq or .bam files for the pipeline.
- index.txt: A text file containing a list of pipeline steps to be executed. Entries in this document match the names of the pipeline directories.

#### 2.5.1 Pipeline Index

The file index.txt contains a list of the pipeline steps to be completed during the analysis, listed in order of completion. An example index.txt would have the following structure:

```
hicseq.analysis-for-hicbench/pipeline$ cat index.txt
   align
   filter
   filter-stats
  tracks
   matrix-filtered
   matrix-prep
   matrix-ic
   matrix-hicnorm
  #matrix-estimated
18
  matrix-stats
20
  compare-matrices
22
  compare-matrices-stats
  boundary-scores
  boundary-scores-pca
26
  domains
  compare-boundaries
  compare-boundaries-stats
  #diff-domains
  hicplotter
  interactions
```

```
annotations annotations—stats
```

Each entry in the index.txt file matches the name of the pipeline step to be completed, represented by the corresponding name of the step's sub-directory in the pipeline directory. One entry is allowed per line in the index.txt file. Entries that begin with a '#' character will be ignored, and pipeline steps that are not included in the index.txt file will not be included in the analysis pipeline.

## 2.5.2 Example Pipeline Step Directory Structure

Each step in the pipeline is represented by a sub-directory in the pipeline directory. An example sub-directory for a pipeline step would have the following structure:

```
hicseq.analysis-for-hicbench/pipeline/align$
                      15 Oct 28 12:10 clean.tcsh -> code/clean.tcsh
7 Sep 29 13:31 code -> ../code
Irwxrwxrwx
               at570
              at570
Irwxrwxrwx
_rw_r__r_
            1
               at570
                       0 Jan 25 11:12 error.log
drwxr-xr-x
            2 at570
                      24 Feb 15 16:47 inpdirs
Irwxrwxrwx
            1
               at570
                       9 Sep 29 13:31 inputs -> ../inputs
            2 at570 77 Dec 28 13:42 params
drwxr-xr-x
drwxr-xr-x
            3 at570
                      62 Feb 16 12:27 results
Irwxrwxrwx
            1 at570 14 Jan 19 15:50 run -> run-align.tcsh
-rwxr-xr-x 1 at570 971 Jan 19 15:49 run-align.tcsh
```

- clean.tcsh: Script for cleaning the directory; remove results and error logs.
- code: Symlink to the directory containing code specific to the analysis type e.g. code.chipseq-standard in this case.
- error.log: File containing errors encountered during execution of the pipeline step, generated at runtime.
- inpdirs: Directory containing symlinks to directories containing input files for use during execution of the pipeline step.
- inputs: Symlink to the directory containing input files.
- params: Directory containing the parameters files associated with the pipeline step files.
- run: Symlink to the 'run' file for the pipeline step.
- run-align.tcsh: 'Run' file for the pipeline step, containing a script that passes pipeline execution information to the wrapper script located in ./code/code.main/pipeline-master-explorer.r.

#### 2.5.3 Example Pipeline Step Results Directory

The base level of a results directory for a pipeline step will have the following structure:

```
hicseq.analysis-for-hicbench/pipeline/align/results/align.by_sample.bowtie2/CD34-HindIII-rep1$

-rw-r-r-- 1 at570 49G Jan 13 01:02 alignments.bam
-rw-r-r-- 1 at570 473 Jan 13 01:02 job.err

-rw-r-r-- 1 at570 47 Jan 12 18:42 job.id
-rw-r--- 1 at570 0 Jan 12 18:42 job.out

-rw-r--- 1 at570 136 Jan 12 18:42 job.sh
-rw-r--- 1 at570 2.3K Jan 13 01:02 job.vars.tsv
```

- alignments.bam: Example alignment output file.
- job.err: File containing the standard error output of the pipeline step.

- job.id: File containing the ID number of the job after submission for execution on the HPC cluster.
- job.out: File containing the standard output of the pipeline step.
- job.sh: File containing the command submitted for execution on the HPC cluster.
- job.vars.tsv: File containing the variables used in the completion of the pipeline step.

# 3 Adding Custom Pipeline Steps

## 3.1 Custom Pipeline Step Overview

The following basic steps should be taken to create a custom pipeline step:

- · Copy an existing step as a template
- Update the new pipeline step name and add it to the entries in the index.txt and as a symlink in the
  parent level of the analysis directory
- · Set the input directories ('inpdirs')
- · Edit the 'run' file and add needed parameter files
- Add a script in the code directory containing the commands needed to run the programs used in the pipeline step

# 3.2 How To Add Custom Pipeline Steps

The steps needed to create a custom pipeline step are explained in detail here:

1. Within the pipeline directory, use a command such as cp -r to make a copy of an existing pipeline step as a template for the new one.

Example pipeline directory:

```
hicseq.analysis-for-hicbench/pipeline$ Is -I
total 818K
drwxr-xr-x 25 at570 at570
                           834 Mar 18 17:14
                           958 Mar 21 19:09
drwxr-xr-x
            5 at570 at570
            5 at570 at570
                           228 Mar 10 16:20 align
drwxr-xr-x
                           234 Mar 21 19:18 annotations
drwxr-xr-x
            5 at 570 at 570
drwxr_xr_x
            5 at 570 at 570
                           240 Mar 21 19:40 annotations-stats
drwxr-xr-x
            5 at 570 at 570
                           238 Mar 18 20:44 boundary-scores
drwxr-xr-x
            5 at 570 at 570
                           242 Mar 21 16:37 boundary-scores-pca
Irwxrwxrwx
            1 at570 at570
                             7 Mar 10 16:20 code -> ../code
Irwxrwxrwx
            1 at570 at570
                            12 Mar 10 16:20 code.main ->
drwxr-xr-x
            5 at570 at570
                           241 Mar 21 16:37 compare-boundaries
                           247 Mar 21 16:37 compare-boundaries-stats
drwxr-xr-x
            5 at 570 at 570
drwxr-xr-x
            5 at570 at570
                           239 Mar 18 20:20 compare-matrices
drwxr-xr-x
            5 at570 at570
                           245 Mar 21 16:37 compare-matrices-stats
drwxr-xr-x
            4 at570 at570
                           245 Mar 21 16:37 diff-domains
drwxr-xr-x
            5 at570 at570
                           230 Mar 18 21:22
                                             domains
                                            filter
drwxr-xr-x
            5 at570 at570
                           229 Mar 10 16:20
drwxr-xr-x
            6 at570 at570
                           256 Mar 10 16:20
                                            filter-stats
drwxr-xr-x
            7 at570 at570
                           323 Mar 21 16:41
                                            hicplotter
-rw-r---r---
            1 at570 at570
                           331 Mar 10 16:20 index txt
Irwxrwxrwx 1 at570 at570
                             9 Mar 10 16:20 inputs ->
drwxr-xr-x
            5 at570 at570
                           235 Mar 10 16:20 interactions
drwxr-xr-x
            4 at570 at570
                           187 Mar 21 16:37 matrix-estimated
drwxr-xr-x
            5 at570 at570
                           238 Mar 10 16:20 matrix-filtered
            5 at570 at570
                           260 Mar 10 16:20 matrix-hicnorm
drwxr-xr-x
drwxr-xr-x
            5 at570 at570
                           232 Mar 10 16:20 matrix-ic
            5 at570 at570
                           234 Mar 18 17:31 matrix-prep
drwxr-xr-x
            5 at570 at570
                           235 Mar 21 16:37 matrix-stats
drwxr-xr-x
Irwxrwxrwx 1 at570 hpchic
                             8 Dec 3 14:56 psync -> ../psync
                           158 Mar 10 16:20
drwxr-xr-x
            4 at570 at570
                                            template
drwxr-xr-x 5 at570 at570
                           229 Mar 10 16:20 tracks
```

2. Adjust the name of the new directory to match the desired name of the new pipeline step. Add this name as an entry in the index.txt file, and make a symlink to this directory from the parent directory in the same style of the existing symlinks to other pipeline steps. The alpha-numeric prefix on the symlink will determine the order in which it will be executed. The command to do this might look like this:

```
hicseq.analysis-for-hicbench$ In -s pipeline/my_new_step __03b-my_new_step
```

#### Example index.txt file contents:

```
hicseq.analysis-for-hicbench/pipeline$ cat index.txt
   align
   filter
   filter-stats
   tracks
  matrix-filtered
  matrix-prep
  matrix-ic
13
   matrix-hicnorm
15
   #matrix-estimated
17
  matrix-stats
19
2
  compare-matrices
   compare-matrices-stats
   boundary-scores
25
  boundary-scores-pca
27
  domains
   compare-boundaries
   compare-boundaries-stats
   #diff_domains
33
   hicplotter
35
   interactions
   annotations
   annotations-stats
```

#### Example parent directory structure:

```
hicseq.analysis-for-hicbench$ Is -I
Irwxrwxrwx
           1 at570 at570 14 Mar 10 16:20
                                            _01a-align -> pipeline/align
Irwxrwxrwx
           1 at570 at570
                          15 Mar 10 16:20
                                            _02a-filter -> pipeline/filter
Irwxrwxrwx
           1 at570 at570
                          21 Mar 10 16:20
                                            _02b-filter-stats -> pipeline/filter-stats
Irwxrwxrwx
           1 at570 at570 15 Mar 10 16:20
                                            _03a-tracks -> pipeline/tracks
Irwxrwxrwx
           1 at570 at570
                          24 Mar 10 16:20
                                            04a-matrix-filtered -> pipeline/matrix-filtered
Irwxrwxrwx 1
             at570 at570
                          20 Mar 10 16:20
                                            05a-matrix-prep -> pipeline/matrix-prep
Irwxrwxrwx
           1 at570 at570
                          18 Mar 10 16:20
                                            06a-matrix-ic -> pipeline/matrix-ic
Irwxrwxrwx 1 at570 at570
                          23 Mar 10 16:20
                                            07a-matrix-hicnorm -> pipeline/matrix-hicnorm
Irwxrwxrwx
           1 at570 at570
                          21 Mar 10 16:20
                                            _08a-matrix-stats -> pipeline/matrix-stats
                                            1 at570 at570
                          25 Mar 10 16:20
Irwxrwxrwx
Irwxrwxrwx
           1 at570 at570
                          31 Mar 10 16:20
Irwxrwxrwx
           1 at570 at570
                          24 Mar 10 16:20
                                            10a-boundary-scores -> pipeline/boundary-scores
           1 at570 at570
                          28 Mar 10 16:20
Irwxrwxrwx
                                            10b-boundary-scores-pca -> pipeline/boundary-scores-pca
Irwxrwxrwx 1 at570 at570
                          16 Mar 10 16:20
                                            11a-domains -> pipeline/domains
           1 at570 at570
                          27 Mar 10 16:20
Irwxrwxrwx
                                            _12a-compare-boundaries -> pipeline/compare-boundaries
Irwxrwxrwx
           1 at570 at570
                          33 Mar 10 16:20
                                            _12b-compare-boundaries-stats -> pipeline/compare-boundaries-
    stats
Irwxrwxrwx
           1 at570 at570
                          19 Mar 10 16:20
                                            13a-hicplotter -> pipeline/hicplotter
             at570 at570
                          21 Mar 10 16:20
                                            14a-interactions -> pipeline/interactions
Irwxrwxrwx
           1 at570 at570
Irwxrwxrwx
                          20 Mar 10 16:20
                                            _15a-annotations -> pipeline/annotations
Irwxrwxrwx
           1 at570 at570
                          26 Mar 10 16:20
                                            _15b-annotations-stats -> pipeline/annotations-stats
Irwxrwxrwx 1 at570 at570
                          30 Mar 14 18:25 code -> code repo/code hicseq-standard
Irwxrwxrwx
           1 at570 at570
                          14 Mar 10 16:20 code main -> code/code main
drwxr-xr-x 10 at570 at570 238 Mar 13 21:45 code.repo
Irwxrwxrwx 1 at570 at570 104 Mar 14 18:25 data -> /ifs/home/at570/disk1/Resources/Code/pipeline-master/
    code/code.main/../../pipelines/hicseq-standard/data
drwxr-xr-x 5 at570 at570 274 Mar 10 16:20 inputs
```

```
drwxr-xr-x 25 at570 at570 834 Mar 18 17:14 pipeline
-rwxr-xr-x 1 at570 at570 211 Mar 11 12:06 psync
-rwxr-xr-x 1 at570 at570 888 Mar 10 16:20 run
-rwxr-xr-x 1 at570 at570 898 Mar 10 16:20 run.dry
```

3. Edit the contents of the directory you have created to hold the information for your new pipeline step. Example pipeline step directory:

```
hicseq.analysis—for—hicbench/pipeline/domains$ Is —I

lrwxrwxrwx 1 at570 at570 15 Mar 10 16:20 clean.tcsh —> code/clean.tcsh
lrwxrwxrwx 1 at570 at570 7 Mar 10 16:20 code —> ../code

-rw—r—r— 1 at570 at570 0 Mar 18 21:22 error.log
drwxr—xr—x 2 at570 at570 155 Mar 10 16:20 inpdirs

Irwxrwxrwx 1 at570 at570 9 Mar 10 16:20 inputs —> ../inputs
drwxr—xr—x 3 at570 at570 146 Mar 10 16:20 params

drwxr—xr—x 6 at570 at570 161 Mar 18 20:48 results
Irwxrwxrwx 1 at570 at570 16 Mar 10 16:20 run —> run—domains.tcsh

-rwxr—xr—x 1 at570 at570 959 Mar 11 14:58 run—domains.tcsh
```

