GenomeFlow User Manual

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1 1D-Functions

1.1 Dependencies

1.1.1 Operating System (OS)

The UNIX environment is required in this protocol.

It is strongly recommended to work under a UNIX-based operating system, such as Ubuntu, Centos/Red Hat, Solaris, or Mac OS X.

If you are using a Windows operating system, install Cygwin first. Cygwin is a free software that provides a UNIX-like environment on Windows. The Cygwin install package can be found at http://www.cygwin.com/.

1.1.2 External Tools

- Install Bowtie2 (http://bowtie-bio.sourceforge.net/index.shtml)
- Install BWA (http://bio-bwa.sourceforge.net/)
- Install Samtools (http://samtools.sourceforge.net/)
- Pick the specific tool for your OS, and follow the installation instructions for each one of them.
- We tested on the following versions for each one of the tools: bowtie2-2.3.4-*, bwa-0.7.17, and samtools-1.6

1.2 Create an index for a reference genome

1.2.1 Purpose

To build index for the reference genome data. Indexing the reference genome makes querying fast, and can also compress the size of the genome data

1.2.2 Input

The reference input FASTA file (usually having extension fa, mfa,.fna or similar). Read more about FASTA files here: https://en.wikipedia.org/wiki/FASTA.

Example:

The human hg19 genome data (.fa) can be downloaded from here:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/hg19_genome/hg19_genome e_FASTA/

1.2.3 Output

It generates a shell script with the name *Indexer_script.sh*.

1.2.4 Output of script:

A list of index files. This varies depending on the tool selected for indexing. BWA output 5 files (NAME.amb, NAME. ann, NAME. bwt, NAME .pac, and NAME.sa), where NAME is a prefix string, and Bowtie outputs 6 files (NAME.1.bt2, NAME.2.bt2, NAME.3.bt2, NAME.4.bt2, NAME.rev.1.bt2, and NAME.rev.2.bt2) where NAME is
bt2_base>.

Example:

Generated index for the hg19 human genome by bowtie2 and bwa tools can be downloaded from here:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/hg19_genome/

1.2.5 Running

- Access the function from the menu toolbar: 1D-Functions/Build index for reference genome
- Generate a script called Indexer_script.sh
- Open a Unix Terminal
- Execute *Indexer_script.sh* in a Unix Terminal



Figure 1: Create an index for a reference genome

Field	Description	Default
Input	A reference genome file having extension. fa, .mfa, .fna or	NA
Reference	similar. For example human genome(GRCh37/hg19)	
Genome file		
Output	The output directory path to output the script	NA
Directory		
Choose tool to	Two options are made available for indexing. Select bwa-	bwa
use	Burrows-Wheeler alignment or Bowtie2.	
Binary file	Browse and select the binary file for the chosen tool	NA
	BWA: Select the bwa binary you compiled from the dependency section Bowtie2: Select the bowtie2-biuild indexer	
Number of	This Option is Only available for the bowtie2-biuild indexer,	8
threads	Specify the number of threads to use for this task. More	
	threads means less processing time taken.	

Generate	This button generates a shell script (.sh) that can be used be	
Scripts	run on a UNIX terminal by the use. This basic parameters	
	required by each tool for genome indexing.	
OS	This button determines if the user's Operating System is	
Compatibility	compatible for running the script directly without a need for a	
	UNIX-like environment for example Cygwin	

1.3 Mapping the raw single or pair read FASTQ files

1.3.1 Purpose

To perform alignment of the index and a set of sequencing read files

1.3.2 Input

A FASTQ read files usually with extension .fq or .fastq. Read more about FASTA files here: https://en.wikipedia.org/wiki/FASTQ_format

Example

Test datasets can be found here:

- MiSeq GM12878 in-situ files:
 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/MiSeq_GM12878/M
 iSeq_GM12878_Data/
- A karyotypically normal human lymphoblastoid cell line (GM06990) from Aiden et al: http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/GM06990
 O_Data/

1.3.3 Output

It generates a shell script with name *Mapper_script_bowtie2.sh* for bowtie2 tool an *Mapper_script_bwa.sh* for bwa tool.

1.3.4 Output of script:

The output will be found in a folder bowtie2_align for bowtie2 and bwa_align for bwa. By default, the output BAM file is named *bwa_mapped.bam* for bwa and named *bowtie2_mapped.bam* for bowtie2.

A BAM binary format (. bam) obtained by converting a SAM file from samtools into a BAM file. Check http://samtools.sourceforge.net for the SAM format specification and the tools for post-processing the alignment.

Example:

The generated bowtie2 and bwa alignment BAM file can be downloaded from the link below for each test data:

- MiSeq GM12878 in-situ files:
 - Bowtie2:
 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/MiSeq_GM1
 2878/bowtie2_align_Miseq/
 - Bwa:
 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/MiSeq_GM1
 2878/bwa_align_Miseq/
- GM06990 Cell line:
 - b Bowtie2:

 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/b

 owtie2_align/
 - b Bwa:

 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/b

 wa_align/

1.3.5 Running

- Access the function from the menu toolbar: 1D-Functions/Map the raw FASTQ files
- Generate a script called *Mapper_script_bowtie2.sh* or *Mapper_script_bwa.sh*
- Open a Unix Terminal
- Execute *Mapper_script_bowtie2.sh* in a Unix Terminal

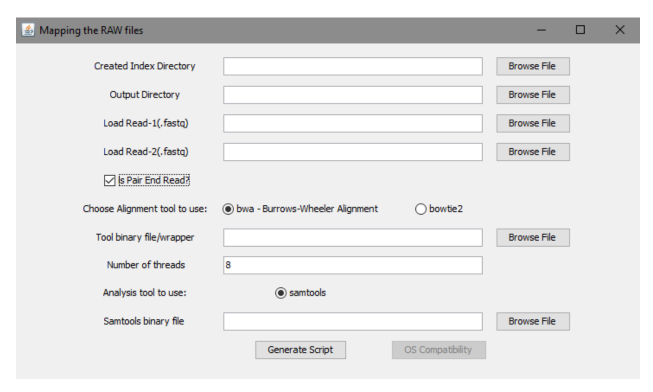


Figure 2: Mapping the raw FASTQ files

Field	Description	Default
Created Index	A path to the index created using bwa or bowtie2	NA
Directory		
Output	The output directory path to output the script	NA
Directory		
Load Read-	The file containing mate 1, or file for a single read e.g	NA
1(.fastq)	HIC003_S2_L001_R1_001.fastq	
Load Read-	The file containing mate 2 e.g	NA
2(.fastq)	HIC003_S2_L001_R2_001.fastq	
Is Pair-End	Check if the data is a pair end read data	unchecked
Read		

