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# ChIA-PET Tool package output illustration

This document explains all the output files, including those used in the visualization report. Figure 1 shows the structure of the DNA constructs generated by ChIA-PET protocol, which is labeled with the terms used in this document. Figure 2 shows the result files along the ChIA-PET data processing steps.

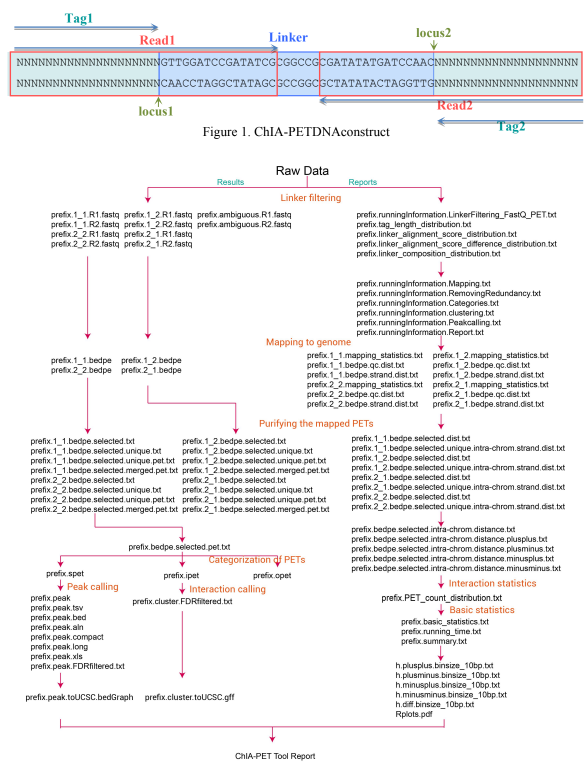


Figure 2. Result files along the ChIA-PET data processing steps

In this file, ChIA-PET data associated with RNA polymerase II (RNAPII) from human breast cancer cell line MCF7 is used for illustration, and "PREFIX" is used as the prefix of the output files. Table 1 contains the file list generated by ChIA-PET Tool, and Table 2 contains the file list used in the visualization report. The files in Table 1 are grouped by the ChIA-PET data processing steps. The formats of the files are elaborated below.

Table 1. Files in the output directory

|                            | Table 1. Files in the o                   |   |
|----------------------------|---|---|
| Processing step            | File name                                 | Description   |
| -                          | PREFIX.1_1.R1.fastq                       | Fastq file of read1, both reads aligned to linker A   |
|                            | PREFIX.1_1.R2.fastq                       | Fastq file of read2, both reads aligned to linker A   |
|                            | PREFIX.1_2.R1.fastq                       | Fastq file of read1, read1 and read2 aligned to linker A and linker B respectively  |
|                            | PREFIX.1_2.R2.fastq                       | Fastq file of read2, read1 and read2 aligned to linker A and linker B respectively  |
| Linker                     | PREFIX.2_1.R1.fastq                       | Fastq file of read1, read1 and read2 aligned to linker B and linker A respectively  |
| Filtering                  | PREFIX.2_1.R2.fastq                       | Fastq file of read2, read1 and read2 aligned to linker B and linker A respectively  |
|                            | PREFIX.2_2.R1.fastq                       | Fastq file of read1, both reads aligned to linker B   |
|                            | PREFIX.2_2.R2.fastq                       | Fastq file of read2, both reads aligned to linker B   |
|                            | PREFIX.ambiguous.R1.fastq                 | Fastq file of read1, with ambiguous linker information  |
|                            | PREFIX.ambiguous.R2.fastq                 | Fastq file of read2, with ambiguous linker information  |
| Mapping to Genome          | PREFIX.1_1.bedpe                          | A bedpe format file of unique mapping PETs converted by BEDtools from mapped BAM file   |
|                            | PREFIX.1_1.bedpe.selected.txt             | File with selected PETs. The mapping scores of both tags are above the mapping quality score threshold, and the optional fields contain the annotation tags: XT:A:U, X0:i:1 and X1:i:0.                       |
|                            | PREFIX.1_1.bedpe.selected.unique.txt      | PETs file after merging PETs that are mapped to the same genomic positions  |
| Purifying the mapped reads | PREFIX.1_1.bedpe.selected.unique.pet.t xt | In this file, the mapping result of a tag is simplified into the format: chromosome, end or start (the genome location that is close to the linker sequence), strand, and sort by chromosome and coordinates. |
|                            | PREFIX.1_1.bedpe.selected.merged.pet.     | PETs file after merging the similarly mapped  |
|                            | txt                                       | PETs into one PET   |
|                            | PREFIX.bedpe.selected.pet.txt             | PETs file after merging both same-linker PET files  |
| PET classification         | PREFIX.ipet                               | File for inter-ligation PETs, including inter-, intra-<br>chromosomal inter-ligation PETs   |