#### First, edit the run file. A sample run file looks like this:

```
hicseq.analysis-for-hicbench/pipeline/domains$ cat run
   #!/bin/tcsh
   source ./code/code.main/custom-tcshrc
                                                   # customize shell environment
  ## USAGE: run-domains.tcsh [--dry-run]
   # this section holds information that will be used in future updates of the software for reporting
  #% This step identifies topologically-associated domains (TADs) using different methods.
   #% TABLES:
12 #% FIGURES:
  # process command—line inputs
   # check to make sure that the proper number of arguments have been passed to the script,
  # if not then print the script lines starting with '##' and exit
   if ($#argv > 1) then
  grep '^##' $0 | scripts-send2err
   endif
  set opt = "$1"
24
  # setup
   # set the 'operation' to be performed, aka name of the pipeline step
26 set op = domains
   # the directories to be used for inputs
28 set inpdirs = "inpdirs/*"
   # an expression which specifies which input branches to include
30 set filter = "*.res 40kb"
                                                   # work only with 40kb resolution
   # the name of the results directory
32 set results = results
34 # create the results directory
   scripts-create-path $results/
  # sends a message to the error logging script
scripts—send2err "=== Operation = $op ===========
  # 'resources' argument to be passed to qsub, referring to CPU cores and GB of RAM to be reserved for the
        iob
   set resources = 1.20G
  # command to be passed to the 'pipeline-master-explorer.r' script
set cmd = "./code/code.main/scripts-qsub-wrapper $resources ./code/hicseq-$op.tcsh"
40
42
   # generate run script
  # the 'pipeline-master-explorer.r' script parses the items set above to create a line of text containing
       the commands to be submitted to qsub
   Rscript ./code/code.main/pipeline-master-explorer.r.-v.-F ~\$filter"~\$cmd"~\$results/\$op~"params/params.\star.
        tcsh"
               "$inpdirs'
                               "sample" 1
  # run and wait until done!
  # if the '--dry-run' argument was not passed to the script
if ("$opt" != "--dry-run") scripts-submit-jobs ./$results/.db/run
```

As listed in the above run file, the following 'setup' items need to be set for the custom pipeline step:

```
set op = <name_of_pipeline_step>
```

The 'operation' to be performed is set as 'op' and should be the name of the pipeline step, as listed in the directory name and in the index.txt file.

```
set inpdirs = "inpdirs/*"
```

The 'inpdirs', or input directories, should be set as the file path to the directory containing symlinks to the input directories. In this case, the contents of inpdirs is as follows:

```
hicseq.analysis-for-hicbench/pipeline/domains$ Is -I inpdirs/
| Irwxrwxrwx 1 at570 at570 22 Mar 10 16:20 matrix-estimated -> ../../matrix-estimated
| Irwxrwxrwx 1 at570 at570 21 Mar 10 16:20 matrix-filtered -> ../../matrix-filtered |
| Irwxrwxrwx 1 at570 at570 20 Mar 10 16:20 matrix-hicnorm -> ../../matrix-hicnorm |
| Irwxrwxrwx 1 at570 at570 15 Mar 10 16:20 matrix-ic -> ../../matrix-ic |
| Irwxrwxrwx 1 at570 at570 17 Mar 10 16:20 matrix-prep -> ../../matrix-prep
```

The setting "inpdirs/\*" will cause all input directories to be used. The entries in the inpdirs directory should be set as needed for the execution of the custom pipeline step.

```
set filter = "*.res_40kb"
```

The 'filter' setting to be used when parsing the 'branches' of the input directory results, for inclusion in the execution of the pipeline step. In this example, only input branches that match the pattern "\*.res\_40kb" will be included. In this example, the following input branches are available:

```
hicseq.analysis—for—hicbench/pipeline/domains$ Is —I inpdirs/matrix—filtered/results/
drwxr—xr—x 3 at570 at570 43 Mar 11 14:43 matrix—filtered.by_sample.res_1000kb
drwxr—xr—x 3 at570 at570 43 Mar 11 14:44 matrix—filtered.by_sample.res_100kb
drwxr—xr—x 3 at570 at570 43 Mar 11 14:44 matrix—filtered.by_sample.res_10kb.maxd_5Mb.rotate45
drwxr—xr—x 3 at570 at570 43 Mar 11 14:45 matrix—filtered.by_sample.res_40kb
```

Based on the given 'filter' setting, only the following branch will be included:

```
drwxr-xr-x 3 at570 at570 43 Mar 11 14:45 matrix-filtered.by_sample.res_40kb
```

This allows for the exclusion of unnecessary analysis branches.

```
set resources = 1,20G
```

This sets the number of computer resources to be reserved by qsub, listed as CPU cores and GB of RAM. If RAM is not a concern, only CPU cores need to be listed. A range of values can be used for CPU cores, such as 8-64, though the utility of this depends on many factors related to your high-performance computing infrastructure and the specifics of the program being run; more cores may not necessarily speed up execution of the task at hand.

```
set cmd = "./code/code.main/scripts-qsub-wrapper $resources ./code/hicseq-$op.tcsh"
```

This line does not need to be modified by the user, but should be noted since it refers to the file in the code directory that will be created later and used to execute the program used in the pipeline step. Importantly, the entry ./code/hicseq-\$op.tcsh in this case refers to the file ./code/hicseq-domains.tcsh.

```
Rscript ./code/code.main/pipeline-master-explorer.r -v -F "$filter" "$cmd" $results/$op "params/params.*.tcsh" "$inpdirs" "" "sample" 1
```

Since this calls the settings that have already been made, this line of the 'run' file does not need to be edited unless the grouping and splitting variables need to be changed. In this case, the command uses the following arguments (as per Section 5.3):

```
pipeline-master-explorer.r [OPTIONS] SCRIPT OUTDIR-PREFIX PARAM-SCRIPTS INPUT-BRANCHES SPLIT-VARIABLE OUTPUT-OBJECT-VARIABLE TUPLES
```

Importantly, the 'split-variable' and 'output-object-variable' come from the headings of columns used as grouping factors in the inputs/sample-sheet.tsv file for the analysis; custom grouping factors can be included in the sample sheet and used here. The output of the pipeline-master-explorer. r script is stored in the file results/.db/run which is created when the run file is executed (the command ./run--dry-run can be used to generate this without running the commands). An example entry will look like this:

```
hicseq.analysis-for-hicbench/pipeline/domains$ head -n 1 results/.db/run
./code/code.main/scripts-qsub-wrapper 1,20G ./code/hicseq-domains.tcsh results/domains.by_sample.
armatus.gamma_0.5/matrix-prep.by_sample.scale/matrix-filtered.by_sample.res_40kb/filter.
by_sample.standard/align.by_sample.bowtie2/CD34-HindIII-rep1 params/params.armatus.gamma_0.5.
tcsh inpdirs/matrix-prep/results/matrix-prep.by_sample.scale/matrix-filtered.by_sample.res_40kb/filter.by_sample.standard/align.by_sample.bowtie2 'CD34-HindIII-rep1'
```

During pipeline step execution, these lines will be submitted to qsub by the script ./code/code.main/scripts-qsub-wrapper.

4. Next, the parameter files must be set for the new pipeline step. These files are contained in the params directory:

```
hicseq.analysis—for—hicbench/pipeline/domains$ Is—I params/
-rwxr—xr—x 1 at570 at570 131 Mar 10 16:20 params.armatus.gamma_0.5.tcsh
-rwxr—xr—x 1 at570 at570 480 Mar 10 16:20 params.hicmatrix.tcsh
-rwxr—xr—x 1 at570 at570 155 Mar 10 16:20 params.topdom.tcsh
```

Importantly, files must use the following naming scheme: params.<name>.tcsh. All files included in the params directory following this naming scheme will be evaluated as a separate 'branch' for analysis. An example parameters file looks like this:

```
hicseq.analysis—for—hicbench/pipeline/domains$ cat params/params.armatus.gamma_0.5.tcsh

#!/bin/tcsh

source ./inputs/params/params.tcsh

set tool = armatus
set chrom_excluded = 'chr[MY]'
set armatus_params = "-g 0.5"
```

Settings that are specific to each analysis branch should be included in these files. Sub-directories in the results directory will be created for each entry in the params directory, as can be seen here:

```
hicseq.analysis—for—hicbench/pipeline/domains$ Is—I results/
drwxr—xr—x 7 at570 at570 246 Mar 18 20:48 domains.by_sample.armatus.gamma_0.5
drwxr—xr—x 7 at570 at570 246 Mar 18 20:48 domains.by_sample.hicmatrix
drwxr—xr—x 7 at570 at570 246 Mar 18 20:49 domains.by_sample.topdom
```

Note that in this case, the full output directory name comes from the following components: step>.by\_<output-object-variable>.<params\_entry>

5. A script containing the commands needed to run the desired program must be created and placed in the code directory, of which a partial list is shown below as an example:

```
hicseq.analysis-for-hicbench/pipeline/domains$ Is -I code/hic*
  -rwxr-xr-x 1 at570 at570 26528 Dec 8 11:42 code/hic-matrix.o
  -rwxr-xr-x 1 at570 at570 117660 Mar 18 15:25 code/hic-matrix.r
 -rwxr-xr-x 1 at570 at570
                           24440 Dec 8 11:42 code/hic-matrix.so
  -rwxr-xr-x 1 at570 at570
                             2471 Mar 10 16:20 code/hicnorm-cis.r
 -rwxr-xr-x 1 at570 at570
                             2542 Mar 10 16:20 code/hicseq-align.tcsh
  -rwxr-xr-x 1 at570 at570
                             2352 Mar 10 16:20 code/hicseq-annotate-tables.tcsh
 -rwxr-xr-x 1 at570 at570
                             3463 Mar 21 19:28 code/hicseq-annotations-enrichments.r
  -rwxr-xr-x 1 at570 at570
                             1547 Mar 21 18:54 code/hicseq-annotations-stats.tcsh
10 -rwxr-xr-x 1 at570 at570
                             1274 Mar 10 16:20 code/hicseq-annotations.tcsh
  -rwxr-xr-x 1 at570 at570
                             2098 Mar 14 17:04 code/hicseq-boundary-scores-pca.tcsh
12 -rwxr-xr-x 1 at570 at570
                             1649 Mar 10 16:20 code/hicseq-boundary-scores.tcsh
  -rwxr-xr-x 1 at570 at570
                             1440 Mar 10 16:20 code/hicseq-compare-boundaries-stats.tcsh
  -rwxr-xr-x 1 at570 at570
                             3138 Mar 10 16:20 code/hicseq-compare-boundaries.tcsh
  -rwxr-xr-x 1 at570 at570
                             1362 Mar 10 16:20 code/hicseq-compare-matrices-stats tcsh
 -rwxr-xr-x 1 at570 at570
                             1596 Mar 10 16:20 code/hicseq-compare-matrices.tcsh
  -rwxr-xr-x 1 at570 at570
                             1902 Mar 10 16:20 code/hicseq-diff-domains.tcsh
```

Importantly, the primary script must follow this naming scheme: hicseq-<pipeline\_step>.tcsh. In this example, this would correspond to hicseq-domains.tcsh. Pre-existing scripts can be used as a template to set up your custom script. This example script has the following contents:

```
hicseq.analysis-for-hicbench/pipeline/domains$ cat code/hicseq-domains.tcsh
   #!/bin/tcsh
  source ./code/code.main/custom-tcshrc
                                               # shell settings
  ##
  ## USAGE: hicseq-domains.tcsh OUTPUT-DIR PARAM-SCRIPT BRANCH OBJECT(S)
  ##
  if ($#argv != 4) then
grep '^##' $0
     exit
   endif
   set outdir = $1
  set params = $2
   set branch = $3
  set objects = ($4)
  # read variables from input branch
   source //code/code.main/scripts-read-job-vars $branch "$objects" "genome genome_dir bin_size"
   # run parameter script
  source $params
23
  # create path
   scripts-create-path $outdir/
27

    MAIN CODE BELOW

29
  # .
31
   # Run domains
  if ((\$tool == armatus) || (\$tool == di) || (\$tool == topdom) || (\$tool == caltads) || (\$tool == hicmatrix)
        ) then
     ./code/hicseq-domains-$tool.tcsh $outdir $params $branch "$objects"
35 else
```

The preamble of the script should require little user intervention, while the bulk of the user's custom pipeline code should be inserted between the 'MAIN CODE' blocks specified within the document. For reference on how to structure your custom code, compare the 'USAGE' entry with the evaluated command to be passed to the script in the results/.db/run file. For convenience, the sample entry is repeated below:

```
hicseq.analysis-for-hicbench/pipeline/domains$ head -n 1 results/.db/run
./code/code.main/scripts-qsub-wrapper 1,20G ./code/hicseq-domains.tcsh results/domains.by_sample.armatus.
gamma_0.5/matrix-prep.by_sample.scale/matrix-filtered.by_sample.res_40kb/filter.by_sample.standard/
align.by_sample.bowtie2/CD34-HindIII-rep1 params/params.armatus.gamma_0.5.tcsh inpdirs/matrix-prep/
results/matrix-prep.by_sample.scale/matrix-filtered.by_sample.res_40kb/filter.by_sample.standard/
align.by_sample.bowtie2 'CD34-HindIII-rep1'
```

While this primary script should be in the .tcsh format, subsequent scripts in the user's preferred language can be called. They should follow the same naming conventions as shown in the code directory example above.

