RNA-Hi-C-tools Documentation

Release 0.3.2

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CONTENTS

RNA-Hi-C-tools 0.3 documentation	3
	3
	4
1.3 Support	4
Analysis pipeline	5
2.1 Overview	5
2.2 Pipeline	5
2.3 Other functions	13
Visualization of local RNA-RNA interactions	15
3.1 Prerequirement	15
3.2 Run the program to generate visualization	15
3.3 Example of result graph	16
Visualization of global RNA-RNA interactome	17
4.1 Prerequirement	17
4.2 Run the program to generate visualization	17
4.3 Example of result graph	17
Python APIs created for this project	19
5.1 Annotation module	19
5.2 "annotated_bed" data class	20
5.3 "RNAstructure" class	21
Updates	25
Indices and tables	27
ython Module Index	29
	1.1 Overview 1.2 Installation 1.3 Support Analysis pipeline 2.1 Overview 2.2 Pipeline 2.3 Other functions Visualization of local RNA-RNA interactions 3.1 Prerequirement 3.2 Run the program to generate visualization 3.3 Example of result graph Visualization of global RNA-RNA interactome 4.1 Prerequirement 4.2 Run the program to generate visualization 4.3 Example of result graph Python APIs created for this project 5.1 Annotation module 5.2 "annotated_bed" data class 5.3 "RNAstructure" class Updates Indices and tables

Contents:

CONTENTS 1

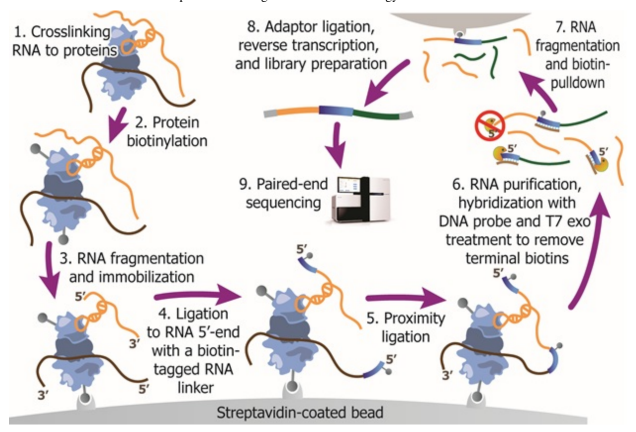
2 CONTENTS

RNA-HI-C-TOOLS 0.3 DOCUMENTATION

1.1 Overview

RNA-Hi-C-tools is a set of bioinformatic tools for analysis of a novel DNA sequencing based technology to detect RNA-RNA interactome and RNA-chromatin interactome (RNA-chromatin interactome is coming soon).

Below is a illustration for the experimental design of this new technology



See also:

Offline documentation.

Download a copy of RNA-Hi-C-tools documentation:

- PDF
- Epub

1.2 Installation

1.2.1 step 1: Install the dependent prerequisites:

- 1. Python libraries [for python 2.x]:
- Biopython
- Pysam
- BAM2X
- Numpy, Scipy
- Parallel python (Only for Select_strongInteraction_pp.py)
- 2. The Boost.Python C++ library
- 3. Other softwares needed:
- Bowtie (not Bowtie 2)
- samtools
- NCBI blast+ (use blastn)

1.2.2 Step 2: Download the package

Clone the package from GitHub:

```
git clone http://github.com/yu68/RNA-Hi-C.git
```

1.2.3 Step 3: Add library source to your python path

Add these lines into your ~/.bash_profile or ~/.profile

```
Location="/path/of/RNA-Hi-C-tools" # change accordingly export PYTHONPATH="$Location/src:$PYTHONPATH" export PATH="$PATH:$Location/bin"
Loc_lib="/path/of/boost_1_xx_0/lib/" # change accordingly export LD_LIBRARY_PATH="$Loc_lib:$LD_LIBRARY_PATH"
```

1.3 Support

For issues related to the use of RNA-Hi-C-tools, or if you want to **report a bug or request a feature**, please contact Pengfei Yu <p3yu at ucsd dot edu>

CHAPTER

TWO

ANALYSIS PIPELINE

2.1 Overview

The next generation DNA sequencing based technology utilize RNA proximity ligation to transfrom RNA-RNA interactions into chimeric DNAs. Through sequencing and mapping these chimeric DNAs, it is able to achieve high-throughput mapping of nearly entire interaction networks. RNA linkers were introduced to mark the junction of the ligation and help to split the chimeric RNAs into two interacting RNAs. This bioinformatic pipeline is trying to obtain the strong interactions from raw fastq sequencing data. The major steps are:

- Step 1: Remove PCR duplicates.
- Step 2: Split library based on barcode.txt.
- Step 3: Recover fragments for each library.
- Step 4: Split partners and classify different types of fragments.
- Step 5: Align both parts of "Paired" fragment to the genome.
- Step 6: Determine strong interactions.
- Step 7: Visualization of interactions and coverages.

Other functions:

- 1. Determine the RNA types of different parts within fragments.
- 2. Find linker sequences within the library.
- 3. Find intersections between two different interaction sets.