Choose tool to	Two options are made available for indexing. Select bwa-	bwa
use	<u>Burrows-Wheeler alignment</u> or <u>Bowtie2</u> .	
	Important : Only select the tool which was used to generated	
	the reference genome Index. bwa can only be used to Map	
	bwa index, and bowtie2 can only be used to map bowtie2	
	index.	
Binary file	Browse and select the binary file for the chosen tool	NA
	BWA: Select the bwa binary you compiled from the	
	dependency section	
	Bowtie2: Select the bowtie2 file to align	
Number of		0
	This Option is Only available for the bowtie2-biuild indexer,	8
threads	Specify the number of threads to use for this task. More	
	threads means less processing time taken.	
Samtools	SAMtools is a collection of tools for manipulating and	NA
binary file	analyzing SAM and BAM alignment files. Using these tools	
	together allows you to get from alignments in SAM format	
	Browse and select the samtools binary file	
Generate	This button generates a shell script (.sh) that can be used be	
Scripts	run on a Unix terminal by the user. This scripts contains the	
	basic parameters required by each tool for Mapping the pair	
	reads to a reference index.	
OS	This button determines if the user's Operating System is	
Compatibility	compatible for running the script directly without a need for a	
	UNIX-like environment for example <u>Cygwin</u>	

1.4 Filter a BAM alignment file

1.4.1 Purpose

To perform filtering of a BAM file to remove low quality map reads, and unmapped reads among others.

1.4.2 Input

The BAM file generated from the mapping step above. For example, by default bwa BAM files is named bwa_mapped.bam and bowtie2 BAM files is named bowtie2_mapped.bam

Example:

The generated bowtie2 alignment BAM file can be downloaded from the link below for each test data:

- MiSeq GM12878 in-situ files:
 - o Bowtie2:

o Bwa:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/MiSeq_GM1 2878/bwa align_Miseq/

- GM06990 Cell line:
 - o Bowtie2:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/bowtie2_align/

o Bwa:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/bwa_align/

1.4.3 Output

It generates a shell script with name Filter_script_samtools.sh

1.4.4 Output of the script:

A BAM binary format (. bam) named bowtie2_mapped.filtered.bam for bowtie2 and bwa_mapped.filtered.bam for bwa

Example:

The generated filtered bowtie2 and bwa alignment BAM file can be downloaded from the link below for the MiSeq GM12878 in-situ test data

- MiSeq GM12878 in-situ files:
 - Bowtie2:
 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/MiSeq_GM1
 2878/bowtie2 Miseq_mapped.filtered.bam
 - Bwa:
 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/MiSeq_GM1
 2878/bwa_Miseq_mapped.filtered.bam
- GM06990 Cell line:
 - Bowtie2:
 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/b
 owtie2 mapped.filtered.bam
 - Bwa:
 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/b
 wa mapped.filtered.bam

1.4.5 Running

- Access the function from the menu toolbar: 1D-Functions/Filter a BAM alignment file
- Generate a script called *Filter_script_samtools.sh*

- Open a Unix Terminal
- Run Filter_script_samtools.sh in a Unix Terminal

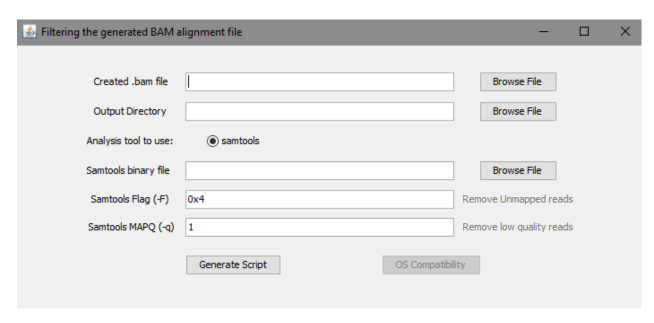


Figure 3: Filter a BAM alignment file

Field	Description	Default
Created .bam	Select the BAM file generated using either BWA or bowtie2.	NA
file	Select the BAM file named bwa_mapped.bam for bwa and	
	BAM file named bowtie2_mapped.bam for bowtie2	
Output	The output directory path to output the script	NA
Directory		
Samtools	samtools is a collection of tools for manipulating and	NA
binary file	analyzing SAM and BAM alignment files. Using these tools	
	together allows you to get from alignments in SAM format	
	Browse and select the samtools binary file	
C . 1 Fl	ř	0.4
Samtools Flag	samtools allows you to sort based on certain flags that are	0x4
(-F)	specified on page 5 on the SAM format specification	

Samtools	An integer value to Skip alignments with MAPQ smaller	1
MAPQ (-q)	than INT. The lowest score is a mapping quality of zero,	
	or mq0 for short. The reads map to multiple places on the	
	genome, and we can't be sure of where the reads originated.	
	To improve the quality of our data, we can remove these low	
	quality reads. Generally, we select reads with MAPQ > 1.	
Generate	This button generates a shell script (.sh) that can be used be	
Scripts	run on a Unix terminal by the user. This scripts contains the	
	basic parameters required by each tool for filtering.	
OS	This button determines if the user's Operating System is	
Compatibility	compatible for running the script directly without a need for a	
	Unix-like environment for example <u>Cygwin</u>	

1.5 Convert a BAM file to a HiC input file format

1.5.1 Purpose

To generate a <u>medium file format</u>, a text file describing mapped Hi-C reads that can be used as input to create a *.hic* file. A hic format file is a binary file containing contact matrices at different resolutions and normalized by different methods.

1.5.2 Input

A filtered BAM alignment file. e.g bwa_mapped.filtered.bam

Example:

The generated filtered bowtie2 and bwa alignment BAM file for the MiSeq GM12878 insitu test data can be downloaded from the link below:

• MiSeq GM12878 in-situ files:

o Bowtie2:

o Bwa:

• GM06990 Cell line:

o Bowtie2:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/bowtie2_mapped.filtered.bam

o Bwa:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/bwa_mapped.filtered.bam

1.5.3 Output

It generates a shell script with name Format_script_samtools.sh

1.5.4 Output of the script:

A medium input file format with 11 columns that can be used to create a .hic file This file format is explained in details here: <u>HiC Input Medium File Format</u>

Example:

The generated input medium file format file for the input test datasets can be downloaded from the link below:

MiSeq GM12878 in-situ files:

o Bowtie2:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/MiSeq_GM1 2878/GenomeFlow_Miseq_formatted.bowtie2.input

o Bwa:

- GM06990 Cell line:
 - o Bowtie2:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/GenomeFlow_formatted.bowtie2.input

o Bwa:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/GenomeFlow formatted.bwa.input

1.5.5 Running

- Access the function from the menu toolbar: 1D-Functions/Convert a BAM file to a HiC input file format
- Generate a script called *Format_script_samtools.sh*
- Open a Unix Terminal
- Execute Format_script_samtools.sh in a Unix Terminal



Figure 4: Convert to HiC Input File Format

Field	Description	Default
Created .bam	Select the BAM file generated from the filtering.	NA
file		
	By default, bwa filtered BAM file is named	
	bwa_mapped.filtered.bam and bowtie2 filtered BAM file is	
	named bowtie2_mapped.filtered.bam	
Output	The output directory path to output the script	NA
Directory		
Samtools	samtools is a collection of tools for manipulating and	NA
binary file	analyzing SAM and BAM alignment files. Using these tools	
	together allows you to get from alignments in SAM format	
	Browse and select the samtools binary file	
Generate	This button generates a shell script (.sh) that can be used be	
Scripts	run on a Unix terminal by the user. This scripts contains the	
	basic parameters required by each tool for filtering.	
OS	This button determines if the user's Operating System is	
Compatibility	compatible for running the script directly without a need for a	
	Unix-like environment for example Cygwin	

1.6 HiC-Express

1.6.1 Purpose

To generate a <u>medium file format</u>, a text file describing mapped Hi-C reads that can be used as input to create a *.hic* file from a raw fastq files derived from a Hi-C experiment.. A hic format file is a binary file containing contact matrices at different resolutions and normalized by different methods.

