# SPORTS1.0

Small non-coding RNA annotation Pipeline Optimized for rRNA- and tRNA-Derived Small RNAs

Requirements

Installation

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Copyright and licensing information

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Disclaimer

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# Requirements

Linux system, enough disk space and Ram depending on the size of RNA deep sequencing data. (Tested system: ubuntu 12.04 LTS, ubuntu 16.04 LTS)

# Installation

1. Download SPORTS1.0 pipeline package.wget

```
https://github.com/junchaoshi/sports1.0/archive/master.zip
```

- 2. Download necessary software, packages and reference databases as listed below:
  - 1. Perl 5 (https://www.perl.org) (Tested version: v5.14.2, v5.22.1); Perl 5 might be already installed in the linux system.
  - 2. Bowtie [1] (http://bowtie-bio.sourceforge.net/index.shtml) (Tested version: 1.1.2, 1.2.1.1)
  - 3. SRA Toolkit (https://ncbi.github.io/sra-tools/) (Tested version: 2.8.2)
  - 4. cutadapt [2] (http://cutadapt.readthedocs.io/en/stable/index.html) (Tested version: 1.11)
  - 5. R (https://www.r-project.org/) (Tested version: 3.2.3, 3.2.5)
  - 6. Reference database (See lists and download link of all pre-compiled species' databases in appendix)
- 3. Installation tutorial for software and packages.
  - 1. Install SPORTS1.0
    - 1. Unpack SPORTS1.0 package.unzip sports1.0-master.zip
    - 2. Attach the SPORTS directory to your PATH:echo 'export
       PATH=\$PATH:your\_path\_to\_sports1.0-master/source' >> ~/.bashrcchmod 755
       your path to sports1.0-master/source/sports.pl
  - 2. Install Bowtie
    - 1. Unpack bowtie-1.x.x-linux-x86\_64.zip.unzip bowtie-1.x.x-linux-x86 64.zip
    - 2. Attach the bowtie directory to your PATH:echo 'export

```
PATH=$PATH:your path to bowtie' >> ~/.bashrc
```

3. If you are administrator user, type the following command and password to easily install bowtie:

sudo apt-get install bowtie

- 4. Install SRA Toolkit
  - 1. Unpack SRA toolkit files.
  - 2. Attach the SRA Toolkit executable path to your PATH: echo 'export PATH=\$PATH:your path to sra-toolkit/bin' >> ~/.bashrc
- 5. Install cutadapt
  - 1. Use pip on the command line to install latest version of cutadapt: pip install --user --upgrade cutadapt
  - 2. Attach the cutadapt directory to your PATH: echo 'export PATH=\$PATH:\$HOME/.local/bin' >> ~/.bashrc
- 6. Install R and R package
  - 1. Unpack R-x.y.z.tar.gz with: tar -xf R-x.y.z.tar.gz
  - 2. Enter into the R-x.y.z directory: cd R-x.y.z
  - 3. Type following command in terminal: ./configure

```
make check
make install
```

4. Install R packages by typing following command in terminal: R

```
install.packages('ggplot2', dependencies=TRUE, repos='http://cran.rstudio.com/')
install.packages('data.table', dependencies=TRUE,
repos='http://cran.rstudio.com/')
install.packages('stringr', dependencies=TRUE, repos='http://cran.rstudio.com/')
q()
```

- 4. Start a new shell session to apply changes to environment variables: source ~/.bashrc
- 5. Test if everything is installed properly: perl -v

```
sports.pl -h
bowtie
fastq-dump
cutadapt -h
R --version
If you get any error messages you should install the software once again.
```

# Script description

#### sports.pl

- 1. Input query format:
  - 1. .sra files.
  - 2. .fastq/.fq, .fasta/.fa files of deep sequencing reads.
- 2. Attention: compressed files need to be unpacked before input!
- 3. Options: --Input: -i <file> Input could be:

```
a .sra, .fastq/.fq or .fasta/.fa file;
a directory (will run all qualified files in the directory recursively);
a text document with absolute path information for each file/folder (when processing multiple data, input each file/folder path per line)
--Output: -o <str> output address of annotation results (default: input address)
-k keep all the intermediate files generated during the running progress
--Alignment: -1 <int> the minimal length of the output sequences (default = 15)
-L <int> the maximal length of the output sequences (default = 45)
-M <int> the total number of mismatches in the entire alignment (default = 0)
-a Remove 5' / 3' adapters
-x <str> (if -a applied) 5' adapter sequence. Default =
"GTTCAGAGTTCTACAGTCCGACGATC"
-y <str> (if -a applied) 3' adapter sequence. Default = "TGGAATTCTCGGGTGCCAAGG"
--Others: -v print version information
```

- -h print this usage message
- 4. Example
  - Example use 1:
- 5. The user wants to map a single fasta file against rat reference genome to get the mapping genome annotation only. (No output figures) Type following command in terminal: sports.pl -i reads.fa -g /foo/bar/Rattus\_norvegicus/UCSC/rn6/Sequence/BowtieIndex/genome

   Example use 2:
- 6. The user wants to map several already trimed human sequencing files to human reference genome, miRNA database, tRNA database, rRNA database and piRNA database by using 4 CPU threads, then to output the result to the address: '/foo/bar/output/'. Write all the fastq files' addresses into a text document, e.g.: seq\_address.txt

- Example use 3:
- 7. The user wants to map several untrimmed mouse sequencing files downloaded from NCBI or somewhere else to mouse reference genome, miRNA database, tRNA database, rRNA database, piRNA database, ensembl noncoding RNA database and Rfam database by using 4 CPU threads, then to output the result to the address: '/foo/bar/output/' and keep all the intermediate files generated during the running progress. Put all the sequencing files into a folder, e.g.:

```
Type following command in terminal: sports.pl -i /foo/bar/download_seq/ -p 4 -a -x GTTCAGAGTCCGACGATC -y TGGAATTCTCGGGTGCCAAGG -g /foo/bar/Mus_musculus/UCSC/mm10/Sequence/BowtieIndex/Genome -m /foo/bar/Mus_musculus/miRBase_21/miRbase_21-mmu -r /foo/bar/Mus_musculus/rRNA_db/mouse_rRNA -t /foo/bar/Mus_musculus/GtRNAdb/mm10-tRNAs -w /foo/bar/Mus_musculus/piRBase/piR_mouse -e /foo/bar/Mus_musculus/Ensembl/Mus_musculus.GRCm38.ncrna -f /foo/bar/Mus_musculus/Rfam_12.3/Rfam-12.3-mouse -o /foo/bar/output/ -k
```

8. Example output file structure for 1 query file input (e.g. SeqFile):

```
Output folder structure

-1_SeqFile
-1_SeqFile
-1_SeqFile_fa (if -k applied)
---unique seqs with reads number
---unique seqs with reads number
---seqs that cannot pass
adapter removing process
---seqs that are lower than
required minimal length threshold
```

```
SeqFile too long reads.fa ---seqs that are higher than
required maximal length threshold
             -SeqFile match_genome.fa
                                             ---seqs that can match to
reference genome
            SeqFile unmatch genome.fa ---seqs that cannot match to
reference genome
             SeqFile_match_<X>_match_genome.fa ---seqs that can match to
both <X> database and reference genome
             SeqFile match <X> unmatch genome.fa ---seqs that can match to
<X> database but not reference genome
             SeqFile unmatch <X> match genome.fa ---seqs that cannot match
to <X> database but can match to reference genome
             LSeqFile_unmatch_<X>_unmatch_genome.fa ---seqs that match to
<X> rfam database nor reference genome
          -SeqFile processed (if -k applied)
             ---seqs that match to reference
genome in BOWTIE format
             SegFile output <X> match genome ---segs that match to both
miRNA database and reference genome in BOWTIE format
             LSeqFile output <X>_unmatch_genome ---seqs that match to miRNA
database but not reference genome in BOWTIE format
          LSeqFile result
             SeqFile output.txt ---6 column table file including
annotation information for every unique sequence
                                           ---3 column table file including
             -SeqFile summary.txt
reads number of each major- (e.g. rRNA) and sub- (e.g. 5S rRNA) classes
             -SeqFile length distribution.txt
including reads number of each length distribution of each major class
             SeqFile mismatch summary.txt ---11 column table file
             SeqFile sncRNA distribution.pdf ---figure of length
distribution of miRNA, rsRNA, tsRNA, piRNA and other RNAs, if sequence matches
existed
             SeqFile rRNA distribution.pdf ---figure of length
distribution of different types of rRNAs, if sequence matches existed
             SeqFile_rRNA mapping.pdf ---figure of rsRNAs mapping
against different types of rRNAs, if sequence matches existed
             LSeqFile_tRNA_mapping.pdf
                                        ---figure of length
```

```
Some output folders only exist when '-k' parameter is applied in sports.pl;

SeqFile_mismatch_summary.txt only exist when '-M' parameter is lager than 0 in sports.pl;

Some output files might not exist if the file size is zero.
```

## fastq2fasta.pl

Extracted from miRDeep2 [3] (https://github.com/rajewsky-lab/mirdeep2)

- 1. Description:Parses fastq format files into fasta format.
- 2. Input: A fastq file.
- 3. Output: A fasta file, one sequence per line (the sequences are expanded).
- 4. options:-
- 5. Example usage:fastq2fasta.pl reads.fq > reads.fa

## fastaparse.pl

Extracted from miRDeep2 [3] (https://github.com/rajewsky-lab/mirdeep2)

- 1. Description:Performs simple filtering of entries in a fasta file.
- 2. Input: A fasta file
- 3. Output: A filtered fasta file
- 4. Options:-a only output entries where the sequence is minimum int nts long-b remove all entries that have a sequence that contains letters other than a,c,g,t,u,n,A,C,G,T,U,N.-s output progress
- 5. Example usage:fastaparse.pl reads.fa -a 15 -s > reads\_no\_short.fa 2>
   reads\_discarded.fa

#### combine\_reads.pl

- 1. Description:Combine reads in the fasta file to get unique sequence and its read number.
- 2. Input: A fasta file
- 3. Output:

```
the highest.
'1234567' represents the reads number of sequence
'TCCCTGGTGGTCTAGTGGTTAGGATTCGGCGC'
```

- 1. Options:-
- 2. Example usage:combine\_reads.pl reads.fa > combined\_reads.fa

### tRNA\_db\_processing.pl

- 1. Description:This script adds CCA end to original genomic tRNA 3' end and add G to Histidine tRNA 5' end.
- 2. Input:The reference tRNA database file in .fa formatA fasta file
- 3. Options: -
- 4. Example usage:tRNA\_db\_processing.pl input\_tRNA\_file.fa

## annotation.pl

- 1. Description:Combine the annotation information generated from sports.pl
- 2. Input:sports.pl output folder address:
- 3. Output:
  - 1. \_output.txt: A 6 column table file including annotation information for every unique sequence.