2.2 Pipeline

2.2.1 Step 1: Remove PCR duplicates.

Starting from the raw pair-end sequencing data, PCR duplicates should be removed as the first step if both the 10nt random indexes and the remaining sequences are exactly the same for two pairs. It is achieved by remove_dup_PE.py

```
usage: remove_dup_PE.py [-h] reads1 reads2

Remove duplicated reads which have same sequences for both forward and reverse reads. Choose the one appears first.

positional arguments:
   reads1    forward input fastq/fasta file
```

```
reads2 reverse input fastq/fasta file
optional arguments:
   -h, --help show this help message and exit
Library dependency: Bio, itertools
```

The program will generate two fastq/fasta files after removind PCR duplicates and report how many read pairs has been removed. The output are prefixed with 'Rm_dupPE'

Note: One pair is considered as a PCR duplicate only when the sequences of both two ends (including the 10nt random index) are the exactly same as any of other pairs.

2.2.2 Step 2: Split library based on barcode.txt.

After removing PCR duplicates, the libraries from different samples are separated based on 4nt barcodes in the middle of random indexes ("RRRBBBBRRR"; R: random, B: barcode). It is implemented by split_library_pairend.py

```
usage: split_library_pairend.py [-h] [-f | -q] [-v] [-b BARCODE]
                                [-r RANGE [RANGE ...]] [-t] [-m MAX_SCORE]
                                input1 input2
Example: split_library_pairend.py -q Rm_dupPE_example.F1.fastq
        Rm_dupPE_example.R1.fastq -b barcode.txt
positional arguments:
                       input fastq/fasta file 1 for pairend data (contain
 input1
                       barcodes)
 input2
                       input fastq/fasta file 2 for pairend data
optional arguments:
 -h, --help
                      show this help message and exit
 -f, --fasta
                      add this option for fasta input file
  -q, --fastq
                       add this option for fastq input file
                show program's version number and exit
 -v, --version
 -b BARCODE, --barcode BARCODE
                       barcode file
 -r RANGE [RANGE ...], --range RANGE [RANGE ...]
                        set range for barcode location within reads, default is
                       full read
 -t, --trim
                       trim sequence of 10nt index
 -m MAX_SCORE, --max_score MAX_SCORE
                        max(mismatch+indel) allowed for barcode match,
                        otherwise move reads into 'unassigned' file
                        default: 2.
Library dependency: Bio
Here is a example for barcode.txt
```

The output of this script are several pairs of fastq/fasta files prefixed with the 4nt barcode sequences, together with another pair of fastq/fasta files prefixed with 'unassigned'.

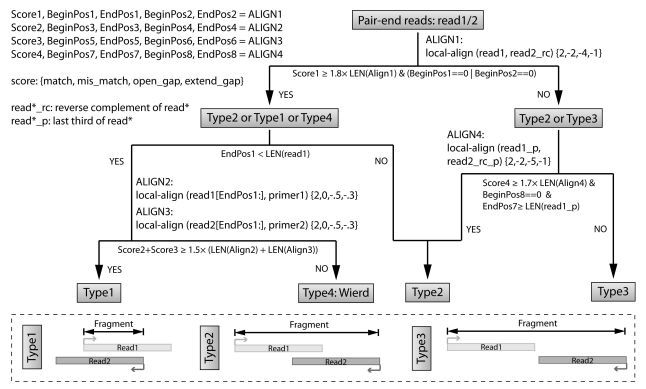
ACCT CCGG GGCG For example, if the input fastq/fasta files are Rm_dupPE_example.F1.fastq and Rm_dupPE_example.R1.fastq, and the barcode file is the same as above, then the output files are:

- ACCT_Rm_dupPE_example.F1.fastq
- ACCT_Rm_dupPE_example.R1.fastq
- CCGG_Rm_dupPE_example.F1.fastq
- CCGG_Rm_dupPE_example.R1.fastq
- GGCG_Rm_dupPE_example.F1.fastq
- GGCG_Rm_dupPE_example.R1.fastq
- unassigned_Rm_dupPE_example.F1.fastq
- unassigned_Rm_dupPE_example.R1.fastq

2.2.3 Step 3: Recover fragments for each library.

After splitting the libraries, the later steps from here (Step 3-7) need to be executed parallelly for each sample.

In this step, we are trying to recover the fragments based on local alignment. The fragments are classifed as several different types as shown in the figure below. The flow chart is also clarified at the top.