|               | PREFIX.spet                     | File for self-ligation PETs                   |  |  |  |  |
|---------------|---------------------------------|---|--|--|--|--|
|               | PREFIX.opet                     | File for other PETs with short distance       |  |  |  |  |
| Interaction   | PREFIX cluster FDR filtered txt | File with statistically-significant chromatin |  |  |  |  |
| calling       | FREFIX.cluster.fDRIIItered.txt  | interactions                                  |  |  |  |  |
| Daala aallina | PREFIX.peak                     | File with peaks                               |  |  |  |  |
| Peak calling  | PREFIX.peaks.FDRfiltered.txt    | File with statistically-significant peaks     |  |  |  |  |
|               | DDEFIV aslata LICCC had Coords  | Peak file which could be visualized with UCSC |  |  |  |  |
| Visualization | PREFIX.peak.toUCSC.bedGraph     | browser                                       |  |  |  |  |
| visualization | DDEELV algester to LICEC off    | Cluster file which could be visualized        |  |  |  |  |
|               | PREFIX.cluster.toUCSC.gff       | with UCSC browser                             |  |  |  |  |

Table 2. Files used in the visualization report

| Table 2. Files used in the vi                              |   |  |  |  |
|--|---|--|--|--|
| File name  | Description                                     |  |  |  |
| PREFIX.tag_length_distribution.txt                         | Distribution of tag length                      |  |  |  |
| PREFIX.linker alignment score distribution.txt             | Distribution of best alignment scores from the  |  |  |  |
| TREFTA.miker_anginnent_score_distribution.txt              | designed linker sequences to the reads          |  |  |  |
|  | Distribution of score differences between the   |  |  |  |
| PREFIX.linker_alignment_score_difference_distribution.txt  | best-aligned linker and the second-best aligned |  |  |  |
|  | linker  |  |  |  |
| PREFIX.linker_composition_distribution.txt                 | Distribution of linker composition              |  |  |  |
| PREFIX.runningInformation.LinkerFiltering_FastQ_PET.tx     | Running information of Linker filtering         |  |  |  |
| t  | Running information of Emiker intering          |  |  |  |
| PREFIX.runningInformation.Mapping.txt                      | Running information of Mapping to genome        |  |  |  |
| PREFIX.runningInformation.RemovingRedundancy.txt           | Running information of Removing redundancy      |  |  |  |
| PREFIX.runningInformation.Categories.txt                   | Running information of Classification           |  |  |  |
| PREFIX.runningInformation.clustering.txt                   | Running information of Peak calling             |  |  |  |
| PREFIX.runningInformation.Peakcalling.txt                  | Running information of Clustering               |  |  |  |
| PREFIX.runningInformation.Report.txt                       | Running information of Generating reports       |  |  |  |
| PREFIX.1 1.mapping statistics.txt                          | Statistics of mapping mode (Non-mappable,       |  |  |  |
| TREFTA.1_1.mapping_statistics.txt                          | Uniquely-mapped and Others)                     |  |  |  |
| PREFIX.1_1.bedpe.qc.dist.txt                               | Distribution of mapping quality scores          |  |  |  |
| PREFIX.1_1.bedpe.strand.dist.txt                           | Distribution of strands information             |  |  |  |
| PREFIX.1_1.bedpe.selected.dist.txt                         | Distribution of counts of PETs that are mapped  |  |  |  |
| TREE TALL T. Decape. Solected. district                    | to the same genomic position                    |  |  |  |
| PREFIX.1_1.bedpe.selected.unique.intra-chrom.strand.dist.t | Distribution of strands information after       |  |  |  |
| xt   | removing redundancy                             |  |  |  |
| PREFIX.bedpe.selected.intra-chrom.distance.txt             | Span and strands information of                 |  |  |  |
| 1  | intra-chromosomal PETs                          |  |  |  |
| PREFIX.bedpe.selected.intra-chrom.distance.plusplus.txt    | Span of PETs with both tags mapped on plus      |  |  |  |
|  | strand in intra-chromosomal PETs                |  |  |  |

|  | Span of PETs with two tags in                     |  |  |  |  |  |  |
|--|---|--|--|--|--|--|--|
| PREFIX.bedpe.selected.intra-chrom.distance.plusminus.txt | intra-chromosomal PETs mapped on plus and         |  |  |  |  |  |  |
|  | minus strand respectively                         |  |  |  |  |  |  |
|  | Span of PETs with two tags in                     |  |  |  |  |  |  |
| PREFIX.bedpe.selected.intra-chrom.distance.minusplus.txt | intra-chromosomal PETs mapped on minus and        |  |  |  |  |  |  |
|  | plus strand respectively                          |  |  |  |  |  |  |
| DDEELY hadro colored inter-shown distance minuscrimeter  | Span of PETs with both tags in                    |  |  |  |  |  |  |
| PREFIX.bedpe.selected.intra-chrom.distance.minusminus.tx | intra-chromosomal PETs mapped on minus            |  |  |  |  |  |  |
| t  | strand in intra-chromosomal PETs                  |  |  |  |  |  |  |
| PREFIX.PET_count_distribution.txt                        | Statistics file of PET counts' distribution       |  |  |  |  |  |  |
| PREFIX.basic_statistics.txt                              | Basic statistics of PETs number                   |  |  |  |  |  |  |
| PREFIX.running_time.txt                                  | The running time of each section                  |  |  |  |  |  |  |
| Chia DET Tool Donort                                     | A html report containingstatistics of information |  |  |  |  |  |  |
| ChIA-PET_Tool_Report                                     | during data processing with ChIA-PET Tool         |  |  |  |  |  |  |