4.

5.

```
ID Sequence Match_Genome Annotation
               Sequence
                                                    Length Reads
   t00000001 TCCCTGGTGGTCTAGTGGTTAGGATTCGGCGC 32 1234567 Yes
tRNA-Glu-CTC 5 end
    -ID: t00000001
                                                  --Represents the unique ID of the
sequence, represents the abundance ranking among all the sequences. In this case,
the abundance of this sequence is the highest.
    -Sequence: TCCCTGGTGGTCTAGTGGTTAGGATTCGGCGC --Represents the sequence.
   -Length: 32
                                                  --Length of the sequence.
   -Reads: 1234567
                                                  --Reads number of the sequence.
    -Match Genome: Yes
                                                  --If the sequence can match the
reference genome.
    -Annotation: tRNA-Glu-CTC 5 end
                                                  -- The annotation of the sequence.
This sequence mapped against the 5' end of tRNA-Glu-CTC sequence.
```

1. \<seq\_fold>\_summary.txt: A 3 column table file including reads number of each major- and sub- classes.

```
Class Sub_Class Reads
tRNAdb-tRNA_5_end_Match_Genome - 123456
- tRNA-Glu-CTC 5 end 78910
```

-----

```
-Class: tRNAdb-tRNA_5_end_Match_Genome --The major class name.
-Sub_Class: tRNA-Glu-CTC_5_end --The sub class name.
-Reads: 123456 --The reads number of the class.
```

1. \<seq\_fold>\_length\_distribution.txt: A 3 column table file including reads number of each length distribution of each major class.

```
6.
```

```
Class Length Reads
tRNAdb-tRNA_5_end_Match_Genome 30 1234
tRNAdb-tRNA_5_end_Match_Genome 31 23456
tRNAdb-tRNA_5_end_Match_Genome 32 34567
tRNAdb-tRNA_5_end_Match_Genome 33 4567
.....

-Class: tRNAdb-tRNA_5_end_Match_Genome --The major class name.
-Length: 30 --Length of the sequence.
-Reads: 1234 --The reads number of the class.
```

- 7. Options: -
- 8. Example usage: annotation.pl <SPORTS\_output\_fold\_address>

# tRNA\_mapping.pl

- 1. Description: Perl script calculates expression level of genomic loci by bowtie 1 output information.
- 2. Input: bowtie 1 output file
- 3. Output: expression summary file
- 4. Options: -
- 5. Example usage: trna mapping.pl bowtie\_output\_file summary\_file

# mismatch\_summary.pl

- 1. Description: Perl script summarizes mismatch information generated by Bowtie 1 if parameter '-M' >0 applied in sports.pl
- 2. Input: bowtie 1 output file and threshold of reads number.
- 3. Output: mismatch summary file
- 4. Options: -
- 5. Example usage: mismatch summary.pl bowtie output file 10 > mismatch summary.txt

# overall\_RNA\_length\_distribution.R

1. Description: Generate figure of length distribution of miRNA, rsRNA, tsRNA, piRNA and other RNAs, if sequence matches exists.

- 2. Input: Files generated by annotation.pl
- 3. Output: \<seq\_fold>\_sncRNA\_distribution.pdf Generate figure of length distribution of different types of rRNAs, if sequence matches exists.
- 4. Options: -

# rRNA\_length\_distribution.R

- 1. Description: Generate figure of length distribution of different types of rRNAs, if sequence matches exists. (e.g. 4.5S, 5S, 5.3S, 5.8S, 12S, 16S, 18S, 28S, 45S ...)
- 2. Input: Files generated by annotation.pl
- 3. Output: \<seq\_fold>\_rRNA\_distribution.pdf
- 4. Options: -

# rRNA\_mapping.R

- 1. Description: Generate figure of rsRNAs mapping against different types of rRNAs if sequence matches existed. (e.g. 4.5S, 5S, 5.3S, 5.8S, 12S, 16S, 18S, 28S, 45S ...)
- 2. Input: Files generated by annotation.pl
- 3. Output: \<seq\_fold>\_rRNA\_mapping.pdf
- 4. Options: -

## tRNA\_mapping.R

- 1. Description: Generate figure of rsRNAs mapping against different types of tRNAs if sequence matches existed.
- 2. Input: Files generated by tRNA\_mapping.pl
- 3. Output: \<seq\_fold>\_tRNA\_mapping.pdf
- 4. Options: -

#### mismatch\_stat.R

- 1. Description: Generate statistics result for mismatch information generated from mismatch\_summary.pl
- 2. Input: File generated by mismatch\_summary.pl and base-calling error
- 3. Output: \<seq\_fold>\_mismatch\_summary.txt
- 4. Options: -

# **Appendix**

Available species lists of bowtie-index based reference database

```
To build bowtie-index of your own:
    cd /foo/bar/your_reference_database/
    bowtie-build <your_reference_database_name>.fa
<your_reference_database_name>
    The built bowtie index will be:
/foo/bar/your_reference_database/your_reference_database_name

Unpack reference genome (e.g. human genome):
    tar zxvf Homo_sapiens_UCSC_hg38.tar.gz
```

Main database source:

```
-mirbase 21 database [4] (Original source: http://www.mirbase.org/index.shtml)

-rRNA database (Original source: https://www.ncbi.nlm.nih.gov/nuccore)

-GtRNAdb 2.0 database [5] (Original source: http://gtrnadb.ucsc.edu/)

-piRBase database [6] (Original source: http://www.regulatoryrna.org/database/piRNA/)

-piRNABank [7] (Original source: http://pirnabank.ibab.ac.in/index.shtml)

-ensembl ncRNA database [8] (Original source: http://www.ensembl.org/index.html)

-rfam 12.3 database [9] (Original source: http://rfam.xfam.org/)
```

- 1. Homo sapiens (Human)
  - 1. annotation databases: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0773ed3d5f6b74f35bbd643e1af221c31&authkey=AcRxf8walnGUIEhgl--8CDc)

```
-ensembl ncRNA database with bowtie-index (Original source:
ftp://ftp.ensembl.org/pub/release-89/fasta/homo_sapiens/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta_files/)
```

1. SPORTS1.0 related parameters if you download recommended reference database:

- 4. Gorilla gorilla (Gorilla)
  - 1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=03a9a8d26cca14b458007e9c6ee4541f7&authkey=Aag33OX-ljvagRWePhYNF3k)

```
-genome with bowtie-index (UCSC gorGor5) (Original source: http://hgdownload.soe.ucsc.edu/goldenPath/gorGor5/bigZips/gorGor5.fa.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source: http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Acaro2/anoCar2-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source: ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

```
/<your_defined_address>/Gorilla_gorilla/UCSC/gorGor5/Sequence/BowtieIndex/genome
-m /<your_defined_address>/Gorilla_gorilla/miRBase_21/miRBase_21-ggo
-t /<your_defined_address>/Gorilla_gorilla/GtRNAdb/gorGor3-tRNAs
-f /<your_defined_address>/Gorilla_gorilla/Rfam_12.3/Rfam-12.3-gorilla
```

- 5. Pan paniscus (Bonobo)
  - annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=03a74e9f6c2594f1e86a31acd8e554621&authkey=AYrgOm8rrAY7hrFYQ03gmTA)

```
-genome with bowtie-index (UCSC panPan1) (Original source: http://hgdownload.soe.ucsc.edu/goldenPath/panPan1/bigZips/panPan1.fa.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-rfam 12.3 database with bowtie-index (Original source: ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

```
/<your_defined_address>/Pan_paniscus/UCSC/panPan1/Sequence/BowtieIndex/genome
-m /<your_defined_address>/Pan_paniscus/miRBase_21/miRBase_21-ppa
-f /<your_defined_address>/Pan_paniscus/Rfam 12.3/Rfam-12.3-Bonobo
```

- 6. Pan troglodytes (Chimp)
  - annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=09c13507ee7414365843de3450aa9ad3e&authkey=AdsLOHA5q2--SbiP2C6Qjpc)

```
-genome with bowtie-index (Ensembl CHIMP2.1.4)
(ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Pan_troglodytes/UCSC/panTro4/Pan_tr
oglodytes_UCSC_panTro4.tar.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Ptrog4/panTro4-tRNAs.fa)
-ensembl ncRNA database with bowtie-index (Original source:
ftp://ftp.ensembl.org/pub/release-89/fasta/pan_troglodytes/ncrna/)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta_files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -q

```
/<your defined address>/Pan troglodytes/UCSC/panTro4/Sequence/BowtieIndex/genome
-m /<your defined address>/Pan troglodytes/miRBase 21/miRBase 21-ptr
-t /<your defined address>/Pan troglodytes/GtRNAdb/panTro4-tRNAs
/<your defined address>/Pan troglodytes/Ensembl/Pan_troglodytes.CHIMP2.1.4.ncrna
-f /<your defined address>/Pan troglodytes/Rfam 12.3/Rfam-12.3-chimp
```

- 7. Pongo abelii (Orangutan)
  - 1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=037168296877942ecb9735f26afcb450a&authkey=AZkU5ib3A0KOiT4KxrhgTGQ)

```
-genome with bowtie-index (UCSC ponAbe2) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/ponAbe2/bigZips/chromFa.tar.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Ppygm2/ponAbe2-tRNAs.fa)
-ensembl ncRNA database with bowtie-index (Original source:
ftp://ftp.ensembl.org/pub/release-89/fasta/pongo abelii/ncrna/)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

```
/<your defined address>/Pongo abelii/UCSC/ponAbe2/Sequence/BowtieIndex/genome
-m /<your defined address>/Pongo abelii/miRBase 21/miRBase 21-ppy
-t /<your defined address>/Pongo abelii/GtRNAdb/ponAbe2-tRNAs
```

- -e /<your defined address>/Pongo abelii/Ensembl/Pongo abelii.PPYG2.ncrna
- -f /<your defined address>/Pongo abelii/Rfam 12.3/Rfam-12.3-orangutan
- 8. Nomascus leucogenys (Gibbon) 6.1. annotation database: (We provide a download link for all databases listed below:

https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.aspx? docid=068ef6629d31b4fb28100f667050be1d1&authkey=AWEPi3HUmOVD PPPnkMhdvs)

```
-genome with bowtie-index (UCSC nomLeu3) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/nomLeu3/bigZips/nomLeu3.fa.gz)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Nleuc3/nomLeu3-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

6.2 SPORTS1.0 related parameters if you download recommended reference database:

```
/<your defined address>/Nomascus leucogenys/UCSC/nomLeu3/Sequence/BowtieIndex/genom
-t /<your defined address>/Nomascus leucogenys/GtRNAdb/nomLeu3-tRNAs
-f /<your_defined_address>/Nomascus_leucogenys/Rfam_12.3/Rfam-12.3-gibbon
```

- 9. Macaca mulatta (Rhesus)
  - annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=07fededf7468444ba9b863b74316b8504&authkey=Aa6X06l4ExLvKtH8ml-CESs)

```
-genome with bowtie-index (UCSC rheMac8) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/rheMac8/bigZips/rheMac8.fa.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Mmula3/rheMac3-tRNAs.fa)
-rRNA database with bowtie-index (Original source:
https://www.ncbi.nlm.nih.gov/nuccore)
```

```
-rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
    /<your defined address>/Macaca mulatta/UCSC/rheMac8/Sequence/BowtieIndex/genome
    -m /<your defined address>/Macaca mulatta/miRBase 21/miRBase 21-mml
    -r /<your defined address>/Macaca mulatta/rRNA db/rhesus rRNA
    -t /<your defined address>/Macaca mulatta/GtRNAdb/rheMac3-tRNAs
    -f /<your defined address>/Macaca mulatta/Rfam 12.3/Rfam-12.3-rhesus
10. Papio anubis (Baboon)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=03a22a2092c7b46fb93a8fc49cf234720&authkey=Ad_hzH3MUIMh0-9BLoH_Vmw)
    -genome with bowtie-index (UCSC papAnu2) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/papAnu2/bigZips/papAnu2.fa.gz)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Panub2/papAnu2-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
     /<your defined address>/Papio anubis/UCSC/papAnu2/Sequence/BowtieIndex/genome
     -t /<your defined address>/Papio anubis/GtRNAdb/papAnu2-tRNAs
    -f /<your defined address>/Papio anubis/Rfam 12.3/Rfam-12.3-baboon
11. Callithrix jacchus (Marmoset)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=077698888fa8d40408df8c979e91146e4&authkey=AeoDYl5a3lKyF-CWgupu6lA)
    -genome with bowtie-index (UCSC calJac3) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/calJac3/bigZips/calJac3.fa.gz)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Cjacc3/calJac3-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -α
     /<your defined address>/Callithrix jacchus/UCSC/calJac3/Sequence/BowtieIndex/genom
    -m /<your defined address>/Callithrix jacchus/rRNAdb/marmoset rRNA
    -t /<your defined address>/Callithrix jacchus/GtRNAdb/calJac3-tRNAs
     -f /<your defined address>/Callithrix jacchus/Rfam 12.3/Rfam-12.3-marmoset
12. Carlito syrichta (Tarsier)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0c761313792b64396b87a66a6e04101be&authkey=AZBsjTA5-hXdASOIimxNL4I)
    -genome with bowtie-index (UCSC tarSyr2) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/tarSyr2/bigZips/tarSyr2.fa.gz)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Tsyri2/tarSyr2-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -q

-f /<your defined address>/Carlito syrichta/Rfam 12.3/Rfam-12.3-tarsier

-t /<your defined address>/Carlito syrichta/GtRNAdb/tarSyr2-tRNAs

/<your defined address>/Carlito syrichta/UCSC/tarSyr2/Sequence/BowtieIndex/genome

13. Rattus norvegicus (rat)

1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0b2cacd8453104b2abb60298863fc4c16&authkey=AZAYeCOsLKuc\_ml-QMqBJoQ) -genome with bowtie-index (UCSC rn6) (Original source: ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Rattus norvegicus/UCSC/rn6/Rattus no rvegicus UCSC rn6.tar.gz) -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml) -rRNA database with bowtie-index (Original source: https://www.ncbi.nlm.nih.gov/nuccore) -tRNA database with bowtie-index (Original source: http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Rnorv5/rn5-tRNAs.fa) -piRNA database with bowtie-index (Original source: http://www.regulatoryrna.org/database/piRNA/ && http://pirnabank.ibab.ac.in/request.html) -ensembl ncRNA database with bowtie-index (Original source: ftp://ftp.ensembl.org/pub/release-89/fasta/rattus norvegicus/ncrna/) -rfam 12.3 database with bowtie-index (Original source: ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/) 2. SPORTS1.0 related parameters if you download recommended reference database: -α /<your defined address>/Rattus norvegicus/UCSC/rn6/Sequence/BowtieIndex/genome -m /<your defined address>/Rattus norvegicus/miRBase 21/miRBase 21-rno -r /<your defined address>/Rattus norvegicus/rRNA db/mouse rRNA -t /<your defined address>/Rattus norvegicus/GtRNAdb/rn5-tRNAs -w /<your defined address>/Rattus norvegicus/piRBase/piR rat /<your defined address>/Rattus norvegicus/Ensembl/Rattus norvegicus.Rnor 6.0.ncrna -f /<your defined address>/Rattus norvegicus/Rfam 12.3/Rfam-12.3-rat 14. Mus musculus (mouse) 1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0833653a140eb47f098267d7a23d3b63c&authkey=Ab8aoYC8paqFl2yRablo7Ok) -genome with bowtie-index (UCSC mm10) (Original source: ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Mus musculus/UCSC/mm10/Mus musculus UCSC mm10.tar.gz) -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml) -rRNA database with bowtie-index (Original source: https://www.ncbi.nlm.nih.gov/nuccore) -tRNA database with bowtie-index (Original source: http://gtrnadb.ucsc.edu/genomes/eukaryota/Mmusc10/mm10-tRNAs.fa) -piRNA database with bowtie-index (Original source: http://www.regulatoryrna.org/database/piRNA/ && http://pirnabank.ibab.ac.in/request.html) -ensembl ncRNA database with bowtie-index (Original source: ftp://ftp.ensembl.org/pub/release-89/fasta/mus musculus/ncrna/) -rfam 12.3 database with bowtie-index (Original source: ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/) 2. SPORTS1.0 related parameters if you download recommended reference database: -g /<your defined address>/Mus musculus/UCSC/mm10/Sequence/BowtieIndex/genome -m /<your\_defined\_address>/Mus\_musculus/miRBase 21/miRBase 21-mmu

-r /<your\_defined\_address>/Mus\_musculus/rRNA\_db/mouse\_rRNA
-t /<your\_defined\_address>/Mus\_musculus/GtRNAdb/mm10-tRNAs

 $-w \ / < \verb|your_defined_address| > / \verb|Mus_musculus/piRBase/piR_mouse| \\$ 

- -e /<your\_defined\_address>/Mus\_musculus/Ensembl/Mus\_musculus.GRCm38.ncma
- -f /<your\_defined\_address>/Mus\_musculus/Rfam\_12.3/Rfam-12.3-mouse

#### 15. Cricetulus griseus (Hamster)

1. annotation database: (We provide a download link for all databases listed below:

```
https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as px?docid=0b0ac3830026f4007958774cbdb421632&authkey=AZnURBxU1PYzUO1yyrcoZ M)
```

```
-genome with bowtie-index (UCSC criGri1) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/criGri1/bigZips/criGri1.fa.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-rRNA database with bowtie-index (Original source:
https://www.ncbi.nlm.nih.gov/nuccore)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Cgris1/criGri1-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -q

 $/ < your\_defined\_address > / Cricetulus\_griseus / UCSC / criGri1 / Sequence / Bowtie Index / genome$  e

- -m /<your\_defined\_address>/Cricetulus\_griseus/miRBase\_21/miRBase\_21-cgr
- -r /<your\_defined\_address>/Cricetulus\_griseus/rRNA\_db/hamster\_rRNA
- -t /<your defined address>/Cricetulus\_griseus/GtRNAdb/criGril-tRNAs
- -f /<your\_defined\_address>/Cricetulus\_griseus/Rfam\_12.3/Rfam-12.3-hamster

#### 16. Cavia porcellus (Guinea pig)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0d27261e6ae9c4402bbc9465addb544de&authkey=Ad1xlk56DNm0StozUWDqCYw)

```
-genome with bowtie-index (UCSC cavPor3) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/cavPor3/bigZips/cavPor3.fa.gz)
-rRNA database with bowtie-index (Original source:
https://www.ncbi.nlm.nih.gov/nuccore)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Cporc3/cavPor3-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

/<your\_defined\_address>/Cavia\_porcellus/UCSC/cavPor3/Sequence/BowtieIndex/genome
-r /<your\_defined\_address>/Cavia\_porcellus/rRNA\_db/guinea\_rRNA

- -t /<your defined address>/Cavia porcellus/GtRNAdb/cavPor3-tRNAs
- -f /<your defined address>/Cavia porcellus/Rfam 12.3/Rfam-12.3-guinea
- 17. Heterocephalus glaber (Naked mole-rat)
  - annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0ceff76c620884395899bcd385bfaa098&authkey=ATiBTnSjWKqY0zPKQOmM6kU)

```
-genome with bowtie-index (UCSC hetGla2) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/hetGla2/bigZips/hetGla2.fa.gz)
-rRNA database with bowtie-index (Original source:
https://www.ncbi.nlm.nih.gov/nuccore)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Hglab2/hetGla2-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta_files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

/<your\_defined\_address>/Heterocephalus\_glaber/UCSC/hetGla2/Sequence/BowtieIndex/ge
nome

- -r /<your\_defined\_address>/Heterocephalus\_glaber/rRNA\_db/mole\_rRNA
- -t /<your\_defined\_address>/Heterocephalus\_glaber/GtRNAdb/hetGla2-tRNAs
- -f /<your defined address>/Heterocephalus glaber/Rfam 12.3/Rfam-12.3-mole

#### 18. Ictidomys tridecemlineatus (Squirrel)

1. annotation database: (We provide a download link for all databases listed below:

https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0419ea57bd00548cc94574d8ce9717fcd&authkey=Aa1pkb9R7850Ss3GFmB6GzM)

```
-genome with bowtie-index (UCSC speTri2) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/speTri2/bigZips/speTri2.fa.gz)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Strid2/speTri2-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -q

 $/ < your\_defined\_address > / Ictidomys\_tridecemlineatus / UCSC / speTri2 / Sequence / Bowtie Index / genome$ 

- -t /<your\_defined\_address>/Ictidomys\_tridecemlineatus/GtRNAdb/speTri2-tRNAs
- -f /<your defined address>/Ictidomys tridecemlineatus/Rfam 12.3/Rfam-12.3-squirrel