1.6.2 Input

A FASTQ read files usually with extension, .fq or .fastq. Read more about FASTA files here: https://en.wikipedia.org/wiki/FASTQ_format

Example:

Test datasets can be found here:

- MiSeq GM12878 in-situ files:
 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/MiSeq_GM12878/M
 iSeq_GM12878_Data/
- A karyotypically normal human lymphoblastoid cell line (GM06990) from Aiden et al: http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/GM0699

 Data/

1.6.3 Output

It generates a shell script with name *HiC-Express.sh*

1.6.4 Output of the script:

An input Medium file format with 11 columns that can be used to create a *.hic* file This file format is explained in details here: <u>HiC Input Medium File Format</u>

Example:

The generated input medium file format file for the input test datasets can be downloaded from the link below:

- MiSeq GM12878 in-situ files:
 - o Bowtie2:

http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/MiSeq_GM1 2878/GenomeFlow Miseq_formatted.bowtie2.input

o Bwa:

- GM06990 Cell line:
 - Bowtie2:
 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/G
 enomeFlow formatted.bowtie2.input
 - Bwa:
 http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/G

1.6.5 Running

• Access the function from the menu toolbar: 1D-Functions/HiC-Express

enomeFlow_formatted.bwa.input

- Generate a script called HiC-Express.sh
- Open a Unix Terminal
- Execute *HiC-Express.sh* in a Unix Terminal.

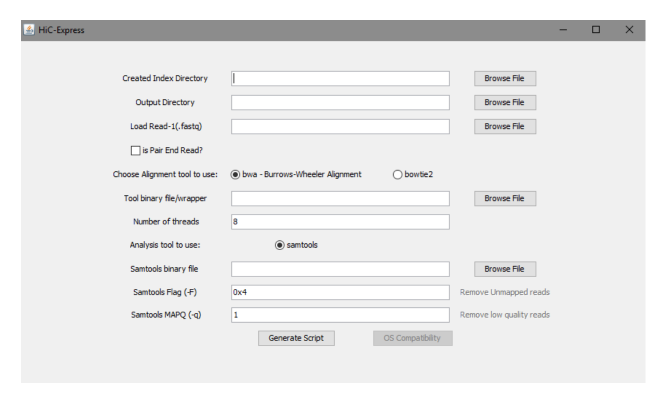


Figure 5: HiC-Express

Field	Description	Default
Created Index	A path to the index created using bwa or bowtie2	NA
Directory		
Output	The output directory path to output the script	NA
Directory		
Load Read-	The file containing mate 1, or file for a single read e.g	NA
1(.fastq)	HIC003_S2_L001_R1_001.fastq	
Load Read-	The file containing mate 2 e.g	NA
2(.fastq)	HIC003_S2_L001_R2_001.fastq	
Is Pair-End	Check if the data is a pair end read data	unchecked
Read		
Choose tool to	Two options are made available for indexing. Select bwa-	bwa
use	<u>Burrows-Wheeler alignment</u> or <u>Bowtie2</u> .	

	Important: Only select the tool which was used to generated	
	the reference genome Index. bwa can only be used to Map	
	bwa index, and bowtie2 can only be used to map bowtie2	
	index.	
Binary file	Browse and select the binary file for the chosen tool	NA
	BWA: Select the bwa binary you compiled from the	
	dependency section	
	Bowtie2: Select the bowtie2 file to align	
Number of	This Option is Only available for the bowtie2-biuild indexer,	8
threads	Specify the number of threads to use for this task. More	
	threads means less processing time taken.	
Samtools	SAMtools is a collection of tools for manipulating and	NA
binary file	analyzing SAM and BAM alignment files. Using these tools	
	together allows you to get from alignments in SAM format	
	Browse and select the samtools binary file	
Samtools Flag	samtools allows you to sort based on certain flags that are	0x4
(-F)	specified on page 5 on the SAM format specification	
Samtools	An integer value to Skip alignments with MAPQ smaller	1
MAPQ (-q)	than INT. The lowest score is a mapping quality of zero,	
	or mq0 for short. The reads map to multiple places on the	
	genome, and we can't be sure of where the reads originated.	
	To improve the quality of our data, we can remove these low	
	quality reads. Generally, we select reads with MAPQ > 1.	
Generate	This button generates a shell script (.sh) that can be used be	
Scripts	run on a Unix terminal by the user. This scripts contains the	
	basic parameters required by each tool for Mapping the pair	
	reads to a reference index.	

OS	This button determines if the user's Operating System is	
Compatibility	compatible for running the script directly without a need for a	
	UNIX-like environment for example Cygwin	

2 2D-Functions

2.1 Convert mapped Hi-C reads to hic format file

2.1.1 Purpose

To create a binary hic format file containing contact matrices at different resolutions and normalized by different methods from a text file describing mapped Hi-C reads

2.1.2 Input file format

Five formats are acceptable: short format, short format with score, medium format, long format and 4DN DCIC format. A sample file is executable/sample_data/GSM1551688_HIC143_merged_nodups.zip (unzip it before use) Another test data is the GM06990 cell line data, that can be downloaded from link below:

- GM06990 Cell line:

 - Bwa:
 <a href="http://sysbio.rnet.missouri.edu/bdm_download/GenomeFlow/GM06990/GenomeFlow-genome

2.1.2.1 Short format

A whitespace separated file that contains, on each line

- str = strand (0 for forward, anything else for reverse)
- chr = chromosome (must be a chromosome in the genome)
- pos = position
- frag = restriction site fragment

If not using the restriction site file option, frag will be ignored, but please see above note on dummy values. readname and strand are also not currently stored within *.hic* files.

2.1.2.2 Short with score format

This format is useful for reading in already processed files, e.g. those that have been already binned and/or normalized; this format can be easily used in conjunction with the -r flag to create a .hic file that contains a single resolution.