# Files in the output directory:

(1) PREFIX.1\_1.bedpe (These explanation is from

http://bed tools.read the docs.org/en/latest/content/general-usage.html)

This file is used to illustrate the information of pair-end tag sequence mapped to genome. This file is getting from unique-mapping PETs.

| chrom1 | start1    | end1      | chrom2 | start2    | end2      | name          | score | strand1 | strand2 |
|--------|-----------|-----------|--------|-----------|-----------|---------------|-------|---------|---------|
| chr5   | 175964094 | 175964115 | chr5   | 175964271 | 175964291 | SRR372741.25  | 25    | -       | +       |
| chr1   | 166569355 | 166569376 | chr12  | 56223587  | 56223608  | SRR372741.36  | 37    | +       | +       |
| chr14  | 71969478  | 71969499  | chr19  | 16530595  | 16530616  | SRR372741.94  | 37    | -       | -       |
| chr10  | 108878463 | 108878484 | chr2   | 184995039 | 184995059 | SRR372741.143 | 37    | -       | -       |
| chr10  | 29904926  | 29904946  | chr11  | 32678927  | 32678947  | SRR372741.167 | 37    | -       | +       |
| chr9   | 127735487 | 127735507 | chr9   | 127736244 | 127736264 | SRR372741.176 | 37    | -       | +       |
| chr2   | 191497466 | 191497487 | chr4   | 83532329  | 83532350  | SRR372741.241 | 37    | +       | -       |
| chr17  | 79680058  | 79680079  | chr17  | 79680152  | 79680172  | SRR372741.247 | 37    | -       | +       |
| chr16  | 10275488  | 10275508  | chr6   | 150597540 | 150597561 | SRR372741.251 | 37    | -       | +       |
| chr1   | 152844798 | 152844818 | chr1   | 152880900 | 152880921 | SRR372741.263 | 37    | -       | +       |

Meaning of the columns:

chrom1: The name of the chromosome on which tag1 exists.

• Use "." for unknown.

**start1**: The zero-based starting position of the tag1 on chrom1.

- The first base in a chromosome is numbered 0.
- As with BED format, the start position in each BEDPE feature is therefore interpreted to be 1 greater than the start position listed in the feature. This column is required.
- Use -1 for unknown.

end1: The one-based ending position of tag1 on chrom1.

- The end position in each BEDPE feature is one-based.
- Use -1 for unknown.

**chrom2**: The name of the chromosome on which tag2 exists.

• Use "." for unknown.

**start2**: The zero-based starting position of tag2 on chrom2.

- The first base in a chromosome is 0.
- As with BED format, the start position in each BEDPE feature is therefore interpreted to be 1 greater than the start position listed in the feature. This column is required.
- Use -1 for unknown.

end2: The one-based ending position of tag2 on chrom2.

- The end position in each BEDPE feature is one-based.
- Use -1 for unknown.

name: Definingthe name of the BEDPE feature.

• In this file, name is the reads id gotten from raw fastq format file.

**score**: The "score" field is the mapping quality score from the BAM alignment.

**strand1**: Definingthe strand for the first end of the feature. Either '+' or '-'.

• Use "." for unknown.

strand2: Defines the strand for the second end of the feature. Either '+' or '-'.

• Use "." for unknown.

(2) PREFIX.1\_1.bedpe.selected.txt and PREFIX.1\_1.bedpe.selected.unique.txt PREFIX.1\_1.bedpe.selected.txt is converted from the bedpe format file PREFIX.1\_1.bedpe described above. The changes are at two point: 1) the name of the BEDPE feature is replaced by '.'; and 2) the mapping quality score of each PET is replaced by the mapping cutoff.