# 4 HiC-Seq Pipeline

## 4.1 Pipeline Steps

Within the parent directory of an analysis, the default pipeline steps are listed as symlinks, in alpha-numeric order starting with "\_\_\_", as seen here:

```
14 Feb
Irwxrwxrwx
             at570
                                      01a-align -> pipeline/align
Irwxrwxrwx
              at570
                    15 Feb
                            7 17:06
                                      _02a-filter -> pipeline/filte:
                                      _02b-filter-stats -> pipeline/filter-stats
Irwxrwxrwx
             at570
                    21 Feb
                            7 17:06
             at570
                    15 Feb 7 17:06
                                      _03a—tracks —> pipeline/tracks
Irwxrwxrwx
             at570
                    24 Feb
                           7 17:06
                                      _04a-matrix-filtered -> pipeline/matrix-filtered
Irwxrwxrwx
             at570
                    20 Feb 7 17:06
                                      _05a-matrix-prep -> pipeline/matrix-prep
Irwxrwxrwx
                    18 Feb 7 17:06
Irwxrwxrwx
             at570
                                      _06a-matrix-ic -> pipeline/matrix-ic
             at570
                    23 Feb 7 17:06
                                      ____07a-matrix-hicnorm -> pipeline/matrix-hicnorm
Irwxrwxrwx
                    21 Feb
Irwxrwxrwx
             at570
                           7 17:06
                                      _08a-matrix-stats -> pipeline/matrix-stats
Irwxrwxrwx
             at570
                    25 Feb 7 17:06
                                      ____09a-compare-matrices -> pipeline/compare-matrices
                    31 Feb 7 17:06
Irwxrwxrwx
             at570
                                      _09b-compare-matrices-stats -> pipeline/compare-matrices-stats
                                      Irwxrwxrwx
             at570
                    24 Feb 7 17:06
             at570
                    28 Feb 7 17:06
Irwxrwxrwx
                                    __11a-domains -> pipeline/domains
             at570
                    16 Feb 7 17:06
Irwxrwxrwx
Irwxrwxrwx
             at570
                    27 Feb 7 17:06
                                      _12a-compare-boundaries -> pipeline/compare-boundaries
             at570
                    33 Feb 7 17:06
Irwxrwxrwx
                                      12b-compare-boundaries-stats -> pipeline/compare-boundaries-stats
Irwxrwxrwx
             at570
                    19 Feb 7 17:06
                                      _
13a-hicplotter -> pipeline/hicplotter
Irwxrwxrwx
             at570
                    21 Feb 7 17:06
                                    __14a-interactions -> pipeline/interactions
             at570
                    20 Feb 7 17:06
Irwxrwxrwx
                                      _15a—annotations —> pipeline/annotations
                    30 Feb 8 17:47 code -> code.repo/code.hicseq-standard
           1 at570
Irwxrwxrwx
             at570
                    14 Nov 12 11:55 code.main -> code/code.main
Irwxrwxrwx
           9 at570 209 Jan 9 10:16 code.repo
drwxr-xr-x
Irwxrwxrwx
             at570 103 Feb 8 17:47
                                    data -> /ifs/home/.../data
           6 at 570 258 Jan 5 09:24 inputs
drwxr-xr-x
drwxr-xr-x 24 at570 799 Feb 7 17:06 pipeline
             at570 210 Dec 2 14:23 psync
-rwxr-xr-x 1
             at570 981 Jan 5 19:40 run
-rwxr-xr-x
           1
           1 at570 554 Dec 18 17:02 run.dry
-rwxr-xr-x
             at570 988 Jan 24 14:42 run.subset
           1
-rwxr-xr-x
-rwxr-xr-x 1 at570 165 Dec 26 08:09 run.usage
```

This functions in informing the user of the order of pipeline steps. Each symlink points back to a directory in the pipeline directory for the corresponding pipeline step, as shown here:

```
pipeline$
total 814K
             4 at570 203 Jan 19 15:50 align
drwxr-xr-x
             5 at 570 207 Jan 19 22:12 annotations
drwxr-xr-x
            5 at570 387 Feb 6 17:46 boundary-scores
5 at570 243 Feb 7 14:17 boundary-scores-pca
drwxr-xr-x
drwxr-xr-x
            1 at570 7 Dec 2 12:39 code -> ../code
1 at570 12 Dec 2 12:39 code.main -> ...
Irwxrwxrwx
Irwxrwxrwx
                                                       ./code.main
            5 at 570 390 Jan 19 22:08 compare-boundaries
drwxr-xr-x
drwxr-xr-x
             5 at570 396 Jan 20 10:59 compare-boundaries-stats
drwxr-xr-x
            5 at 570 435 Jan 19 16:04 compare-matrices
drwxr-xr-x
             6 at570 419 Jan 19 16:05 compare-matrices-stats
drwxr-xr-x
             5 at 570 445 Jan 19 22:10 diff -domains
drwxr-xr-x
             5 at 570 379 Jan 20 13:06 domains
drwxr-xr-x 5 at570 229 Jan 19 16:19 filter
drwxr{-}xr{-}x
             6 at570 256 Jan 19 15:53 filter-stats
drwxr-xr-x
             5 at 570 206 Jan 19 22:11 hicplotter
             1 at570 297 Feb 6 17:52 index.txt
-rw-r---r
Irwxrwxrwx 1
               at570
                       9 Dec 2 12:39 inputs ->
             5 at 570 208 Jan 19 22:12 interactions
drwxr-xr-x
             5 at570 239 Jan 19 16:01 matrix-estimated
drwxr-xr-x
drwxr-xr-x
             5 at570 238 Jan 19 16:08 matrix-filtered
drwxr-xr-x
            5 at570 232 Jan 19 16:00 matrix-ic
             5 at570 234 Jan 19 15:59 matrix-prep
drwxr-xr-x
             5 at570 208 Jan 19 16:03 matrix-stats
drwxr-xr-x
            1 at570
                       8 Dec 3 14:56 psync -> ../psync
Irwxrwxrwx
            4 at570 158 Dec 22 17:44 template
drwxr-xr-x
            5 at570 229 Jan 19 15:55 tracks
drwxr-xr-x
```

The pipeline directory contains files and symlinks needed for each step in the pipeline. The steps to be exectuted are defined in two ways:

- 1. A file called 'index.txt' lists the names of each step in the pipeline, in the order in which they will be completed. This file is located in the 'pipeline' directory.
- 2. A subdirectory within the 'pipeline' directory with the same name as its corresponding entry in the 'index.txt' file must be included to hold the parameters and commands to be run, and the results produced.

#### Index file:

```
pipeline/index.txt$
  align
   filter
   filter-stats
  tracks
   matrix-filtered
  matrix-prep
   matrix-ic
  #matrix-estimated
  matrix-stats
  compare-matrices
  compare-matrices-stats
  boundary-scores
  boundary-scores-pca
  domains
26
  compare-boundaries
  compare-boundaries-stats
  #diff-domains
30
  hicplotter
32
  interactions
34
  annotations
```

Steps listed in 'index.txt' which have been commented out (i.e. start with a # character) will not be included in the analysis. Custom pipeline steps can be easily included by adding the corresponding entry to the 'index.txt' and creating a subdirectory within the 'pipeline' directory.

For details on adding custom pipeline steps, see Section ??

#### 4.1.1 Default Parameters

These parameters are used by default across pipeline steps.

```
/inputs/params/params.tcsh$
#!/bin/tcsh

# load basic tools
module unload samtools
module unload java
module unload gcc
module unload python
module load samtools/1.2.1
module load bedtools/2.22.0
module load java/1.7
module load picard—tools

# load tools required for each step of the pipeline (this can be overriden in local param scripts)
module load bowtie2/2.2.6
module load armatus/2014—05—19
module load caltads/0.1.0
```

```
module load ghmm/0.9

# sample sheet file
set sheet = inputs/sample—sheet.tsv
```

# 4.1.2 Pipeline Step Execution Flowchart

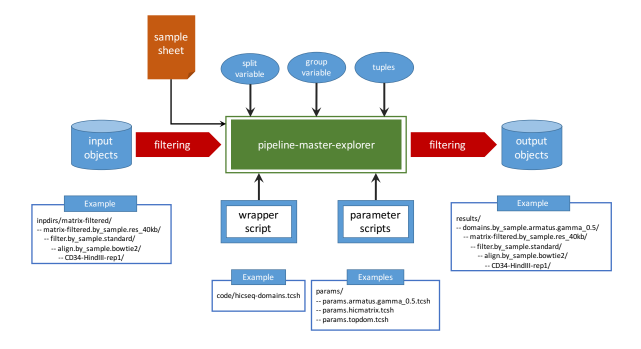


Figure 2: Overview of pipeline step execution for default analysis steps. See Section 4.1.2.

# 4.2 Alignment

## 4.2.1 Input

Raw data in fastq or fastq.gz files (Section 2.4).

## 4.2.2 Analysis

## Default parameters:

```
params.bowtie2.tcsh$
#!/bin/tcsh

source ./inputs/params/params.tcsh

set aligner = bowtie2

set genome = `./code/read-sample-sheet.tcsh $sheet $object genome`
set genome_index = inputs/genomes/$genome/bowtie2.index/genome
set align_params = "--very-sensitive-local --local"
```

## **4.2.3** Output

```
-rw-r - 1 at570 49G Jan 13 01:02 alignments.bam
-rw-r - 1 at570 473 Jan 13 01:02 job.err

-rw-r - 1 at570 47 Jan 12 18:42 job.id
-rw-r - 1 at570 0 Jan 12 18:42 job.out
-rw-r - 1 at570 136 Jan 12 18:42 job.sh
-rw-r - 1 at570 2.3K Jan 13 01:02 job.vars.tsv
```

# 4.3 Filter

## 4.3.1 Input

Data from the pipeline align step is used as input (Section 4.2).

## 4.3.2 Analysis

## Default parameters:

```
params.standard.tcsh$

#!/bin/tcsh

source ./inputs/params/params.tcsh

set filter_params = "—mapq 30 —min-dist 25000 —max-offset 500 —filter-dups"
```

# **4.3.3 Output**

```
-rw-r-r- 1 at570 1.7G Jan 13 14:26 filtered.reg.gz
2 -rw-r-r- 1 at570 65K Jan 13 14:25 job.err
-rw-r-r- 1 at570 47 Jan 13 13:15 job.id
4 -rw-r-r- 1 at570 0 Jan 13 13:15 job.out
-rw-r-r- 1 at570 195 Jan 13 13:15 job.sh
6 -rw-r-r- 1 at570 2.1K Jan 13 14:26 job.vars.tsv
-rw-r-r- 1 at570 378 Jan 13 14:24 stats.tsv
```

# 4.4 Filter Stats

## 4.4.1 Input

Data from the pipeline filter step is used as input (Section 4.3).

## 4.4.2 Analysis

## Default parameters:

```
params.standard.tcsh$
#!/bin/tcsh
source ./inputs/params/params.tcsh
```

## **4.4.3 Output**

See Figure 3 and Figure 4. Default output:

```
-rw-r - 1 at570 6.5K Feb 11 15:27 counts.pdf

2 -rw-r - 1 at570 34 Feb 11 15:27 job.err
-rw-r - 1 at570 47 Feb 11 15:27 job.id

4 -rw-r - 1 at570 52 Feb 11 15:27 job.out
-rw-r - 1 at570 226 Feb 11 15:27 job.sh

6 -rw-r - 1 at570 6.7K Feb 11 15:27 percent.pdf
```



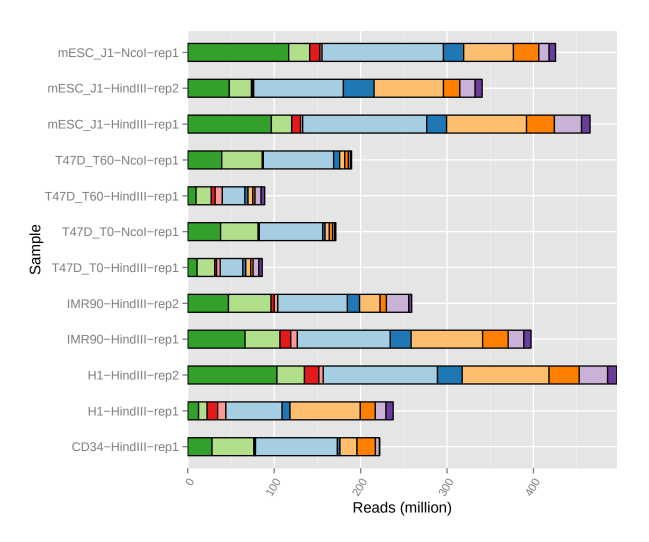


Figure 3: Filter Stats counts sample output

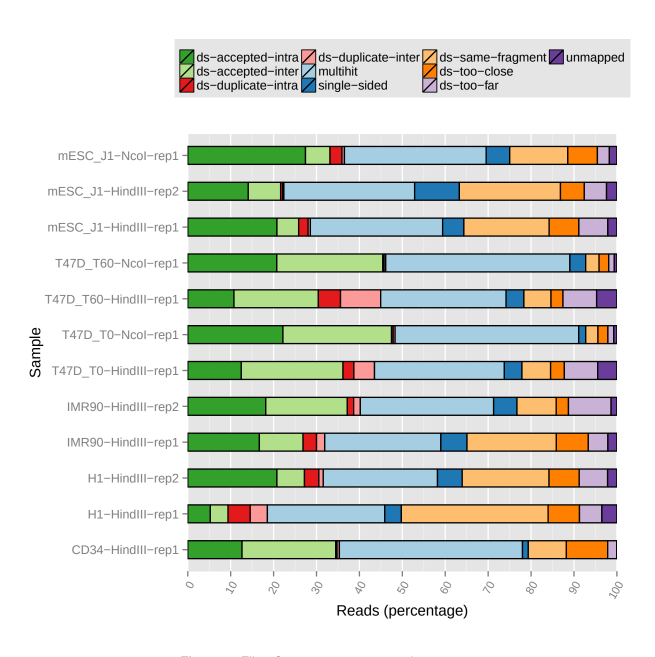


Figure 4: Filter Stats percentage sample output

## 4.5 Tracks

## 4.5.1 Input

Data from the pipeline filter step is used as input (Section 4.3).