#### 19. Ochotona princeps (Pika)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=068d8da8388b74c38943e50237a259a88&authkey=AemP73yDCDiObexlk-DcRVQ)
 ``-genome with bowtie-index (UCSC ochPri3) (Original source:

http://hgdownload.soe.ucsc.edu/goldenPath/ochPri3/bigZips/ochPri3.fa.gz) -tRNA database with bowtie-index (Original source:

http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Oprin3/ochPri3-tRNAs.fa) -rfam 12.3 database with bowtie-index (Original source:

ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta\_files/)

2. SPORTS1.0 related parameters if you download recommended reference database: -q

 $/< your\_defined\_address > / Ochotona\_princeps / UCSC / ochPri3 / Sequence / Bowtie Index / genome - t / < your\_defined\_address > / Ochotona\_princeps / GtRNAdb / ochPri3 - tRNAs$ 

-f /<your defined address>/Ochotona princeps/Rfam 12.3/Rfam-12.3-pika

#### 20. Oryctolagus cuniculus (Rabbit)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=01b2d69333c97448196cc69b212e92fc9&authkey=ARbwyo0-WJX10nDdXwvNygc)

```
-genome with bowtie-index (UCSC oryCun2) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/oryCun2/bigZips/oryCun2.fa.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-rRNA database with bowtie-index (Original source:
https://www.ncbi.nlm.nih.gov/nuccore)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Ocuni2/oryCun2-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta_files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

/<your\_defined\_address>/Oryctolagus\_cuniculus/UCSC/oryCun2/Sequence/BowtieIndex/ge
nome

```
-m /<your defined address>/Oryctolagus cuniculus/miRBase_21/miRBase_21-ocu
```

- -r /<your defined address>/Oryctolagus cuniculus/rRNA db/rabbit rRNA
- -t /<your defined address>/Oryctolagus cuniculus/GtRNAdb/oryCun2-tRNAs
- -f /<your defined address>/Oryctolagus cuniculus/Rfam 12.3/Rfam-12.3-rabbit

#### 21. Ovis aries (Sheep)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0e880ae29a16747bebf5b9afdb5956921&authkey=AYIP0UkTwtVuiul45XA7mYE)

```
-genome with bowtie-index (UCSC oviAri3) (Original source:
http://hgdownload.cse.ucsc.edu/goldenPath/oviAri3/bigZips/oviAri3.fa.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
```

```
-tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Zmays5/zeaMay5-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
    /<your defined address>/Ovis aries/UCSC/oviAri3/Sequence/BowtieIndex/genome
    -m /<your defined address>/Ovis aries/miRBase 21/miRBase 21-oar
    -t /<your defined address>/Ovis aries/GtRNAdb/oviAri1-tRNAs
    -f /<your defined address>/Ovis aries/Rfam 12.3/Rfam-12.3-sheep
22. Bos taurus (Cow)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0a58e56dc7cb743299631fec15b72e69d&authkey=AeosGslpVMdvtZa6gtKJBCE)
    -genome with bowtie-index (UCSC bosTau8) (Original source:
    ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Bos taurus/UCSC/bosTau8/Bos taurus U
    CSC bosTau8.tar.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Btaur8/bosTau8-tRNAs.fa)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensembl.org/pub/release-89/fasta/bos taurus/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
    /<your defined address>/Bos taurus/UCSC/bosTau8/Sequence/BowtieIndex/genome
    -m /<your defined address>/Bos taurus/miRBase 21/miRBase 21-bta
    -r /<your defined address>/Bos taurus/rRNA db/cow rRNA
    -t /<your_defined_address>/Bos_taurus/GtRNAdb/bosTau8-tRNAs
    -e /<your defined address>/Bos taurus/Ensembl/Bos taurus.UMD3.1.ncrna
    -f /<your defined address>/Bos taurus/Rfam 12.3/Rfam-12.3-cow
23. Sus scrofa (Pig)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0e15d39ac05b24a9b899e7a9dfcf96773&authkey=AYaNWQ9KLlkqq7f2qbWzchc)
    -genome with bowtie-index (UCSC susScr3) (Original source:
    ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Sus scrofa/UCSC/susScr3/Sus scrofa U
    CSC susScr3.tar.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Sscro3/susScr3-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
    /<your defined address>/Sus scrofa/UCSC/susScr3/Sequence/BowtieIndex/genome
    -m /<your defined address>/Sus scrofa/miRBase 21/miRBase 21-ssc
    -r /<your defined address>/Sus scrofa/rRNA db/pig rRNA
    -t /<your defined address>/Sus scrofa/GtRNAdb/susScr3-tRNAs
    -f /<your defined address>/Sus scrofa/Rfam 12.3/Rfam-12.3-pig
```

24. Tursiops truncatus (Dolphin)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=08396ad3619a247d695508aee771e3991&authkey=AVKuiimDuoOGVKfGvZcK\_ik)

```
-genome with bowtie-index (UCSC turTru2) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/turTru2/bigZips/turTru2.fa.gz)
     -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Ttrun2/turTru2-tRNAs.fa)
     -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
     /<your defined address>/Tursiops truncatus/UCSC/turTru2/Sequence/BowtieIndex/genom
     -t /<your defined address>/Tursiops truncatus/GtRNAdb/turTru2-tRNAs
     -f /<your defined address>/Tursiops truncatus/Rfam 12.3/Rfam-12.3-dolphin
25. Balaenoptera acutorostrata (Minke whale)
  1. annotation database: (We provide a download link for all databases listed below:
     https://ncrnainfo-my.sharepoint.com/personal/sports ncrna info/ layouts/15/guestaccess.as
     px?docid=0d27ad173ffdb4dcc8a4954f9ba5426eb&authkey=AUdYvbA- g0lzVEEBlth8V8)
     -genome with bowtie-index (UCSC balAcul) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/balAcu1/bigZips/balAcu1.fa.gz)
     -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Bacut1/balAcu1-tRNAs.fa)
     -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
     /<your defined address>/Balaenoptera acutorostrata/UCSC/balAcu1/Sequence/BowtieInd
    ex/genome
     -t /<your defined address>/Balaenoptera acutorostrata/GtRNAdb/balAcu1-tRNAs
     -f /<your defined address>/Balaenoptera acutorostrata/Rfam 12.3/Rfam-12.3-whale
26. Erinaceus europaeus (Hedgehog)
  1. annotation database: (We provide a download link for all databases listed below:
     https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
     px?docid=0633e2d7781ab4aefb59fc03f1347657b&authkey=AXrP4XnyiHQqkC5WOUhhS5w)
     -genome with bowtie-index (UCSC eriEur2) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/eriEur2/bigZips/eriEur2.fa.gz)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Eeuro2/eriEur2-tRNAs.fa)
     -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
     /<your defined address>/Erinaceus europaeus/UCSC/eriEur2/Sequence/BowtieIndex/geno
     -t /<your defined address>/Erinaceus europaeus/GtRNAdb/eriEur2-tRNAs
     -f /<your defined address>/Erinaceus europaeus/Rfam 12.3/Rfam-12.3-hedgehog
27. Sorex araneus (Shrew)
  1. annotation database: (We provide a download link for all databases listed below:
     https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
     px?docid=0399046c56fb246c39630c84db61a23ad&authkey=AWE7pAlGt1TAAUylD4qoE9A)
     -genome with bowtie-index (UCSC sorAra2) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/sorAra2/biqZips/sorAra2.fa.gz)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Saran2/sorAra2-tRNAs.fa)
     -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
     /<your defined address>/Sorex araneus/UCSC/sorAra2/Sequence/BowtieIndex/genome
    -t /<your defined address>/Sorex araneus/GtRNAdb/sorAra2-tRNAs
```

-f /<your defined address>/Sorex araneus/Rfam 12.3/Rfam-12.3-shrew

- 28. Canis familiaris (Dog)
  - 1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=06a45d60105a04796b5e96a9417d86f4c&authkey=AY37yxKX-C5u9DG71tzrFEI)

```
-genome with bowtie-index (UCSC canFam3) (Original source:
```

ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Canis\_familiaris/UCSC/canFam3/Canis\_ familiaris UCSC canFam3.tar.gz)

-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)

-tRNA database with bowtie-index (Original source:

http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Cfami3/canFam3-tRNAs.fa)

-rfam 12.3 database with bowtie-index (Original source:

ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)

- 2. SPORTS1.0 related parameters if you download recommended reference database: -g
  - /<your\_defined\_address>/Canis\_familiaris/UCSC/canFam3/Sequence/BowtieIndex/genome
  - -m /<your\_defined\_address>/Canis\_familiaris/miRBase\_21/miRBase\_21-cfa
  - -t /<your defined address>/Canis familiaris/GtRNAdb/canFam3-tRNAs
  - -f /<your defined address>/Canis familiaris/Rfam 12.3/Rfam-12.3-dog
- 29. Mustela putorius furo (Ferret)
  - annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0ef4e067d42f241359e2e606b14e0d8f9&authkey=AUW2yr7SM356KpD5uhBNOeM)

```
-genome with bowtie-index (UCSC musFur1) (Original source:
```

http://hgdownload.soe.ucsc.edu/goldenPath/musFur1/bigZips/musFur1.fa.gz)

-tRNA database with bowtie-index (Original source:

http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Mputo1/musFur1-tRNAs.fa)

-rfam 12.3 database with bowtie-index (Original source:

ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)

2. SPORTS1.0 related parameters if you download recommended reference database: -g

/<your defined address>/Mustela furo/UCSC/musFur1/Sequence/BowtieIndex/genome

- -t /<your defined address>/Mustela furo/GtRNAdb/musFur1-tRNAs
- -f /<your defined address>/Mustela furo/Rfam 12.3-ferret
- 30. Ailuropoda melanoleuca (Panda)
  - 1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0215799b172f94c909c5251061e317540&authkey=AdE82hkEi1MHl3OL1vY92b8)

-genome with bowtie-index (UCSC ailMel1) (Original source:

http://hgdownload.soe.ucsc.edu/goldenPath/ailMel1/bigZips/ailMel1.fa.gz)

-tRNA database with bowtie-index (Original source:

http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Amela1/ailMel1-tRNAs.fa)

-rfam 12.3 database with bowtie-index (Original source:

ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta\_files/)

2. SPORTS1.0 related parameters if you download recommended reference database: -g

/<your\_defined\_address>/Ailuropoda\_melanoleuca/UCSC/ailMel1/Sequence/BowtieIndex/g
enome

- -t /<your defined address>/Ailuropoda melanoleuca/GtRNAdb/ailMel1-tRNAs
- -f /<your defined address>/Ailuropoda melanoleuca/Rfam-12.3-panda
- 31. Felis catus (Cat)
  - annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0203fe1a0c8954879878ef8cabcfe37cf&authkey=AZ32jdZI7FoFU\_t8\_NFb9o0)