PREFIX.1\_1.bedpe.selected.unique.txt is a file generated from PREFIX.1\_1.bedpe.selected.txtby keeping only one PET from those PETs with both tags mapped at the same positions.

| chrom1 | start1    | end1      | chrom2 | start2    | end2      | cutoff | strand1 | strand2 |
|--------|-----------|-----------|--------|-----------|-----------|--------|---------|---------|
| chr5   | 175964094 | 175964115 | chr5   | 175964271 | 175964291 | 20     | -       | +       |
| chr1   | 166569355 | 166569376 | chr12  | 56223587  | 56223608  | 20     | +       | +       |
| chr14  | 71969478  | 71969499  | chr19  | 16530595  | 16530616  | 20     | -       | -       |
| chr10  | 108878463 | 108878484 | chr2   | 184995039 | 184995059 | 20     | -       | -       |
| chr10  | 29904926  | 29904946  | chr11  | 32678927  | 32678947  | 20     | -       | +       |
| chr9   | 127735487 | 127735507 | chr9   | 127736244 | 127736264 | 20     | -       | +       |
| chr2   | 191497466 | 191497487 | chr4   | 83532329  | 83532350  | 20     | +       | -       |
| chr17  | 79680058  | 79680079  | chr17  | 79680152  | 79680172  | 20     | -       | +       |
| chr16  | 10275488  | 10275508  | chr6   | 150597540 | 150597561 | 20     | -       | +       |
| chr1   | 152844798 | 152844818 | chr1   | 152880900 | 152880921 | 20     | -       | +       |

Meaning of the columns:

**chrom1**: The name of the chromosome on which the tag1 exists.And all the PETs that either side chromosome information unknown are removed.

**start1**: The zero-based starting position of tag1on chrom1.

end1: The one-based ending position of tag1on chrom1.

**chrom2**: The name of the chromosome on which the tag2 exists. And all the PETs that either side chromosome information unknown are removed.

**start2**: The zero-based starting position of tag2 on chrom2.

end2: The one-based ending position of tag2 on chrom2.

/: Using '.' to replace the name of the BEDPE feature. It doesn't have exact meaning.

cutoff: Threshold of score value. Using this value to replace the mapping score.

**strand1**: Defines the strand for the first end of the feature.

**strand2**: Defines the strand for the second end of the feature.

# (3) PREFIX.1\_1.bedpe.selected.unique.pet.txt

This file is used to illustrate the PETs after merging those mapped to the same genome position, which are in a high chance caused by PCR amplification. And the mapped location of the tag sequence is replaced with a point, which is the border between tag and linker.

| chrom1 | locus1    | strand1 | chrom2 | locus2    | strand2 |
|--------|-----------|---------|--------|-----------|---------|
| chr10  | 100003964 | -       | chr10  | 100005553 | -       |
| chr10  | 100012097 | +       | chr10  | 100012839 | -       |
| chr10  | 100004117 | -       | chr10  | 100070385 | -       |
| chr10  | 100022370 | -       | chr10  | 100025881 | -       |
| chr10  | 100029170 | -       | chr10  | 100056212 | -       |
| chr10  | 100059995 | -       | chr10  | 100061239 | -       |
| chr10  | 100069332 | -       | chr10  | 100086674 | -       |
| chr10  | 100072260 | -       | chr10  | 100173581 | -       |
| chr10  | 100072560 | -       | chr10  | 100175402 | -       |
| chr10  | 100092053 | -       | chr10  | 100171865 | -       |

Meaning of the columns:

**chrom1**: The name of the chromosome on which tag1 exists.

**locus1**: Locus1 is the border between tag1 and linker. Using end1 as the value of location if tag1 is aligned to plus strand; in contrast, start1 is used when tag1 is aligned to minus strand.

**strand1**: Defines the strand for tag1.

**chrom2**: The name of the chromosome on which tag2 exists.

**locus2**: Locus2 is the border between tag2 and linker. Using end2 as the value of location if tag2 is aligned to plus strand, in contrast, start2 is used when tag2 is aligned to minus strand.

strand2: Defines the strand for tag2.

#### (4) PREFIX.1 1.bedpe.selected.merged.pet.txt

PREFIX.bedpe.selected.pet.txt

PREFIX.ipet

PREFIX.spet

PREFIX.opet

PREFIX.1\_1.bedpe.selected.merged.pet.txtis used to illustrate the location of PETs after removing similar position caused by PCR amplification and ultrasonic disruption.

PREFIX.bedpe.selected.pet.txtcontains same linker PETs after removing redundancy.

PREFIX.ipet is the file with inter-ligation PETs.

PREFIX.spetis the file with self-ligation PETs.

PREFIX.opet is the file containing the other PETs with short distance.

| chrom1 | locus1    | strand1 | chrom2 | locus2    | strand2 | counts | index |
|--------|-----------|---------|--------|-----------|---------|--------|-------|
| chr10  | 103618745 | -       | chr10  | 103618888 | +       | 1.0    | -1    |

| chr10 | 103619140 | - | chr10 | 103662080 | + | 1.0 | -1 |
|-------|-----------|---|-------|-----------|---|-----|----|
| chr10 | 103633680 | - | chr10 | 103634188 | + | 1.0 | -1 |
| chr10 | 103634897 | - | chr10 | 103753306 | + | 1.0 | -1 |
| chr10 | 103635188 | - | chr10 | 103635449 | + | 2.0 | -1 |
| chr10 | 103640011 | - | chr10 | 103641203 | + | 1.0 | -1 |
| chr10 | 103647998 | - | chr10 | 103648487 | + | 1.0 | -1 |
| chr10 | 103657683 | - | chr10 | 103657935 | + | 1.0 | -1 |
| chr10 | 103661110 | - | chr10 | 103661482 | + | 1.0 | -1 |
| chr10 | 103661281 | - | chr10 | 103661712 | + | 1.0 | -1 |

**chrom1**: The name of the chromosome on which tag1 exists.

locus1: The location of tag1 after merging similar PETs.