## 4.5.2 Analysis

### Default parameters:

```
params.standard.tcsh$

#!/bin/tcsh

source ./inputs/params/params.tcsh

set bin_size = 40000 # this is a commonly used bin size
```

# 4.5.3 **Output**

```
-rw-r-r-- 1 at570 4.1K Jan 13 15:55 job.err

-rw-r-r-- 1 at570 47 Jan 13 15:10 job.id

-rw-r-r-- 1 at570 0 Jan 13 15:11 job.out

-rw-r-r-- 1 at570 242 Jan 13 15:10 job.sh

-rw-r-r-- 1 at570 2.6K Jan 13 15:55 job.vars.tsv

-rw-r-r-- 1 at570 1.1G Jan 13 15:54 track.washu.tsv.gz

-rw-r-r-- 1 at570 789K Jan 13 15:55 track.washu.tsv.gz.tbi
```

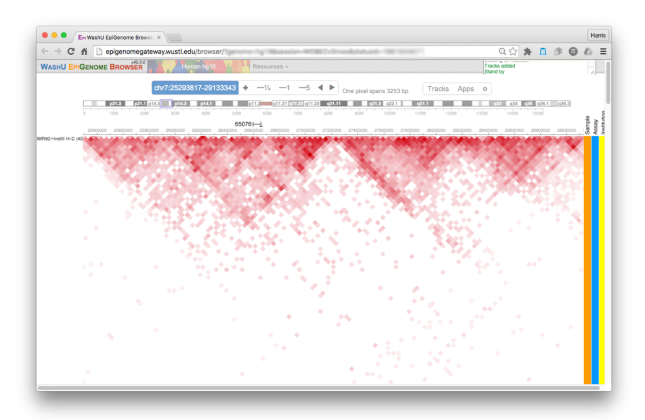


Figure 5: WashU tracks loaded in browser.

## 4.6 Matrix Filtered

#### 4.6.1 Input

Data from the pipeline filter step is used as input (Section 4.3).

#### 4.6.2 Analysis

Default parameters files:

```
-rwxr-xr-x 1 at570 195 Nov 24 11:20 params.res_1000kb.tcsh
-rwxr-xr-x 1 at570 194 Nov 25 15:11 params.res_100kb.tcsh
-rwxr-xr-x 1 at570 210 Dec 1 12:41 params.res_10kb.maxd_5Mb.rotate45.tcsh
-rwxr-xr-x 1 at570 193 Nov 30 16:22 params.res_10kb.tcsh
5 rwxr-xr-x 1 at570 193 Nov 24 11:20 params.res_40kb.tcsh
```

#### Default parameters:

```
params.res_1000kb.tcsh$
#!/bin/tcsh

source ./inputs/params/params.tcsh

set bin_size = 1000000
7 set max_dist = 0
set ref = $genome_dir/genome.bed
9 set matrix_params = "—bin-size $bin_size —max-dist $max_dist -R $ref"
```

```
params.res_100kb.tcsh$
#!/bin/tcsh

source ./inputs/params/params.tcsh

set bin_size = 100000

rest max_dist = 0
    set ref = $genome_dir/genome.bed
    set matrix_params = "—bin_size $bin_size —max_dist $max_dist -R $ref"
```

```
params.res_10kb.maxd_5Mb.rotate45.tcsh$

#!/bin/tcsh

source ./inputs/params/params.tcsh

set bin_size = 10000

set max_dist = 5000000

set ref = $genome_dir/genome.bed

set matrix_params = "__bin_size $bin_size __max_dist __rotate45 _R $ref"
```

```
params.res_10kb.tcsh
#!/bin/tcsh

source ./inputs/params/params.tcsh

set bin_size = 10000

set max_dist = 0
    set ref = $genome_dir/genome.bed
    set matrix_params = "__bin_size $bin_size __max_dist = R $ref"
```

```
params.res_40kb.tcsh$
#!/bin/tcsh

source ./inputs/params/params.tcsh
```

```
5
    set bin_size = 40000
7    set max_dist = 0
    set ref = $genome_dir/genome.bed
9    set matrix_params = "__bin_size $\int bin_size \text{-max_dist } \name R $\reft{ref}"
```

## 4.6.3 Output

```
at570 56K Jan 13 15:57 ignored_loci.txt
at570 9.6K Jan 13 16:02 job.err
-rw-r---r-
-rw-r---r-
-rw-r---r-
              at570
                       47 Jan 13 15:57 job.id
_rw_r __r __
              at570
                       0 Jan 13 15:57
                                       job.out
                     266 Jan 13 15:57
              at570
-rw-r---r--
                                       job.sh
-rw-r---r--
              at570 2.7K Jan 13 16:02 job.vars.tsv
                      75M Jan 13 16:01 matrix.chr1.tsv
-rw-r---r--
              at570
-rw-r---r--
              at570
                      23M Jan 13 16:01 matrix.chr10.tsv
-rw-r---r--
              at570
                      22M Jan 13 16:01 matrix.chr11.tsv
              at570
                      22M Jan 13 16:01 matrix.chr12.tsv
_rw_r__r_
                      16M Jan 13 16:01 matrix.chr13.tsv
-rw-r---r--
              at570
              at570
                      14M Jan 13 16:01 matrix.chr14.tsv
-rw-r---r--
                      13M Jan 13 16:01 matrix.chr15.tsv
-rw-r---r--
              at570
              at570 9.9M Jan 13 16:01 matrix.chr16.tsv
-rw-r---r--
-rw-r--r-
              at570 8.0M Jan 13 16:01 matrix.chr17.tsv
-rw-r---r--
              at570 7.4M Jan 13 16:01 matrix.chr18.tsv
-rw-r---r--
              at570 4.3M Jan 13 16:01 matrix.chr19.tsv
-rw-r---r--
              at570
                     71M Jan 13 16:01 matrix.chr2.tsv
-rw-r---r--
              at570 4.9M Jan 13 16:01 matrix.chr20.tsv
-rw-r---r--
              at570 2.9M Jan 13 16:01 matrix.chr21.tsv
-rw-r---r--
              at570 3.3M Jan 13 16:01 matrix.chr22.tsv
-rw-r---r--
              at570
                      48M Jan 13 16:01 matrix.chr3.tsv
-rw-r--r--
              at570
                      44M Jan 13 16:01 matrix.chr4.tsv
-rw-r---r--
              at570
                      40M Jan 13 16:01 matrix.chr5.tsv
-rw-r--r--
               at570
                      36M Jan 13 16:02 matrix.chr6.tsv
-rw-r---r--
              at570
                      31M Jan 13 16:02 matrix.chr7.tsv
-rw-r--r--
               at570
                      26M Jan 13 16:02 matrix.chr8.tsv
-rw-r---r
               at570
                      24M Jan 13 16:02 matrix.chr9.tsv
-rw-r--r--
               at570
                       29 Jan 13 16:02 matrix.chrM.tsv
-rw-r---r
              at570
                     29M Jan 13 16:02 matrix.chrX.tsv
              at570 4.3M Jan 13 16:02 matrix.chrY.tsv
```

# 4.7 Matrix Prep

#### 4.7.1 Input

Data from the pipeline matrix-filtered step is used as input (Section 4.6).

### 4.7.2 Analysis

#### Default parameters:

```
params.scale_impute.tcsh$
#!/bin/tcsh

source ./inputs/params/params.tcsh

set chrom_excluded = 'chr[MY]'  # excluded chromosomes
set prep_params = "—scale —impute"
```

#### **4.7.3** Output

```
drwxr-xr-x 2 at570 3.4K Jan 13 16:16
  -rw-r---r-
              1 at570 4.0K Jan 13 16:18 job.err
  -rw-r---r-
              1
                at570
                        47 Jan 13 16:14
                                        job.id
  -rw-r---r-
                at570
                         0 Jan 13 16:15 job.out
  -rw-r-r-
              1 at570 345 Jan 13 16:14 job.sh
  -rw-r--- 1
                at570 3.3K Jan 13 16:18 job.vars.tsv
  -rw-r---r--
                at570 371M Jan 13 16:17 matrix chr1 tsv
  -rw-r--- 1
                at570 110M Jan 13 16:16 matrix.chr10.tsv
  -rw-r---r
                at570 109M Jan 13 16:15 matrix.chr11.tsv
  -rw-r--- 1
                at570 107M Jan 13 16:16 matrix.chr12.tsv
                at570
                       80M Jan 13 16:16 matrix.chr13.tsv
  -rw-r---r
  -rw-r---r
                at570
                       69M Jan 13 16:16 matrix.chr14.tsv
  -rw-r---r
                at570
                       63M Jan 13 16:16 matrix.chr15.tsv
                at570
                       49M Jan 13 16:16 matrix.chr16.tsv
  -rw-r---r-
                at570
                       40M Jan 13 16:16 matrix.chr17.tsv
15
                at570
                       37M Jan 13 16:16 matrix.chr18.tsv
   -rw-r---r
17
  -rw-r---r
                at570
                       21M Jan 13 16:16 matrix.chr19.tsv
  -rw-r---r
                at570 353M Jan 13 16:17 matrix.chr2.tsv
                       24M Jan 13 16:16 matrix.chr20.tsv
  -rw-r---r--
                at570
19
                       14M Jan 13 16:16 matrix.chr21.tsv
  -rw-r---r-
                at570
                       16M Jan 13 16:16 matrix.chr22.tsv
  -rw-r---r--
                at570
                at570 234M Jan 13 16:17 matrix.chr3.tsv
  -rw-r---r-
                at570 219M Jan 13 16:17 matrix.chr4.tsv
  -rw-r--r--
  -rw-r---r--
                at570 196M Jan 13 16:17 matrix.chr5.tsv
  -rw-r-r-
                at570 175M Jan 13 16:17 matrix.chr6.tsv
  -rw-r---r--
                at570 152M Jan 13 16:17 matrix.chr7.tsv
                at570 128M Jan 13 16:17 matrix.chr8.tsv
  -rw-r---r--
                at570 120M Jan 13 16:17 matrix.chr9.tsv
  _rw_r__r_
  -rw-r-r 1 at570 144M Jan 13 16:17 matrix.chrX.tsv
```

### 4.8 Matrix IC

#### 4.8.1 Input

Data from the pipeline matrix-filtered step is used as input (Section 4.6).

### 4.8.2 Analysis

#### Default parameters:

```
params.standard.tcsh$
#!/bin/tcsh

source ./inputs/params/params.tcsh

module unload gcc  # this is necessary in order to take care of module conflicts in our system module unload python module load python/2.7.3

set chrom_excluded = 'chr[MY]'  # excluded chromosomes set cutoff = 0.05
```

### 4.8.3 Output

```
drwxr-xr-x 2 at570 3.4K Jan 13 16:50
   -rw-r---r
                at570 1.4K Jan 13 16:52 job.err
  -rw-r---r-
                at570
                        47 Jan 13 16:47
                                        job.id
  -rw-r---r-
                at570
                         0 Jan 13 16:47 job.out
  -rw-r---r-
                at570
                       333 Jan 13 16:47
                                        job.sh
  -rw-r--- 1
                at570 3.5K Jan 13 16:52 job.vars.tsv
                at570 371M Jan 13 16:50 matrix.chr1.tsv
                at570 110M Jan 13 16:49 matrix.chr10.tsv
  -rw-r---r-
                at570 109M Jan 13 16:49 matrix.chr11.tsv
                at570 107M Jan 13 16:49 matrix.chr12.tsv
                       80M Jan 13 16:49 matrix.chr13.tsv
  -rw-r---r
                at570
11
                at570
                       69M Jan 13 16:49 matrix.chr14.tsv
  -rw-r---r
13
  -rw-r---r
                at570
                       63M Jan 13 16:49 matrix.chr15.tsv
  _rw_r__r
                at570
                       49M Jan 13 16:49 matrix.chr16.tsv
                       40M Jan 13 16:49 matrix.chr17.tsv
  -rw-r---r-
                at570
15
                       37M Jan 13 16:49 matrix.chr18.tsv
  -rw-r---r
                at570
                       21M Jan 13 16:49 matrix.chr19.tsv
  -rw-r---r--
                at570
                at570 353M Jan 13 16:52 matrix.chr2.tsv
  -rw-r---r-
                at570
                       24M Jan 13 16:49 matrix.chr20.tsv
19
  -rw-r--r--
  -rw-r---r--
                at570 14M Jan 13 16:50 matrix.chr21.tsv
  -rw-r-r-
                at570
                       16M Jan 13 16:50 matrix.chr22.tsv
  -rw-r---r-
                at570 234M Jan 13 16:51 matrix.chr3.tsv
                at570 219M Jan 13 16:51 matrix.chr4.tsv
  -rw-r---r--
23
  -rw-r---r-
                at570 196M Jan 13 16:51 matrix.chr5.tsv
                at570 175M Jan 13 16:51 matrix.chr6.tsv
  -rw-r--r-
                at570 152M Jan 13 16:51 matrix.chr7.tsv
  -rw-r---r-
                at570 128M Jan 13 16:51 matrix.chr8.tsv
  -rw-r---r--
                at570 120M Jan 13 16:51 matrix.chr9.tsv
  -rw-r---r--
  -rw-r-r- 1 at570 144M Jan 13 16:51 matrix.chrX.tsv
```

### 4.9 Matrix HiCNorm

#### 4.9.1 Input

Data from the pipeline matrix-filtered step is used as input (Section 4.6).

