

**NAME**

vsearch — chimera detection, clustering, dereplication, masking, pairwise alignment, searching, shuffling and sorting of amplicons from metagenomic projects.

**SYNOPSIS**

Chimera detection:

**vsearch** --uchime\_denovo *fastafile* (--chimeras | --nonchimeras | --uchimealns | --uchimeout) *outputfile* [*options*]

**vsearch** --uchime\_ref *fastafile* (--chimeras | --nonchimeras | --uchimealns | --uchimeout) *outputfile* --db *fastafile* [*options*]

Clustering:

**vsearch** (--cluster\_fast | --cluster\_size | --cluster\_smallmem) *fastafile* (--alnout | --blast6out | --centroids | --clusters | --msaout | --uc | --userout) *outputfile* --id *real* [*options*]

Dereplication:

**vsearch** --derep\_fulllength *fastafile* (--output | --uc) *outputfile* [*options*]

Masking:

**vsearch** --maskfasta *fastafile* --output *outputfile* [*options*]

Pairwise alignment:

**vsearch** --allpairs\_global *fastafile* (--alnout | --blast6out | --matched | --notmatched | --uc | --userout) *outputfile* (--acceptall | --id *real*) [*options*]

Searching:

**vsearch** --usearch\_global *fastafile* --db *fastafile* (--alnout | --blast6out | --uc | --userout) *outputfile* --id *real* [*options*]

Shuffling:

**vsearch** --shuffle *fastafile* --output *outputfile* [*options*]

Sorting:

**vsearch** (--sortbylength | --sortbysize) *fastafile* --output *outputfile* [*options*]

**DESCRIPTION**

Environmental or clinical molecular studies generate large volumes of amplicons (e.g. SSU-rRNA sequences) that need to be dereplicated, masked, sorted, searched, clustered or compared to reference sequences. The aim of **vsearch** is to offer a all-in-one open source tool to perform these tasks, using optimized algorithm implementations and harvesting the full potential of modern computers, thus providing fast and accurate data processing.

Nucleotide sequence comparisons is at the core of **vsearch**. To speed up comparisons, **vsearch** implements an extremely fast implementation of the Needleman-Wunsch algorithm, making use of the Streaming SIMD Extensions (SSE2) of modern x86-64 CPUs. If SSE2 instructions are not available, **vsearch** exits with an error message. For comparisons involving sequences longer than 5,000 nucleotides, **vsearch** uses a slower alignment method with smaller memory requirements.

**Input**

**vsearch** input is a fasta file containing one or several nucleotide sequences. For each sequence, the sequence identifier is defined as the string comprised between the ">" symbol and the first space, or the end of the line, whichever comes first. Additionally, if the line starts with ">[:size=integer;label]", contains ">label;size=integer;label" or ends with ">label;size=integer[:]", **vsearch** will remove the pattern [:size=integer[:]] from the header and interpret *integer* as the number of occurrences (or abundance) of the sequence in the study. That abundance information is used or created during chimera detection, dereplication, sorting and searching.

The nucleotide sequence is defined as a string of IUPAC symbols (ACGTURYSWKMDBHVN), starting after the end of the identifier line and ending before the next identifier line, or the file end. **vsearch** silently ignores ascii characters 9 to 13, and exits with an error message if ascii characters 0 to 8, 14 to 31, "." or "-" are present. All other ascii or non-ascii characters are stripped and complained about in a non-blocking

warning message.

**vsearch** operations are case insensitive, except when soft masking is activated. For `--usearch_global` (searching), `--cluster_fast` and `--cluster_smallmem` (clustering), and `--maskfasta` (masking) commands, the case is important if soft masking is used. Soft masking is specified with the options `--dbmask soft` (for searching) or `--qmask soft` (for searching, clustering and masking). When using soft masking, lower case letters indicate masked symbols, while upper case letters indicate regular symbols. Masked symbols are never included in the unique *k*-mers used in searching. When soft masking is not activated, all letters are converted to upper case internally and used in result files.