We will use a complied program recoverFragment to do that

recoverFragment - recover fragment into 4 different categories from pair-end seq data

SYNOPSIS

DESCRIPTION

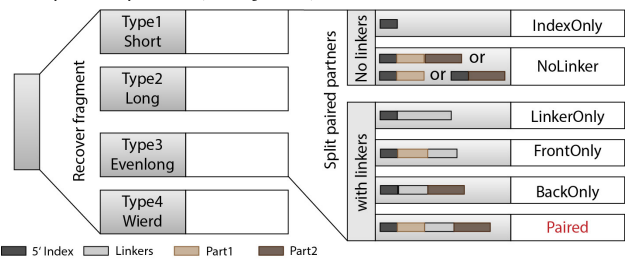
-h, --help

2.2. Pipeline 7

```
Displays this help message.
    --version
          Display version information
    -I, --inputs STR
          input of forward and reverse fastq file, path of two files separated by SPACE
    -p, --primer STR
          fasta file contianing two primer sequences
    -v, --verbose
          print alignment information for each alignment
EXAMPLES
    recoverFragment -I read_1.fastq read_2.fastq -p primer.fasta
          store fragment using fasta/fastg into 4 output files
          'short_*', 'long_*','evenlong_*','wierd_*'
VERSION
    recoverFragment version: 0.1
    Last update August 2013
```

2.2.4 Step 4: Split partners and classify different types of fragments.

When we recovered the fragments, the next we are goting to do is to find parts that are separated by the linkers, and from here, we will be able to classify the fragments into different types: "IndexOnly", "NoLinker", "LinkerOnly", "BackOnly", "FrontOnly", "Paired". (see the figure below).



This will be done by split_partner.py

```
type3_2
                        read_2 for evenlong (type3) fastq file
optional arguments:
 -h, --help
                        show this help message and exit
  -e EVALUE, --evalue EVALUE
                        cutoff evalues, only choose alignment with evalue less
                        than this cutoffs (default: 1e-5).
 --linker_db LINKER_DB
                        BLAST database of linker sequences
  --blast_path BLAST_PATH
                        path for the local blast program
  -o OUTPUT, --output OUTPUT
                        output file containing sequences of two sepatated
                        part.s
 -t TRIM, --trim TRIM trim off the first this number of nt as index,
                        default:10
 -b BATCH, --batch BATCH
                        batch this number of fragments for BLAST at a time.
                        default: 200000
  -r, --release
                        set to allow released criterion for Paired fragment in
                        Type 3, include those ones with no linker in two reads
 -1 LENGTH, --length LENGTH
                        shortest length to be considered for each part of the
                        pair, default: 15
Library dependency: Bio, itertools
```

Note: New option added in version 0.3.1, which could allow two different strategies for selection of "Paired" fragments from the Type3 fragments. The --release option will allow a read pair to be called as "Paired" fragment even when the linker are not detected in both reads.

The linker fasta file contain sequences of all linkers

```
>L1
CTAGTAGCCCATGCAATGCGAGGA
>L2
AGGAGCGTAACGTACCCGATGATC
```

The output fasta files will be the input file name with different prefix ("NoLinker", "LinkerOnly", "BackOnly", "FrontOnly", "Paired") for different types. The other output file specified by $-\circ$ contains information of aligned linker sequences for each Type1/2 fragment.

For example, if the commend is

```
split_partner.py fragment_ACCT.fasta evenlong_ACCTRm_dupPE_stitch_seq_1.fastq
    evenlong_ACCTRm_dupPE_stitch_seq_2.fastq
    -o fragment_ACCT_detail.txt --linker_db linker.fa
```

Then, the output files will be:

- backOnly_fragment_ACCT.fasta
- NoLinker_fragment_ACCT.fasta
- frontOnly_fragment_ACCT.fasta
- Paired1_fragment_ACCT.fasta
- Paired2_fragment_ACCT.fasta

2.2. Pipeline 9

fragment_ACCT_detail.txt

The format of the last output file fragment_ACCT_detail.txt will be "Name | linker_num | linker_loc | Type | linker order". Here are two examples:

```
HWI-ST1001:238:H0NYEADXX:1:1101:10221:1918 L1:2;L2:1 19,41;42,67;68,97 None L2;L1;L1 HWI-ST1001:238:H0NYEADXX:1:1101:4620:2609 L1:2 28,46;47,79 Paired L1;L1
```

In the **first** fragment, there are three regions can be aligned to linkers, 2 for L1 and 1 for L2, the order is L2, L1, L1. And they are aligned in region [19,41], [42,67], [68,97] of the fragment. "None" means this fragment is either 'LinkerOnly' or 'IndexOnly' (in this case it is 'LinkerOnly'). This fragment won't be written to any of the output fasta files.

In the **second** fragment, two regions can be aligned to linkers, and they are both aligned to L1. The two regions are in [28,46], [47,79] of the fragment. the fragment is "Paired" because on both two sides flanking the linker aligned regions, the length is larger than 15nt. The left part will be writen in Paired1_fragment_ACCT.fasta and the right part in Paired2_fragment_ACCT.fasta