### 4.9.2 Analysis

### Default parameters:

```
params.standard.tcsh$
#!/bin/tcsh

source ./inputs/params/params.tcsh

set chrom_excluded = 'chr[MY]' # excluded chromosomes
```

#### 4.9.3 Output

```
2 at570 3.4K Feb 8 17:10
drwxr-xr-x
                                       _jdata
-rw-r---r--
             at570
                    13K Feb
                            8 17:25
                                      job.err
-rw-r--r 1
              at570
                     47 Feb 8 17:06 job.id
-rw-r---r--
              at570
                      0 Feb
                             8 17:06
                                     job.out
-rw-r-r-
              at570
                    343 Feb
                            8 17:06 job.sh
-rw-r---r
              at570 3.2K Feb
                             8 17:25 job.vars.tsv
-rw-r-r-- 1
              at570 86M Feb 8 17:19 matrix.chr1.tsv
-rw-r---r
              at570
                    28M Feb
                             8 17:10 matrix.chr10.tsv
-rw-r--r--
              at570
                    28M Feb
                             8 17:12 matrix.chr11.tsv
-rw-r---r
              at570
                    28M Feb
                             8 17:10 matrix.chr12.tsv
-rw-r---r--
              at570
                    21M Feb
                            8 17:09 matrix.chr13.tsv
              at570
                    18M Feb
                             8 17:09 matrix.chr14.tsv
-rw-r---r
              at570
                    16M Feb
                             8 17:09 matrix.chr15.tsv
-rw-r---r
              at570
                     13M Feb
                             8 17:09 matrix.chr16.tsv
              at570 9.7M Feb
                             8 17:09 matrix.chr17.tsv
-rw-r---r
              at570
                    11M Feb
                             8 17:09 matrix.chr18.tsv
-rw-r---r
              at570 5.3M Feb
                             8 17:08 matrix.chr19.tsv
-rw-r---r
              at570
                    85M Feb
                             8 17:25 matrix.chr2.tsv
-rw-r-r-
              at570 6.6M Feb
                             8 17:10 matrix.chr20.tsv
                             8 17:09 matrix.chr21.tsv
-rw-r---r
              at570 3.6M Feb
-rw-r-r-
              at570 3.8M Feb
                             8 17:09 matrix.chr22.tsv
                    58M Feb
                             8 17:17 matrix.chr3.tsv
-rw-r---r-
              at570
-rw-r---r-
              at570
                    55M Feb
                             8 17:20 matrix.chr4.tsv
              at570
                    49M Feb
                             8 17:15 matrix.chr5.tsv
-rw-r---r--
-rw-r---r-
              at570
                    44M Feb
                             8 17:15 matrix.chr6.tsv
-rw-r---r
              at570
                    38M Feb
                             8 17:17 matrix.chr7.tsv
-rw-r--r-
              at570
                    33M Feb
                             8 17:14 matrix.chr8.tsv
                    29M Feb
                             8 17:13 matrix.chr9.tsv
              at570
-rw-r---r--
                    34M Feb
-rw-r---r--
              at570
                             8 17:15 matrix.chrX.tsv
```

## 4.10 Matrix Stats

#### 4.10.1 Input

Data from the pipeline steps matrix-filtered (Section 4.6), matrix-hicnorm (Section 4.9), matrix-prep (Section 4.7), and matrix-ic (Section 4.8) are used as input.

# 4.10.2 Analysis

## Default parameters:

```
params.standard.tcsh$
#!/bin/tcsh

source ./inputs/params/params.tcsh

set chrom_excluded = 'chr[MY]' # excluded chromosomes
```

## 4.10.3 Output

## See Figure 6. Default output:

```
1 -rw-r - 1 at570 39K Feb 11 16:11 job.err

-rw-r - 1 at570 47 Feb 11 15:48 job.id

3 -rw-r - 1 at570 0 Feb 11 15:48 job.out

-rw-r - 1 at570 480 Feb 11 15:48 job.sh

5 -rw-r - 1 at570 5.3K Feb 11 16:11 job.vars.tsv

-rw-r - 1 at570 59K Feb 11 16:11 stats.pdf
```

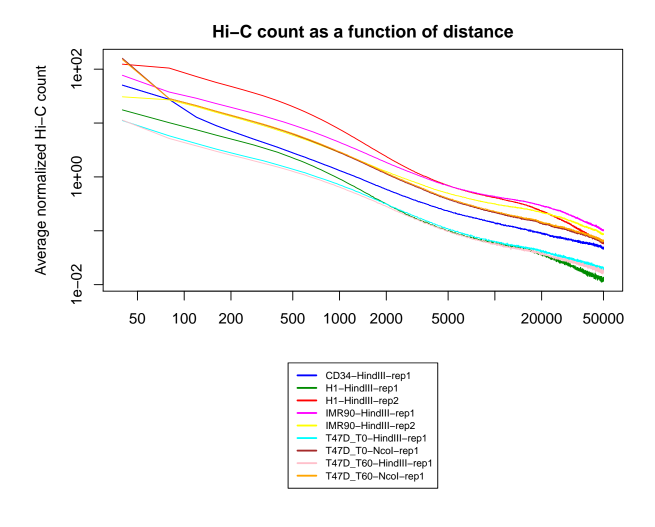


Figure 6: Matrix Stats sample output

## 4.11 Compare Matrices

### 4.11.1 Input

Data from the pipeline steps matrix-filtered (Section 4.6), matrix-hicnorm (Section 4.9), matrix-prep (Section 4.7), and matrix-ic (Section 4.8) are used as input.

## 4.11.2 Analysis

#### Default parameters:

```
params.standard.tcsh$

#!/bin/tcsh

source ./inputs/params/params.tcsh

set chrom_excluded = 'chr[MYX]'  # excluded chromosomes

set max_dist = `echo 10000000/$bin_size | bc`  # number of bins (max distance = 10Mb)

set compare_params = "—max-dist=$max_dist —n-dist=1 —min-lambda=0.0 —max-lambda=1.0 —n-lambda=6 —gamma=0"
  # only used if estimation was done with max-lambda=Inf
```

#### 4.11.3 Output

```
at570
                                9 01:26 chr1.cor.pearson.tsv
-rw-r---r--
                        17 Feb
                        17 Feb
             1 at570
                                9 01:26 chr1.cor.spearman.tsv
-rw-r---r--
                        17 Feb
-rw-r---r--
               at570
                                9 01:27 chr10.cor.pearson.tsv
             1 at570
                        17 Feb
                                9 01:27 chr10.cor.spearman.tsv
_rw_r ___ r ___
                        17 Feb
-rw-r---r--
               at570
                                9 01:28 chr11.cor.pearson.tsv
                       17 Feb
                                9 01:28 chr11.cor.spearman.tsv
_rw_r__r__
             1 at570
_rw_r__r_
               at570
                       17 Feb 9 01:28 chr12.cor.pearson.tsv
-rw-r---r--
             1 at 570
                       17 Feb
                               9 01:28 chr12.cor.spearman.tsv
               at570
                       17 Feb 9 01:29 chr13.cor.pearson.tsv
_rw_r__r_
                       17 Feb
-rw-r---r---
             1 at 570
                               9 01:29 chr13.cor.spearman.tsv
_rw_r__r_
               at570
                       17 Feb
                                9 01:29 chr14.cor.pearson.tsv
                       17 Feb
_rw_r ___ r ___
             1 at 570
                                9 01:29 chr14.cor.spearman.tsv
                       17 Feb
17 Feb
_rw_r__r_
               at570
                               9 01:30 chr15.cor.pearson.tsv
_rw_r ___ r ___
             1 at570
                                9 01:30 chr15.cor.spearman.tsv
_rw_r__r
               at570
                       17 Feb
                                9 01:30 chr16.cor.pearson.tsv
             1
-rw-r---r--
             1 at 570
                        17 Feb
                                9 01:30 chr16.cor.spearman.tsv
_rw_r__r_
             1
               at570
                       17 Feb 9 01:31 chr17.cor.pearson.tsv
                       17 Feb
-rw-r---r--
             1 at570
                                9 01:31 chr17.cor.spearman.tsv
-rw-r---r--
               at570
                       17 Feb
                                9 01:31 chr18.cor.pearson.tsv
               at570
                       17 Feb
                                9 01:31 chr18.cor.spearman.tsv
-rw-r---r--
-rw-r---r--
               at570
                       17 Feb
                                9 01:31 chr19.cor.pearson.tsv
                       17 Feb
_rw_r__r
             1 at570
                                9 01:31 chr19.cor.spearman.tsv
-rw-r---r
               at570
                       17 Feb
                                9 01:33 chr2.cor.pearson.tsv
               at570
                        17 Feb
                                9 01:33 chr2.cor.spearman.tsv
-rw-r---r
                                9 01:33 chr20.cor.pearson.tsv
-rw-r---r
               at570
                       17 Feb
-rw-r---r
                                9 01:33 chr20.cor.spearman.tsv
             1 at570
                       17 Feb
               at570
                       17 Feb
                                9 01:33 chr21 cor pearson tsv
               at570
                        17 Feb
                                9 01:33 chr21.cor.spearman.tsv
               at570
                       17 Feb
                                9 01:33 chr22.cor.pearson.tsv
                                9 01:33 chr22.cor.spearman.tsv
               at570
                        17 Feb
-rw-r---r--
             1 at570
                       17 Feb
                                9 01:35 chr3.cor.pearson.tsv
                                9 01:35 chr3.cor.spearman.tsv
-rw-r---r
               at570
                        17 Feb
               at570
                       17 Feb
                                9 01:36 chr4.cor.pearson.tsv
                        17 Feb
                                9 01:36 chr4.cor.spearman.tsv
-rw-r--r-
               at570
_rw_r__r
               at570
                       17 Feb
                                9 01:37 chr5.cor.pearson.tsv
-rw-r---r--
               at570
                        17 Feb
                                9 01:37 chr5.cor.spearman.tsv
                       17 Feb
                                9 01:38 chr6.cor.pearson.tsv
_rw_r__r_
               at570
                       17 Feb
                                9 01:38 chr6.cor.spearman.tsv
-rw-r---r--
               at570
-rw-r--r-
             1 at570
                       17 Feb
                                9 01:39 chr7.cor.pearson.tsv
-rw-r---r--
               at570
                        17 Feb
                                9 01:39 chr7.cor.spearman.tsv
               at570
                       17 Feb
-rw-r---r
                                9 01:40 chr8.cor.pearson.tsv
-rw-r---r
               at570
                        17 Feb
                                9 01:40 chr8.cor.spearman.tsv
-rw-r---r--
             1 at570
                        17 Feb
                               9 01:41 chr9.cor.pearson.tsv
-rw-r---r--
             1 at570
                        17 Feb
                                9 01:41 chr9.cor.spearman.tsv
             1 at570 1.9K Feb 9 01:41 cor.pearson.tsv
-rw-r---r--
```

# 4.12 Compare Matrices Stats

## 4.12.1 Input

Data from the pipeline compare-matrices step is used as input (Section 4.11).

### 4.12.2 Analysis

### Default parameters:

```
params.standard.tcsh$
#!/bin/tcsh

source ./inputs/params/params.tcsh
```

# 4.12.3 Output

See Figure 7, and See Figure 8. Default output:

```
spearman/$
-rw-r — 1 at570 161K Feb 12 11:25 cor.spearman.tsv
-rw-r — 1 at570 8.9K Feb 12 11:25 correlograms.pdf
-rw-r — 1 at570 12K Feb 12 11:25 summary.tsv
```

```
pearson/$
2 -rw-r-r- 1 at570 159K Feb 12 11:25 cor.pearson.tsv
-rw-r-r- 1 at570 9.0K Feb 12 11:25 correlograms.pdf
4 -rw-r-r- 1 at570 12K Feb 12 11:25 summary.tsv
```

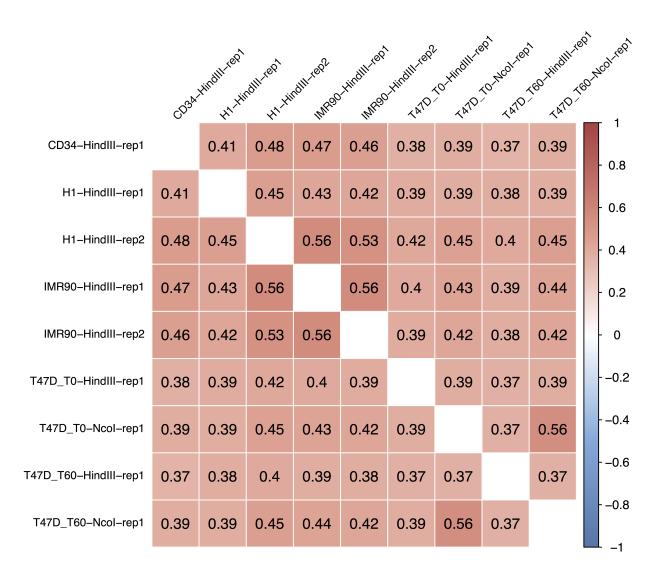


Figure 7: Compare Matrices Stats Spearman sample correlograms. See Section 4.12.

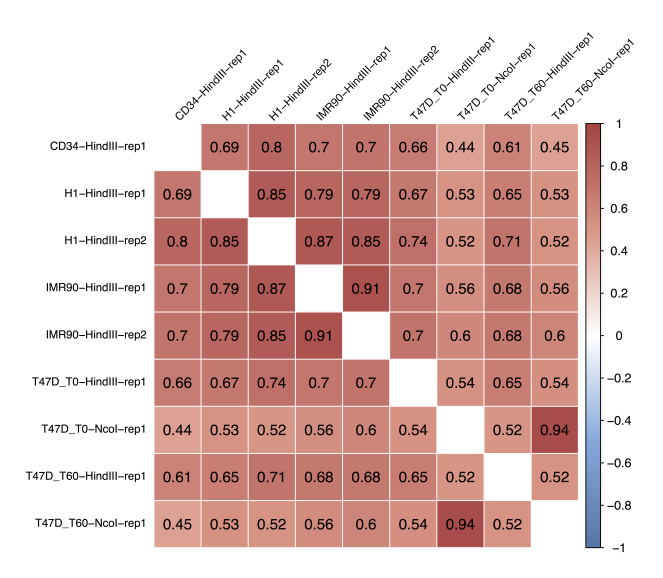


Figure 8: Compare Matrices Stats Pearson sample correlograms. See Section 4.12.