A whitespace separated file that contains, on each line

```
<str1> <chr1> <pos1> <frag1> <str2> <chr2> <pos2> <frag2> <score>
```

- str = strand (0 for forward, anything else for reverse)
- chr = chromosome (must be a chromosome in the genome)
- pos = position
- frag = restriction site fragment
- score = the score imputed to this read

If not using the restriction site file option, frag will be ignored, but please see above note on dummy values. readname and strand are also not currently stored within .hic files.

2.1.2.3 Medium format

A whitespace separated file that contains, on each line

```
<readname> <str1> <chr1> <pos1> <frag1> <str2> <chr2> <pos2> <frag2> <mapq1> <mapq2>
```

• str = strand (0 for forward, anything else for reverse)

- chr = chromosome (must be a chromosome in the genome)
- pos = position
- frag = restriction site fragment
- mapq = mapping quality score

If not using the restriction site file option, frag will be ignored, but please see above note on dummy values. If not using mapping quality filter, mapq will be ignored, readname and strand are also not currently stored within .hic files.

2.1.2.4 Long format

The long format is used by <u>Juicer</u> and takes in directly the *merged_nodups.txt* file.

A whitespace separated file that contains, on each line

```
<str1> <chr1> <pos1> <frag1> <str2> <chr2> <pos2> <frag2> <mapq1> <cigar1>
<sequence1> <mapq2> <cigar2> <sequence2> <readname1> <readname2>
```

- str = strand (0 for forward, anything else for reverse)
- chr = chromosome (must be a chromosome in the genome)
- pos = position
- frag = restriction site fragment
- mapq = mapping quality score
- cigar = cigar string as reported by aligner
- sequence = DNA sequence

If not using the restriction site file option, frag will be ignored, but please see above note on dummy values. If not using mapping quality filter, mapq will be ignored. readname, strand, cigar, and sequence are also not currently stored within .hic files.

2.1.2.5 4DN DCIC format

A file that follows the 4DN DCIC format specification (the 4DN DCIC format specification). See the link for more information. Briefly, there should be a header with the first seven columns reserved:

```
## pairs format v1.0
#columns: readID chr1 position1 chr2 position2 strand1 strand2
```

If the columns line contains (in any field after field 7) both frag1 and frag2, those will also be read in; otherwise Pre will set frag1=0 and frag2=1, so that no reads are discarded. Other fields are ignored.

2.1.3 Output

A binary .hic file containing contact matrices

2.1.4 Running

Access the function from the menu toolbar: 2D-Functions/Convert to HiC

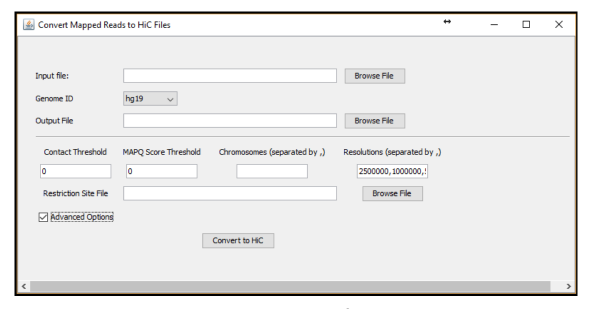


Figure 6: Convert to HiC function

Field	Description	Default
Input file	A text file describes mapped Hi-C reads (format described	NA
	above)	
Genome ID	Version genome of Hi-C data	hg19
Output File	A name of the output hic format file	NA
Contact	Number of interaction threshold for contacts to be used in	0
Threshold	creating contact matrices.	
MAPQ Score	Mapping quality score threshold for reads to be considered in	0
Threshold	creating contact matrices.	
Chromosomes	Chromosomes for which their contact matrices need to be	All (when
	created. When left blank, all chromosomes will be considered.	left blank)
	Chromosomes must be separated by a comma (,).	
Resolutions	List of resolutions of contact matrices to be created.	2500000, 1000000, 500000, 250000,
	Resolutions are separated by a comma (,)	100000, 50000, 25000,10000,5000
Restriction Site	Each line starts with a chromosome number followed by	blank
File	positions of restriction sites on that chromosome, in numeric	
	order, and ending with the size of the chromosome. When	
	provided, 8 additional fragment-delimited resolutions are	
	added: 500f, 250f, 100f, 50f, 20f, 5f, 2f, 1f	

2.2 Extract contact matrices from a hic format

2.2.1 Purpose

To extract a contact matrix from a hic format into a sparse matrix format in a text file

2.2.2 Input

A local path to a hic format or an online link to a hic format. A link to a hic file: https://www.encodeproject.org/files/ENCFF219YOB/@@download/ENCFF219YOB.hic

2.2.3 Output

A contact matrix in sparse matrix format (each line represents a contact by three numbers separated by whitespaces: position1 postion2 interaction_frequency)

2.2.4 Running

Access the function from the menu toolbar: 2D-Functions/Extract HiC

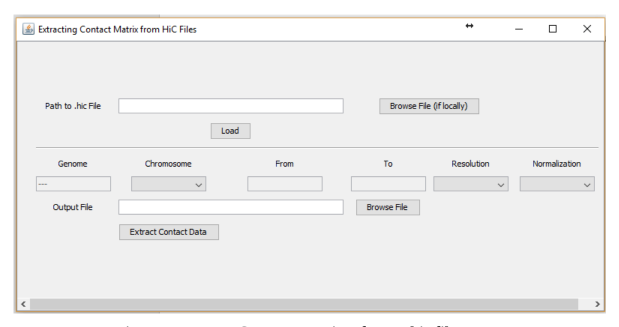


Figure 7: Extract Contact Matrices from a hic file

Field /	Description	Default
Button		
Path to .hic	An online link or local path to a hic format file	NA
File		
Load	Click this button to fetch information from the header of the hic	NA
	file.	

Genome	Genome version of the hic file	NA
Chromosomes	List of resolutions of contact matrices in the hic file	NA
From	Start of a fragment (to extract its contact matrix). When From and	Blank
	To are left blank, the whole chromosome is considered.	
То	End of a fragment (to extract its contact matrix). When From and	Blank
	To are left blank, the whole chromosome is considered.	
Resolution	List of resolutions of contact matrices in the hic file	NA
Normalization	List of normalization methods used to normalize contact matrices	NA
Extract	Click this button to initiate extracting contact data	NA
Contac Data		

2.3 Normalize HiC contact matrices

2.3.1 Purpose

To normalize contact matrices in sparse matrix format.