When comparing sequences during chimera detection, dereplication, searching and clustering, T and U are considered identical, regardless of their case. If two symbols are non-identical, their alignment will result in the negative mismatch score (default -4), except if one or both of the symbols are ambiguous (RYSWK-MDBHVN) in which case the score is zero. Alignment of two identical ambiguous symbols (e.g. R vs R) also receives a score of zero.

Optionally, **vsearch** can be compiled to accepted compressed fasta files as input (gz and bzip2 formats).

## Options

**vsearch** recognizes a large number of command-line options. For easier navigation, options are grouped below by theme (chimera detection, clustering, dereplication, masking, shuffling, sorting, and searching). We start with general options that apply to all themes.

General options:

- help** Display a short help and exit.
- version** Output version information and exit.
- fasta\_width** *positive integer*  
Fasta files produced by **vsearch** are wrapped (sequences are written on lines of *integer* nucleotides, 80 by default). Set that value to 0 to eliminate the wrapping.
- maxseqlength** *positive integer*  
All **vsearch** operations will discard sequences of length equal or greater than *integer* (50,000 nucleotides by default).
- minseqlength** *positive integer*  
All **vsearch** operations will discard sequences of length smaller than *integer* (1 nucleotide by default for sorting or shuffling, 32 nucleotides for clustering, dereplication or searching).
- notrunclabels**  
Do not truncate sequence labels at first space, use the full header in output files.

Chimera detection options:

Chimera detection is based on a scoring function controlled by five options (`--dn`, `--mindiffs`, `--mindiv`, `--minh`, `--xn`). Sequences are first sorted by decreasing abundance (if available), and compared on their *plus* strand only (case insensitive).

In *de novo* mode, input fasta file should present abundance annotations (pattern `[;]size=integer[;]` in the fasta header). The input order influences the chimera detection, we recommend to sort sequences by decreasing abundance (default of `--derep_fulllength` command). If your sequence set needs to be sorted, please see the `--sortbysize` command in the sorting section.

**--abskew** *real*

When using `--uchime_denovo`, the abundance skew is used to distinguish in a 3-way alignment which sequence is the chimera and which are the parents. The assumption is that chimeras appeared later in the PCR amplification process and are therefore less abundant than their parents. The default value is 2.0, which means that the parents should be at least 2 times more abundant than their chimera. Any positive value greater than 1.0 can be used.

- alignwidth** *positive integer*  
Width of 3-way alignments in `--uchimealns` output. The default value is 80. Set to 0 to eliminate wrapping.
- chimeras** *filename*  
Output chimeric sequences to *filename*, in fasta format. Output order may vary when using multiple threads.
- db** *filename*  
When using `--uchime_ref`, detect chimeras using the fasta-formatted reference sequences contained in *filename*. Reference sequences are assumed to be chimera-free. Chimeras will not be detected if their parents (or sufficiently close relatives) are not present in the database.
- dn** *real* No vote pseudo-count (parameter *n* in the chimera scoring function) (1.4).
- mindiffs** *positive integer*  
Minimum number of differences per segment (3).
- mindiv** *real*  
Minimum divergence from closest parent (0.8).
- minh** *real*  
Minimum score (h). Increasing this value tends to reduce the number of false positives and to decrease sensitivity. Default value is 0.28. (value ranging from 0.0 to 1.0 included).
- nonchimeras** *filename*  
Output non-chimeric sequences to *filename*, in fasta format. Output order may vary when using multiple threads.
- self** When using `--uchime_ref`, ignore a reference sequence when its label matches the label of the query sequence (useful to estimate false-positive rate in reference sequences).
- selfid** When using `--uchime_ref`, ignore a reference sequence when its nucleotide sequence is strictly identical with the query sequence.
- threads** *positive integer*  
Number of computation threads to use (1 to 256) with `uchime_ref`. The number of threads should be lesser or equal to the number of available CPU cores. The default is to launch one thread per available logical core.
- uchime\_denovo** *filename*  
Detect chimeras present in the fasta-formatted *filename*, without external references (i.e. *de novo*). Automatically sort the sequences in *filename* by decreasing abundance beforehand. Multithreading is not supported.
- uchime\_ref** *filename*  
Detect chimeras present in the fasta-formatted *filename* by comparing them with reference sequences (option `--db`). Multithreading is supported.
- uchimealns** *filename*  
Write 3-way global alignments (parentA, parentB, chimera) to *filename* using a human-readable format. Use `--alignwidth` to modify alignment length. Output order may vary when using multiple threads.
- uchimeout** *filename*  
Write chimera detection results to *filename* using the `uchime` tab-separated format of 18 fields (see the list below). Use `--uchimeout5` to use a format compatible with `usearch v5` and earlier versions. Rows output order may vary when using multiple threads.
1. score: higher score means a more likely chimeric alignment.