2.2.5 Step 5: Align both parts of "Paired" fragment to the genome.

In this step, we will use the Paired1* and Paired2* fasta files output from the previous step. The sequences of part1 and part2 are aligned to the mouse genome mm9 with Bowtie and the pairs with both part1 and part2 mappable are selected as output. We also annotate the RNA types of each part in this step. All of these are implemented using script Stitch-seq_Aligner.py.

```
usage: Stitch-seq_Aligner.py [-h] [-s samtool_path] [-a ANNOTATION]
                            [-A DB_DETAIL]
                            miRNA_reads mRNA_reads bowtie_path miRNA_ref
                            mRNA_ref
Align miRNA-mRNA pairs for Stitch-seq. print the alignable miRNA-mRNA pairs
with coordinates
positional arguments:
 part1_reads paired part1 fasta file
 part2_reads
                     paired part2 fasta file
 bowtie_path
                     path for the bowtie program
                    reference genomic seq for part1
 part1_ref
 part2_ref
                      reference genomic seg for part2
optional arguments:
                   show this help message and exit
 -h, --help
 -b, --bowtie2
                      set to use bowtie2 (--sensitive-local) for alignment,
                      need to change reference index and bowtie_path
 -u, --unique
                      set to only allow unique alignment
 -s samtool_path, --samtool_path samtool_path
                       path for the samtool program
 -a ANNOTATION, --annotation ANNOTATION
                       If specified, include the RNA type annotation for each
                       aligned pair, need to give bed annotation RNA file
 -A DB_DETAIL, --annotationGenebed DB_DETAIL
                       annotation bed12 file for lincRNA and mRNA with intron
                       and exon
```

Library dependency: Bio, pysam, itertools

An annotation file for different types of RNAs in mm9 genome (bed format, 'all_RNAs-rRNA_repeat.txt.gz') was included in Data folder. The annotation bed12 file for lincRNA and mRNA ('Ensembl_mm9.genebed.gz') was

also included in Data folder. One can use the option -a ../Data/all_RNAs-rRNA_repeat.txt.gz -A ../Data/Ensembl mm9.genebed.gz for annotation.

Here is a example:

```
Stitch-seq_Aligner.py Pairedl_fragment_ACCT.fasta Paired2_fragment_ACCT.fasta ~/Software/bowtie-0.12.7/bowtie mm9 mm9 -s samtools -a ../Data/all_RNAs-rRNA_repeat.txt.gz -A ../Data/Ensembl_mm9.genebed.gz > ACCT_fragment_paired_align.txt
```

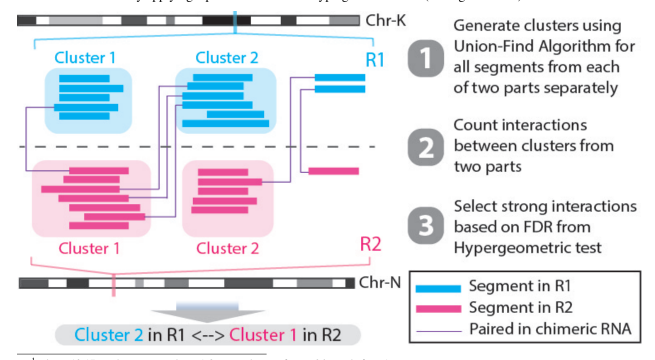
The format for the output file ACCT_fragment_paired_align.txt will be:

Column ¹	Description
1	chromosome name of part1
2,3	start/end position of part1
4	strand information of part1
5	sequence of part1
6	RNA type for part1
7	RNA name for part1
8	RNA subtype ² for part1
9	name of the pair

Note: Bowtie2 ("-sensitive-local" mode) option is added in version 0.3.1 for the user to choose, the reference index and bowtie_path need to be changed accordingly if you use bowtie2 instead of bowtie. User can also choose unique aligned reads or not by setting --unique option.

2.2.6 Step 6: Determine strong interactions.

In this step, we will generate clusters with high coverage separately for all part1 (R1) an part2 (R2) segments. Then based on the pairing information, we count the interactions between clusters from part1 and part2. The strong interactions can be selected by applying a p-value cutoff from hypergeometric test. (See figure below)



¹column 10-17 are the same as column 1-8 except they are for part2 instead of part1.

2.2. Pipeline 11

²subtype can be intron/exon/utr5/utr3 for lincRNA and mRNA (protein-coding), '.' for others

We will use the script Select_strongInteraction_pp.py, parallel computing are implemented for clustering parallelly on different chromosomes:

```
usage: Select_strongInteraction_pp.py [-h] -i INPUT [-M MIN_CLUSTERS]
                                      [-m MIN_INTERACTION] [-p P_VALUE]
                                      [-o OUTPUT] [-P PARALLEL] [-F]
find strong interactions from paired genomic location data
optional arguments:
 -h, --help
                        show this help message and exit
 -i INPUT, --input INPUT
                        input file which is the output file of Stitch-seq-
                       Aligner.py
 -M MIN_CLUSTERS, --min_clusterS MIN_CLUSTERS
                        minimum number of segments allowed in each cluster,
                        default:5
 -m MIN_INTERACTION, --min_interaction MIN_INTERACTION
                        minimum number of interactions to support a strong
                        interaction, default:3
 -p P_VALUE, --p_value P_VALUE
                        the p-value based on hypergeometric distribution to
                        call strong interactions, default: 0.05
 -o OUTPUT, --output OUTPUT
                        specify output file
 -P PARALLEL, --parallel PARALLEL
                        number of workers for parallel computing, default: 5
 -F, --FDR
                        Compute FDR if specified
need Scipy for hypergeometric distribution
```

The input of the script is the output of Step 5 (ACCT_fragment_paired_align.txt in the example). "annotated_bed" class is utilized in this script.