```
-genome with bowtie-index (UCSC felCat8) (Original source:
```

http://hgdownload.soe.ucsc.edu/goldenPath/felCat8/bigZips/felCat8.fa.gz)

-rRNA database with bowtie-index (Original source:

https://www.ncbi.nlm.nih.gov/nuccore)

-tRNA database with bowtie-index (Original source:

```
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Ocuni2/oryCun2-tRNAs.fa) -rfam 12.3 database with bowtie-index (Original source: ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta_files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

 $/\!<\! \texttt{your\_defined\_address}\!\!>\!/ \texttt{Felis\_catus/UCSC/felCat8/Sequence/BowtieIndex/genome}$ 

- -r /<your\_defined\_address>/Felis\_catus/rRNA\_db/cat\_rRNA
- -t /<your\_defined\_address>/Felis\_catus/GtRNAdb/felCat5-tRNAs
- -f /<your defined address>/Felis catus/Rfam 12.3/Rfam-12.3-cat

#### 32. Equus caballus (Horse)

1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0e7011a73d57545ea8be3c71335b3cf4c&authkey=AWHkBfbzVmwz\_HjmQXo8lJU)

```
-genome with bowtie-index (UCSC equCab2) (Original source:
ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Equus_caballus/UCSC/equCab2/Equus_ca
ballus_UCSC_equCab2.tar.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Ecaba2/equCab2-tRNAs.fa)
-rRNA database with bowtie-index (Original source:
https://www.ncbi.nlm.nih.gov/nuccore)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

```
/<your_defined_address>/Equus_caballus/UCSC/equCab2/Sequence/BowtieIndex/genome
-m /<your_defined_address>/Equus_caballus/miRBase_21/miRBase_21-eca
-r /<your_defined_address>/Equus_caballus/rRNA_db/horse_rRNA
-t /<your_defined_address>/Equus_caballus/GtRNAdb/equCab2-tRNAs
-f /<your_defined_address>/Equus_caballus/Rfam 12.3/Rfam-12.3-horse
```

- 33. Ceratotherium simum (White rhinoceros)
  - 1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0f836e7106664459bacda6f94dc15e22d&authkey=AVSh1b1dr84q53sLTqD9sRA)

```
-genome with bowtie-index (UCSC cerSim1) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/cerSim1/bigZips/cerSim1.fa.gz)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Csimu1/cerSim1-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

 $/ < your\_defined\_address > / Ceratotherium\_simum/UCSC/cerSim1/Sequence/BowtieIndex/genome$ 

- -t /<your defined address>/Ceratotherium simum/GtRNAdb/cerSim1-tRNAs
- -f /<your\_defined\_address>/Ceratotherium\_simum/Rfam\_12.3/Rfam-12.3-rhinoceros
- 34. Myotis lucifugus (Microbat)
  - 1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=03dcc351bbc274cfc8692e5f2b1f5d0f3&authkey=AbjlxSozqu1c-2sRUrTGn7k)

```
-genome with bowtie-index (UCSC myoLuc2) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/myoLuc2/bigZips/myoLuc2.fa.gz)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Mluci2/myoLuc2-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta_files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g /<your defined address>/Myotis lucifugus/UCSC/myoLuc2/Sequence/BowtieIndex/genome

```
-t /<your_defined_address>/Myotis_lucifugus/GtRNAdb/myoLuc2-tRNAs
-f /<your defined address>/Myotis lucifugus/Rfam 12.3/Rfam-12.3-bat
```

#### 35. Trichechus manatus (Manatee)

1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=06448172c574b4b9cbe0d906cf75bc68b&authkey=AX\_fEuPmr18NZPEFbcg9nEQ)

```
-genome with bowtie-index (UCSC triMan1) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/triMan1/bigZips/triMan1.fa.gz)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Tmana1/triMan1-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

 $/ < your\_defined\_address > / Trichechus\_manatus / UCSC / triMan1 / Sequence / Bowtie Index / genome e$ 

- $\verb|-t|/<your_defined_address|| Trichechus_manatus/GtRNAdb/triMan1-tRNAs||$
- -f /<your defined address>/Trichechus manatus/Rfam 12.3/Rfam-12.3-manatee

#### 36. Loxodonta africana (Elephant)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0e0146e32fe4745bba50bbede409efddc&authkey=AT8bNP2DhjKysDaGD4Qy-7s)

```
-genome with bowtie-index (UCSC loxAfr3) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/loxAfr3/bigZips/loxAfr3.fa.gz)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Lafri3/loxAfr3-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

/<your\_defined\_address>/Loxodonta\_africana/UCSC/loxAfr3/Sequence/BowtieIndex/genom
e

- -t /<your defined address>/Loxodonta africana/GtRNAdb/loxAfr3-tRNAs
- -f /<your defined address>/Loxodonta africana/Rfam 12.3/Rfam-12.3-elephant
- 37. Dasypus novemcinctus (Armadillo)
  - 1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0c68adc6b2afc447a9dfe45a1a5eedd49&authkey=AUPTS51dr88E3AMvcZtljrk)

```
-genome with bowtie-index (UCSC dasNov3) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/dasNov3/bigZips/dasNov3.fa.gz)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Dnove3/dasNov3-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta_files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

 $/ < your\_defined\_address > / \texttt{Dasypus\_novemcinctus/UCSC/dasNov3/Sequence/BowtieIndex/genome} \\$ 

- -t /<your\_defined\_address>/Dasypus\_novemcinctus/GtRNAdb/dasNov3-tRNAs
- -f /<your defined address>/Dasypus novemcinctus/Rfam 12.3/Rfam-12.3-armadillo

#### 38. Notamacropus eugenii (Wallaby)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=076db8740d2e34caabb25669fd6297e36&authkey=AWv-dFi1Mm7C0QH9K00WHH4)

```
-genome with bowtie-index (UCSC macEug2) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/macEug2/bigZips/macEug2.fa.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source:
```

```
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Meuge2/macEug2-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
    /<your defined address>/Notamacropus eugenii/UCSC/macEug2/Sequence/BowtieIndex/gen
    ome
    -m /<your defined address>/Notamacropus eugenii/miRBase 21/miRBase 21-meu
    -t /<your defined address>/Notamacropus eugenii/GtRNAdb/macEug2-tRNAs
    -f /<your_defined_address>/Notamacropus_eugenii/Rfam_12.3/Rfam-12.3-wallaby
39. Sarcophilus harrisii (Tasmanian devil)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0469dc3788cbb40bab7472ee70a230dc0&authkey=AXA1nDb4QpYeuI|FC-D6mL0)
    -genome with bowtie-index (UCSC sarHarl) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/sarHar1/bigZips/sarHar1.fa.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Sharr1/sarHar1-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
    /<your defined address>/Sarcophilus harrisii/UCSC/sarHar1/Sequence/BowtieIndex/gen
    -m /<your defined address>/Sarcophilus harrisii/miRBase 21/miRBase 21-sha
    -t /<your defined address>/Sarcophilus harrisii/GtRNAdb/sarHar1-tRNAs
     -f /<your defined address>/Sarcophilus harrisii/Rfam 12.3/Rfam-12.3-tasmanian
40. Monodelphis domestica (Opossum)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=05e9cd2f3891a4761ade95d98aa3ebc78&authkey=Ae jBKldBC7HypVTN7S98Rw)
    -genome with bowtie-index (UCSC monDom5) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/monDom5/bigZips/chromFa.tar.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Mdome5/monDom5-tRNAs.fa)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensembl.org/pub/release-89/fasta/monodelphis domestica/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
    /<your defined address>/Monodelphis domestica/UCSC/monDom5/Sequence/BowtieIndex/ge
    -m /<your_defined_address>/Monodelphis_domestica/miRBase_21/miRBase_21-mdo
    -t /<your defined address>/Monodelphis domestica/GtRNAdb/monDom5-tRNAs
    /<your defined address>/Monodelphis domestica/Ensembl/Monodelphis domestica.BROADO
```

-f /<your\_defined\_address>/Monodelphis\_domestica/Rfam\_12.3/Rfam-12.3-opossum
41. Ornithorhynchus anatinus (Platypus)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=07f0f4ed088844a86afa5db763407699c&authkey=AUFx9yWXHtg1CQc-wfpJ81M)

```
-genome with bowtie-index (UCSC ornAna2) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/ornAna2/bigZips/ornAna2.fa.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
```

```
-tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Oanat1/ornAna1-tRNAs.fa)
     -piRNA database with bowtie-index (Original source:
    http://pirnabank.ibab.ac.in/request.html)
     -ensembl ncRNA database with bowtie-index (Original source:
     ftp://ftp.ensembl.org/pub/release-89/fasta/ornithorhynchus anatinus/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
     /<your defined address>/Ornithorhynchus anatinus/UCSC/ornAna2/Sequence/BowtieIndex
     /genome
     -m /<your defined address>/Ornithorhynchus anatinus/miRBase 21/miRBase 21-oan
     -t /<your defined address>/Ornithorhynchus anatinus/GtRNAdb/ornAnal-tRNAs
     -w /<your defined address>/Ornithorhynchus anatinus/piRBase/piR platypus
     /<your defined address>/Ornithorhynchus anatinus/Ensembl/Ornithorhynchus anatinus.
     -f /<your defined address>/Ornithorhynchus anatinus/Rfam 12.3/Rfam-12.3-platypus
42. Taeniopygia guttata (Zebra finch)
  1. annotation database: (We provide a download link for all databases listed below:
     https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
     px?docid=0defbf712045f4e7f85b373b0eba4cd1b&authkey=AbieXkg6akbsD8tY1oGctDI)
     -genome with bowtie-index (UCSC taeGut2) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/taeGut2/bigZips/taeGut2.fa.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Tgutt2/taeGut2-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
     /<your defined address>/Taeniopygia guttata/UCSC/taeGut2/Sequence/BowtieIndex/geno
     -m /<your defined address>/Taeniopygia quttata/miRBase 21/miRBase 21-tqu
     -t /<your defined address>/Taeniopygia guttata/GtRNAdb/taeGut2-tRNAs
     -f /<your defined address>/Taeniopygia guttata/Rfam 12.3/Rfam-12.3-finch
43. Melopsittacus undulatus (Budgerigar)
  1. annotation database: (We provide a download link for all databases listed below:
     https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
     px?docid=0e0046ab3b1a9458a9a183b45507ea0c3&authkey=AWiCroVnVDQcPENZDX_76VM)
     -genome with bowtie-index (UCSC melUnd1) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/melUnd1/bigZips/melUnd1.fa.gz)
     -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Mundu1/melUnd1-tRNAs.fa)
     -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
     /<your defined address>/Melopsittacus undulatus/UCSC/melUnd1/Sequence/BowtieIndex/
    genome
     -t /<your defined address>/Melopsittacus undulatus/GtRNAdb/melUnd1-tRNAs
     -f /<your defined address>/Melopsittacus undulatus/Rfam 12.3/Rfam-12.3-budgerigar
44. Gallus gallus (Chicken)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
     px?docid=0abafb326c4074fe9971d60a26497126c&authkey=AcRw 9ltjRXbdNAfmLRl gg)
```