2.3.2 Input

A contact matrix in sparse matrix format (each line represents a contact by three numbers separated by whitespaces: position1 position2 interaction_frequency)

2.3.3 Output

A normalized contact matrix in sparse matrix format. The matrix is normalized by the Iterative Correction and Eigenvector decomposition (ICE) method

2.3.4 Running

Access the function from the menu toolbar: 2D-Functions/Normalized HiC Data

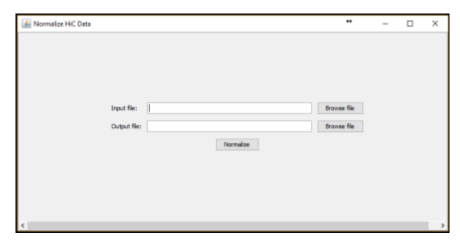


Figure 8: Normalize HiC contact matrices

2.4 Visualizing Dataset in 2D format

2.4.1 Purpose

To create a two dimensional (2D) graphical representation of a contact matrix from an input file.

2.4.2 Input

A sparse matrix format (each line represents a contact by three numbers separated by whitespaces: position1 postion2 interaction_frequency(IF)) or an input file in square matrix format (a full matrix representing all the contact regions). Mark the *Is Square Matrix?* box if the input is a square matrix.

An example sparse matrix file can be found here:

/executable/sample_data/ contact_matrices/ chr11_10kb_gm12878_list_125mb_135mb.txt Examples of square matrix files can be found here:

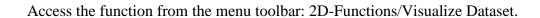
/executable/sample_data/ contact_matrices/square_matrices/

Note: Resolution for square matrices = 40000

2.4.3 Output

A Heatmap which is a graphical representation of contact data where numeric values in the input contact matrix are represented as colors based according to a selected color gradient.

2.4.4 Running



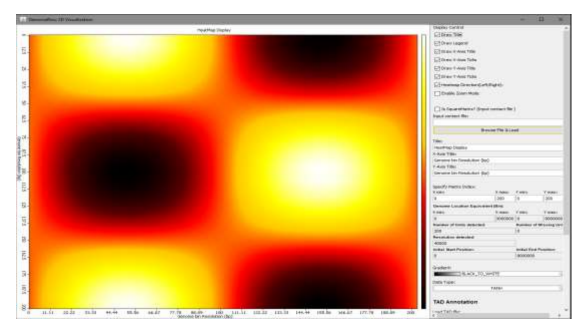


Figure 9: Visualize Dataset in 2D Format

2.4.5 Display Controls

The description of the display controls on the display window is given below.

Field	Description	Default
Draw Title	It shows or hides the Heatmap title	checked
Draw Legend	It shows or hides the color legend	checked
Draw X-Axis Title	It shows or hides the X-axis title label on the	checked
	2D display window	
Draw X-Axis Ticks	It shows or hides the X-axis ticks label on	checked
	the 2D display window	
Draw Y-Axis Title	It shows or hides the Y-axis title label on the	checked
	2D display window	

Draw Y-Axis Ticks	It shows or hides the Y-axis ticks label on	checked
	the 2D display window	
Heatmap Direction(Left/Right)	It changes the Y-axis origin of the heatmap	checked
	matrix from the Bottom-Left to Top-Left and	
	vice versa	
Enable Zoom Mode	It allows the user to zoom in/out of the heat	unchecked
	map matrix	
Is Square Matrix?(Input contact file)	Allows the user to specify if the input is a	unchecked
	Square matrix (a full matrix) or a sparse	
	matrix. If checked, it displays a textbox for	
	the user to specify the matrix resolution.	
Specify Resolution	It is visible only if <i>Is SquareMatrix?</i> is	NA
	checked. It allows user specify resolution for	
	the input matrix.	
Input contact file	A text file containing a contact matrix in any	NA
	of the format described above.	
Title	Allows user to specify the title of the	Heatmap
	heatmap	Display
X-Axis Title	Allows user to specify the X-Axis title for	Genome bin
	the heatmap	Resolution (bp)
Y-Axis Title	Allows user to specify the Y-Axis title for	Genome bin
	the heatmap	Resolution (bp)
X min	It allows the user to specify the minimum X-	0
	axis Tick for the heatmap	
X max	It allows the user to specify the maximum X-	200
	axis Tick for the heatmap	
Y min	It allows the user to specify the minimum Y-	0
	axis Tick for the heatmap	
Y max	It allows the user to specify the maximum Y-	200
	axis Tick for the heatmap	

X min [Genome Location	It shows the genomic position equivalent for	0
Equivalent]	the minimum X-axis Tick for the heatmap	
X max[Genome Location Equivalent]	It shows the genomic position equivalent for	8000000
	the maximum X-axis Tick for the heatmap	
Y min [Genome Location	It shows the genomic position equivalent for	0
Equivalent]	the minimum Y-axis Tick for the heatmap	
Y [Genome Location Equivalent]	It shows the genomic position equivalent for	8000000
	the maximum Y-axis Tick for the heatmap	
Number of Units detected	It shows the number of regions found in the	200
	input matrix	
Number of Missing Units	It shows the number of gaps or missing	0
	regions noted from the input matrix	
Resolution detected	It displays the resolution of the input matrix	40000
Initial Start Position	It shows the minimum genome position	0
	observed from the input matrix	
Initial End Position	It shows the maximum genome position	8000000
	observed from the input matrix	
Gradient	An array of Color used as a gradient. One	НОТ
	color is used as the bottom gradient and	
	another color is used as the top gradient.	
	Hence, it produces a gradient from one color	
	to the other. The Gradient Colors are	
	explained below	
GRADIENT_BLACK_TO_WHITE	Produces a gradient from black (low) to	
	white (high)	
GRADIENT_BLUE_TO_RED	Produces a gradient from blue (low) to red	
	(high)	
GRADIENT_HEAT	Produces a different gradient for hot things	
	(black, brown, orange, white)	

GRADIENT_HOT	Produces a gradient for hot things (black,	
	red, orange, yellow, white)	
GRADIENT_MAROON_TO_GOLD	Produces a gradient from maroon (low) to	
	gold (high)	
GRADIENT_RAINBOW	Produces a gradient through the rainbow:	
	violet, blue, green, yellow, orange, red	
GRADIENT_RED_TO_GREEN	Produces a gradient from red (low) to green	
	(high)	
GRADIENT_ROY	Produces a gradient through red, orange,	
	yellow	
Data Type	It determines the type of data to be	TANH
	displayed. The types available are the raw	
	input data, a Tanh of input data, a Pearson	
	correlation of input data, and a Spearman	
	correlation of the input data.	

2.4.6 TAD Annotation

The description of the display controls on the display window for TAD annotation is given below.

Field	Description	Default
Load TAD file	Browse and Load a .bed format file	NA
	containing the TADs identified for the input	
	matrix	
Identified TAD	It shows the TADs in the input file	NA
Show TAD on Heatmap	It marks the boundary of the TADs	
	identified on the displayed heatmap	
Display Multiple TADs	Once checked, allows TADs from different	unchecked
	method to be overlapped on the same display	
	window. This function is useful for	

	comparing TADs identified by different	
	methods for a dataset.	
Choose Display Color	Choose the color for the TAD boundary	Color 1
	marks	

2.4.7 Demonstration

Figure 5 below shows the TAD annotation for the TADs identified by two TAD identification algorithms (ClusterTAD and DI) for mESC Chromosome 17 from Ren Lab.

