



shear Documentation

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GETTING STARTED

1.1 What is SHEAR?

SHEAR (Simply Handler for Error and Adapter Removal) is a short-read trimmer for high-throughput sequencing fastq files. SHEAR first scans the fastq file(s) and automatically detects likely adapter and primer contaminants. Then it calls Scythe (<https://github.com/vsbuffalo/scythe>), which removes reads using a Bayesian error-tolerant approach that effectively removes adapters even if the adapter itself contains a sequence variation due to sequencing error. SHEAR then trims and filters reads based on minimum quality and content cutoffs. Additionally, SHEAR is designed to automate the simultaneous trimming and concatenation of multiple paired read files into a single trimmed file ready for mapping or assembly.

1.2 Requirements

- Python 2.7.x or 3.x (3.x recommended)

1.2.1 Optional

- Scythe: <https://github.com/vsbuffalo/scythe> (**Strongly recommended**)

1.3 Installation

No installation is necessary, simply clone the repository from GitHub. Scythe should be installed according to its instructions.:

```
git clone https://www.github.com/jbpease/shear
```

1.4 Preparing your data

Standard fastq files with four lines per entry (header, sequence, gap, quality) should be used. ABI solid colorspace reads are not currently supported. When using paired-end mode reads only need to be sorted and

contain the same number of reads if the `-U/--filter-unpaired` mode is used, since this will remove both reads from a pair when either of them is filtered out.

1.5 Usage

1.5.1 Paired-end

```
python shear.py --fq1 FASTQ.SAMPLE1.p1.fastq FASTQ.SAMPLE2.p1.fastq .  
↪ .. --fq2 FASTQ.SAMPLE1.p2.fastq FASTQ.SAMPLE2.p2.fastq ... --out1 ↪  
↪ FASTQ.sheared.p1.fq --out2 FASTQ.sheared.p2.fq
```

1.5.2 Single-read

```
python shear.py --fq1 FASTQ.SAMPLE1.p1.fastq FASTQ.SAMPLE2.p1.fastq .  
↪ .. --fq2 FASTQ.SAMPLE1.p2.fastq FASTQ.SAMPLE2.p2.fastq ... --out1 ↪  
↪ FASTQ.sheared.p1.fq --out2 FASTQ.sheared.p2.fq
```

1.5.3 Config file alternative

Alternatively to a full set of command line arguments you can enter a single positional argument that points in a text file. You can then specify command line arguments more neatly over several lines as in the example:

```
--fq1  
FASTQ.SAMPLE1.p1.fastq  
FASTQ.SAMPLE2.p1.fastq  
--fq2  
FASTQ.SAMPLE1.p2.fastq  
FASTQ.SAMPLE2.p2.fastq  
--out1 FASTQ.sheared.p1.fq  
--out2 FASTQ.sheared.p2.fq
```

1.6 Version History

1.6.1 v. 2017-06-21

Important fix for `--trim_pattern_3` option. Removal of the `-trim-pattern-5` option. Minor fixes to manual, readme and some parameters.

1.6.2 v. 2017-06-20

Major upgrade, fixed issues with compatibility with Scythe, added automatic adapter finding with `adapt.py`, included new manual and documentation with Sphinx. WARNING: program options have changed signifi-

cantly to be more conventional and consistent in format. Please be advised that old commands will need to be updated in terms of flag names.

1.6.3 v. 2015-09-13

Fixes for Python3 compatibility , add gzip capability

1.6.4 v. 0.007

Fixes to default parameters and option processing

1.6.5 v. 0.006

Minor fixes to default parameters

1.6.6 v. 0.005

Major update, fixed filtered output, clean-up, removed GC content filter

1.6.7 v. 0.004

Added support for combining multiple pairs of input files

1.6.8 v. 0.003

Alpha Release

PROGRAM PARAMETER DESCRIPTIONS

2.1 adapt

2.1.1 Description

This program (as part of SHEAR) searches adapter sequences and generates an adapter file for use with SHEAR/Scythe.

2.1.2 Parameters

-h/--help

Description: show this help message and exit

Type: boolean flag

--fq1 (required)

Description: one or more fastq file paths, separated by spaces

Type: file path; **Default:** None

-o/--out (required)

Description: output FASTA of adapters detected

Type: file path; **Default:** None

--fq2

Description: one or more fastq file paths separated by spaces, only use this for paired-end fastq files and enter these files in the same order as their counterparts in **-fq1**

Type: file path; **Default:** None

-k/--end-klength

Description: Length of end kmer to tabulate for possible adapter matches.

Type: integer; **Default:** 16

-m/--mode

Description: known=only use list of known adapters;endmer=search for common 3'end sequences;both=both known and endmers

Type: None; **Default:** known

Choices: ('known', 'endmer', 'both')

--quiet

Description: Suppress progress messages

Type: boolean flag

-E/--end-min-match

Description: Minimum proportion of read match required to report the endmer as a possible match.

Type: float; **Default:** 0.0001

-M/--known-min-match

Description: Minimum proportion of read match required to report the endmer as a possible match.Set to -1 (default) to accept all matches

Type: float; **Default:** -1

-N/--number-of-reads

Description: Number of reads to search in each fastq

Type: integer; **Default:** 200000

2.2 shear

2.2.1 Description

SHEAR is a read trimmer that coordinates the automatic finding of adapter sequences, removes adapters using Scythe, implements various trimming and filtering options for high-throughput short read sequences, and allows coordinated removal of paired end sequence files.

