

# SMALT Manual

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Version 0.7.4

## Abstract

*SMALT* is a pairwise sequence alignment program for the efficient mapping of DNA sequencing reads onto genomic reference sequences. It uses a combination of short-word hashing and dynamic programming. Most types of sequencing platforms are supported including paired-end sequencing reads.

## 1 Synopsis

*smalt* *TASK* [**OPTIONS**] [*INDEX SEQFIL-A* [*SEQFIL-B*]]

### Available tasks

*smalt index* [**INDEX-OPTIONS**] *INDEX REFSEQ-FILE*

builds a hash index of k-mer words in the reference sequences and stores it on disk. Two files are written to disk: *INDEX.smi*

*smalt map* [**MAP-OPTIONS**] *INDEX READ-FILE* [*MATE-FILE*]

loads the index into memory and aligns single or paired-end reads against the reference sequences.

*smalt sample* [**SAMPLE-OPTIONS**] *INDEX READ-FILE* [*MATE-FILE*]

samples distribution of insert lengths for paired reads. A subset of the pairs is aligned with a reference using the index.

*smalt check* *READ-FILE* [*MATE-FILE*]

checks whether read input files conform to FASTA/FASTQ format specification.

*smalt help*

for a brief summary of this software.

*smalt version*

prints version information.

### Help on individual tasks

*smalt TASK -H*

e.g. *smalt index -H* for help on options influencing the generation of the hash index.

## 2 Description

Running *SMALT* involves two steps. First, an index of short words has to be built (smalt index). Then the sequencing reads are mapped onto the reference (smalt map).

*SMALT* uses a hash table of words of fixed-length sampled along the genomic reference sequence in the file *REFSEQ-FILE* at equidistant steps. The sequencing reads in the file *READ-FILE* (and *MATE-FILE*) are then mapped against the genomic reference sequences one-by-one.

First, exactly matching seeds are identified in the reference sequences by looking up the *k*-mer words of the read in the hash index. Based on these seeds, potentially matching sequence segments are selected for alignment by a Smith-Waterman algorithm.

## 3 Options

### 3.1 INDEX-OPTIONS

- k** *wordlen* Sets the length of the hashed words. *wordlen* is an integer with  $2 < \textit{wordlen} \leq 20$  (default: 13).
- s** *skipstep* Sampling step size, i.e. the distance between successive words that are hashed along the genomic reference sequence. With the option **-s 1** every word is hashed, with **-s 2** every second word, with **-s 3** every third etc. By default *skipstep* is set equal to *wordlen*.

### 3.2 MAP-OPTIONS

- a** When this flag is set, explicit alignments are output along with the mappings.
- c** *mincover* Only consider mappings where the *k*-mer word seeds cover the query read to a minimum extent. If *mincover* is an integer or floating point value  $> 1.0$ , at least this many bases of the read must be covered by *k*-mer word seeds. If *mincover* is a floating point value  $\leq 1.0$ , it specifies the fraction of the query read length that must be covered by *k*-mer word seeds. This option can be used only when the **-x** flag is also set.
- d** *scorediff* Set a threshold of the Smith-Waterman alignment score relative to the maximum score. When mapping single reads, all alignments resulting in Smith-Waterman scores within *scorediff* of the maximum are reported. Mappings with scores lower than this value are skipped. If *scorediff* is set to a value  $< 0$ , no threshold is set and all alignments are reported (possibly limited by the **-m** *minscor* and **-y** *minid* options).

For paired reads, only a value of 0 is supported. With the option **-d 0** all alignments or alignment pairings with the best score are printed. For reads that have multiple best mappings the default is to select a read/pair at random depending on the option **-r**. With **-r -1** reads (mate pairs) with multiple best mappings are reported as 'not mapped'.

**-f *format*** Specifies the output format. *format* can be one of the following strings:

***bam*** (<http://samtools.sourceforge.net>)

Optional extensions: *bam:x,clip* (see *sam* output format). The bam header is always written.