2. Q: query sequence label.
3. A: parent A sequence label.
4. B: parent B sequence label.
5. T: top parent sequence label (i.e. parent most similar to the query). That field is removed when using `--uchimeout5`.
6. idQM: percentage of similarity of query (Q) and model (M) constructed as a part of parent A and a part of parent B.
7. idQA: percentage of similarity of query (Q) and parent A.
8. idQB: percentage of similarity of query (Q) and parent B.
9. idAB: percentage of similarity of parent A and parent B.
10. idQT: percentage of similarity of query (Q) and top parent (T).
11. LY: yes votes in the left part of the model.
12. LN: no votes in the left part of the model.
13. LA: abstain votes in the left part of the model.
14. RY: yes votes in the right part of the model.
15. RN: no votes in the right part of the model.
16. RA: abstain votes in the right part of the model.
17. div: divergence, defined as (idQM - idQT).
18. YN: query is chimeric (Y), or not (N), or is a borderline case (?).

**--uchimeout5**

When using `--uchimeout`, write chimera detection results using a tab-separated format of 17 fields (drop the 5th field of `--uchimeout`), compatible with usearch version 5 and earlier versions.

**--xn** *real* No vote weight (parameter beta) (8.0).

Clustering options:

**vsearch** implements a single-pass, greedy star-clustering algorithm, similar to the algorithms implemented in usearch, DNAClust and sumacust. Important parameters are the global clustering threshold (`--id`) and the pairwise identity definition (`--iddef`).

**--centroids** *filename*

Output cluster centroid sequences to *filename* file, in fasta format. The centroid is the sequence that seeded the cluster (i.e. the first sequence of the cluster).

**--cluster\_fast** *filename*

Clusterize the fasta sequences in *filename*, automatically perform a sorting by decreasing sequence length beforehand.

**--cluster\_size** *filename*

Clusterize the fasta sequences in *filename*, automatically perform a sorting by decreasing sequence abundance beforehand.

**--cluster\_smallmem** *filename*

Clusterize the fasta sequences in *filename* without automatically modifying their order beforehand. Sequence are expected to be sorted by decreasing sequence length, unless `--usersort` is used.

**--clusters** *string*

Output each cluster to a separate fasta file using the prefix *string* and a ticker (0, 1, 2, etc.) to construct the path and filenames.

