# Genome-analysis-tools manual version 1.0.0

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#### 1 Overview

These tools are designed for analyzing the peaks and/or read distributions derived from next-generation sequencing such as ChIP-seq, MNase-seq or RNA-seq. For

peaks, BED format as well as any tab-delimited table with chromosome name, start position and end position are available. For reads, compressed WIG format (.wig.gz, either one file for whole chromosomes or each file for each chromosome is OK) and bedGraph (.bedgraph) are supported. To make comparison between samples possible, these reads should be normalized by using softwares such as DROMPA or BEDTools. Each tool and option are explained in detail with sample files in the following section.

## 2 Requirement

Genome-analysis-tools are written in ANSI C and can be executed on Linux OS. Genome-analysis-tools requires the following libraries:

- GCC compiler (<u>http://gcc.gnu.org/</u>)
- GNU C Library(glibc) (<a href="http://www.gnu.org/software/libc/">http://www.gnu.org/software/libc/</a>)
- zlib (<u>http://www.zlib.net/</u>)
- (optional) R (<a href="http://www.r-project.org/">http://www.r-project.org/</a>)

### 3 Installation

Install through git:

- \$ git clone git@github.com:KojiMasuda/genome-analysis-tools.git
- \$ cd genome-analysis-tools
- \$ make

Install through source file:

- \$ wget https://github.com/KojiMasuda/genome-analysis-tools/archive/master.zip
- \$ unzip master.zip
- \$ cd genome-analysis-tools-master
- \$ make

If you get an installation error, make sure that all required libraries are installed.

Add the directory you installed the tools to your PATH environment variable. If you installed it in "\$HOME/genome-analysis-tools" directory, add the following line to your Bash startup file such as \$HOME/.bashrc;

\$ export PATH=\$PATH:HOME/genome-analysis-tools

## 4 Genome-analysis-tools

In general, <command> without any argument or <command> -h/--help shows command usage. Also, <command> -v shows version information. In most tools, the first line of input file is considered as header if --header option (default:off) is specified. To support as many format as possible, you can specify the number of column. For example, if you have tab-delimited file of which start position is on 5th column, set --col start 4 (note that column number option is zero-based).

#### 4.1. ga overlap

ga\_overlap reports the overlaps of two peaks/summits.

Among output files;

```
peak1_over_peak2.txt is peak1 which overlap peak2,
peak1_nonover_peak2.txt is peak1 which doesn't overlap peak2,
peak1_vs_peak2.txt is peak1 with over/nonover flags,
peak1_vs_peak2_summary.txt is overlapping summary.
```

If you need overlapping of peak1 to peak2 with header:

```
$ ga_overlap -1 sample/peak1.txt -2 sample/peak2.txt --header
```

If the input files don't have headers, just remove --header option.

In addition, if you need *the number* of overlapping peak2, specify --count option:

```
$ ga_overlap -1 sample/peak1.txt -2 sample/peak2.txt --header --count
```

You may want information of overlapping peak2, specify --preserve2 option:

```
$ ga overlap -1 sample/peak1.txt -2 sample/peak2.txt --header --preserve2
```

Or you may want to compare *the summit* of peak1, specify the start and the end column to the summit (summit is on 4th column in this case):

```
$ ga_overlap -1 sample/peak1.txt -2 sample/peak2.txt --header --col_start1 3 --col_end1 3
```

Note that the column number is zero-based.

In some case, you may want to use the fixed length as a peak width. In that case, specify --hw <int> option as a half-width (total width is 2xhw):

```
$ ga_overlap -1 sample/peak1.txt -2 sample/peak2.txt --header --col_start1 3 --col_end1 3 --hw 250 ##total width is 500 bp
```

Type ga overlap -h for the detail explanation of options.

## 4.2. ga\_reads\_summit

**ga\_reads\_summit** reports the average distribution of reads around summits or specific positions such as transcription start sites (TSS) or transcription end sites (TES).

```
Output file is made in the directory of --sig file. In the output file;

1st column <relative_pos>: distance from peak summit

2nd column <smt_mean>: average read values around summit

3rd column <CI95.00percent_U>: Upper 95% confidence interval

4th column <CI95.00percent_L>: Lower 95% confidence interval

5th column <smtNb>: summit number

6th column <Centered>: centered peak name

7th column <Signal>: signal(read) name
```

If you have peak sets and want to calculate the average ChIP read distribution;

- \$ ga\_reads\_summit --smt sample/peak1.txt --sig sample/chip1/chip1 --sigfmt sepwiggz --col start 3 --col end 3 --header
- \$ Rscript sample/R/script1.r