**strand1**: Defines the strand for tag1.

**chrom2**: The name of the chromosome on which tag2 exists.

locus2: The location of tag2 after merging similar PETs.

strand2: Defines the strand for tag2.

**counts**: Counts of similar position PETs mapped to genome.

index: Indexis reserved for future use.

# (5) PREFIX.cluster.FDRfiltered.txt

This file contains the statistically-significant interaction clusters and related information.

| chrom1   | start1          |     | end1           | chi   | rom2              |                   | start1   |       | end2        | ipe  | t counts  | type    |
|----------|-----------------|-----|----------------|-------|-------------------|-------------------|----------|-------|-------------|------|-----------|---------|
| chr10    | 5454640         |     | 5455156        | chr10 |                   |                   | 5469890  |       | 5470405     |      | 2         | 1       |
| chr10    | 5487515         |     | 5489041        | cł    | hr10              |                   | 5587424  |       | 5589742     |      | 9         | 1       |
| chr10    | 5488183         |     | 5488970        | cł    | hr10              |                   | 5655080  |       | 5655943     |      | 2         | 1       |
| chr10    | 5488356         |     | 5489176        | cł    | hr10              |                   | 5534701  |       | 5535565     |      | 2         | 1       |
| chr10    | 5488670         |     | 5489194        | cł    | hr10              |                   | 5531320  |       | 5531883     |      | 2         | 1       |
| chr10    | 5489401         |     | 5490133        | cł    | hr10              |                   | 5638857  |       | 5639501     |      | 2         | 1       |
| chr10    | 5490050         |     | 5491267        | cł    | hr10              |                   | 5587813  |       | 5589259     |      | 4         | 1       |
| chr10    | 5492324         |     | 5492945        | cł    | hr10              |                   | 5586500  |       | 5587151     |      | 2         | 1       |
| chr10    | 5504728         |     | 5505495        | cł    | hr10              |                   | 5588057  |       | 5588979     |      | 2         | 1       |
| chr10    | 5505491         |     | 5506273        | cł    | hr10              |                   | 5589580  |       | 5590267     |      | 2         | 1       |
| distance | anchor1 tag cou | nts | anchor2 tag co | unts  | p-va              | lue               | p.adjust |       | -log10(p-va | lue) | -log10(p. | adjust) |
| 15249    | 18              |     | 5              | 2.926 |                   | 2-11              | 1.46e-10 |       | 10.53       |      | 9.8       | 4       |
| 100305   | 182             |     | 390            |       | 7.56e-26 3.71e-24 |                   | 3.71e-24 |       | 25.12       |      | 23.4      | 13      |
| 166935   | 159             |     | 14             |       | 2.186             | e-08              | 3.32e-08 |       | 7.66        |      | 7.4       | 8       |
| 46367    | 159             |     | 50             |       | 2.94              | e-07              | 3.40e-07 | 6.53  |             |      | 6.4       | 7       |
| 42669    | 82              |     | 35             | 35    |                   | e-08              | 5.35e-08 |       | 7.42        |      | 7.2       | 7       |
| 149412   | 41              |     | 22             |       | 3.61              | 3.61e-09 7.35e-09 |          | 8.44  |             |      | 8.1       | 3       |
| 97877    | 63              |     | 289            |       | 3.70€             | 70e-13 3.38e-12   |          | 12.43 |             |      | 11.4      | 17      |
| 94191    | 12              |     | 57             |       | 2.016             | -09               | 4.54e-09 | 8.70  |             |      | 8.3       | 4       |
| 83406    | 20              |     | 187            |       | 6.30€             | -08               | 8.40e-08 |       | 7.20        |      | 7.0       | 8       |

| 84041 18 | 92 | 1.22e-08 | 2.03e-08 | 7.91 | 7.69 |
|----------|----|----------|----------|------|------|
|----------|----|----------|----------|------|------|

**chrom1**: The name of the chromosome on which cluster anchor1 exists.

**start1**: The start location of cluster anchor1.

end1: The end location of cluster anchor1.

**chrom2**: The name of the chromosome on which cluster anchor2exists.

start2: The start location ofcluster anchor2.

end2: The end location of cluster anchor2.

ipet counts: Counts of PETs between two anchors of an interaction cluster.

**type**: Interactive type. 1 represents intra-chromosomal interaction, 0 for inter-chromosomal interaction.

**distance**: Distance between both two anchors of intra-chromosomal PET. And if the two anchors are located on different chromosomes, the value of distance is 2,147,483,647 (as the largest integer with 32bit, 2^31-1).

anchor1 tag counts: Counts of tag loci fall into cluster anchor1.

anchor2 tag counts: Counts of tag loci fall into cluster anchor2.