***cigar*** Compact Idiosyncratic Gapped Alignment Report

(see <http://www.sanger.ac.uk/resources/software/ssaha2>)

***gff*** General Feature Format version 2

(see <http://www.sanger.ac.uk/resources/software/gff/spec.html>)

***sam*** (default) Sequence Alignment/Map format (<http://samtools.sourceforge.net>)

Optional extension: *sam* followed by a colon and one or more comma separated keywords, e.g. *sam:nohead,x,clip*:

***nohead*** for output without the SAM header

***clip*** for hard clipped sequences

***x*** for CIGAR alignment strings containing the letter 'X' to indicate a mismatch

No spaces are allowed in the format string. By default, sequences are soft clipped and, when writing to a file, the header is present.

***samsoft*** like *sam* (for backwards compatibility)

***ssaha*** native output format of the SSAHA2 software package

(<http://www.sanger.ac.uk/resources/software/ssaha2>)

**-F *inform*** Specifies the input format. *inform* can be either 'fastq' (default), 'sam' or 'bam' (see: [samtools.sourceforge.net](http://samtools.sourceforge.net)). SAM/BAM input formats require additional libraries to be installed and may generate temporary files. A directory for those temporary files may be specified with the option **-T**.

**-g *insfil*** Use the distribution of insert sizes stored in the file *insfil*. This file is in ASCII format and can be generated using the *smalt sample* task.

**-H** Print instructions on screen.

**-i *insertmax*** Maximum insert size for paired-end reads. *insertmax* is a positive integer (default 500).

**-j *insertmin*** Minimum insert size for paired-end reads *insertmax* is a positive integer (default 0).

- l *pairtyp* Type of read pair library used. This specifies the expected relative orientation of mate-pairs. *pairtyp* can be either *pe*, i.e. for the Illumina paired-end library for short inserts (orientation:  $\rightarrow \leftarrow$ ), *mp* for the Illumina mate pair library (long inserts,  $\leftarrow \rightarrow$ ) or *pp* when both mates are on the same strand ( $\rightarrow \rightarrow$ ) as in 454 read-pair libraries.
- m *minscor* Sets an absolute threshold of the Smith-Waterman scores. Mappings with scores below that threshold will not be reported. *minscor* is a positive integer (default *minscor* = *wordlen* + *skipstep* - 1).
- n *nthreads* Run *SMALT* using multiple threads. *nthread* is the number of additional threads forked for mapping. The order of the reads is not preserved for the output unless the flag -O is specified.
- o *oufilnam* Write mapping output (e.g. SAM lines) to a separate file named *oufilnam*. If this option is not specified, mappings are written to standard output together with other messages.
- O Output mappings the order of the reads in the input files when using multiple threads (option -n *nthreads*).
- p Report partial alignments if they are complementary on the query read (split or chimeric reads). A maximum of two partial alignments are output per read. The second alignment is labelled 'P' ('-f ssaha' or '-f cigar' formats) or has the 'secondary alignment' bit-flag (0x100) of the SAM FLAG field raised ('-f sam' or '-f samsoft').
- q *minbasq* Sets a base quality threshold  $0 \leq \text{minbasq} \leq 10$  (default *minbasq* = 0). k-mer words of the read with base pairs that have a base quality below this threshold are not looked up in the hash index.
- r *seed* Determines how reads or mate pairs with multiple best mappings are reported. If *seed*  $\geq 0$  report an alignment (pair) selected at random. With *seed* = 0 (default) a seed is derived from the current calendar time. If *seed* < 0 reads with multiple best mappings are reported as 'not mapped'.
- S *scorspec* Specify alignment penalty scores for a match or mismatch (substitution), or for opening or extending a gap. *scorspec* is a comma separated list of integer assignments to one or more of the following variables: match, subst, gapopen, gapext. *scorspec* must not contain spaces. Example: -S *gapopen=-5,gapext=-4*. Default:-S *match=1,subst=-2,gapopen=-4,gapext=-3*.
- T *tmpdir* Write temporary files to directory *tmpdir* (used when reading sequence input in SAM/BAM format).
- w Output complexity weighted Smith-Waterman scores.
- x This flag triggers a more exhaustive search for alignments at the cost of decreased speed. In paired-end mode each mate is mapped independently. (By default the mate with fewer hits in the hash index is mapped first and the vicinity is searched for its mate.)

**-y *minid*** Filters output alignments by a threshold in the number of exactly matching nucleotides. If  $0 \leq \textit{minid} \leq 1$  denotes the number of matching bases as a fraction of the read length. *minid* = 1 labels as 'unmapped' all reads that do not have an exact match.