# 4.13 Boundary Scores

### 4.13.1 Input

Data from the pipeline steps matrix-filtered (Section 4.6), matrix-hicnorm (Section 4.9), matrix-prep (Section 4.7), and matrix-ic (Section 4.8) are used as input.

#### 4.13.2 Analysis

#### Default parameters:

```
params.standard.tcsh$
#!/bin/tcsh
source ./inputs/params/params.tcsh
set chrom_excluded = 'chr[MYX]'
                                                                                      # excluded chromosomes
set boundary_scores_params = ( \
 -min-lambda=0.0 -max-lambda=1.0 -n-lambda=6 -gamma=0 \
--preprocess=none \
-distance=`echo 500000/$bin_size | bc` \
-distance2=`echo 500000/$bin_size | bc` \
  -skip-distance=0 \
—flank-dist=`echo 500000/$bin_size | bc` \
 -tolerance=0.01 \
--alpha=0.50 \
 -track-dist=`echo 2000000/$bin_size | bc` \
--presentation=none \
```

# 4.13.3 Output

```
-rw-r - 1 at570 9.0M Feb 15 14:25 all_scores.k=001.tsv
-rw-r - 1 at570 17K Feb 15 14:25 job.err

-rw-r - 1 at570 47 Feb 15 14:07 job.id
-rw-r - 1 at570 0 Feb 15 14:11 job.out
-rw-r - 1 at570 345 Feb 15 14:07 job.sh
-rw-r - 1 at570 3.2K Feb 15 14:25 job.vars.tsv
```

# 4.14 Boundary Scores PCA

### 4.14.1 Input

Data from the pipeline boundary-scores step is used as input (Section 4.13).

### 4.14.2 Analysis

### Default parameters:

```
params.standard.tcsh

#!/bin/tcsh

source ./inputs/params/params.tcsh

set chrom_excluded = 'chr[MYX]'  # excluded chromosomes

set group_var = 'cell_type'  # grouping variable (from sample sheet) to be used for color assignment)
```

# 4.14.3 Output

### See Figure 9. Default output:

```
-rw-r-r- 1 at570 4.1K Feb 15 15:20 job.err
-rw-r-r-1 at 570
                   47 Feb 15 15:18 job.id
-rw-r-r- 1 at570 936 Feb 15 15:20
-rw-r-r- 1 at570 564 Feb 15 15:18 job.sh
-rw-r-r- 1 at570 4.8K Feb 15 15:20 job.vars.tsv
-rw-r-r- 1 at570 211 Feb 15 15:18 labels.tsv
-rw-r-r- 1 at570 4.4K Feb 15 15:19 pca.Dl.k=001.pdf
-rw-r-r 1 at570 4.4K Feb 15 15:19 pca.Dl2.k=001.pdf
-rw-r-r- 1 at570 4.4K Feb 15 15:19 pca.diff.k=001.pdf
-rw-r-r 1 at570 4.4K Feb 15 15:20 pca. diffratio.k=001.pdf
-rw-r-r- 1 at570 4.4K Feb 15 15:19 pca.inter.k=001.pdf
-rw-r-r- 1 at570 4.4K Feb 15 15:18 pca.intra-left.k=001.pdf
-rw-r-r- 1 at570 4.4K Feb 15 15:19 pca.intra-max.k=001.pdf
-rw-r-r- 1 at570 4.4K Feb 15 15:19 pca.intra-min.k=001.pdf
-rw-r-r- 1 at570 4.4K Feb 15 15:18 pca.intra-right.k=001.pdf
-rw-r-r- 1 at570 4.4K Feb 15 15:20 pca.novel-max.k=001.pdf
-rw-r-r- 1 at570 4.4K Feb 15 15:20 pca.novel-min.k=001.pdf
-rw-r-r- 1 at570 4.4K Feb 15 15:19 pca.ratio.k=001.pdf
```

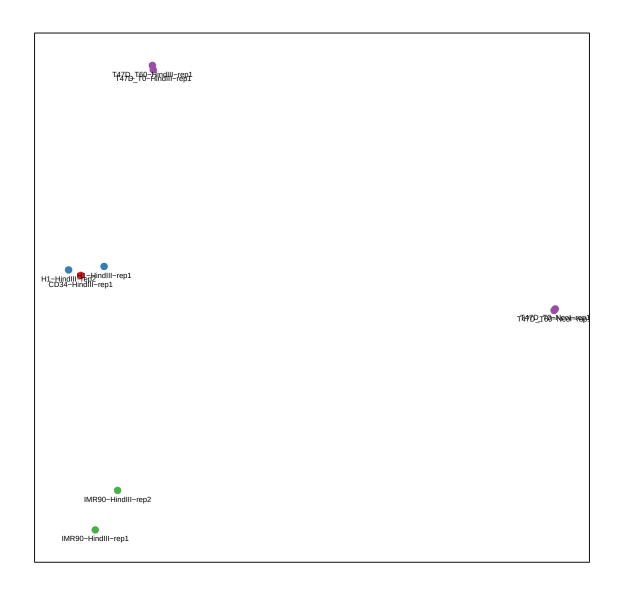


Figure 9: Boundary Scores PCA sample output. See Section 4.14.

### 4.15 Domains

#### 4.15.1 Input

Data from the pipeline steps matrix-filtered (Section 4.6), matrix-hicnorm (Section 4.9), matrix-prep (Section 4.7), and matrix-ic (Section 4.8) are used as input.

# 4.15.2 Analysis

### Default parameters:

```
params.armatus.gamma_0.5.tcsh$

#!/bin/tcsh

source ./inputs/params/params.tcsh

set tool = armatus
set chrom_excluded = 'chr[MY]'
set armatus_params = "-g 0.5"
```

```
params.hicmatrix.tcsh$
  #!/bin/tcsh
  source ./inputs/params/params.tcsh
  set tool = hicmatrix
  set chrom_excluded = 'chr[MY]'
10 set hicmatrix_params = ( \
   -min-lambda=0.0 --max-lambda=1.0 --n-lambda=6 --gamma=0 \
  ---preprocess=none \
   -method=ratio \
  --distance='echo 500000/$bin size | bc' \
   -distance2=`echo 500000/$bin_size | bc` \
  --skip-distance=0 \
   -flank-dist=`echo 500000/$bin_size | bc` \
  --tolerance=0.01 \
  —alpha=0.25 \
—track-dist=`echo 2000000/$bin_size | bc` \
    -presentation=none \
```

```
params.topdom.tcsh$

#!/usr/bin/tcsh

source ./inputs/params/params.tcsh

set tool = topdom
    set topdompath = "./code/TopDom.R"

set chrom_excluded = 'chr[MY]'
    set winsize = 5
```

# 4.15.3 Output

```
-rw-r - 1 at570 288K Feb 15 16:31 domains.k=001.bed
-rw-r - 1 at570 28K Feb 15 16:31 job.err

-rw-r - 1 at570 47 Feb 15 16:13 job.id
-rw-r - 1 at570 5.6K Feb 15 16:31 job.out

-rw-r - 1 at570 347 Feb 15 16:13 job.sh
-rw-r - 1 at570 2.7K Feb 15 16:31 job.vars.tsv
```

# 4.16 Compare Boundaries

## 4.16.1 Input

Data from the pipeline domains step is used as input (Section 4.15).

### 4.16.2 Analysis

## Default parameters:

```
params.standard.tcsh$
2 #!/bin/tcsh

4 source ./inputs/params/params.tcsh
6 set flank_dist = $bin_size
    set black_lists = ($genome_dir/centrotelo.bed)
```

#### 4.16.3 Output

```
-rw-r---r--
            1 at570 573K Feb 16 00:18 boundaries1.k=001.bed
-rw-r---r-
            1 at570 573K Feb 16 00:18 boundaries2.k=001.bed
-rw-r---r--
            1 at570 268K Feb 16 00:18 common_boundaries.k=001.bed
-rw-r---r-
           1 at570 154 Feb 16 00:18 comparison.tsv
-rw-r-r-
            1 at570 268K Feb 16 00:18 intersection k=001.bed
-rw-r-r-
           1 at570 70 Feb 16 00:18 job.err
-rw-r--r--
            1 at570
                     47 Feb 16 00:18 job.id
-rw-r-r-
            1 at570
                     0 Feb 16 00:18 job.out
            1 at570 456 Feb 16 00:18 job.sh
           1 at570 4.5K Feb 16 00:18 job.vars.tsv
           1 at570 268K Feb 16 00:18 union.k=001.bed
```

# 4.17 Compare Boundaries Stats

# 4.17.1 Input

Data from the pipeline compare-boundaries step is used as input (Section 4.16).

### 4.17.2 Analysis

### Default parameters:

```
params.standard.tcsh$
#!/bin/tcsh
source ./inputs/params/params.tcsh
```

# 4.17.3 Output

## See Figure 11 and Figure 10. Default output:

```
-rw-r - 1 at570 6.9K Feb 12 12:52 comparisons.tsv
2 -rw-r - 1 at570 27K Feb 12 12:52 correlograms.pdf
-rw-r - 1 at570 238 Feb 12 12:52 job.err
4 -rw-r - 1 at570 47 Feb 12 12:51 job.id
-rw-r - 1 at570 52 Feb 12 12:52 job.out
6 -rw-r - 1 at570 3.5K Feb 12 12:51 job.sh
-rw-r - 1 at570 51K Feb 12 12:52 job.vars.tsv
8 -rw-r - 1 at570 5.6K Feb 12 12:52 raw_comparisons.pdf
```

# **Number of boundaries**

	CD34-HindIII-rep1	H1-HindIII-rep1	H1-HindIII-rep2	IMR90-HindIII-rep1	IMR90-HindIII-rep2	T47D_T0-HindIII-rep1	T47D_T0-Ncol-rep1	T47D_T60-HindIII-rep1	T47D_T60-Ncol-rep1
CD34-HindIII-rep1	9158	5231	5277	4045	4157	5006	3532	4869	3546
H1-HindIII-rep1	5231	7258	4724	3860	4059	4481	3215	4481	3232
H1-HindIII-rep2	5277	4724	6564	3615	3719	4186	2860	4149	2890
IMR90-HindIII-rep1	4045	3860	3615	5464	4291	3539	2666	3576	2678
IMR90-HindIII-rep2	4157	4059	3719	4291	5800	3767	2813	3749	2823
T47D_T0-HindIII-rep1	5006	4481	4186	3539	3767	7364	3252	5027	3264
T47D_T0-Ncol-rep1	3532	3215	2860	2666	2813	3252	5417	3205	4467
T47D_T60-HindIII-rep1	4869	4481	4149	3576	3749	5027	3205	7215	3234
T47D_T60-Ncol-rep1	3546	3232	2890	2678	2823	3264	4467	3234	5401

Figure 10: Example raw comparisons. See Section 4.17.

# Overlap (%)

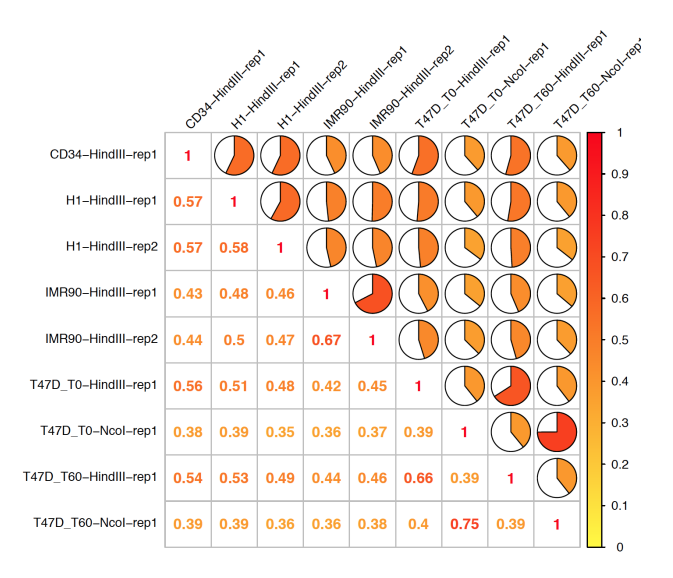


Figure 11: Example correlograms. See Section 4.17.

### 4.18 HiC Plotter

### 4.18.1 Input

Data from the pipeline steps matrix-filtered (Section 4.6), matrix-hicnorm (Section 4.9), matrix-prep (Section 4.7), and matrix-ic (Section 4.8) are used as input.

#### 4.18.2 Analysis

#### Default parameters:

```
params.standard.tcsh$
  #!/bin/tcsh
  source ./inputs/params/params.tcsh
  # HiCplotter path
  set hicplotter_path = ./code/HiCPlotter2.py
  # create bedgraphs for boundary scores
  set bscores_branch = ../boundary-scores/results/boundary-scores.by_sample.standard/`echo $branch | sed 's/.*
       results\///'
  set cell_type = `echo $objects[1] | cut -d'-' -f1`
  set f = $bscores_branch/$objects[1]/all_scores.k=001.tsv
  set methods = (intra-max DI ratio)
  set bedgraphs = ()
  set bedgraph_labels = ($methods)
  18
    set bedgraphs = ($bedgraphs $outdir/bscores.$m.bedGraph)
20
  # add CTCF ChIP-seq
  if (-e inputs/data.external/$cell_type/CTCF.bedGraph) then
    set bedgraphs = ($bedgraphs inputs/data.external/$cell_type/CTCF.bedGraph)
   set bedgraph_labels = ($bedgraph_labels CTCF)
  endif
26
  # regions to plot
set regions = "chr8:125000000-133000000"
28
  set tiles = "params/regions.bed"
set tiles_labels = "regions"
  set highlight = 1
  set highlight_bed = "params/highlight.bed"
set fileheader = 0  # Either 1 or 0
                            # Either 1 or 0 (header / no header)
  set insulation_score = 0  # Either 1 or 0 (include insulation index or not)
```

# 4.18.3 Output

#### See Figure 12. Default output:

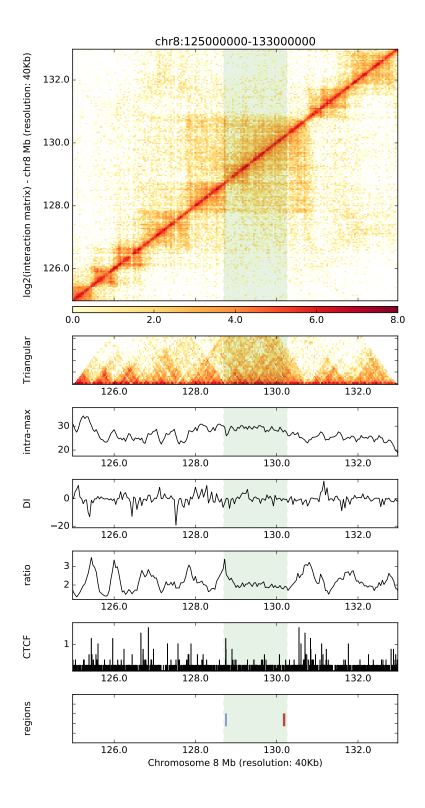


Figure 12: HiCPlotter sample output Page 50 of 62

### 4.19 Interactions

#### 4.19.1 Input

Data from the pipeline matrix-filtered step is used as input (Section 4.6).