2.2.2 Parameters

`-h/--help`

Description: show this help message and exit

Type: boolean flag

`--fq1 (required)`

Description: one or more fastq file paths, separated by spaces

Type: file path; **Default:** None

`--out1 (required)`

Description: Output fastq file path. Note this is a single output file that concatenates the processed outputs from all files in `-fq1`

Type: file path; **Default:** None

`-a/--adapters`

Description: Skip adapter finding and use these adapter files. Either enter (1) a single file to use for all fastq files, (2) one file per single end file or pair of paired-end files. Adapter file(s) should be in FASTA format.

Type: file path; **Default:** None

`--clean-header`

Description: removes any additional terms from the header line (useful after STAR)

Type: boolean flag

`-f/--trim-fixed`

Description: Trim a fixed number of bases from the FRONT:END of the sequence (NOT recommended).

Type: None; **Default:** 0:0

`--filt1`

Description: Output fastq file path for sequences that were filtered out. Note this is a single output file that concatenates the rejected outputs from all files in `-fq1`. Default is `[-out1]_filtered_1.fastq`

Type: file path; **Default:** None

`--filt2`

Description: Output fastq file path for sequences that were filtered out. Note this is a single output file that concatenates the rejected outputs from all files in `-fq2`. Default is `[-out2]_filtered_2.fastq`

Type: file path; **Default:** None

`--fq2`

Description: one or more fastq file paths separated by spaces, only use this for paired-end fastq files and enter these files in the same order as their counterparts in `-fq1`

Type: file path; **Default:** None

`-k/--adapter-end-klength`

Description: (Adapter finding) Length of 3'-end kmer to tabulate for possible adapter matches.

Type: integer; **Default:** 16

`--log-path`

Description: Manually specify log file path, default is 'shear_TIMESTAMP'

Type: file path; **Default:** None

`-m/--adapter-mode`

Description: (Adapter finding) known=only use list of known adapters;endmer=search for common 3' end sequences;both=both known and endmers

Type: None; **Default:** known

Choices: ('known', 'endmer', 'both')

`-n/--retain-ambig`

Description: By default ambiguous nucleotides (N) are removed from both ends of each read. If this flag is specified, N's are retained.

Type: boolean flag

`--out2`

Description: output fastq file path. Note this is a single output file that concatenates the processed outputs from all files in `-fq2`

Type: file path; **Default:** None

-p/--trim-poly

Description: Trim poly-A or poly-T repeats of at least this length from the front or end.

Type: integer; **Default:** 12

-q/--trim-qual

Description: Trim bases below this quality score from the FRONT:END of each read.

Type: None; **Default:** 20:20

--quality-scale

Description: Quality scale is usually automatically determined, but use this to set manually.

Type: None; **Default:** None

Choices: ('sanger', 'illumina', 'phred', 'solexa')

--quiet

Description: Suppress progress messages

Type: boolean flag

--retain-temp

Description: Retain temporary files (none=remove all; tempfastq=remove temporary fastq files from scythe; exceptadapters=remove temp fastq and log files from Scythe but keep adapter file; all=keep all temporary file)

Type: None; **Default:** none

Choices: ['none', 'tempfastq', 'exceptadapters', 'all']

-s/--scythe-prior

Description: Bayesian prior for proportion of adapters expected to be sampled in Scythe.

Type: float; **Default:** 0.1

--scythe-match

Description: Minimum number of bases required for a match in Scythe.

Type: integer; **Default:** 5

--scythe-skip

Description: Skip scythe 3' adapter removal.

Type: boolean flag

-t/--platform

Description: Sequencing Platform

Type: None; **Default:** TruSeq

Choices: ('TruSeq', 'TruSeqDualIndex')

--temp-dir

Description: directory to use for temporary files

Type: file path; **Default:** .

--trim-qual-pad

Description: Trim additional bases next to low-quality bases specified by --trim-qual from the FRONT:END of each read.

Type: None; **Default:** 0:0

-z/--trim-pattern-3

Description: Comma-separated list of specific sequences to trim from the 3' end. Can be used for extra stringent adapter trimming.

Type: None; **Default:** None

-A/--filter-ambig

Description: Filter reads with more than this number of ambiguous nucleotides (N's; set as 0 to skip

Type: integer; **Default:** 5

-E/--adapter-end-min-match

Description: (Adapter finding) Minimum proportion of read matches required to report the 3'-end-mer as a possible match.

Type: float; **Default:** 0.0001

-I/--filter-low-info

Description: Filter out reads with mutual information scores exceeding this value (ADVANCED, removes highly repetitive reads).

Type: float; **Default:** 0.0

-L/--filter-length

Description: Filter out reads that contain fewer than this many characters after trimming.

Type: integer; **Default:** 30

-M/--adapter-known-min-match

Description: (Adapter finding) Minimum proportion of read matches required to report a known contaminant as a possible match. Use -1 (default) to accept all matches.

Type: float; **Default:** -1

-N/--adapter-number-of-reads

Description: (Adapter finding) Number of reads to search in each fastq

Type: integer; **Default:** 200000

-Q/--filter-quality

Description: Filter out reads with a mean quality score below this value (before trimming).

Type: integer; **Default:** 3

-U/--filter-unpaired

Description: If either read in a read pair is filtered out, the counterpart reads is also filtered out regardless of quality.

Type: boolean flag

-X/--scythe-executable

Description: Set the path of the scythe executable manually.

Type: None; **Default:** scythe

INDICES AND TABLES

- genindex
- modindex
- search