**p-value**: This value represents the statistical significance of the interaction. It is calculated by hyper-geometric distribution.

**p.adjust**: P.adjust means p-value adjustment with Benjamini-Hockberg method (1995)

-log10(p-value): The negative logarithm of p-value.

-log10(p.adjust): The negative logarithm of adjusted p-value.

### (6) PREFIX.peak

This file is used to illustrate the primary peak information obtained from self-ligation PETs.

| chrom | start  | end    | summit start | summit end | peak coverage |
|-------|--------|--------|--------------|------------|---------------|
| chr1  | 840639 | 840651 | 840639       | 840651     | 14            |
| chr1  | 841447 | 841511 | 841447       | 841511     | 7             |
| chr1  | 843472 | 843495 | 843472       | 843495     | 5             |
| chr1  | 846703 | 846732 | 846703       | 846732     | 16            |
| chr1  | 847869 | 847883 | 847869       | 847883     | 5             |
| chr1  | 849284 | 849285 | 849284       | 849285     | 20            |
| chr1  | 852786 | 852788 | 852786       | 852788     | 46            |
| chr1  | 853677 | 853746 | 853677       | 853679     | 58            |
| chr1  | 854750 | 854812 | 854750       | 854763     | 48            |
| chr1  | 858117 | 858423 | 858117       | 858149     | 9             |

Meaning of the columns:

**chrom**: The name of the chromosome on which peak exists.

**start**: The start position of peak. This column is used in broad peak.

end: The end position of peak. This column is used in broad peak.

**summit start**: The start position of peak summit. This column is used in narrow peak.

summit end: The end position of peak summit.

**peak coverage**: The coverage of the overlapped peaks.

### (7) PREFIX.peak.FDRfiltered.txt

This file is used to illustrate the high potential peaks which passed the threshold of adjusted p-value.

| chrom | start  | end    | peak coverage | p-value  | p.adjust |
|-------|--------|--------|---------------|----------|----------|
| chr1  | 852786 | 852788 | 46            | 1.73e-07 | 1.8e-06  |
| chr1  | 853677 | 853679 | 58            | 1.44e-13 | 2.31e-12 |
| chr1  | 854750 | 854763 | 48            | 1.2e-07  | 1.26e-06 |
| chr1  | 860735 | 860737 | 37            | 9.32e-08 | 1.03e-06 |
| chr1  | 877184 | 877186 | 23            | 1.25e-09 | 1.6e-08  |
| chr1  | 902065 | 902068 | 53            | 1.08e-40 | 4.55e-39 |
| chr1  | 935164 | 935172 | 61            | 4.6e-30  | 1.36e-28 |
| chr1  | 936330 | 936335 | 46            | 8.52e-16 | 1.51e-14 |
| chr1  | 955915 | 955938 | 31            | 2.29e-17 | 4.29e-16 |
| chr1  | 994919 | 994929 | 25            | 5.89e-09 | 7.26e-08 |

Meaning of the columns:

**chrom**: The name of the chromosome on which the peak exists

**start**: The start position of peak **end**: The end position of peak

peak coverage: The coverage of the overlapped peaks.

**p-value**: This value represents the reliability of a real peak. It is calculated by Poisson distribution.

**p.adjust**: P.adjust means p-value adjustment with Benjamini-Hockberg method.

# Files used in the visualization report:

(1) PREFIX.linker alignment score distribution.txt

This file is used to illustrate the distribution of best alignment scores from the designed linker sequences to the reads.

| alignment score | counts of read1 | counts of read2 |
|-----------------|-----------------|-----------------|
| 0               | 0               | 99987           |
| 1               | 2815            | 9249            |
| 2               | 27337           | 46442           |
| 3               | 24444           | 46457           |
| 4               | 67866           | 81805           |
| 5               | 87995           | 95125           |
| 6               | 103164          | 105683          |
| 7               | 100254          | 123026          |
| 8               | 98584           | 125184          |
| 9               | 84207           | 120977          |
| 10              | 124532          | 162743          |
| 11              | 186583          | 256884          |
| 12              | 135917          | 204520          |
| 13              | 808328          | 1041042         |
| 14              | 1521765         | 1875530         |

| 15 | 33572085 | 32416084 |
|----|----------|----------|
| 16 | 43485247 | 43744265 |
| 17 | 967484   | 909505   |
| 18 | 100999   | 65062    |
| 19 | 57964    | 28000    |

**alignment score**: Local alignment score of reads with the best matched linker by dynamic programming.

**counts of read1**: Counts of read1 that are aligned to best matched linker with certain local alignment score.

**counts of read2**: Counts of read2 that are aligned to best matched linker with certain local alignment score.

### (2) PREFIX.linker\_alignment\_score\_difference\_distribution.txt

This file is used to illustrate the score difference distribution between the best-aligned linker and the second-best aligned linker. It shows how the barcode distinguishes different linkers.

| score difference | counts of read1 | counts of read2 |
|------------------|-----------------|-----------------|
| 0                | 237941          | 381176          |
| 1                | 247817          | 243882          |
| 2                | 282467          | 347700          |
| 3                | 877610          | 787925          |
| 4                | 406169          | 562655          |
| 5                | 79505566        | 79234232        |

Meaning of the columns:

**score difference**: Reads mapped to different linkers will get a set of local alignment scores. This column is the score difference between the best aligned and second-best aligned one.

counts of read1: Counts of read1 with certain score difference.

counts of read2: Counts of read2 with certain score difference.