Here is a example:

Select_strongInteraction.py -i ACCT_fragment_paired_align.txt -o ACCT_interaction_clusters.txt

The column description for output file ACCT_interaction_clusters.txt is:

Column	Description
1	chromosome name of cluster in part1
2,3	start/end position of cluster in part1
4	RNA type for cluster in part1
5	RNA name for cluster in part1
6	RNA subtype for cluster in part1
7	# of counts for cluster in part1
8-14	Same as 1-7, but for cluster in part2
15	# of interactions between these two clusters
16	log(p-value) of the hypergeometric testing

2.2.7 Step 7: Visualization of interactions and coverages.

There are two ways of visulization provided (LOCAL and GLOBAL):

- Visualization of local interactions.
- Visualization of global interactome.

2.3 Other functions

2.3.1 Determine the RNA types of different parts within fragments.

2.3.2 Find linker sequences within the library.

2.3.3 Find intersections between two different interaction sets.

The script tool intersectInteraction.py could be used to identify overlap of interactions between two interaction set from independent experiments (two replicates or treatment v.s. control)

```
usage: intersectInteraction.py [-h] -a FILEA -b FILEB [-s START] [-n NBASE]
                               [-o OUTPUT] [-c]
find intersections (overlaps) between two interaction sets
optional arguments:
 -h, --help
                        show this help message and exit
 -a FILEA, --filea FILEA
                        file for interaction set a
 -b FILEB, --fileb FILEB
                        file for interaction set b
 -s START, --start START
                        start column number of the second part in each
                        interaction (0-based), default:7
  -n NBASE, --nbase NBASE
                        number of overlapped nucleotides for each part of
                      interactions to call intersections, default: 1
 -o OUTPUT, --output OUTPUT
                        specify output file
                        calculate p-values based on 100times permutations
 -p, --pvalue
require 'random'&'numpy'&'scipy' module if set '-p'
```

if "-p" option is set, then the program will do permutation for 100 times by shuffling the two partners of interactions in set a. A p-value will be calculate based on permutation distribution.

2.3.4 RNA structure prediction by adding digestion site information

The script will take selfligated chimeric fragments from given snoRNA (ID) and predict secondary structures with and without constraints of digested single strand sites. It is also able to compare the known structure in dot format if the known structure is available and specified by "-a". The script needs RNAStructure software for structure prediction ("-R") and and VARNA command line tool for visualization ("-v").

2.3. Other functions 13

```
linkedPair
                        file for information of linked pairs, which is output
                        of 'Stitch-seq_Aligner.py'
optional arguments:
 -h, --help
                        show this help message and exit
 -g GENOMEFA, --genomeFa GENOMEFA
                        genomic sequence, need to be fadix-ed
 -R RNASTRUCTUREEXE, --RNAstructureExe RNASTRUCTUREEXE
                        folder of RNAstrucutre suite excutable
 -a ACCEPTDOT, --acceptDot ACCEPTDOT
                        accepted structure in dot format, for comparing of
                        accuracy, no comparison if not set
 -o OUTPUT, --output OUTPUT
                        output distribution of digested sites with dot
                        structures, can be format of eps, pdf, png,...
 -s samtool_path, --samtool_path samtool_path
                        path for the samtool program
  -v VARNA, --varna VARNA
                        path for the VARNA visualization for RNA
  -c COLORMAPSTYLE, --colorMapStyle COLORMAPSTYLE
                        style of color map, choose from: "red", "blue",
                        "green", "heat", "energy", and "bw", default: "heat"
```

Here is a example:

```
python RNA_structure_prediction.py \
    ENSMUSG00000064380 \
    /data2/sysbio/UCSD-sequencing/2013-11-27-Bharat_Tri_Shu/Undetermined_indices/Sample_lane8/ACCT_GGCG
-a Snora73_real_dot.txt \
    -o Snora73_distribution.pdf
```

Here "Snora73_real_dot.txt" is dot format of known Snora73 structure This will generate three eps files with secondary structures ("Predict", "Refine", "Accepted (known)". Also the output pdf file contains the distribution of digested sites in whole RNA molecule.