Step 1:

To run this demonstration, load a sample square matrix as the input contact file.

The example file can be found here: /executable/sample_data/

contact_matrices/square_matrices/mESC_nij.chr17. Resolution for the square matrix = 40000

Load the contact file as instructed here: <u>Visualizing Dataset in 2D format</u>

Step 2:

Modify the highlighted fields on the display window. The table below shows the values set for each field in the display control.

Field	Value
Draw Title	checked
Draw Legend	checked
Draw X-Axis Title	checked
Draw X-Axis Ticks	checked
Draw Y-Axis Title	checked
Draw Y-Axis Ticks	checked
Heatmap Direction(Left/Right)	checked
Enable Zoom Mode	unchecked
Is SquareMatrix?(Input contact file)	checked
Specify Resolution	<mark>40000</mark>

Input contact file	Path/to/chr17/inputfile
Title	HeatMap Display
X-Axis Title	Number of Bins
Y-Axis Title	Number of Bins
X min	500
X max	<mark>700</mark>
Y min	<u>500</u>
Y max	<mark>700</mark>
X min [Genome Location Equivalent]	20000000
X max[Genome Location Equivalent]	28000000
Y min [Genome Location Equivalent]	20000000
Y [Genome Location Equivalent]	28000000
Number of Units detected	2382
Number of Missing Units	0
Resolution detected	40000
Initial Start Position	0
Initial End Position	95240000
Gradient	НОТ
Data Type	TANH

Step 3:

- a. Browse & Load the ClusterTAD file found here:
 - ClusterTAD: /executable/sample_data/TAD_annotation/mESC_TAD_bed/ ClusterTAD /chr17.bed.
- b. Select a Unique from Color 1 to 4. (Ex: Color 1 for ClusterTAD and Color 2 for DI)
- c. Click the *Show TAD on Heatmap* button.

Step 4:

To display multiple TADs on the Heatmap, Mark/Check the *Display Multiple TADs* then Repeat <u>Step 3</u> with the DI file found here:

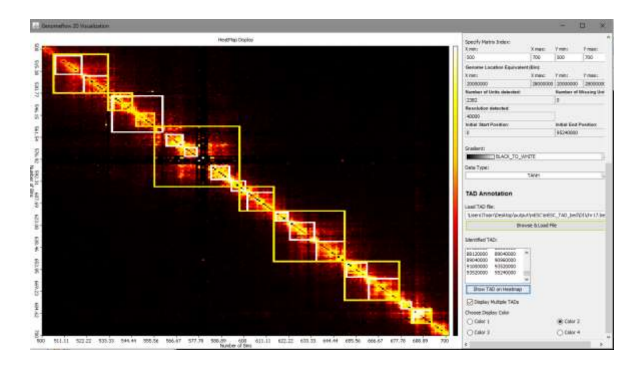


Figure 10: Demonstration of TAD Annotation on 2D Heatmap

2.5 Identify TAD

2.5.1 Purpose

To identify Topological Associated domains from input contact matrix.

2.5.2 Input

An input file in square matrix format (a full matrix representing all the contact regions) or a sparse matrix format (each line represents a contact by three numbers separated by whitespaces: position1 postion2 interaction_frequency(IF)).

An example sparse matrix file can be found here:

/executable/sample_data/ contact_matrices/ chr11_10kb_gm12878_list_125mb_135mb.txt

2.5.3 Output

A TAD with the best quality will be generated prefixed with **BestTAD_** in **bed** format. This file will be found here: /Selected_output_directory_from_GUI/Output/TADs/.

2.5.4 Running

Access the function from the menu toolbar: 2D-Functions/Identify TAD.

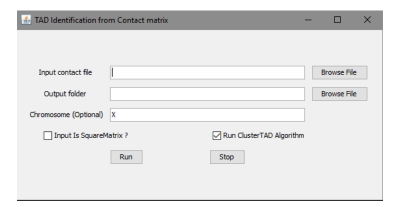


Figure 11: Identifying TADs on a contact matrix

Field	Description	Default
Input contact file	An input file in any of the format described	NA
	above	
Output folder	Directory to output the comparison report	NA
Is SquareMatrix?(Input contact file)	Allows the user to specify if the input is a	unchecked
	Square matrix (a full matrix) or a sparse	
	matrix. If checked, it displays a textbox for	
	the user to specify the matrix resolution.	
Data Resolution	It is visible only if <i>Is SquareMatrix?</i> is	40000
	checked. It allows user specify resolution for	
	the input matrix.	
Chromosome (optional)	Allows user to specify the chromosome data	X

Run ClusterTAD Algorithm	The default algorithm used for TAD	checked
	identification from the input contact Matrix	
Run	To start the identification process. A	NA
	progress bar is displayed to show the steps	
	taken by the TAD identification algorithm,.	
Stop	During the identification, if this button is	NA
	pressed, the program will stop.	

2.6 Check TAD consistency between two TADs from different methods

2.6.1 Purpose

To compare two TADs from two different Topological Associated domains identification method.

2.6.2 Input

A file containing TADs in .bed format. The method whose TADs consistency is to be checked is termed Method-1, and the methods whose TADs is to be compared with is termed Method-2. Choose the same chromosome for different methods. For example, to compare TAD from ClusterTAD with DI for chromosome 17,

Method-1 = /executable/sample_data/TAD_annotation/mESC_TAD_bed/ ClusterTAD /chr17.bed.

Method-2 = /executable/sample_data/TAD_annotation/mESC_TAD_bed/ DI /chr17.bed

2.6.3 Output

A report of the consistency of the Method-1 with Method-2. The output reports the following cases:

Case	Description
------	-------------

Case 1	The number of Exact TADs found in both
	Method-1 and Method-2
Case 2	The number of Sub-TADs that exist between
	Method-1 and Method-2
Case 3	The number of Conflicting TADs.
Case 4	The number of TADs in Method-1 but not found
	in Method-2

2.6.4 Running

Access the function from the menu toolbar: 2D-Functions/Check TAD Consistency.

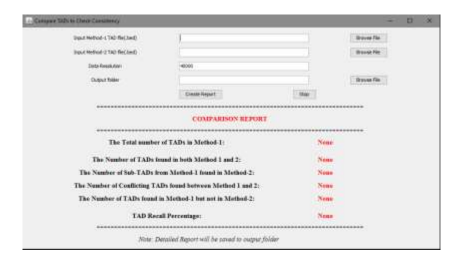


Figure 12: Comparing two TADs for a consistency check

Field	Description	Default
Input Method-1 TAD file(.bed)	Browse the .bed format file containing the	NA
	TADs identified by Method-1	
Input Method-2 TAD file(.bed)	Browse the .bed format file containing the	NA
	TADs identified by Method-2	
Data Resolution	The Resolution of the dataset the TADs were	40000
	identified from.	