When you specify --sigfmt sepwiggz (separated WIG.GZ file for each chromosome), set file name before "\_chr\*.wig.gz" as read file. In this case it's "sample/chip1/chip1". In the above example, set --col\_start 3 and --col\_end 3 to calculate the read distribution around summit positions, which is stored on 4th column in peak1.txt file. The second command creates png file of the output (Figure 1)

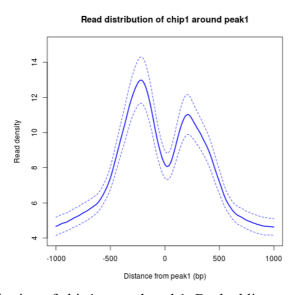


Figure 1. Read distribution of chip1 around peak1. Dashed lines are upper and lower 95 % confidence intervals.

If you have input file as a denominator, signals from --sig file are divided by signals from --sig d file by specifying --sig d option;

- \$ ga\_reads\_summit --smt sample/peak1.txt --sig sample/chip1/chip1 --sigfmt sepwiggz --col start 3 --col end 3 --header --sig d sample/input1/input1
- \$ Rscript sample/R/script2.r

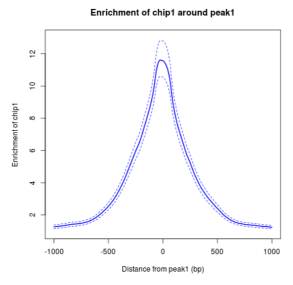


Figure 2. Distribution of chip1 enrichment around peak1. Dashed lines are upper and lower 95 % confidence intervals.

The second command creates png file of the output file (Figure 2).

In some case, you may want to know the read distribution throughout the genome. If you specify --rand <int> option, the positions from 1 to the genome length is randomly picked up (the number of the position is same as the peak numbers), then read density is calculated around the picked-up positions. This process is iterated for <int> times then the mean values are reported;

\$ ga\_reads\_summit --smt sample/peak1.txt --sig sample/chip1/chip1 --sigfmt sepwiggz --col\_start 3 --col\_end 3 --header --sig\_d sample/input1/input1 --rand 1000 --gt sample/genome table.txt

## \$ Rscript sample/R/script3.r

Don't forget setting genome table by --gt option for random simulation because the tool would know the genome length from this file. The second command creates png file of the output file (Figure 3).

#### Enrichment of chip1 around peak1

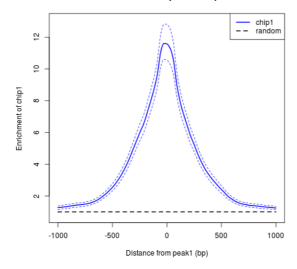


Figure 3. Distribution of chip1 enrichment (blue) around peak1 with enrichment from randomly picked up positions throughout genome (dashed black line). Dashed blue lines are upper and lower 95 % confidence intervals.

If you want to know the read distribution around TSS, specify proper start and end positions by --col\_start and --col\_end options, respectively. Generally speaking, each gene has its strandness. Thus let the tool know the column for the strandness of annotation file by --col\_strand option;

\$ ga\_reads\_summit --smt sample/anno.txt --sig sample/nuc.wig.gz --sigfmt onewiggz --col chr 2 --col start 4 --col end 5 --col strand 3

\$ Rscript sample/R/script4.r

Note that "onewiggz" is specified for --sigfmt argument because "sample/nuc.wig.gz" is WIG.GZ file which contains data for whole chromosome. When you specify --col\_strand, the tool checks whether the gene has '+' or '-'. The second command creates png file of the output file (Figure 4).

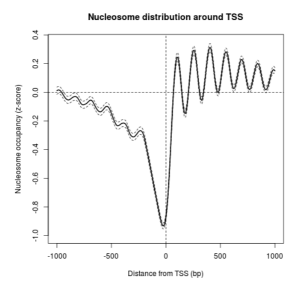


Figure 4. Nucleosome occupancy around TSS. Dashed lines are upper and lower 95 % confidence intervals.

Set appropriate half-window size (--hw), step size (--step) and window size for read calculation (--win) for your sample (default is suitable for yeast sample). Type ga\_reads\_summit -h for the detail explanation of options.

### 4.3. ga reads summit all

**ga\_reads\_summit\_all** reports the read distributions around ALL summits/peaks. The usage is quite similar to **ga\_reads\_summit** tool (see section 4.2).