VISUALIZATION OF LOCAL RNA-RNA INTERACTIONS

3.1 Prerequirement

This program require python modules: xplib, matplotlib, numpy, bx-python

3.2 Run the program to generate visualization

The script "Plot_interaction.py" will be used for this purpose,

```
usage: Plot_interaction.py [-h] [-n N] [-s START [START ...]] [-d DISTANCE]
                           [-g GENEBED] [-w PHYLOP_WIG] [-p PAIR_DIST] [-S]
                           [-o OUTPUT]
                           interaction linkedPair
plot linked pairs around a given interaction. information of linked pairs are
stored in file '*_fragment_paired_align.txt'
positional arguments:
                        Interaction file from output of
 interaction
                        'Select_strongInteraction_pp.py'
 linkedPair
                        file for information of linked pairs, which is output
                        of 'Stitch-seq_Aligner.py'
optional arguments:
 -h, --help
                        show this help message and exit
 -n N
                        Choose region to plot, it can be a number (around n-th
                        interaction in the interaction file) or one/two
                        regions with format 'chr:start-end', default=1
 -s START [START ...], --start START [START ...]
                        start column number of the second region in
                        interaction file and linkedPair file, default=(7,8)
 -d DISTANCE, --distance DISTANCE
                        the plus-minus distance (unit: kbp) flanking the
                        interaction regions to be plotted, default=10
 -g GENEBED, --genebed GENEBED
                        the genebed file from Ensembl, default:
                        ../Data/Ensembl_mm9.genebed
 -w PHYLOP_WIG, --phyloP_wig PHYLOP_WIG
                        the bigWig file for phyloP scores, defualt:
                        mouse.phyloP30way.bw
 -p PAIR_DIST, --pair_dist PAIR_DIST
                        two interacted parts within this distance are
```

```
considered as self-ligated and they are marked or eliminated (see option -s for slim mode), default:
200bp
-S, --Slim set slim mode to eliminate self ligated interactions
-o OUTPUT, --output OUTPUT output plot file, can be format of emf, eps, pdf, png, ps, raw, rgba, svg, svgz
```

Note: linkedPair file is the output *_fragment_paired_align.txt from *Step5:Stitch-seq_Aligner.py* of the pipeline; Interaction txt file is the output of *Step6:Select_strongInteraction_pp.py*.

3.3 Example of result graph

Example code:

```
python Plot_interaction.py
    ACCT_interaction_clusters_rmrRNA.txt \
    ACCT_fragment_paired_align_rmRNA_sort.txt.gz \
    -n 2412 \
    -d 5 \
    -o local_interaction.pdf
```

5796000

Result figure:

5797000

chr19:5796284-5796519 <-> chr19:5796822-5797016

Explanation:

5795000

chr19

5798000

5799000

CHAPTER

FOUR

VISUALIZATION OF GLOBAL RNA-RNA INTERACTOME

4.1 Prerequirement

This program is powered by RCircos.

Required R packages (our program will check for the presence of these packages and install/load them automatically if not present):

• argparse, RCircos, biovizBase, rtracklayer

The program also require a python script "bam2tab.py" (already in /bin/ folder) to call coverage from BAM2X

4.2 Run the program to generate visualization

We will use the script "Plot_Circos.R" for this purpose.

```
usage: Plot_Circos.R [-h] [-g GENOME] [-b BIN] [-o OUTPUT]
                  interaction part1 part2
positional arguments:
 interaction
                       the interaction file, [required]
 part1
                       aligned BAM file for part1, [required]
 part2
                       aligned BAM file for part2, [required]
optional arguments:
 -h, --help
                        show this help message and exit
 -g GENOME, --genome GENOME
                        genome information, choice: mm9/mm10/hg19 et.al.,
                        [default: mm9]
 -b BIN, --bin BIN
                       window size for the bins for coverage calling, [default: 100000.0]
 -o OUTPUT, --output OUTPUT
                        output pdf file name, [default: Interactome_view.pdf]
```

Note: part1, part2 BAM files are the ones generated from *Step5:Stitch-seq_Aligner.py* of the pipeline; Interaction txt file is the output of *Step6:Select_strongInteraction_pp.py*.

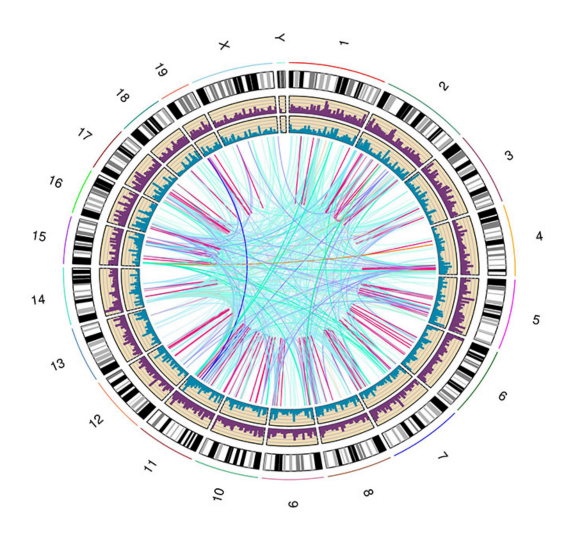
4.3 Example of result graph

Example code:

Rscript Plot_Circos.R GGCG_interaction_clusters.txt
 sort_Paired1_fragment_GGCG.bam sort_Paired2_fragment_GGCG.bam
 -b 100000 -o Interactome_GGCG.pdf

Result figure:

RNA-RNA interactome



Explanation:

CHAPTER

FIVE

PYTHON APIS CREATED FOR THIS PROJECT

5.1 Annotation module

For the purpose of annotating RNA types for genomic regions.

```
Annotation.overlap(bed1, bed2)
```

This function compares overlap of two Bed object from same chromosome

Parameters

- bed1 A Bed object from xplib.Annotation.Bed (BAM2X)
- bed2 A Bed object from xplib.Annotation.Bed (BAM2X)

Returns boolean - True or False

Example:

```
>>> from xplib.Annotation import Bed
>>> from Annotation import overlap
>>> bed1=Bed(["chr1",10000,12000])
>>> bed2=Bed(["chr1",9000,13000])
>>> print overlap(bed1,bed2)
True
```

Annotation. Subtype (bed1, genebed)

This function determines intron or exon or utr from a BED12 file.