Output folder	Directory to output the comparison report	NA
Create Report	Once this button is pressed, a progress bar is	NA
	displayed to show the steps taken by the	
	TAD identification algorithm,.	
Stop	During the check, if this button is pressed,	NA
	the program will stop.	

3 3D-Functions

3.1 3D model reconstruction by LorDG

3.1.1 Purpose

To build 3D chromosomes and genome models

3.1.2 Input

A contact matrix in sparse matrix format

3.1.3 Output

3D models in. gss format file and .pdb format file

3.1.4 Running

Access the function from the menu toolbar: 3D-Functions/LorDG-3D Modeller

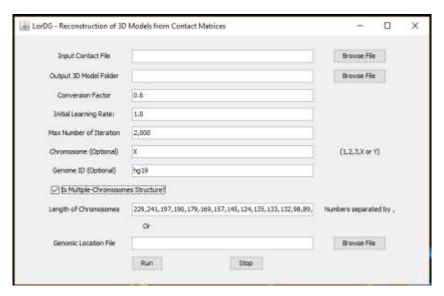


Figure 13: 3D Model reconstruction by LorDG

Field /Button	Description	Default
Conversion	α in the formula $d_{ij} = \frac{1}{IF_{ij}^{\alpha}}$, where IF_{ij} is interaction	1.0
Factor	frequency between <i>i</i> and <i>j</i> . When the field is left blank,	
	the program will search for the best value in the range	
	[0.1-3.0] with a step size of 0.1. Users can also specify a	
	range to search by put 2 numbers separated by a hyphen	
	(e.g. 0.5-1.0). During the searching, the right-top corner	
	of the main screen displays information about the current	
	value being tested.	
Initial Learning	Initial learning rate of the optimization. Higher learning	1.0
Rate	rate can speed up the reconstruction process but can	
	cause the process to fail as well	
Max number of	Maximum number of iterations for the optimization	1000
Iteration		
Chromosome	Chromosome name of the contact matrix in the input. If	X
	the input contains contact matrix of the whole genome,	
	leave this field blank.	
Genome ID	Genome version of the contact matrix in the input.	hg19

Is Multiple-	if the input contains both inter-and intra-chromosomal	unchecked
Chromosomes	contacts data, this checkbox should be checked.	
Structure?	contacts data, this eneckbox should be enecked.	
Length of	This field contains a list of lengths of chromosomes in	
Chromosomes	increasing order of chromosome names and separated by	
	commas, if "Is Multiple-Chromosomes Structure" is	
	checked. Please note that these lengths should not	
	contain omitted regions (e.g. centromeres) in the input of	
	chromosomes.	
Run	To start the reconstruction process. The main screen	NA
	displays how models are being formed from initially	
	random models. The information about the	
	reconstruction is displayed in the top-right corner of the	
	main screen. The conversion factor is being used to build	
	model and the current value of the objective function	
	(higher is better). After the reconstruction is finished, the	
	score of the model is displayed in the top-right corner of	
	the main screen (the lower the value is, the better the	
	model is).	
Stop	During the reconstruction, if this button is pressed, the	NA
	program will stop and output the currently best structure.	
	If the program is searching for the best conversion factor,	
	it will stop the searching and use the best-found	
	conversion factor to build models.	

3.2 3D model reconstruction by 3DMax

3.2.1 Purpose

To build 3D chromosomes and genome models

3.2.2 Input

A contact matrix in sparse matrix format

3.2.3 Output

3D models in .gss format file and .pdb format file

3.2.4 Running

Access the function from the menu toolbar: 3D-Functions/3DMax-3D Modeller

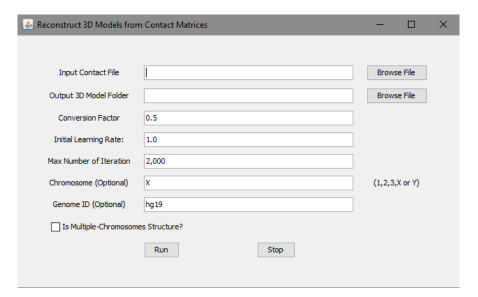


Figure 14: 3D Model reconstruction by 3DMax

Field /Button	Description	Default
Conversion	α in the formula $d_{ij} = \frac{1}{IF_{ij}^{\alpha}}$, where IF_{ij} is interaction	0.5
Factor	frequency between i and j . When the field is left blank,	
	the program will search for the best value in the range	
	[0.1-2.0] with a step size of 0.1. Users can also specify a	
	range to search by put 2 numbers separated by a hyphen	
	(e.g. 0.5-1.0). During the searching, the right-top corner	
	of the main screen displays information about the current	
	value being tested.	

Initial learning rate of the optimization. Higher learning	1.0
rate can speed up the reconstruction process but can	
cause the process to fail as well	
Maximum number of iterations for the optimization	2000
Chromosome name of the contact matrix in the input. If	X
the input contains contact matrix of the whole genome,	
leave this field blank.	
Genome version of the contact matrix in the input.	hg19
if the input contains both inter-and intra-chromosomal	unchecked
contacts data, this checkbox should be checked.	
This field contains a list of lengths of chromosomes in	
increasing order of chromosome names and separated by	
commas, if "Is Multiple-Chromosomes Structure" is	
checked. Please note that these lengths should not	
contain omitted regions (e.g. centromeres) in the input of	
chromosomes.	
To start the reconstruction process. The main screen	NA
displays how models are being formed from initially	
random models. The information about the	
reconstruction is displayed in the top-right corner of the	
main screen. The conversion factor is being used to build	
model and the current value of the objective function	
(higher is better). After the reconstruction is finished, the	
score of the model is displayed in the top-right corner of	
the main screen (the lower the value is, the better the	
model is).	
During the reconstruction, if this button is pressed, the	NA
program will stop and output the currently best structure.	
If the program is searching for the best conversion factor,	
	rate can speed up the reconstruction process but can cause the process to fail as well Maximum number of iterations for the optimization Chromosome name of the contact matrix in the input. If the input contains contact matrix of the whole genome, leave this field blank. Genome version of the contact matrix in the input. if the input contains both inter-and intra-chromosomal contacts data, this checkbox should be checked. This field contains a list of lengths of chromosomes in increasing order of chromosome names and separated by commas, if "Is Multiple-Chromosomes Structure" is checked. Please note that these lengths should not contain omitted regions (e.g. centromeres) in the input of chromosomes. To start the reconstruction process. The main screen displays how models are being formed from initially random models. The information about the reconstruction is displayed in the top-right corner of the main screen. The conversion factor is being used to build model and the current value of the objective function (higher is better). After the reconstruction is finished, the score of the model is displayed in the top-right corner of the main screen (the lower the value is, the better the model is). During the reconstruction, if this button is pressed, the program will stop and output the currently best structure.

it will stop the searching and use the best-found conversion factor to build models.