### 4.19.2 Analysis

### Default parameters:

```
#!/bin/tcsh
source ./inputs/params/params.tcsh
set chrom_excluded = 'chr[MYX]'  # excluded chromosomes
set loop_params = "--bin-size=$bin_size --lambda-id=6 --rpk2b-cutoff=1.0 --loop-cutoff=4.0 --min-distance=40000"
# parameters for identifying significant interactions
```

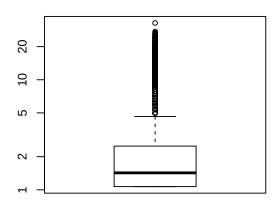
### 4.19.3 Output

### See Figure 13. Default output:

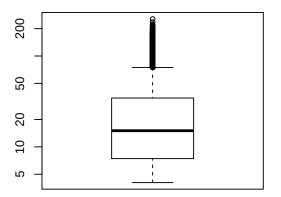
```
drwxr-xr-x 2 at570 3.3K Feb
                              5 10:12
                                        _jdata
-rw-r---r
              at570 4.8K Feb
                              5 10:17 job.err
-rw-r---r--
              at570
                     47 Feb
                             5 10:11 job.id
-rw-r---r
              at570
                      0 Feb
                              5 10:11 job.out
-rw-r-r-
           1
              at570 375 Feb
                             5 10:11 job.sh
-rw-r---r
              at570 3.6K Feb
                              5 10:17 job.vars.tsv
drwxr-xr-x 2 at570
                     54 Feb
                             5 10:15 matrix.chr1
drwxr-xr-x
            2 at570
                     54 Feb
                              5 10:14 matrix.chr10
drwxr-xr-x 2 at570
                     54 Feb
                             5 10:14 matrix.chr11
           2 at570
                     54 Feb
                              5 10:14 matrix.chr12
drwxr-xr-x
            2 at570
                     54 Feb
                              5 10:13 matrix.chr13
drwxr-xr-x
drwxr-xr-x
            2 at570
                     54 Feb
                              5 10:13 matrix.chr14
drwxr-xr-x 2 at570
                     54 Feb
                             5 10:13 matrix.chr15
drwxr-xr-x
           2 at570
                     54 Feb
                              5 10:13 matrix.chr16
           2 at570
                     54 Feb
                              5 10:13 matrix.chr17
drwxr-xr-x
drwxr-xr-x
           2 at570
                     54 Feb
                              5 10:13 matrix.chr18
drwxr-xr-x
           2 at570
                     54 Feb
                             5 10:13 matrix.chr19
drwxr-xr-x
           2 at570
                     54 Feb
                              5 10:16 matrix.chr2
drwxr-xr-x
            2 at570
                     54 Feb
                              5 10:13 matrix.chr20
drwxr-xr-x 2 at570
                     54 Feb
                              5 10:12 matrix.chr21
drwxr-xr-x 2 at570
                     54 Feb
                              5 10:13 matrix.chr22
drwxr-xr-x
           2 at570
                     54 Feb
                              5 10:15 matrix.chr3
drwxr-xr-x
            2 at570
                     54 Feb
                              5 10:15 matrix.chr4
drwxr-xr-x
           2 at570
                     54 Feb
                              5 10:15 matrix.chr5
drwxr-xr-x
            2 at570
                     54 Feb
                              5 10:15 matrix.chr6
           2 at570
                     54 Feb
drwxr-xr-x
                              5 10:14 matrix.chr7
            2 at570
                      54 Feb
                              5 10:14 matrix.chr8
drwxr-xr-x
drwxr-xr-x 2 at570
                             5 10:14 matrix.chr9
                     54 Feb
```

```
matrix.chr1$
-rw-r-r- 1 at570 4.7M Feb 5 10:15 loops.tsv
-rw-r- 1 at570 27K Feb 5 10:16 plots.pdf
```

# Scaled Hi-C counts



# Distance-normalized Hi-C scores



# Hi-C scaled count per distance

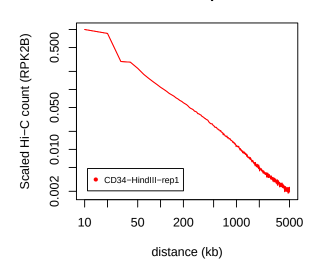


Figure 13: Interactions sample output

### 4.20 Annotations

### 4.20.1 Input

Data from the pipeline interactions step is used as input (Section 4.19).

### 4.20.2 Analysis

### Default parameters:

### 4.20.3 Output

```
-rw-r-r 1 at570 5.9M Feb 5 17:33 bin.annotated.tsv
-rw-r---r--
            at570 3.8M Feb 5 17:33 bin.gene.tsv
-rw-r-r- 1 at570 7.5M Feb 5 17:33 bin.loci.tsv
-rw-r---r--
           1 at570 8.7M Feb 5 17:33 bin.reg
-rw-r-r- 1 at570 5.4K Feb 5 17:33 job.err
-rw-r-r- 1 at570 47 Feb 5 17:32 job.id
-rw-r-r- 1 at 570
                     0 Feb
                           5 17:33 job.out
-rw-r-r- 1 at570 434 Feb
                           5 17:32 job.sh
-rw-r-r 1 at570 3.1K Feb 5 17:33 job.vars.tsv
           1 at570 42M Feb 5 17:33 loci.reg
-rw-r---r
-rw-r-r- 1 at570 45M Feb 5 17:33 table.annotated.tsv
```

### 4.21 Annotations Stats

### 4.21.1 Input

Data from the pipeline annotations step is used as input (Section 4.20).

### 4.21.2 Analysis

### Default parameters:

```
params.standard.tcsh$
#!/bin/tcsh

source ./inputs/params/params.tcsh

set nbest = 10000  # choose top-scoring interactions to calculate enrichments
```

## 4.21.3 Output

## See Figure 14. Default output:

```
-rw-r-r- 1 at570 77 Feb 16 17:26 counts.tsv
-rw-r-r- 1 at570 350 Feb 16 17:26 enrich.tsv
-rw-r-r- 1 at570 7.0K Feb 16 17:26 enrichment.pdf
-rw-r-r- 1 at570 121 Feb 16 17:26 job.err
-rw-r-r- 1 at570 47 Feb 16 17:25 job.id
-rw-r-r- 1 at570 62 Feb 16 17:25 job.out
-rw-r-r- 1 at570 507 Feb 16 17:25 job.sh
-rw-r-r- 1 at570 3.1K Feb 16 17:26 job.vars.tsv
-rw-r-r- 1 at570 184 Feb 16 17:25 top_counts.tsv
```

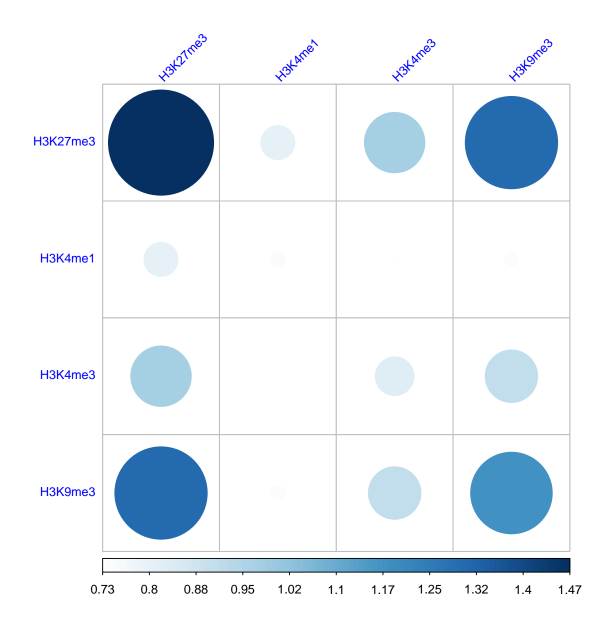


Figure 14: Annotation Stats enrichment sample output. See Section 4.21.

# 5 Appendix

# 5.1 Error Logs

Errors encountered during pipeline execution can be viewed with:

```
| project_directory >$ code.main/pipeline -errors
```

Analysis results can be removed with:

# 5.2 Other Pipeline Software: gtools-hic

```
code.repo/bin/gtools-hic$
USAGE:
  gtools-hic OPERATION [OPTIONS] <REGION-SET>
VERSION:
  genomic-tools 3.0.0
DESCRIPTION:
  Pipeline for HiC-seq data analysis. For detailed description and list of options choose an operation and use
    the —help option.
OPERATION:
  align
                  Iteratively aligns HiC-seq read pairs to reference genome using bowtie2.
  classify
                   Classifies and computes various metrics for HiC-seq aligned read pairs.
  filter
                  Filters HiC-seq aligned read pairs for common experimental artifacts.
  bin
                  Bins filtered read pairs to genomic bins of desired resolution.
  matrix
                   Create Hi-C count matrix
  convert
                  Convert contact matrix into WashU Epigenome Browser format.
```

### 5.2.1 gtools-hic align

```
code.repo/bin/gtools-hic align —help
  USAGE:
    gtools-hic align [OPTIONS] READ1-FASTQ READ2-FASTQ
  DESCRIPTION:
    Iteratively aligns HiC-seq read pairs to reference genome using bowtie2.
      Input: FASTQ files
     * Output: aligned reads in SAM format (same order as in fastq files)
12
  OPTIONS:
      -help
14
                                                                                                    [true]
    -h
                                help
                                                                                                    [true]
                                verbose mode
                                                                                                    [false]
16
    ---work-dir
                                working directory (required)
                                minimum truncated read length
                                                                                                    [30]
    -\!\!-\!\!min-\!len
18
    --len-diff
                                read truncation step
                                                                                                    [10]
                                number of threads for bowtie2 run
20
                                full bowtie2 path (version >= 2.1.0)
                                                                                                    [bowtie2]
     -bowtie-path
      -bowtie-index
                                full bowtie2 index prefix path
                                                                                                    [genome/bowtie2.index
       /genome]
```

### 5.2.2 gtools-hic classify

```
code.repo/bin/gtools-hic classify ---help$
  USAGE:
    gtools-hic classify [OPTIONS] <ALIGNED-READS>
    Classifies and computes various metrics for HiC-seq aligned read pairs.
    * Input: aligned reads in SAM format (sorted by read-id, at most one alignment per read)

    Output: tab-separated table

12
  OPTIONS:
      -help
14
                                                                                                   [true]
    -h
                                help
                                                                                                   [true]
                                verbose mode
                                                                                                   [false]
16
    -\mathsf{E}
                                enzyme fragments (BED/GFF/SAM/REG)
                                                                                                   [3.000000e+01]
18
    --mapq
                                minimum mapping quality (MAPQ)
    --min-dist
                                miminum allowed distance between 5's of reads in read pair
                                                                                                   [500]
                                maximum allowed offset of 5's of reads from fragment ends
    --max-offset
                                                                                                   [500]
```

#### 5.2.3 gtools-hic filter

```
code.repo/bin/gtools-hic filter ---help$
     gtools-hic filter [OPTIONS] <ALIGNED-READS>
     Filters HiC-seq aligned read pairs for common experimental artifacts.
     * Input: aligned reads in SAM format (sorted by read-id, at most one alignment per read)
     * Output: filtered read pairs in REG format
   OPTIONS:
13
     --help
                                   help
                                                                                                            [true]
    -h
15
                                   help
                                                                                                            [true]
                                   verbose mode
    -v
                                                                                                            [false]
    -E
                                   enzyme fragments (BED/GFF/SAM/REG)
                                   minimum mapping quality (MAPQ)
miminum allowed distance between 5's of reads in read pair
                                                                                                            [3.000000e+01]
    --mapq
    --min-dist
                                                                                                            [500]
19
                                   maximum allowed offset of 5's of reads from fragment ends
    --max-offset
                                                                                                            [500]
                                   filter duplicate read pairs as PCR artifacts output statistics file (default=stderr)
21
     ---filter-dups
                                                                                                            [false]
     -stats
                                                                                                            []
```

#### 5.2.4 gtools-hic bin

```
code.repo/bin/gtools-hic bin ---help$
  USAGE:
    gtools-hic bin [OPTIONS] <FILTERED-READ-PAIRS>
  DESCRIPTION:
    Bins filtered read pairs to genomic bins of desired resolution.
    * Input: filtered read pairs in REG format
10
    * Output: binned read pairs
  OPTIONS:
14
    --help
                                help
                                                                                                  [true]
    -h
                                help
                                                                                                  [true]
    -v
                                verbose mode
                                                                                                  [false]
                                genomic bin size
                                                                                                  [1000000]
    --bin-size
                                genome region file (BED/REG)
    -g
```