Parameters

- **bed1** A Bed object defined by xplib.Annotation.Bed (BAM2X)
- genebed A Bed12 object representing a transcript defined by xplib Annotaton.Bed with information of exon/intron/utr from an BED12 file

Returns str – RNA subtype. "intron"/"exon"/"utr3"/"utr5"/"."

Example:

```
>>> from xplib.Annotation import Bed
>>> from xplib import DBI
>>> from Annotation import Subtype
>>> bed1=Bed(["chr13",40975747,40975770])
>>> a=DBI.init("../../Data/Ensembl_mm9.genebed.gz","bed")
>>> genebed=a.query(bed1).next()
>>> print Subtype(bed1,genebed)
"intron"
```

Annotation.annotation(bed, ref_allRNA, ref_detail, ref_repeat)

This function is based on <code>overlap()</code> and <code>Subtype()</code> functions to annotate RNA type/name/subtype for any genomic region.

Parameters

- **bed** A Bed object defined by xplib.Annotation.Bed (in BAM2X).
- ref_allRNA the DBI.init object (from BAM2X) for bed6 file of all kinds of RNA
- ref_detail the DBI.init object for bed12 file of lincRNA and mRNA with intron, exon, UTR
- ref_detail the DBI.init object for bed6 file of mouse repeat

Returns list of str – [type,name,subtype]

Example:

```
>>> from xplib.Annotation import Bed
>>> from xplib import DBI
>>> from Annotation import annotation
>>> bed=Bed(["chr13",40975747,40975770])
>>> ref_allRNA=DBI.init("../../Data/all_RNAs-rRNA_repeat.txt.gz","bed")
>>> ref_detail=DBI.init("../../Data/Ensembl_mm9.genebed.gz","bed")
>>> ref_repeat=DBI.init("../../Data/mouse.repeat.txt.gz","bed")
>>> print annotation(bed,ref_allRNA,ref_detail,ref_repeat)
["protein_coding","gcnt2","intron"]
```

5.2 "annotated bed" data class

```
class data structure.annotated bed(x=None, **kwargs)
```

To store, compare, cluster for the genomic regions with RNA annotation information. Utilized in the program Select_stronginteraction_pp.py

Cluster(c)

Store cluster information of self object

Parameters c – cluster index

Example:

```
>>> a=annotated_bed(chr="chr13", start=40975747, end=40975770)
>>> a.Cluster(3)
>>> print a.cluster
3
```

Note: a.cluster will be the count information when a become a cluster object in *Select_stronginteraction_pp.py*

Update (S, E)

Update the upper and lower bound of the cluster after adding segments using Union-Find.

Parameters

- \mathbf{S} start loc of the newly added genomic segment
- E end loc of the newly added genomic segment

Example:

```
>>> a=annotated_bed(chr="chr13", start=40975747, end=40975770)
    >>> a.Update(40975700,40975800)
    >>> print a.start, a.end
    40975700 40975800
___init__(x=None, **kwargs)
    Initiation example:
    >>> str="chr13 40975747
                                      40975770
                                                               ATTAAG...TGA protein_coding gcnt
    >>> a=annotated_bed(str)
    >>> a=annotated_bed(chr="chr13",start=40975747,end=40975770,strand='+',type="protein_coding"
 _lt___(other)
    Compare two objects self and other when they are not overlapped
        Parameters other - another annotated_bed object
        Returns boolean – "None" if overlapped.
    Example:
    >>> a=annotated_bed(chr="chr13", start=40975747, end=40975770)
    >>> b=annotated_bed(chr="chr13",start=10003212,end=10005400)
    >>> print a>b
    False
 _str__()
    Use print function to output the cluster information (chr, start, end, type, name, subtype, cluster)
    Example:
    >>> str="chr13 40975747
                                      40975770
                                                       +
                                                               ATTAAG...TGA
                                                                                 protein_coding gcnt
    >>> a=annotated_bed(str)
    >>> a.Cluster(3)
    >>> a.Update(40975700,40975800)
    >>> print a
    "chr13 40975700
                             40975800
                                              protein_coding gcnt2 intron 3"
overlap (other)
    Find overlap between regions
        Parameters other - another annotated_bed object
        Returns boolean
```

5.3 "RNAstructure" class

```
class RNAstructure .RNAstructure (exe path=None)
```

Interface class for RNAstructure executable programs.

```
DuplexFold (seq1=None, seq2=None, dna=False)
```

Use "DuplexFold" program to calculate the minimum folding between two input sequences