- consout** *filename*  
Output cluster consensus sequences to *filename*. For each cluster, a multiple alignment is computed, and a consensus sequence is constructed by taking the majority symbol (nucleotide or gap) from each column of the alignment. Columns containing a majority of gaps are skipped, except for terminal gaps. Use **--construncate** to take terminal gaps into account (not implemented yet).
- id** *real*  
Do not add the target to the cluster if the pairwise identity with the centroid is lower than *real* (value ranging from 0.0 to 1.0 included). The pairwise identity is defined as the number of (matching columns) / (alignment length - terminal gaps). That definition can be modified by **--iddef**.
- iddef** 0/1/2/3/4  
Change the pairwise identity definition used in **--id**. Values accepted are:
0. CD-HIT definition using shortest sequence as numerator.
  1. edit distance: (matching columns) / (alignment length).
  2. edit distance excluding terminal gaps (default value).
  3. Marine Biological Lab definition counting each extended gap as a single difference.
  4. BLAST definition, equivalent to **--iddef** 2 in a context of global pairwise alignment.
- msaout** *filename*  
Output a multiple sequence alignment and a consensus sequence for each cluster to *filename*, in fasta format. The consensus sequence is constructed by taking the majority symbol (nucleotide or gap) from each column of the alignment. Columns containing a majority of gaps are skipped, except for terminal gaps.
- qmask** *none/dust/soft*  
Mask simple repeats and low-complexity regions in sequences using the *dust* or the *soft* algorithms, or do not mask (*none*). Warning, when using *soft* masking, clustering becomes case sensitive. The default is to mask using *dust*.
- sizein**  
Take into account the abundance annotations present in the input fasta file (search for the pattern "[>]size=*integer*[:]" in sequence headers).
- sizeout**  
Add abundance annotations to the output fasta files (add the pattern ";size=*integer*;" to sequence headers). If **--sizein** is specified, abundance annotations are reported to output files, and each cluster centroid receives a new abundance value corresponding to the total abundance of the amplicons included in the cluster (**--centroids** option). If **--sizein** is not specified, input abundances are set to 1 for amplicons, and to the number of amplicons per cluster for centroids.
- strand** *plus/both*  
When comparing sequences with the cluster seed, check the *plus* strand only (default) or check *both* strands.
- threads** *positive integer*  
Number of computation threads to use (1 to 256). The number of threads should be less or equal to the number of available CPU cores. The default is to launch one thread per available logical core.
- uc** *filename*  
Output clustering results in *filename* using a uclust-like format. See <http://www.drive5.com/usearch/manual/ucout.html> for a description of the format.

**--usersort**

When using `--cluster_smallmem`, allow any sequence input order, not just a decreasing length ordering.

Most searching options also apply to clustering:

`--alnout`, `--blast6out`, `--userout`, `--userfields`, `fastapairs`, `--matched`, `--notmatched`, `--max-accept`, `--maxreject`, score filtering, gap penalties, masking. (see the Searching section).

Dereplication options:

**--derep\_fulllength** *filename*

Merge strictly identical sequences contained in *filename*. Identical sequences are defined as having the same length and the same string of nucleotides (case insensitive, T and U are considered the same). As **vsearch** needs to read *filename* twice, *filename* must be a real file, not a stream.

**--maxuniquesize** *positive integer*

Discard sequences with an abundance value greater than *integer*.

**--minuniquesize** *positive integer*

Discard sequences with an abundance value smaller than *integer*.

**--output** *filename*

Write the dereplicated sequences to *filename*, in fasta format and sorted by decreasing abundance. Identical sequences receive the header of the first sequence of their group. If `--sizeout` is used, the number of occurrences (i.e. abundance) of each sequence is indicated at the end of their fasta header using the pattern `";size=integer;"`.

**--sizein**

Take into account the abundance annotations present in the input fasta file (search for the pattern `"[>]size=integer[;]"` in sequence headers).

**--sizeout**

Add abundance annotations to the output fasta file (add the pattern `";size=integer;"` to sequence headers). If `--sizein` is specified, each unique sequence receives a new abundance value corresponding to its total abundance (sum of the abundances of its occurrences). If `--sizein` is not specified, input abundances are set to 1, and each unique sequence receives a new abundance value corresponding to its number of occurrences in the input file.

**--strand** *plus/both*

When searching for strictly identical sequences, check the *plus* strand only (default) or check *both* strands.