3.3 Chromatin loop identification

3.3.1 Purpose

To identify chromatin loop in 3D models

3.3.2 Input

A 3D model to visualize

3.3.3 Output

A list of chromatin loops in a bed format file (optional) and highlighted in the 3D model

3.3.4 Running

Access the function from the menu toolbar: 3D-Functions/Loop Detection

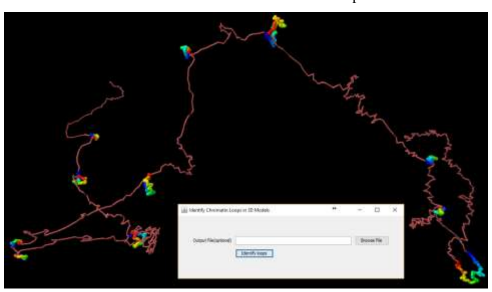


Figure 15: Chromatin loops

The function identifies chromatin loops and highlights them in the 3D model. The loops can also be outputted into a bed format file specified in the Output File field. The top-right corner of the main screen displays the number of chromatin loops identified.

Loops are colored in spectrum (from blue to red). To highlight loops better, color the model by a single color (right-click on the main screen, choose color/structure/chain)

3.4 Model annotation

3.4.1 Purpose

To annotate 3D models with genomic elements

3.4.2 Input

A 3D model (e.g. in executable/sample_data/models) and genomic elements in bed format files (e.g. in executable/track_files)

3.4.3 Output

3D model is annotated with data from bed format files

3.4.4 Running

Access the function from the menu toolbar: 3D-Functions/Model Annotation

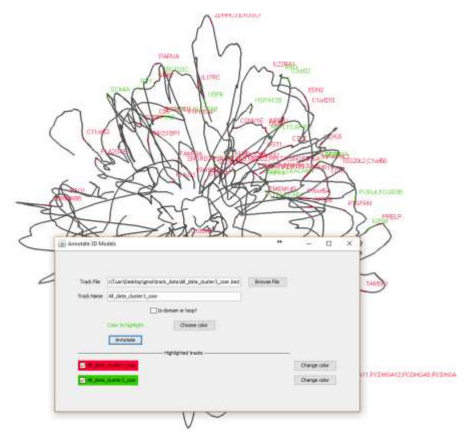


Figure 16: Function to annotate 3D models

To better highlight track data, change the color of the model to a sing color (right-click on the main screen, Color/Structure/Reset). The background can be changed to white to (Color/Background/White)

Field /	Description	Default
Button		
Track file	A file in bed format (see executable/track_files for example) to annotate the model	NA
Track name	A unique name associated with the above input file	Name of track file
Is domain or loop?	Indicate if the track file contains domains or loops. Adjacent domains/loops will be colored in red/blue alternatively.	Unchecked
Choose	To pick a color to label annotation and points overlapped by genomic elements in the track file.	Random
Change color	To change color of the corresponding track	NA
	Checking corresponding track names will display or hidden the content of tracks.	

To get the genomic coordinate of a point, left-click or mouse-over to the point as shown in **Figure**.

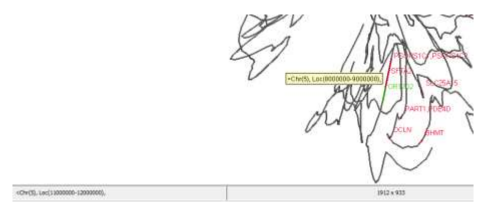


Figure 17: Coordinate of a point in the model

3.5 Gene expression data visualization (a special case of model annotation)

3.5.1 Purpose

To display gene expression level along a 3D model

3.5.2 Input

- a) A 3D model in GSS format (e.g. in executable/sample_data/models/chr11_10kb_gm12878_list_60mb_70mb_1514493462531.gss) to visualize,
- A gene expression data file in GCT format (
 http://software.broadinstitute.org/cancer/software/genepattern/file-formats-guide#GCT
), an example file is executable/sample_data/gene_expression/allaml.dataset.gct .
- c) And a text file to specify genomic coordinates of probes/genes in the GCT format file (each line consists of 4 elements separated by space or tab, e.g.: probe_or_gene_name chr_number start end). A sample is executable/sample_data/gene_expression/probe_coordinates.txt

These 3 following files are prepared for demo: executable/sample_data/models/chr11_10kb_gm12878_list_60mb_70mb_1514493462531.gss,

executable/sample_data/gene_expression/allaml.dataset.gct and executable/sample_data/gene_expression/ probe_coordinates.txt.

3.5.3 Output

Expression levels of genes/probes are annotated in the 3D model. Usually, the GCT file contains several samples and therefore, the median value (across all samples) together with minimum and maximum values (in brackets) are displayed next to probe/gene names. If the 3D model and the gene expression data file have no overlap, no annotation will be added to the 3D model.

3.5.4 Running

Access the function from the menu toolbar: 3D-Functions/Model Annotation. A GCT file must be filled in the "Track File" field.

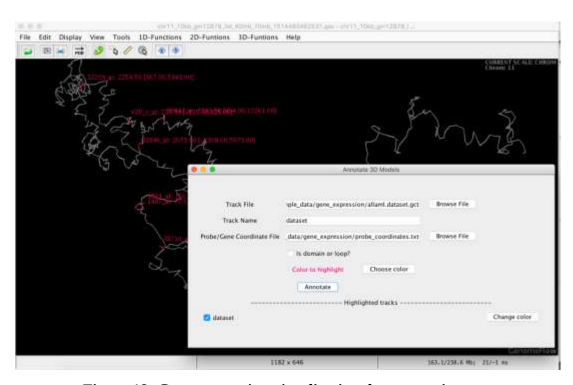


Figure 18: Gene expression visualization demonstration

3.6 Comparing 2 models

3.6.1 Purpose

To superimpose and compare two 3D-models in GSS format.

3.6.2 Input

Two chromosome models in GSS format.

3.6.3 Output

The two models are scaled, superimposed and visualized. Spearman's correlation and RMSE between pairwise distances of the two models are calculated.

3.6.4 Running

Access the function from the menu toolbar: 3D-Functions/Compare Models

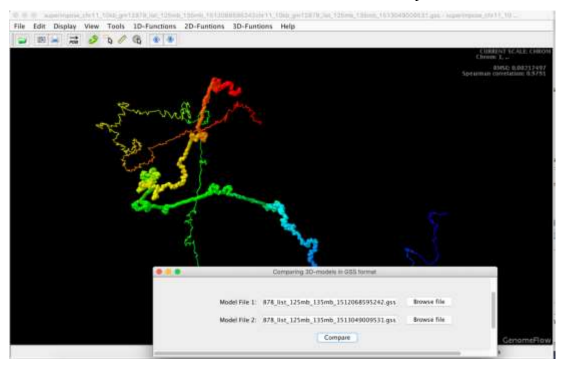


Figure 19: Comparing two constructed models