-genome with bowtie-index (UCSC galGal5) (Original source:

```
ftp://igenome2:u7NMwVkm@ftp.illumina.com/Gallus gallus/UCSC/galGal5/Gallus gallus
    UCSC galGal5.tar.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Ggall4/galGal4-tRNAs.fa)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -piRNA database with bowtie-index (Original source:
    http://www.regulatoryrna.org/database/piRNA/)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensembl.org/pub/release-89/fasta/gallus gallus/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. PORTS1.0 related parameters if you download recommended reference database: -q
     /<your defined address>/Gallus gallus/UCSC/galGal5/Sequence/BowtieIndex/genome
    -m /<your defined address>/Gallus gallus/miRBase 21/miRBase 21-gga
    -r /<your defined address>/Gallus gallus/rRNA db/chicken rRNA
    -t /<your defined address>/Gallus gallus/GtRNAdb/galGal4-tRNAs
    -w /<your_defined_address>/Gallus_gallus/piRBase/piR_gga_v1.0
    /<your defined address>/Gallus gallus/Ensembl/Gallus gallus.Gallus gallus-5.0.ncrn
    -f /<your defined address>/Gallus gallus/Rfam 12.3/Rfam-12.3-chicken
45. Meleagris gallopavo (Turkey)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0c88e1661f08e4b6d945d87e8120bdf07&authkey=AWlCjj414nXNuNbrNM7mbE4)
    -genome with bowtie-index (UCSC melGall) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/melGal1/bigZips/melGal1.fa.gz)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Mgall1/melGal1-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
    /<your defined address>/Meleagris gallopavo/UCSC/melGal1/Sequence/BowtieIndex/geno
    -t /<your_defined_address>/Meleagris_gallopavo/GtRNAdb/melGall-tRNAs
    -f /<your defined address>/Meleagris gallopavo/Rfam 12.3/Rfam-12.3-turkey
46. Chrysemys picta (Painted Turtle)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0fbd62d91eb4442a88525f89684a74242&authkey=AY3fgBdvPRzWpqqD4yd_Sqw)
    -genome with bowtie-index (UCSC chrPic1) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/chrPic1/bigZips/chrPic1.fa.gz)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Ocuni2/oryCun2-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
     /<your defined address>/Chrysemys picta/UCSC/chrPic1/Sequence/BowtieIndex/genome
    -t /<your defined address>/Chrysemys picta/GtRNAdb/chrPic1-tRNAs
    -f /<your defined address>/Chrysemys picta/Rfam 12.3/Rfam-12.3-turtle
```

47. Anolis carolinensis (Lizard)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=06986df31c45c4a00a6a2b47ce4ee7af2&authkey=AaE-i58-1fjJeqcDHZUpLLo)

```
-genome with bowtie-index (UCSC anoCar2) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/anoCar2/bigZips/anoCar2.fa.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Acaro2/anoCar2-tRNAs.fa)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensembl.org/pub/release-89/fasta/anolis carolinensis/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
    /<your defined address>/Anolis carolinensis/UCSC/anoCar2/Sequence/BowtieIndex/geno
    -m /<your defined address>/Anolis carolinensis/miRBase 21/miRBase 21-aca
    -t /<your defined address>/Anolis carolinensis/GtRNAdb/anoCar2-tRNAs
    /<your defined address>/Anolis carolinensis/Ensembl/Anolis carolinensis.AnoCar2.0.
     -f /<your defined address>/Anolis carolinensis/Rfam 12.3/Rfam-12.3-lizard
48. Xenopus laevis (Frog)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0d9b3d45b3bf3483c826c36d2b340f0fd&authkey=AdGZ969RHd1lpwBpm7lsGEQ)
    -genome with bowtie-index (UCSC xenTro7) (Original source:
    ftp://hqdownload.soe.ucsc.edu/goldenPath/xenTro7/biqZips/xenTro7.fa.qz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Xtrop3/xenTro3-tRNAs.fa)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -piRNA database with bowtie-index (Original source:
    http://www.regulatoryrna.org/database/piRNA/)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensembl.org/pub/release-89/fasta/xenopus tropicalis/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
    /<your defined address>/Xenopus laevis/UCSC/xenTro7/Sequence/BowtieIndex/genome
    -m /<your defined address>/Xenopus laevis/miRBase 21/miRBase 21-xtr
    -r /<your defined address>/Xenopus laevis/rRNA db/frog rRNA
    -t /<your defined address>/Xenopus laevis/GtRNAdb/xenTro3-tRNAs
    -w /<your defined address>/Xenopus laevis/piRBase/piR xtr v1.0
    -e /<your defined address>/Xenopus laevis/Ensembl/Xenopus tropicalis.JGI 4.2.ncrna
    -f /<your defined address>/Xenopus laevis/Rfam 12.3/Rfam-12.3-frog
49. Latimeria chalumnae (Coelacanth)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0be8104aeb59d4ff89909d0c62d0a2f4e&authkey=AasK4DZcbB12a8wB8CNz6Ak)
    -genome with bowtie-index (UCSC latChal) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/latCha1/bigZips/latCha1.fa.gz)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Lchal1/latChal-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
     ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -q

```
/<your defined address>/Latimeria chalumnae/UCSC/latChal/Sequence/BowtieIndex/geno
    -r /<your defined address>/Latimeria chalumnae/rRNA db/coelacanth rRNA
    -t /<your defined address>/Latimeria chalumnae/GtRNAdb/latChal-tRNAs
    -f /<your defined address>/Latimeria chalumnae/Rfam 12.3/Rfam-12.3-coelacanth
50. Tetraodon nigroviridis (Tetraodon)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=06252eaadd5894a3795afce052716fd17&authkey=AWZ9FVt-iphiOFRDzuKoltA)
    -genome with bowtie-index (UCSC tetNig2) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/tetNig2/bigZips/chromFa.tar.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Tnigr2/tetNig2-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
     /<your defined address>/Tetraodon nigroviridis/UCSC/tetNig2/Sequence/BowtieIndex/g
    enome
    -m /<your defined address>/Tetraodon nigroviridis/miRBase 21/miRBase 21-tni
    -t /<your defined address>/Tetraodon nigroviridis/GtRNAdb/tetNig2-tRNAs
     -f /<your defined address>/Tetraodon nigroviridis/Rfam 12.3/Rfam-12.3-tetraodon
51. Takifugu rubripes (Fugu)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=026501a5c4aa54930a00289acf9691f5f&authkey=AWUaDYW0 VZx-1dy5wnCfdO)
    -genome with bowtie-index (UCSC fr3) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/fr3/bigZips/fr3.fa.gz)
    -tRNA database with bowtie-index (Original source:
    http://qtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Trubr3/fr3-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
    /<your defined address>/Takifugu rubripes/UCSC/fr3/Sequence/BowtieIndex/genome
    -t /<your defined address>/Takifugu rubripes/GtRNAdb/fr3-tRNAs
     -f /<your defined address>/Takifugu rubripes/Rfam_12.3/Rfam-12.3-fugu
52. Gasterosteus aculeatus (Stickleback)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0268db779d3654712b39d5450bac55894&authkey=AQwh0wJeAcvF78JRu4RHddM)
    -genome with bowtie-index (UCSC gasAcu1) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/gasAcul/bigZips/chromFa.tar.gz)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Gacul1/gasAcu1-tRNAs.fa)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensembl.org/pub/release-89/fasta/gasterosteus aculeatus/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
     /<your defined address>/Gasterosteus aculeatus/UCSC/gasAcu1/Sequence/BowtieIndex/g
    enome
    -t /<your defined address>/Gasterosteus aculeatus/GtRNAdb/gasAcu1-tRNAs
     /<your defined address>/Gasterosteus aculeatus/Ensembl/Gasterosteus aculeatus.BROA
```

DS1.ncrna

-f /<your\_defined\_address>/Gasterosteus\_aculeatus/Rfam\_12.3/Rfam-12.3-stickleback
53. Oryzias latipes (Medaka)

1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0f5089e2cc47245f28dca1a3f8f75343c&authkey=AV8N2wRviWth\_LihlqpB0zk)

```
-genome with bowtie-index (UCSC oryLat2)
(http://hgdownload.soe.ucsc.edu/goldenPath/oryLat2/bigZips/oryLat2.fa.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Olati2/oryLat2-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

```
/<your_defined_address>/Oryzias_latipes/UCSC/oryLat2/Sequence/BowtieIndex/genome
-m /<your_defined_address>/Oryzias_latipes/miRBase_21/miRBase_21-ola
-t /<your_defined_address>/Oryzias_latipes/GtRNAdb/oryLat2-tRNAs
-f /<your_defined_address>/Oryzias_latipes/Rfam_12.3/Rfam-12.3-medaka
```

#### 54. Oreochromis niloticus (Nile tilapia)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0cf791d1d99de4f4c8ed7f8de6d5694f8&authkey=AZZgbe3d1aKb7GDuCDlv81w)

```
-genome with bowtie-index (UCSC oreNil2) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/oreNil2/bigZips/oreNil2.fa.gz)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Onilo2/oreNil2-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

- 2. SPORTS1.0 related parameters if you download recommended reference database: -g
  - /<your\_defined\_address>/Oreochromis\_niloticus/UCSC/oreNil2/Sequence/BowtieIndex/ge
    nome
  - -t /<your defined address>/Oreochromis niloticus/GtRNAdb/oreNil2-tRNAs
  - -f /<your defined address>/Oreochromis niloticus/Rfam 12.3/Rfam-12.3-tilapia

#### 55. Gadus morhua (Atlantic cod)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=08f5a65a279f34d518cc4017cb04fe469&authkey=ASMQLlwmWReD\_bVW922N0lo)