Parameters

- seq1,seq2 two DNA/RNA sequences as string, or existing fasta file name
- dna boolean input. Specify then DNA parameters are to be used

Returns minimum binding energy, (unit: kCal/Mol)

Example:

```
>>> from RNAstructure import RNAstructure
>>> RNA_prog = RNAstructure(exe_path="/home/yu68/Software/RNAstructure/exe/")
>>> seq1 = "TAGACTGATCAGTAAGTCGGTA"
>>> seq2 = "GACTAGCTTAGGTAGGATAGTCAGTA"
>>> energy=RNA_prog.DuplexFold(seq1, seq2)
>>> print energy
```

Fold (seq=None, ct_name=None, sso_file=None, Num=1)

Use "Fold" program to predict the secondary structure and output dot format.

Parameters

- seq one DNA/RNA sequence as string, or existing fasta file name
- ct_name specify to output a ct file with this name, otherwise store in temp, default: None
- sso_file give a single strand offset file, format see http://rna.urmc.rochester.edu/Text/File Formats.html#Offset
- Num choose Num th predicted structure

Returns dot format of RNA secondary structure and RNA sequence.

Example:

Parameters exe_path – the folder path of the RNAstructure executables

Example:

```
>>> from RNAstructure import RNAstructure
>>> RNA_prog = RNAstructure(exe_path="/home/yu68/Software/RNAstructure/exe/")
scorer(ct_name1, ct_name2)
```

Use 'scorer' pogram to compare a predicted secondary structure to an accepted structure. It calculates two quality metrics, sensitivity and PPV

Parameters

- ct_name1 The name of a CT file containing predicted structure data.
- ct_name2 The name of a CT file containing accepted structure data, can only store one structure.

Returns sensitivity, PPV, number of the best predicted structure.

Example:

```
>>> ct_name1 = "temp_prediction.ct"
>>> ct_name2 = "temp_accept.ct"
>>> from RNAstructure import RNAstructure
>>> RNA_prog = RNAstructure(exe_path="/home/yu68/Software/RNAstructure/exe/")
>>> sensitivity, PPV, Number = RNA_prog.scorer(ct_name1,ct_name2)
```

Note: RNA Hi-C tools benifits a lot from BAM2X, a convenient python interface for most common NGS datatypes. Try BAM2X now!

UPDATES

Version 0.3.2 (2014-05-07):

• change the name into RNA-Hi-C

2014-05-06:

- In "Select_strongInteraction_pp.py" function, now annotations are updated after doing clustering and for strong interaction. The indexing of annotation files may take some time.
- New "RNA_structure_prediction.py" function to refine RNA structure prediction based on empirical offset of free energies for single strand nucleotide.

New features in 0.3.1 (2014-05-02):

- Add "-release" option in "*split_partner.py*" function. Allow a Type3 read-pair considered to be a "Paired" chimeric fragment even linker does not show up.
- Fix bugs in "Select_strongInteraction_pp.py" function when the number of mapped pairs is low and some chromosomes don't have any mapped read in part1 or part2.
- Add bowtie 2 option and Unique-align option in "Stitch-seq Aligner.py" function.
- Different colors for different types of interactions in the *visualization of interactome*.
- New API for folding energies of two RNA molecules, see "RNAstructure".
- Allow permutation-based strategies to calculate the p-value for the overlap between two independent interaction sets in "intersectInteraction.py" function

New features in 0.2.2:

- "Plot_interaction.py" function to plot local RNA-RNA interactions.
- "intersectInteraction.py" function to call overlap between two independent interaction sets.

26 Chapter 6. Updates

CHAPTER

SEVEN

INDICES AND TABLES

- genindex
- modindex
- search

PYTHON MODULE INDEX

а

Annotation, 19

30 Python Module Index

Symbols	split_par
init() (RNAstructure.RNAstructure method), 22 init() (data_structure.annotated_bed method), 21	Stitch-se Subtype(
lt() (data_structure.annotated_bed method), 21 str() (data_structure.annotated_bed method), 21	U Undata()
A	Update()
annotated_bed (class in data_structure), 20 Annotation (module), 19 annotation() (in module Annotation), 19	
С	
Cluster() (data_structure.annotated_bed method), 20	
D	
DuplexFold() (RNAstructure.RNAstructure method), 21	
F	
Fold() (RNAstructure.RNAstructure method), 22	
1	
intersectInteraction.py, 13	
0	
overlap() (data_structure.annotated_bed method), 21 overlap() (in module Annotation), 19	
P	
Plot_Circos.R, 17 Plot_interaction.py, 15	
R	
recoverFragment, 7	
remove_dup_PE.py, 5 RNA_structure_prediction.py, 13	
RNAstructure (class in RNAstructure), 21	
S	
scorer() (RNAstructure.RNAstructure method), 22 Select_strongInteraction_pp.py, 11	

split_library_pairend.py, 6

```
split_partner.py, 8
Stitch-seq_Aligner.py, 10
Subtype() (in module Annotation), 19

Update() (data_structure.annotated_bed method), 20
```