#### (3) PREFIX.tag length distribution.txt

This file is used to illustrate the tag length distribution. Tags refer to the DNA fragments that are digested by enzyme *Mme*Ifar away from linkers. Such fragments are gotten by removing linker from sequencing reads. The selected tags will be kept for further processing.

| tag length | counts of read1 | counts of read2 |
|------------|-----------------|-----------------|
| 0          | 0               | 99987           |
| 1          | 111             | 188             |
| 2          | 1974            | 1805            |
| 3          | 3808            | 4744            |
| 4          | 939             | 2204            |
| 5          | 1480            | 2448            |
| 6          | 1923            | 2922            |
| 7          | 2533            | 3813            |
| 8          | 7913            | 8360            |

| 9  | 103553   | 102500   |
|----|----------|----------|
| 10 | 41470    | 45188    |
| 11 | 34000    | 37377    |
| 12 | 22210    | 25660    |
| 13 | 25898    | 28738    |
| 14 | 30130    | 31580    |
| 15 | 27110    | 27857    |
| 16 | 44094    | 39730    |
| 17 | 73781    | 50902    |
| 18 | 88013    | 76375    |
| 19 | 1010611  | 1006545  |
| 20 | 45191493 | 45902278 |
| 21 | 34479577 | 33648812 |
| 22 | 222849   | 252044   |
| 23 | 37024    | 42881    |
| 24 | 15408    | 17240    |
| 25 | 12873    | 13127    |
| 26 | 14795    | 15981    |
| 27 | 14197    | 15407    |
| 28 | 17627    | 18741    |
| 29 | 13407    | 14184    |
| 30 | 8027     | 8469     |
| 31 | 5226     | 5380     |
| 32 | 2649     | 2447     |
| 33 | 778      | 1045     |
| 34 | 89       | 523      |
| 35 | 0        | 88       |
|    |          |          |

tag length: The length of tag sequence.

**counts of read1**: Counts of tag1 with certain tag length. **counts of read2**: Counts of tag2 with certain tag length.

# (4) PREFIX.linker\_composition\_distribution.txt

This file is used to illustrate the proportion of each linker combination aligned to the reads.

|            | A_A      | A_B     | B_A     | B_B      | Ambiguous | Total    |
|------------|----------|---------|---------|----------|-----------|----------|
| Numbers    | 33822895 | 4744854 | 4748192 | 32998692 | 5242937   | 81557570 |
| Percentage | 41.47%   | 5.82%   | 5.82%   | 40.46%   | 6.43%     | 100%     |

Meaning of the columns:

**A** A: "A A" means the category of PETs that both tags optimally aligned to linker A.

**A\_B**: "A\_B" means the category of PETs that tag1 and tag2 optimally aligned to linker A and linker B respectively.

**B\_A**: "B\_A" means the category of PETs that tag1 and tag2 optimally aligned to linker B and

linker A respectively.

**B\_B**: "B\_B" means the category of PETs that both tags optimally aligned to linker B.

Ambiguous: "Ambiguous" means the category of PETs that not satisfy one of the criteria below:

- 1) the best linker alignment score is equal to or larger than the **minimum\_linker\_alignment\_score**,
- 2) the difference between the best and the second best linker alignment scoresis equal to or larger than the **minSecondBestScoreDiff**, 3) tag length should conform to the specified range, and 4) the barcodes in the PETs must completely match the designed barcodes in the linkers.

**Total**: Counts of the whole PETs.

**Attention**: A\_A and B\_B should constitute the majority of the PETsin a good ChIA-PET library as the sample shows.

# (5) PREFIX.1\_1.mapping\_statistics.txt

This file contains the statistics of the pair-end reads mapping results. Each end has three possible mapping results: Non-mappable, Uniquely-mapped and Others, and there are nine combinations from the two ends.

| read1 read2     | Non-mappable | Uniquely-mapped | Others   |
|-----------------|--------------|-----------------|----------|
| Non-mappable    | 46           | 4146            | 8459     |
| Uniquely-mapped | 6254         | 4757858         | 7807698  |
| Others          | 11679        | 7163264         | 14063491 |

The meanings of the different categories are as follows:

**Non-mappable**: Mappingpaired-end reads to a reference genome using BWA can obtain a SAM format file. There are two conditions for Non-mappable reads. 1) If the mapping result of a tag in the SAM file doesn't have optional fields (less than 12 columns), it is classified into non-mappabletag. 2) If the mapping result of a tag in the SAM file has optional fields and there is a label XT:A:N, this tag is classified into "non-mappable" category.