```
-genome with bowtie-index (UCSC gadMor1) (Original source:
http://hgdownload.soe.ucsc.edu/goldenPath/gadMor1/bigZips/gadMor1.fa.gz)
-rRNA database with bowtie-index (Original source:
https://www.ncbi.nlm.nih.gov/nuccore)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Gmorh1/gadMor1-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

```
/<your_defined_address>/Gadus_morhua/UCSC/gadMor1/Sequence/BowtieIndex/genome
-r /<your defined address>/Gadus morhua/rRNA db/cod rRNA
```

- -t /<your\_defined\_address>/Gadus\_morhua/GtRNAdb/gadMor1-tRNAs
- -f /<your defined address>/Gadus morhua/Rfam 12.3/Rfam-12.3-cod

#### 56. Danio rerio (Zebrafish)

1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=07d3448d6cd29485498e70f8f067a619d&authkey=Aa1-VjshX-GKLZI7limHFlo)

```
-genome with bowtie-index (UCSC danRer10) (Original source: ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Danio rerio/UCSC/danRer10/Danio reri
```

```
o UCSC danRer10.tar.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Dreri v8/danRer6-tRNAs.fa)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -piRNA database with bowtie-index (Original source:
    http://www.regulatoryrna.org/database/piRNA/)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensembl.org/pub/release-89/fasta/danio rerio/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
    /<your defined address>/Danio rerio/UCSC/danRer10/Sequence/BowtieIndex/genome
    -m /<your defined address>/Danio rerio/miRBase 21/miRBase 21-dre
    -r /<your defined address>/Danio rerio/rRNA db/zebrafish rRNA
    -t /<your defined address>/Danio rerio/GtRNAdb/danRer6-tRNAs
    -w /<your defined address>/Danio rerio/piRBase/piR dre v1.0
    -e /<your defined address>/Danio rerio/Ensembl/Danio rerio.GRCz10.ncrna
    -f /<your defined address>/Danio rerio/Rfam 12.3/Rfam-12.3-zebrafish
57. Callorhinchus milii (Elephant shark)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=053976e5c17b9435982d22223b9389eba&authkey=ASL1VUPU4Ol2PskRXpj94t4)
    -genome with bowtie-index (UCSC calMill) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/calMil1/bigZips/calMil1.fa.gz)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Cmili1/calMil1-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
    /<your defined address>/Callorhinchus milii/UCSC/calMil1/Sequence/BowtieIndex/geno
    me
    -t /<your defined address>/Callorhinchus milii/GtRNAdb/calMil1-tRNAs
    -f /<your defined address>/Callorhinchus milii/Rfam 12.3/Rfam-12.3-shark
58. Petromyzon marinus (Lamprey)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=013a1f7a5585b462b801f1cfe3faf2cdd&authkey=AVN94xcHd1_Aa2ofYovsw8Q)
    -genome with bowtie-index (UCSC petMar2) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/petMar2/bigZips/petMar2.fa.gz)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Pmari2/petMar2-tRNAs.fa)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
    /<your defined address>/Petromyzon marinus/UCSC/petMar2/Sequence/BowtieIndex/genom
    -r /<your defined address>/Petromyzon marinus/rRNA db/lamprey rRNA
    -t /<your defined address>/Petromyzon marinus/GtRNAdb/petMar2-tRNAs
    -f /<your_defined_address>/Petromyzon_marinus/Rfam_12.3/Rfam-12.3-lamprey
59. Strongylocentrotus purpuratus (Sea urchin)
```

1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as

```
px?docid=0840567689dbe4ff49852c8744056f172&authkey=AWhmaOmPgN5fuB1RXIKc69U)
    -genome with bowtie-index (UCSC strPur2) (Original source:
    http://hgdownload.soe.ucsc.edu/goldenPath/strPur2/bigZips/strPur2.fa.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Spurp/Spurp-tRNAs.fa)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensemblgenomes.org/pub/metazoa/release-36/fasta/strongylocentrotus purpu
    ratus/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
    /<your defined address>/Strongylocentrotus purpuratus/UCSC/strPur2/Sequence/Bowtie
    Index/genome
    -m /<your defined address>/Strongylocentrotus purpuratus/miRBase 21/miRBase 21-spu
    -r /<your defined address>/Strongylocentrotus purpuratus/rRNA db/urchin rRNA
    -t /<your defined address>/Strongylocentrotus purpuratus/GtRNAdb/Spurp-tRNAs
    /<your defined address>/Strongylocentrotus purpuratus/Ensembl/Strongylocentrotus p
    urpuratus.GCA 000002235.2.ncrna
    -f
     /<your defined address>/Strongylocentrotus purpuratus/Rfam 12.3/Rfam-12.3-urchin
60. Drosophila melanogaster (Drosophila)

    annotation database: (We provide a download link for all databases listed below:

    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=06d15ec2032e141648ce4eedd413b0e0c&authkey=ARejQLC8ofAhQq9lwwoB0Pw)
    -genome with bowtie-index (UCSC dm6) (Original source:
    ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Drosophila melanogaster/UCSC/dm6/Dro
    sophila melanogaster UCSC dm6.tar.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/genomes/eukaryota/Dmela6/dm6-tRNAs.fa)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -piRNA database with bowtie-index (Original source:
    http://www.regulatoryrna.org/database/piRNA/)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensemblgenomes.org/pub/metazoa/release-36/fasta/drosophila melanogaster/
    ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
    /<your defined address>/Drosophila melanogaster/UCSC/dm6/Sequence/BowtieIndex/geno
    -m /<your defined address>/Drosophila melanogaster/miRBase 21/miRBase 21-dme
    -r /<your defined address>/Drosophila melanogaster/rRNA db/drosophila rRNA
    -t /<your defined address>/Drosophila melanogaster/GtRNAdb/dm6-tRNAs
    -w /<your defined address>/Drosophila melanogaster/piRBase/piR dme
     /<your_defined_address>/Drosophila_melanogaster/Ensembl/Drosophila_melanogaster.BD
    -f /<your defined address>/Drosophila melanogaster/Rfam 12.3/Rfam-12.3-drosophila
61. Anopheles gambiae (Mosquito)
  1. annotation database: (We provide a download link for all databases listed below:
```

# https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=003bfc542d06b42edb24a94969bb12b36&authkey=AdgdWQ5emWRWTYvXcgPT0Fo)

```
-genome with bowtie-index (UCSC anoGam1) (Original source: http://hgdownload.soe.ucsc.edu/goldenPath/anoGam1/bigZips/chromFa.zip)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source: http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Agamb/Agamb-tRNAs.fa)
-ensembl ncRNA database with bowtie-index (Original source: ftp://ftp.ensemblgenomes.org/pub/metazoa/release-36/fasta/anopheles_gambiae/ncrna/)
-rfam 12.3 database with bowtie-index (Original source: ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

```
/<your_defined_address>/Anopheles_gambiae/UCSC/anoGam1/Sequence/BowtieIndex/genome
-m /<your_defined_address>/Anopheles_gambiae/miRBase_21/miRBase_21-aga
-t /<your_defined_address>/Anopheles_gambiae/GtRNAdb/Agamb-tRNAs
-e
/<your_defined_address>/Anopheles_gambiae/Ensembl/Anopheles_gambiae.AgamP4.ncrna
-f /<your_defined_address>/Anopheles_gambiae/Rfam 12.3/Rfam-12.3-mosquito
```

#### 62. Pristionchus pacificus (Roundworm)

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0877c09c8493b47e9b9fb6c2c0bdc6015&authkey=AQJrbZlyZz9MDnPB9uYxMYE)

```
-genome with bowtie-index (UCSC priPac1) (Original source: http://hgdownload.soe.ucsc.edu/goldenPath/priPac1/bigZips/chromFa.tar.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source: http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Ppaci1/priPac1-tRNAs.fa)
-rfam 12.3 database with bowtie-index (Original source: ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

```
/<your_defined_address>/Pristionchus_pacificus/UCSC/priPac1/Sequence/BowtieIndex/g
enome
```

```
-m /<your defined address>/Pristionchus pacificus/miRBase 21/miRBase 21-ppc
```

- -r /<your defined address>/Pristionchus pacificus/rRNA db/roundworm rRNA
- -t /<your\_defined\_address>/Pristionchus\_pacificus/GtRNAdb/priPac1-tRNAs
- -f /<your defined address>/Pristionchus pacificus/Rfam 12.3/Rfam-12.3-roundworm

#### 63. Caenorhabditis elegans (Nematode):

 annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0cffdc7d5a3844567bc5303adca47fb81&authkey=AdA1jCVLRJkz2k8oSVU-pCI)

```
-genome with bowtie-index (UCSC ce10) (Original source:
ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Caenorhabditis_elegans/UCSC/ce10/Cae
norhabditis_elegans_UCSC_ce10.tar.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/genomes/eukaryota/Celeg_WS220/ce10-tRNAs.fa)
-rRNA database with bowtie-index (Original source:
https://www.ncbi.nlm.nih.gov/nuccore)
-piRNA database with bowtie-index (Original source:
http://www.regulatoryrna.org/database/piRNA/)
-ensembl ncRNA database with bowtie-index (Original source:
ftp://ftp.ensemblgenomes.org/pub/metazoa/release-36/fasta/caenorhabditis_elegans/n
crna/)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