**--topn** *positive integer*

Output only the top *integer* sequences (i.e. the most abundant).

**--uc** *filename*

Output dereplication results in *filename* using a uclust-like format. See <http://www.drive5.com/usearch/manual/ucout.html> for a description of the format. In the context of dereplication, the option `--uc_allhits` has no effect.

Masking options:

An input sequence can be composed of lower- or uppercase nucleotides. Lowercase nucleotides are silently set to uppercase before masking, unless the `--qmask` soft option is used. Here are the results of combined masking options `--qmask` (or `--dbmask` for database sequences) and `--hard-mask`, assuming each input sequences contains both lower and uppercase nucleotides:

qmask	hardmask	action
none	off	no masking, all symbols uppercased
none	on	no masking, all symbols uppercased
dust	off	masked symbols lowercased, others uppercased
dust	on	masked symbols changed to Ns, others uppercased

soft	off	lowercase symbols masked, no case changes
soft	on	lowercase symbols masked and changed to Ns

**--hardmask**

Mask low-complexity regions by replacing them with Ns instead of setting them to lower case.

**--maskfasta** *filename*

Mask simple repeats and low-complexity regions in sequences contained in *filename*. The default is to mask using *dust* (use --qmask to modify that behavior).

**--output** *filename*

Write the masked sequences to *filename*, in fasta format.

**--qmask** *none/dust/soft*

Mask simple repeats and low-complexity regions in sequences using the *dust* or the *soft* algorithms, or do not mask (*none*). The default is to mask using *dust*.

**--threads** *positive integer*

Number of computation threads to use (1 to 256). The number of threads should be lesser or equal to the number of available CPU cores. The default is to launch one thread per available logical core.

Pairwise alignment options:

**--allpairs** **global** *filename*

Perform optimal global pairwise alignments of all vs. all fasta sequences contained in *filename*. The results of the  $n * (n-1) / 2$  alignments are written to the result files specified with --alnout, --blast6out, --fastapairs --matched, --notmatched, --uc or --userout (see Searching section below). Specify either the --acceptall option to output all pairwise alignments, or specify an identity level with --id to discard weak alignments. Most other accept/reject options (see Searching options below) may also be used. Sequences are aligned on their *plus* strand only. This command is multi-threaded.

**--acceptall**

Write the results of all alignments to output files. This option overrides all other accept/reject options (e.g. --id).

**--id** *real* Reject the sequence match if the pairwise identity is lower than *real* (value ranging from 0.0 to 1.0 included).

**--threads** *positive integer*

Number of computation threads to use (1 to 256). The number of threads should be lesser or equal to the number of available CPU cores. The default is to launch one thread per available logical core.

Searching options:

**--alnout** *filename*

Write pairwise global alignments to *filename* using a human-readable format. Use --rowlen to modify alignment length. Output order may vary when using multiple threads.

**--blast6out** *filename*

Write search results to *filename* using a blast-like tab-separated format of twelve fields (listed below), with one line per query-target matching (or lack of matching if --output\_no\_hits is used). Output order may vary when using multiple threads. A similar output can be obtained with --userout *filename* and --userfields query+target+id+alnlen+mism+opens+qlo+qhi+tlo+thi+evalue+bits. A complete list and description is available in the section "Userfields" of this manual.

1. *query*: query label.

2. *target*: target (database sequence) label. The field is set to "\*" if there is no alignment.
3. *id*: percentage of identity (real value ranging from 0.0 to 100.0). The percentage identity is defined as  $100 * (\text{matching columns}) / (\text{alignment length} - \text{terminal gaps})$ . See fields *id0* to *id4* for other definitions.
4. *alnlen*: length of the query-target alignment (number of columns). The field is set to 0 if there is no alignment.
5. *mism*: number of mismatches in the alignment (zero or positive integer value).
6. *opens*: number of columns containing a gap opening (zero or positive integer value).
7. *qlo*: first nucleotide of the query aligned with the target. Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
8. *qhi*: last nucleotide of the query aligned with the target. Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
9. *tlo*: first nucleotide of the target aligned with the query. Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
10. *thi*: last nucleotide of the target aligned with the query. Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
11. *eval*: expectancy-value (not computed for nucleotide alignments). Always set to -1.
12. *bits*: bit score (not computed for nucleotide alignments). Always set to 0.