#### 5.2.5 gtools-hic matrix

```
\verb|code.repo/bin/gtools-hic matrix| --- help |
  USAGE:
     gtools-hic matrix [OPTIONS] <FILTERED-READ-PAIRS>
  DESCRIPTION:
     Create Hi-C count matrix.
     * Input: filtered read pairs in REG format
     * Output: contact matrix
12
  OPTIONS:
14
      -help
                                 help
                                                                                                     [true]
    -\mathsf{h}
                                 help
                                                                                                     [true]
                                 verbose mode
16
    --bin-size
                                 genomic bin size (in nucleotides)
                                                                                                     [5000]
                                 maximum distance between bins (in nucleotides; default = no restriction)[0]
18
      -max-dist
     -rotate45
                                 rotate matrix by 45 degrees (applicable if —max-dist > 0)
                                                                                                    [false]
    -R
                                 reference region file (BED/REG)
20
                                 output file prefix
    -р
                                                                                                     ij
```

## 5.2.6 gtools-hic convert

```
code.repo/bin/gtools-hic convert —help
    gtools-hic convert [OPTIONS] <CONTACT-MATRIX>
    Convert contact matrix into WashU Epigenome Browser format.
    * Input: locus-labelled contact matrix
    * Output: WashU Epigenome Browser format
  OPTIONS:
13
    -help
                               help
                                                                                                  [true]
15
                               help
                                                                                                  [true]
                                                                                                  [false]
                               verbose mode
    -v
                                                                                                  [false]
    --col-labels
                               input matrix has column labels
    -t
                               matrix element separator
                                                                                                 []
[1.000000e+00]
                               normalization constant
19
    -c
                               maximum distance between interacting loci (default = no limit) [0]
                                                                                                 [0.000000e+00]
    -min
    -d
```

# 5.3 Other Pipeline Software: pipeline-master-explorer.r

The pipeline-master-explorer.r script, located in the code.main directory, is the driver of combinatorial parameter exploration during the execution of each pipeline step.

```
code.main$ ./pipeline_master_explorer.r —help
Usage: pipeline_master_explorer.r [OPTIONS] SCRIPT OUTDIR_PREFIX PARAM_SCRIPTS INPUT_BRANCHES SPLIT_VARIABLE
OUTPUT_OBJECT_VARIABLE TUPLES
     -v, --verbose
Print more messages.
     -S SAMPLE-SHEET, --sample-sheet=SAMPLE-SHEET
       Sample sheet file name (required) [default "inputs/sample-sheet.tsv"].
11
    -F FILTER-BRANCH, -- filter -branch=FILTER-BRANCH
       Regular expression for filtering input branches [default ""].
13
15
       -exclude-branch=EXCLUDE-BRANCH
       Regular expression for excluding input branches [default ""].
17
       -exclude-obj=EXCLUDE-OBJ
       Regular expression for excluding input objects [default ""].
19
       -exclude-outdir=EXCLUDE-OUTDIR
21
       Regular expression for excluding output directories [default ""].
23
    -h. --help
       Show this help message and exit
```

# 5.4 System and Session Information

This document was created with: LATEX 2 $_{\mathcal{E}}$  2005/12/01

```
system('uname -srv',intern=T)
## [1] "Linux 2.6.32-573.18.1.el6.x86_64 #1 SMP Tue Feb 9 22:46:17 UTC 2016"
sessionInfo()
## R version 3.2.3 (2015-12-10)
## Platform: x86_64-redhat-linux-gnu (64-bit)
## Running under: CentOS release 6.7 (Final)
## locale:
## [1] LC_CTYPE=en_US.UTF-8
                                 LC_NUMERIC=C
                                                             LC_TIME=en_US.UTF-8
   [4] LC_COLLATE=en_US.UTF-8
                                  LC_MONETARY=en_US.UTF-8
                                                             LC_MESSAGES=en_US.UTF-8
                                  LC_NAME=C
  [7] LC_PAPER=en_US.UTF-8
                                                             LC_ADDRESS=C
## [10] LC_TELEPHONE=C
                                  LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
## attached base packages:
## [1] stats graphics grDevices utils datasets methods base
##
## other attached packages:
## [1] knitr_1.12.3
## loaded via a namespace (and not attached):
## [1] magrittr_1.5 formatR_1.2.1 tools_3.2.3
                                                stringi_1.0-1 highr_0.5.1 stringr_1.0.0
## [7] evaluate_0.8
```

#### 5.4.1 LATEX File List

```
article.cls
                      2005/09/16 v1.4f Standard LaTeX document class
     size10.clo
                      2005/09/16 v1.4f Standard LaTeX file (size option)
   graphicx.sty
                      1999/02/16 v1.0 f Enhanced LaTeX Graphics (DPC,SPQR)
                      1999/03/16 v1.13 key=value parser (DPC)
2006/02/20 v1.00 Standard LaTeX Graphics (DPC,SPQR)
     keyval.sty
   graphics.sty
        trig.sty
                      1999/03/16 v1.09 sin cos tan (DPC)
   graphics.cfg
pdftex.def
                      2007/01/18 v1.5 graphics configuration of teTeX/TeXLive
                      2007/01/08 v0.04d Graphics/color for pdfTeX
      color.stv
                      1999/02/16
      color.cfg
                      2007/01/18 v1.5 color configuration of teTeX/TeXLive
    framed.sty
                      2011/10/22 v 0.96: framed or shaded text with page breaks
                      1997/06/16 v2.0g defines alltt environment
2005/04/12 PSNFSS-v9.2a Palatino w/ Pazo Math (D.Puga, WaS)
      alltt.sty
13 mathpazo.sty
                      2005/04/12 PSNFSS-v9.2a (WaS)
     helvet.sty
   fontenc.sty
                      2005/09/27 v1.99g Standard LaTeX file
2002/07/08 v3.2 Page Geometry
      t1enc.def
   geometry.sty
   geometry.cfg
                      2003/11/04 v 4.01
       cite.sty
                      2007/01/07\ v3.0\,k Customising captions (AR) 2007/01/07\ v3.0\,k caption3 kernel (AR)
    caption.sty
   caption3.sty
                      2007/02/07 v6.75r Hypertext links for LaTeX 2007/02/07 v6.75r Hyperref: PDFDocEncoding_definition (HO)
   hyperref.sty
     pd1enc.def
   hyperref.cfg
                      2002/06/06 v1.2 hyperref configuration of TeXLive
   kvoptions.sty
                       2006/08/22 v2.4 Connects package keyval with LaTeX options (HO
                      2007/02/07 v6.75r Hyperref: PDF Unicode definition (HO)
      puenc.def
                      2005/06/27 ver 3.2 Verb mode for urls, etc. 2007/02/07 v6.75r Hyperref driver for pdfTeX
         url.sty
   hpdftex.def
   breakurl.sty
                      2006/08/26 v1.20 Breakable hyperref URLs
   xkeyval.sty
                      2006/11/18 v2.5f package option processing (HA)
                      2006/11/18 v2.5f key=value parser (HA)
    xkeyval.tex
```

```
forloop.sty
                    2006/09/18 v3.0 For Loops for LaTeX
     ifthen.sty
                     2001/05/26 v1.1c Standard LaTeX ifthen package (DPC)
       tikz.sty
                    2006/10/17 \text{ v1.10 (rcs-revision 1.68)}
        pgf.sty
                    2006/10/11 v1.10 (rcs-revision 1.7)
     pgfrcs.sty
37
                    2006/10/26 v1.10 (rcs-revision 1.14)
     pafrcs.code.tex
    pgfcore.sty
                    2006/10/11 v1.10 (rcs-revision 1.4)
                    2006/10/16 v1.10 (rcs-revision 1.19)
     pgfsys.sty
41
     pgfsys.code.tex
                                2006/10/16 (rcs-revision 1.4)
2006/10/16 (rcs-revision 1.4)
   pgfsyssoftpath.code.tex
43
   pgfsysprotocol.code.tex
                    2007/01/21 v2.11 LaTeX color extensions (UK)
     xcolor.sty
      color.cfg
                    2007/01/18 v1.5 color configuration of teTeX/TeXLive
45
    pgfcore.code.tex
   pgfbaseshapes.sty
                          2006/10/16 v1.10 (rcs-revision 1.16)
   pgfbaseshapes.code.tex
                       2006/10/16 v1.10 (rcs-revision 1.5)
   pgfbaseplot.sty
   pgfbaseplot code tex
                         2006/10/16 v1.10 (rcs-revision 1.5)
   pgfbaseimage.sty
   pgfbaseimage.code.tex
                         2006/10/16 v1.10 (rcs-revision 1.5)
   pgfbaselayers.stv
   pgfbaselayers.code.tex
                          2006/10/16 v1.10 (rcs-revision 1.10)
   pgfbasesnakes.sty
   pgfbasesnakes.code.tex
                           2006/10/16 v1.10 (rcs-revision 1.9)
   pgfbasepatterns.stv
57
   pgfbasepatterns code tex
   pgfcomp-version\,-0-65.sty
                                 2006/10/11 v1.10 (rcs-revision 1.4)
59
       calc.sty
                    2005/08/06 v4.2 Infix arithmetic (KKT,FJ)
                    2006/10/16 v1.10 (rcs-revision 1.5)
     pgffor.sty
61
     \verb"pgffor.code.tex"
       tikz.code.tex
63
    amsmath.stv
                    2000/07/18 v2.13 AMS math features
    amstext.sty
                    2000/06/29 v2.01
65
                     1999/11/30 v2.0
     amsgen.sty
                     1999/11/29 v1.2d
67
     amsbsy.sty
                    1999/12/14 \ v2.01 \ operator \ names
     amsopn.sty
                    2001/02/13 v0.1 j Color table columns (DPC) 2005/08/23 v2.4b Tabular extension package (FMi)
69
   colortbl.sty
      array.sty
                    2004/10/17 1.3b (Carsten Heinz)
2004/10/17 1.3b (Carsten Heinz)
   listings.sty
   Istpatch.sty
                    2004/09/07 1.3 (Carsten Heinz)
    Istmisc.sty
                    2004/09/05 1.3 listings configuration
   listings.cfg
                    2004/09/05 1.3 listings language file 2004/09/05 1.3 listings language file
   Istlang1.sty
   Istlang1.sty
    upquote.sty
                    2003/08/11 v1.1 Covington's upright-quote modification to verba
   tim and verb
   textcomp.sty
                    2005/09/27 v1.99g Standard LaTeX package
                    2001/06/05 v3.0e (jk/car/fm) Standard LaTeX file
     ts1enc.def
                    1999/05/25 v2.5h Standard LaTeX font definitions
     ts1cmr.fd
      t1phv.fd
                   2001/06/04 scalable font definitions for T1/phv.
   supp-pdf.tex
   ragged2e.sty
                    2003/03/25 v2.04 ragged2e Package (MS)
   everysel.sty
                     1999/06/08 v1.03 EverySelectfont Package (MS)
    nameref.sty
                    2006/12/27 v2.28 Cross-referencing by name of section
   refcount.sty
                    2006/02/20 v3.0 Data extraction from references (HO)
   hic-manual_base.out
   hic-manual base.out
    ot1pplx.fd
                    2004/09/06 font definitions for OT1/pplx.
    omlzplm.fd
                   2002/09/08 Fontinst v1.914 font definitions for OML/zplm.
                    2002/09/08 Fontinst v1.914 font definitions for OMS/zplm.
    omszplm.fd
                    2002/09/08 Fontinst v1.914 font definitions for OMX/zplm.
    omxzplm.fd
    ot1zplm.fd
                   2002/09/08 Fontinst v1.914 font definitions for OT1/zplm.
   figure/NYU_Langone.jpg
   child / Introduction / install -setup-run.tex
                   1999/05/25 v2.5h Standard LaTeX font definitions
   figure/sample_sheet_screenshot.png
   child / Introduction / dependencies . tex
   child/default-pipeline-components.tex
                   2001/06/04 scalable font definitions for TS1/phv.
     ts1phv.fd
   child/code-structure.tex
   child/auto_report.tex
   child/custom_pipeline_step.tex
   child/HiC/index.tex
   child/HiC/align.tex
   child/HiC/filter.tex
   child/HiC/filter-stats.tex
   figure / filter -stats_counts.pdf
   figure / filter -stats_percent.pdf
   child/HiC/tracks.tex
   child/HiC/matrix-filtered.tex
child/HiC/matrix-prep.tex
```

```
\verb|child/HiC/matrix-ic.tex| \\
          child/HiC/matrix-hicnorm.tex
            \verb|child/HiC/matrix-stats|| state|| to the constant of the co
          figure/matrix-stats_stats.pdf
            child/HiC/compare-matrices.tex
          child/HiC/compare-matrices-stats.tex
             figure / compare-matrices-stats\_correlograms.pdf
          figure/compare-matrices-stats_pearson_correlograms.pdf
            child/HiC/boundary-scores.tex
          child/HiC/boundary-scores-pca.tex
          figure/boundary-scores-pca_pca_Dl_k_001.pdf
child/HiC/domains.tex
            child/HiC/compare-boundaries.tex
          child/HiC/compare-boundaries-stats.tex
          figure/compare-boundaries-stats_raw_comparisons.pdf
figure/compare-boundaries-stats_correlograms.pdf
            child/HiC/hicplotter.tex
          figure / hicplotter_chr8 -125000000-133000000.pdf
131
            child/HiC/interactions.tex
          figure/interactions_plots.pdf
          child/HiC/annotations.tex
child/HiC/annotations—stats.tex
135
           figure / annotations - stats _ enrichment . pdf child / Appendix / appendix . tex
               ts1cmtt.fd
                                                                     1999/05/25 v2.5h Standard LaTeX font definitions
139
```