```
/<your defined address>/Caenorhabditis elegans/UCSC/ce10/Sequence/BowtieIndex/geno
    -m /<your defined address>/Caenorhabditis elegans/miRBase 21/miRBase 21-cel
    -r /<your defined address>/Caenorhabditis elegans/rRNA db/cel rRNA
    -t /<your defined address>/Caenorhabditis elegans/GtRNAdb/ce10-tRNAs
    -w /<your defined address>/Caenorhabditis elegans/piRBase/piR cel v1.0
    /<your defined address>/Caenorhabditis elegans/Ensembl/Caenorhabditis elegans.WBce
    1235.ncrna
    -f /<your defined address>/Caenorhabditis elegans/Rfam 12.3/Rfam-12.3-nematode
64. Saccharomyces cerevisiae (Yeast)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0a974d08ffcb842dea8d18462dcfaf2f4&authkey=AQ6wsr9hNwVfCfgvuScKStE)
    -genome with bowtie-index UCSC sacCer3) (Original source:
    ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Saccharomyces cerevisiae/UCSC/sacCer
    3/Saccharomyces cerevisiae UCSC sacCer3.tar.gz)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Scere3/sacCer3-tRNAs.fa)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensemblgenomes.org/pub/fungi/release-36/fasta/saccharomyces cerevisiae/n
    crna/)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
    /<your defined address>/Saccharomyces cerevisiae/UCSC/sacCer3/Sequence/BowtieIndex
    -r /<your defined address>/Saccharomyces cerevisiae/rRNA db/yeast rRNA
    -t /<your defined address>/Saccharomyces cerevisiae/GtRNAdb/sacCer3-tRNAs
    /<your defined address>/Saccharomyces cerevisiae/Ensembl/Saccharomyces cerevisiae.
    R64-1-1.ncrna
    -f /<your defined address>/Saccharomyces cerevisiae/Rfam 12.3/Rfam-12.3-yeast
65. Zea mays (Corn)
  1. annotation database: (We provide a download link for all databases listed below:
    https://ncrnainfo-my.sharepoint.com/personal/sports ncrna info/ layouts/15/guestaccess.as
    px?docid=0abdce07adf9449e89bdcb89d4d4609a4&authkey=AViNEp3y6Y4hNBYSEuW_sYg)
     -genome with bowtie-index (Ensembl AGPv4) (Original source:
    ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Zea mays/Ensembl/AGPv4/Zea mays Ense
    mbl AGPv4.tar.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Zmays5/zeaMay5-tRNAs.fa)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensemblgenomes.org/pub/plants/release-36/fasta/zea mays/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta_files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
    /<your defined address>/Zea mays/Ensembl/AGPv4/Sequence/BowtieIndex/genome
    -m /<your_defined_address>/Zea_mays/miRBase_21/miRBase_21-zma
    -r /<your defined address>/Zea mays/rRNA db/corn rRNA
    -t /<your defined address>/Zea mays/GtRNAdb/zeaMay5-tRNAs
```

```
-e /<your_defined_address>/Zea_mays/Ensembl/Zea_mays.AGPv4.ncrna
-f /<your defined address>/Zea mays/Rfam 12.3/Rfam-12.3-corn
```

#### 66. Sorghum bicolor (Sorghum)

1. annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0335fb9b05a024e61819c04db7ffd2a51&authkey=AdMT31NWtny7F1NzJNn3zZI)

```
-genome with bowtie-index (Ensembl Sbi1) (Original source:
ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Sorghum_bicolor/Ensembl/Sbi1/Sorghum
_bicolor_Ensembl_Sbi1.tar.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Sbico/Sbico-tRNAs.fa)
-rRNA database with bowtie-index (Original source:
https://www.ncbi.nlm.nih.gov/nuccore)
-ensembl ncRNA database with bowtie-index (Original source:
ftp://ftp.ensemblgenomes.org/pub/plants/release-36/fasta/sorghum_bicolor/ncrna/)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g

```
/<your_defined_address>/Sorghum_bicolor/Ensembl/Sbi1/Sequence/BowtieIndex/genome
-m /<your_defined_address>/Sorghum_bicolor/miRBase_21/miRBase_21-sbi
-r /<your_defined_address>/Sorghum_bicolor/rRNA_db/sorghum_rRNA
-t /<your_defined_address>/Sorghum_bicolor/GtRNAdb/Sbico-tRNAs
-e
/<your_defined_address>/Sorghum_bicolor/Ensembl/Sorghum_bicolor.Sorghum_bicolor_v2.
ncrna
```

- -f /<your defined address>/Sorghum bicolor/Rfam 12.3/Rfam-12.3-sorghum
- 67. Oryza sativa (Rice)
  - annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=0510818bde4a342868928f982420dd07c&authkey=ASfslUL0BX2HYew7xrw2x1A)

```
-genome with bowtie-index (Ensembl IRGSP-1.0) (Original source:
ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Oryza_sativa_japonica/Ensembl/IRGSP-
1.0/Oryza_sativa_japonica_Ensembl_IRGSP-1.0.tar.gz)
-mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
-tRNA database with bowtie-index (Original source:
http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Osati/Osati-tRNAs.fa)
-rRNA database with bowtie-index (Original source:
https://www.ncbi.nlm.nih.gov/nuccore)
-ensembl ncRNA database with bowtie-index (Original source:
ftp://ftp.ensemblgenomes.org/pub/plants/release-36/fasta/oryza_sativa/ncrna/)
-rfam 12.3 database with bowtie-index (Original source:
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta_files/)
```

2. SPORTS1.0 related parameters if you download recommend reference database: -g

```
/<your_defined_address>/Oryza_sativa/Ensembl/IRGSP-1.0/Sequence/BowtieIndex/genome
-m /<your_defined_address>/Oryza_sativa/miRBase_21/miRBase_21-osa
-r /<your_defined_address>/Oryza_sativa/rRNA_db/rice_rRNA
-t /<your_defined_address>/Oryza_sativa/GtRNAdb/Osati-tRNAs
-e /<your_defined_address>/Oryza_sativa/Ensembl/Oryza_sativa.IRGSP-1.0.ncrna
-f /<your_defined_address>/Oryza_sativa/Rfam_12.3/Rfam-12.3-rice
```

- 68. Arabidopsis thaliana (Arabidopsis)
  - annotation database: (We provide a download link for all databases listed below: https://ncrnainfo-my.sharepoint.com/personal/sports\_ncrna\_info/\_layouts/15/guestaccess.as px?docid=08ec312d8fc7d4211952170468dfbb202&authkey=AeVag9p4ifZJG58C5UbXstE)

```
-genome with bowtie-index (Ensembl TAIR10) (Original source: ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Arabidopsis thaliana/Ensembl/TAIR10/
```

```
Arabidopsis thaliana Ensembl TAIR10.tar.qz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Athal10/araThal-tRNAs.fa)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensemblgenomes.org/pub/plants/release-36/fasta/arabidopsis thaliana/ncrn
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -q
    /<your defined address>/Arabidopsis thaliana/Ensembl/TAIR10/Sequence/BowtieIndex/g
    enome
    -m /<your defined address>/Arabidopsis thaliana/miRBase 21/miRBase 21-ath
    -r /<your defined address>/Arabidopsis thaliana/rRNA db/Arabidopsis rRNA
    -t /<your defined address>/Arabidopsis thaliana/GtRNAdb/araThal-tRNAs
    /<your defined_address>/Arabidopsis_thaliana/Ensembl/Arabidopsis_thaliana.TAIR10.n
    -f /<your defined address>/Arabidopsis thaliana/Rfam 12.3/Rfam-12.3-arabidopsis
69. Glycine max (Soybean)

    annotation database: (We provide a download link for all databases listed below:

    https://ncrnainfo-my.sharepoint.com/personal/sports ncrna info/ layouts/15/guestaccess.as
    px?docid=079974655bb2e4f62b669c3d1701fffeb&authkey=AWVriz_LNaTMDsBjbMXDS60)
    -genome with bowtie-index (Ensembl Gm01) (Original source:
    ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Glycine max/Ensembl/Gm01/Glycine max
    Ensembl Gm01.tar.gz)
    -mirbase 21 with bowtie-index (Original source: http://www.mirbase.org/ftp.shtml)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/eukaryota/Gmax2/glyMax2-tRNAs.fa)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -rfam 12.3 database with bowtie-index (Original source:
    ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
  2. SPORTS1.0 related parameters if you download recommended reference database: -g
    /<your defined address>/Glycine max/Ensembl/Gm01/Sequence/BowtieIndex/genome
    -m /<your defined address>/Glycine max/miRBase 21/miRBase 21-gma
    -r /<your defined address>/Glycine max/rRNA db/soybean rRNA
    -t /<your defined address>/Glycine max/GtRNAdb/glyMax2-tRNAs
     -f /<your defined address>/Glycine max/Rfam 12.3/Rfam-12.3-soybean
70. Escherichia coli (E.coli)

    annotation database: (We provide a download link for all databases listed below:

    https://ncrnainfo-my.sharepoint.com/personal/sports_ncrna_info/_layouts/15/guestaccess.as
    px?docid=0645cc2a0024d41fdba5be31a17bd5374&authkey=AaR4ui2QEXqh2-SpOIxaUik)
     -genome with bowtie-index (Ensembl EB1) (Original source:
    ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/Escherichia coli K 12 DH10B/Ensembl/
    EB1/Escherichia coli K 12 DH10B Ensembl EB1.tar.gz)
    -tRNA database with bowtie-index (Original source:
    http://gtrnadb.ucsc.edu/GtRNAdb2/genomes/bacteria/Esch coli/eschColi-tRNAs.fa)
    -rRNA database with bowtie-index (Original source:
    https://www.ncbi.nlm.nih.gov/nuccore)
    -ensembl ncRNA database with bowtie-index (Original source:
    ftp://ftp.ensemblgenomes.org/pub/bacteria/release-36/fasta/bacteria 91 collection/
    escherichia coli/ncrna/)
    -rfam 12.3 database with bowtie-index (Original source:
```

```
ftp://ftp.ebi.ac.uk/pub/databases/Rfam/12.3/fasta files/)
```

2. SPORTS1.0 related parameters if you download recommended reference database: -g
 /<your\_defined\_address>/Escherichia\_coli/Ensembl/EB1/Sequence/BowtieIndex/genome
 -r /<your\_defined\_address>/Escherichia\_coli/rRNA\_db/e\_coli\_rRNA
 -t /<your\_defined\_address>/Escherichia\_coli/GtRNAdb/eschColi-tRNAs
 -e
 /<your\_defined\_address>/Escherichia\_coli/Ensembl/Escherichia\_coli.HUSEC2011CHR1.nc
 rna
 -f /<your\_defined\_address>/Escherichia\_coli/Rfam\_12.3/Rfam-12.3-e\_coli

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# History

1.0.4 - 03/25/2018 1. New function added: if '-M > 0' applied in sports.pl, statistics of mismatch information will generated to predict potential nucleic acid modification loci; 2. Figure of tRNA mapping information will generated if tRNA database is defined; 3. A sample output with the parameters '-M 1' of dataset GSM2304822 (mouse sperm) is uploaded <a href="https://example.com/here/beta/4012612">https://example.com/here/beta/4012612</a>

according to the paper " Juan Pablo Tosar, Carlos Rovira, Alfonso Cayota. Non-coding RNA fragments account for the majority of annotated piRNAs expressed in somatic non-gonadal tissues.

Communications Biology. 2018;1, Article number: 2. doi: 10.1038/s42003-017-0001-7 ".

1.0.2 - 01/22/2018 1. Fixed bugs to generate more accurate mature tRNA database.

1.0.1 - 01/13/2018 1. Fixed bugs that caused unexpected exit. 2. Optimized script to speed up tRNA

annotation process.

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service, corrections, or upgrades to the package.

**Contact information** 

Contact author: Junchao Shi

E-mail: sports.rna@gmail.com; junchao.shi@yahoo.com