### 3.3 SAMPLE-OPTIONS

**-m *minscor*** Sets an absolute threshold of the Smith-Waterman scores. Mappings with scores below that threshold will not be used for the distribution of insert lengths.

**-n *nthreads*** Run *SMALT* using multiple threads.

**-o *oufilnam*** Write output to a separate file named *oufilnam* rather than standard output.

**-q *minbasq*** Sets a base quality threshold  $0 \leq \textit{minbasq} \leq 10$  (default *minbasq* = 0). k-mer words of the read with base pairs that have a base quality below this threshold are not looked up in the hash index.

**-u *nreads*** Map only every *nreads*-th read pair (default 100).

## 4 Input files and formats

The reference sequence file *REFSEQ-FILE* has to be in FASTA or FASTQ format (see [http://en.wikipedia.org/wiki/FASTQ\\_format](http://en.wikipedia.org/wiki/FASTQ_format)). Variations of the FASTQ format are explained in <http://maq.sourceforge.net/fastq.shtml>.

The sequencing read file *READ-FILE* can be in FASTA/FASTQ (default) or, when **-F *bam*** or **-F *sam*** is specified, in SAM/BAM format (see <http://samtools.sourceforge.net>).

If *MATE-FILE* is present on the command line, reads are to be mapped in pairs and the mate files *READ-FILE* and *MATE-FILE* are in FASTA or FASTQ format. Corresponding mates are identified by the position in the respective FASTA/FASTQ file. Read names are *not* checked by the software to see if the respective mates correspond. It is up to the user to make sure the mates of the *i*-th read are the *i*-th sequences in the FASTQ files. As a consequence paired and unpaired reads cannot be mixed in FASTA/FASTQ input files.

*SMALT* can read FASTA/FASTQ files that are compressed using gzip (<http://www.gzip.org>).

### 4.1 Unix pipes and reading from standard input

A hyphen-minus ('-') for *READ-FILE* reads sequence data from standard input.

*SMALT* with FASTQ paired-read input can be integrated in a UNIX pipeline using named pipes (FIFOs). For example, to have *SMALT* read paired sequencing reads from gzipped FASTQ files, *mate\_1.fq.gz* and *mate\_2.fq.gz*, one can have *gzip* pipe its output into two FIFOs:

```
> mkfifo pipe_1
> mkfifo pipe_2
> smalt map -o output.sam test pipe_1 pipe_2 &
> gzip -dc mate_1.fq.gz > pipe_1 &
> gzip -dc mate_2.fq.gz > pipe_2 &
```

## 5 Note on Paired Reads

The **-i** and **-j** options specify the expected insert size range influence the way in which the mates are aligned. The mate with fewer hits in the hash index is mapped first and the vicinity defined by the expected range is searched for its mate. If the **-x** option is specified both mates are aligned independently.

In some output formats the reads are labelled or flagged, e.g. as a *proper* pair in the SAM format, with respect to the insert size range. But the alignments of all mates will be reported regardless of the range specified with the **-i** and **-j** options.

### 5.1 Definition of 'proper' pairs

By default, the mates of a read pair are in *proper* orientation when they map to opposite strands with the 5'-ends on the outside of the double stranded segment spanned by the pair as expected from the Illumina paired-end (PE) libraries with short insert lengths. A *proper* pair has both mates mapped in *proper* orientation within the expected insert range (specified with the **-i** and **-j** options).

Paired reads from the Illumina mate-pair (MP) libraries for long insert lengths will have the 3' ends on the outside of the segment spanned by the pair. Signalling this to the program *via* the **-l mp** option is important for correct labelling, e.g. of 'proper' pairs, and also for assigning the correct mapping quality scores.

### 5.2 Definition of insert size

With version 0.7.1 *SMALT* adopts definition of insert sizes according to SAM/BAM specification versions 1.3 onwards. In earlier *SMALT* versions the insert size of a read pair refers to the distance between the 5'-ends of the mapped reads (in accordance with early SAM/BAM specifications).