**Uniquely-mapped**: If the mapping quality score MAPQ (5th column) of a tag is more than 20 and the optional fields contain the following labels: XT:A:U(12th column), X0:i:1 (16th column), X1:i:0 (17th column), this tagis classified into "uniquely-mapped" category.

**Others**: If the mapping result of a tag doesn't meet the criteria above, it is classified into "Others" category, which includes the tags with low mapping scores and multiply-mapped tags.

**Attention**: In a SAM format file, the mapping results of the PETs appear in pairs. So the first line in the pairs is read1, the second line in pairs is read2.

#### (6) PREFIX.1 1.bedpe.qc.dist.txt

This file contains the distribution of mapping scores from unique-mapping PETs.

| counts  | score |
|---------|-------|
| 15234   | 25    |
| 4669548 | 37    |
| 2       | 50    |
| 73074   | 60    |

Meaning of the columns:

**counts**: Counts of PETs with the corresponding mapping scores.

**score**: Score value(8th column) from the bedpe format file above.

(7) PREFIX.1\_1.bedpe.strand.dist.txt

This file contains the distribution of strand combinations from the uniquely-mapped PETs. There are four types of strand combinations: 1) both tags are mapped to plus strands (++), 2) tag1 is mapped to plus strand and tag2 is mapped to minus strand (+-), 3) tag1 is mapped to minus strand and tag2 is mapped to plus strand (-+), 4) both tags are mapped to minus strands (--).

| counts  | strand1 | strand2 |
|---------|---------|---------|
| 841994  | -       | -       |
| 2227108 | -       | +       |
| 845768  | +       | -       |
| 842988  | +       | +       |

Meaning of the columns:

counts: Counts of certain strand combination.

strand1: Strand information of tag1 (9th column) from the bedpe format file above.

strand2: Strand information of tag2 (10th column) from the bedpe format file above.

### (8) PREFIX.1 1.bedpe.selected.dist.txt

This file is about the summary statistics of the redundancies caused by PCR amplification. The redundancy means the PETs that are mapped exactly to the same genomic position, which are from PCR amplification with high chance.

| counts  | repetitions |
|---------|-------------|
| 3358894 | 1           |
| 566169  | 2           |
| 75928   | 3           |
| 8548    | 4           |
| 841     | 5           |
| 66      | 6           |
| 7       | 7           |

Meaning of the columns:

**counts**: means the number of frequencies appeared in the data.

repetitions: meansthe frequency of PETs mapped to the same genomic position.

### (9) PREFIX.1\_1.bedpe.selected.unique.intra-chrom.strand.dist.txt

This file is used to illustrate strands combination of intra-chromosomal inter-ligation PETs.

| counts  | strand1 | strand2 |
|---------|---------|---------|
| 169000  | -       | -       |
| 1338210 | -       | +       |
| 172396  | +       | -       |
| 169847  | +       | +       |

Meaning of the columns:

**counts**: Counts of PETs with certain strand combinations. The selected are intra-chromosomal inter-ligation PETs.

**strand1**: Strand of tag1 from the intra-chromosomal inter-ligation PETs after removing redundancy.

strand2: Strand of tag2 from the intra-chromosomal inter-ligation PETs after removing

redundancy.

#### (10) PREFIX.bedpe.selected.intra-chrom.distance.txt

This file contains the distance between two tags of the PETs and strand information. Based on the ChIA-PET design and Illumina sequencing principle, there should be more PETs with strand combination -+ at short distance than other combination.

| span  | strand1 | strand2 |
|-------|---------|---------|
| 1589  | -       | -       |
| 1036  | -       | -       |
| 742   | +       | -       |
| 66268 | -       | -       |
| 2505  | -       | -       |
| 3511  | -       | -       |
| 27042 | -       | -       |
| 27157 | -       | -       |
| 1244  | -       | -       |
| 17342 | -       | -       |

Meaning of the columns:

span: Span between the two tags of individual intra-chromosomal PETs.

**strand1**: Strand of tag1 in an intra-chromosomal PETs.

strand2: Strand of tag2 in an intra-chromosomal PETs.

(11) PREFIX.bedpe.selected.intra-chrom.distance.plusplus.txt

PREFIX.bedpe.selected.intra-chrom.distance.plusminus.txt

PREFIX.bedpe.selected.intra-chrom.distance.minusplus.txt

PREFIX.bedpe.selected.intra-chrom.distance.minusminus.txt

These four files are separated from PREFIX.bedpe.selected.intra-chrom.distance.txtaccording to strand combination.

| span     |
|----------|
| 90590128 |
| 672      |
| 527      |
| 62489    |
| 3631     |
| 99731215 |
| 108495   |
| 1440     |
| 52       |
| 1014     |

Meaning of the columns:

**span**: Span between the two tags of individual intra-chromosomal PETs in certain strand combination.