**--db** *filename*

Compare query sequences (specified with `--usearch_global`) to the fasta-formatted target sequences contained in *filename*, using global pairwise alignment.

**--dbmask** *none/dust/soft*

Mask simple repeats and low-complexity regions in target database sequences using the *dust* or the *soft* algorithms, or do not mask (*none*). Warning, when using *soft* masking search commands become case sensitive. The default is to mask using *dust*.

**--dbmatched** *filename*

Write database target sequences matching at least one query sequence to *filename*, in fasta format. If the option `--sizeout` is used, the number of queries that matched each target sequence is indicated using the pattern `";size=integer;"`.

**--dbnotmatched** *filename*

Write database target sequences not matching query sequences to *filename*, in fasta format.

**--fastapairs** *filename*

Write pairwise alignments of query and target sequences to *filename*, in fasta format.

**--fulldp**

Dummy option. To maximize search sensitivity, **vsearch** uses a 8-way 16-bit SIMD vectorized full dynamic programming algorithm (Needleman-Wunsch), whether or not `--fulldp` is specified.

**--gapext** *string*

Set penalties for a gap extension. See `--gapopen` for a complete description of the penalty declaration system. The default is to initialize the six gap extending penalties using a penalty of 2 for extending internal gaps and a penalty of 1 for extending terminal gaps, in both query and target sequences (i.e. 2I/1E).



**--gapopen** *string*

Set penalties for a gap opening. A gap opening can occur in six different contexts: in the query (Q) or in the target (T) sequence, at the left (L) or right (R) extremity of the sequence, or inside the sequence (I). Sequence symbols (Q and T) can be combined with location symbols (L, I, and R), and numerical values to declare penalties for all possible contexts: aQL/bQI/cQR/dTL/eTI/fTR, where abcdef are zero or positive integers, and "/" is used as a separator.

To simplify declarations, the location symbols (L, I, and R) can be combined, the symbol (E) can be used to treat both extremities (L and R) equally, and the symbols Q and T can be omitted to treat query and target sequences equally. For instance, the default is to declare a penalty of 20 for opening internal gaps and a penalty of 2 for opening terminal gaps (left or right), in both query and target sequences (i.e. 20I/2E). If only a numerical value is given, without any sequence or location symbol, then the penalty applies to all gap openings. To declare an infinite penalty value, the symbol "\*" can be used to indicate that gap openings are forbidden in that context.

**vsearch** always initializes the six gap opening penalties using the default parameters (20I/2E). The user is then free to declare only the values he/she wants to modify. The *string* is scanned from left to right, accepted symbols are (0123456789/LIREQT\*), and later values override previous values.

Please note that **vsearch**, in contrast to **usearch**, only allows integer gap penalties. Because the lowest gap penalties are 0.5 by default in **usearch**, all default scores and gap penalties in **vsearch** have been doubled in order to obtain similar alignments.

**--hardmask**

Mask low-complexity regions by replacing them with Ns instead of setting them to lower case. For more information, please see the Masking section.

**--id** *real* Reject the sequence match if the pairwise identity is lower than *real* (value ranging from 0.0 to 1.0 included). The search process sorts target sequences by decreasing number of *k*-mers they have in common with the query sequence, using that information as a proxy for sequence similarity. That efficient pre-filtering will also prevent pairwise alignments with weakly matching targets, as there needs to be at least 6 shared *k*-mers to start the pairwise alignment, and at least one out of every 16 *k*-mers from the query needs to match the target. Consequently, using values lower than **--id** 0.5 is not likely to capture more weakly matching targets. The pairwise identity is by default defined as the number of (matching columns) / (alignment length - terminal gaps). That definition can be modified by **--iddef**.