### 5.3 Extension of the CIGAR output format

The CIGAR output format (option **-f cigar**) produced by *smalt* comprises a label after the GIGAR tag (e.g. label A in GIGAR:A:51). The labels have the following meaning:

- A** mates are in *proper* orientation within the limits specified by the **-i** and **-j** options.
- B** mates in *proper* orientation outside the limits specified by the **-i** and **-j** options but on the same reference sequence (i.e. chromosome or contig).
- C** mates are not in *proper* orientations but on the same chromosome or contig.
- D** mates are mapped to different chromosomes or contigs.
- N** read could not be mapped.
- P** Alignment is the second partial alignment of a split (chimaeric) read (only with **-p** flag).
- R** Read is reported as 'not mapped' because there were multiple possible mappings, but no random assignment was made.
- S** Read was mapped as a single read (sole mapped read of a pair).

## 6 Note on Smith-Waterman Scores

*SMALT* uses 'standard' Smith-Waterman scores:

match: +1; mismatch: -2; gap opening: -4; gap extension: -3.

There is currently no way for the user to modify these settings.

The options **-d scordiff** and **-m minscor** which determine how many alignments are reported, are based on Smith-Waterman scores rather than e.g. edit distance. Calculating *scordiff* from the edit distance is simple if *SMALT* is run without the **-w** flag.

## 7 Memory Requirements

The memory footprint of *SMALT* is determined primarily by the total number  $N$  of base pairs of the genomic reference sequences and by the word length  $k$  (option **-k k**) and the sampling step  $s$  (option **-s s**) with which the hash index is generated. The index itself occupies less than approx.  $4(N/s + \min(4^k, 4^{13} + N/s))$  bytes of memory (disk space). The genomic reference sequences occupy approx.  $2N/5$  bytes.

For example an index of words of length 13 sampled at every 6<sup>th</sup> position (options **-k 13 -s 6**) from the human genome ( $N = 3 \times 10^9$ ) requires 3.3 Gb of memory. An index of the human genome built with options **-k 13 -s 13** (default) requires 2.3 GB. The recommended setting for 100 bp Illumina reads, **-k 20 -s 13**, requires 3.8 GB.

## 8 Index Files

The command  
*smalt index [-k k] [-s s]INDEX REFSEQ-FILE*  
writes 2 files to disk:

**INDEX.sma** Compressed set of reference sequences for which the hash table of k-mer words was generated.  $N * 2/5$  bytes where  $N$  is the total number of base pairs of the genomic reference sequences.

**INDEX.smi** The actual hash index. The file size is about  $4(N/s + \min(4^k, 4^{13+N/s}))$  bytes.

## 9 Sequence File Formats

Sequence input files are expected in (possibly gzipped) FASTA/FASTQ format (see [http://en.wikipedia.org/wiki/FASTQ\\_format](http://en.wikipedia.org/wiki/FASTQ_format)) or in SAM/BAM format (see <http://samtools.sourceforge.net>).

Variations of the FASTQ format are explained in <http://maq.sourceforge.net/fastq.shtml>.

## 10 Version

Version: 0.7.4 of March 19, 2014.

## 11 License and Copyright

**Copyright** © 2010-2013 Genome Research Limited.

**License** The source code will be made available eventually under the GNU General Public License (<http://www.gnu.org/licenses/>).

## 12 Authors

SMALT was written by Hannes Ponstigl [[hp3@sanger.ac.uk](mailto:hp3@sanger.ac.uk)] at the Wellcome Trust Sanger Institute, Cambridge, UK in 2010.

## 13 Examples

### 13.1 Paired-end Illumina-Solexa reads, human genome

#### 13.1.1 Longer reads ( $\approx 100$ bp)

The insert size be 300bp, the reads provided in two FASTQ files: `mate1.fq` contains the 1<sup>st</sup> and `mate2.fq` the 2<sup>nd</sup> read of each pair. The human chromosome

sequences be in the FASTA file `GRCh37.fa`.

**Build the hash index:** `smalt index -k 20 -s 13 hs37k20s13 GRCh37.fa`  
This writes an index file `hs37k20s13.smi` of 2.7 GB and a sequence file `hs37k20s13.sma` of 1.2 GB to the disk using 4.0 GB of memory.

**Map the reads:** `smalt map -f samsoft -o hs37map.sam hs37k13s13 mate1.fq mate2.fq`  
This writes a file `hs37map.sam` with alignments in SAM format using 'soft clipping' which retains the entire read sequence.