If you have mammalian ChIP reads, specifying only --sig argument is recommended because of sparse input reads;

- \$ ga\_reads\_summit\_all --smt sample/anno\_mm10.bed --sig sample/chip2/chip2 --sigfmt sepwiggz --col strand 5 --hw 5000 --step 100 --win 500
- \$ Rscript sample/R/script5.r

Note that --col\_strand argument is specified for strandness of annotation genes. Half-window size (--hw), step size (--step) and window size for read calculation (--win) are set for this sample. The second command creates png file of the ouput file (Figure 5).

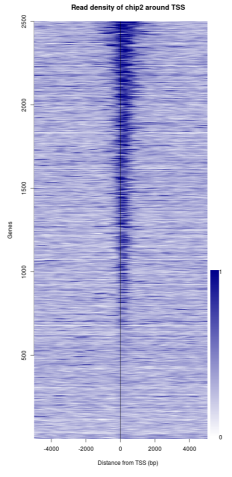


Figure 5. Heat map of chip2 reads around TSS.

If you have yeast ChIP reads, you can specify --sig\_d argument as an input as well as --sig argument as a ChIP;

- \$ ga\_reads\_summit\_all --smt sample/peak1.txt --sig sample/chip1/chip1 --sigfmt sepwiggz --col\_start 3 --col\_end 3 --header --sig\_d sample/input1/input1 --win 300
- \$ Rscript sample/R/script6.r

The second command creates png file of the output file (Figure 6).

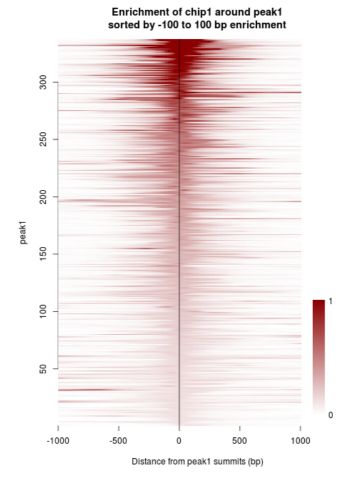


Figure 6. Enrichment of chip1 around peak1.

## 4.4. ga calc dist

**ga\_calc\_dist** reports the distance of two peaks(mode:two) or inter-summit distance(mode:isd). If you have one peak set and want the inter-summit distance;

\$ ga\_calc\_dist -1 sample/peak1.txt -mode isd --header

In some case, you may want to know the distance between two peak sets. When you set "two" mode, **ga\_calc\_dist** calculate the distance from each peak1 to *the closest peak2*;

\$ ga\_calc\_dist -1 sample/peak1.txt -2 sample/peak2.txt -mode two --header

## 4.5. ga\_reads\_region

**ga\_reads\_region** reports the amount of reads inside regions.

You may be interested in read amounts inside gene, up-stream of TSS, down-stream of TES, or around summits. With specific "region" or "smt" modes, you can calculate the read amount inside regions you're interested in. The basic usages are;

<sup>&</sup>quot;isd" is set as -mode argument for inter-summit distance calculation.

\$ ga\_reads\_region [options] --smt <file\_summit> --sig <file\_signal> --sigfmt <sig format:bedgraph | sepwiggz | onewiggz> --mode smt --col\_smt <column of summit>

or

\$ ga\_reads\_region [options] --smt <file\_summit> --sig <file\_signal> --sigfmt <sig format:bedgraph | sepwiggz | onewiggz> --mode <region mode: region | up-tss | tss-dw | up-tss-dw | up-tes | tes-dw | up-tes-dw>

Set --smt, --sig, --sigfmt arguments as explained in section 4.2. You can set suitable --mode argument for your purpose. See scheme in Figure 7 for detail. By specifying --col strand option, the tool consider strandness of region such as gene.

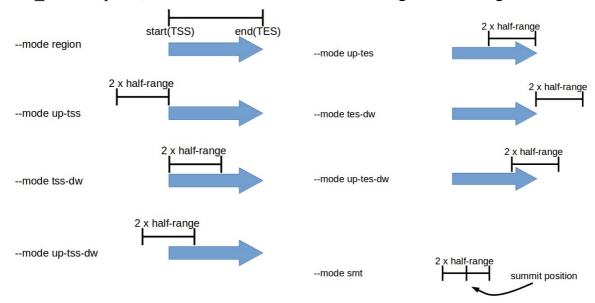


Figure 7: The graphical explanation of mode options. When "region" is specified as --mode argument, start and end position is used and half-range (--hw) is ignored. On the other hand, fixed region length is specified by half-range as --hw argument when other mode is specified.

Note that the amount of reads inside regions is normalized by region length.

- 4.6. ga\_nuc\_region
- 4.7. ga\_nuc\_summit
- 4.8. ga\_deltaG
- 4.9. ga\_RPKM

#### 5 References