### 13.1.2 short reads (36 bp)

Because of the short read length, the hash index should be built with a smaller sampling step size, for example `-s 3` or `-s 2`. Larger step sizes would result in reduced sensitivity and increased error rate.

**Build the hash index:** `smalt index -s 3 hs37k13s3 GRCh37.fa`  
This writes an index file `hs37k13s3.smi` of 3.8 GB and a sequence file `hs37k13s3.sma` of 1.2 GB to the disk. The memory footprint is 5.3 MB.

**Map the reads:** `smalt map -o hs37map.cig hs36k13s3 mate1.fq mate2.fq`  
This writes a file `hs37map.cig` with mappings in CIGAR format.

## 13.2 Single Roche-454 reads (human)

**Build the hash index:** `smalt index -s 4 hs37k13s4 GRCh37.fa`  
This writes an index file `hs37k13s4.smi` of 3.0 GB and a sequence file `hs37k13s4.sma` of 1.1 GB to the disk using 4.4 GB of memory.

**Map the reads:** `smalt map -f ssaha -o hs37map.ssaha hs36k13s4 reads.fq`  
This writes a file `hs37map.ssaha` with alignments in SSAHA2 native format.

## 13.3 Single Illumina-Solexa reads (bacterial genome)

Map 76 bp single reads (FASTQ file `reads.fq` of the bacterium *S. suis* (FASTA file `suis.fa`).

For small genomes, one can often afford using the most sensitive settings for the hash index, i.e. `-s 1`, and possibly reduce the word length, e.g. `-k 11`.

**Build the hash index:** `smalt index -k 11 -s 1 suisk11s1 suis.fa`

**Map the reads:** `smalt map suisk11s1 reads.fq`  
This writes a CIGAR lines to standard output.

platform	genome	length	options	memory
Illumina	<i>H. sapiens</i>	108 bp	-k 20 -s 13	3.8 GB
Illumina	<i>H. sapiens</i>	76 bp	-k 13 -s 6	3.3 GB
Illumina	<i>H. sapiens</i>	54 bp	-k 13 -s 4	4.3 GB
Illumina	<i>H. sapiens</i>	36 bp	-k 13 -s 3	5.3 GB
Illumina	<i>C. elegans</i>	76 bp	-k 13 -s 4	
Illumina	<i>P. falciparum</i>	72 bp	-k 13 -s 2	
Illumina	<i>C. suis</i>	72 bp	-k 13 -s 2	
Roche-454	<i>H. sapiens</i>	200 bp	-k 13 -s 4	4.3 GB
Capillary	<i>H. sapiens</i>	500 bp	-k 13 -s 4	4.3 GB
small RNAs	<i>H. sapiens</i>	$\geq 12$ bp	-k 11 -s 2	7.3 GB

## 14 Tuning performance

By tuning two parameters, the word length (`-k wordlen`) and the step size (`-s stepsiz`) with which the index is built, one can trade sensitivity and accuracy against speed and memory efficiency.

A necessary condition for a read to register a match on a segment of the genomic reference is that there be at least one contiguous stretch of *wordlen* identical nucleotides between the two sequences.

This is, however, not a sufficient criterion. Because hashed words are sampled only every *stepsiz* base pairs along the reference, any particular word in the sequencing read may be missed. But even when there is a contiguous segment of  $\text{wordlen} + \text{stepsiz} - 1$  identical nucleotide between the sequences, a match may be missed on rare occasions because, depending on the command line flags, heuristics are employed to speed up program execution.

Generally *wordlen* should be set to 13 (the default) but can be increased to 20 for Illumina reads of  $> 100$ bp length or reduced to 11 for very short query sequences of 11 – 24 base pairs. The choice of *stepsiz* is more critical and depends on the available computer memory, on the size of the genome and the expected degree of variation between the sequenced genome and the reference, as well as on the sequencing platform with its inherent sequencing error profile.

The table is intended as a guideline for economical choices of `-s stepsiz` for a range of scenarios. Reducing *stepsiz* results in lower error rates at the cost of a reciprocal increase of the memory footprint and, particularly with the `-x` flag, of reduced execution speeds.