**--iddef** 0/1/2/3/4

Change the pairwise identity definition used in **--id**. Values accepted are:

0. CD-HIT definition using shortest sequence as numerator.
1. edit distance.
2. edit distance excluding terminal gaps (default value of **--id**).
3. Marine Biological Lab definition counting each extended gap as a single difference.
4. BLAST definition, equivalent to **--iddef** 2 in a context of global pairwise alignment.

The option **--userfields** accepts the fields id0 to id4, in addition to the field id, to report the pairwise identity values corresponding to the different definitions.

**--idprefix** *positive integer*

Reject the target sequence if the first *integer* nucleotides do not match the query sequence.

- idsuffix** *positive integer*  
Reject the target sequence if the last *integer* nucleotides do not match the query sequence.
- leftjust**  
Reject the target sequence if the alignment begins with gaps.
- match** *integer*  
Score assigned to a match (i.e. identical nucleotides) in the pairwise alignment. The default value is 2.
- matched** *filename*  
Write query sequences matching database target sequences to *filename*, in fasta format.
- maxaccepts** *positive integer*  
Maximum number of hits to accept before stopping the search. The default value is 1. This option works in pair with maxrejects. The search process sorts target sequences by decreasing number of *k*-mers they have in common with the query sequence, using that information as a proxy for sequence similarity. If the first target sequence passes the acceptance criteria, it is accepted as best hit and the search process stops for that query. If maxaccepts is set to a higher value, more hits are accepted. If maxaccepts and maxrejects are both set to 0, the complete database is searched.
- maxdiffs** *positive integer*  
Reject the target sequence if the alignment contains at least *integer* substitutions, insertions or deletions.
- maxgaps** *positive integer*  
Reject the target sequence if the alignment contains at least *integer* insertions or deletions.
- maxhits** *positive integer*  
Maximum number of hits to show once the search is terminated (hits are sorted by decreasing identity). Unlimited by default value. **It applies to alnout, blast6out, uc, userout, fastapairs.**
- maxid** *real*  
Reject the target sequence if its percentage of identity with the query is greater than *real*.
- maxqsize** *positive integer*  
Reject query sequences with an abundance greater than *integer*.
- maxqt** *real*  
Reject if the query/target sequence length ratio is greater than *real*.
- maxrejects** *positive integer*  
Maximum number of non-matching target sequences to consider before stopping the search. The default value is 32. This option works in pair with maxaccepts. The search process sorts target sequences by decreasing number of *k*-mers they have in common with the query sequence, using that information as a proxy for sequence similarity. If none of the first 32 examined target sequences pass the acceptance criteria, the search process stops for that query (no hit). If maxrejects is set to a higher value, more target sequences are considered. If maxaccepts and maxrejects are both set to 0, the complete database is searched.
- maxsizeratio** *real*  
Reject if the query/target abundance ratio is greater than *real*.
- maxsl** *real*  
Reject if the shorter/longer sequence length ratio is greater than *real*.
- maxsubs** *positive integer*  
Reject the target sequence if the alignment contains more than *integer* substitutions.

- mid** *real*  
Reject the alignment if the percentage of identity is lower than *real* (ignoring all gaps, internal and terminal).
- mincols** *positive integer*  
Reject the target sequence if the alignment length is shorter than *integer*.
- minq** *real*  
Reject if the query/target sequence length ratio is lower than *real*.
- minsize** *real*  
Reject if the query/target abundance ratio is lower than *real*.
- minsl** *real*  
Reject if the shorter/longer sequence length ratio is lower than *real*.
- mints** *positive integer*  
Reject target sequences with an abundance lower than *integer*.
- mismatch** *integer*  
Score assigned to a mismatch (i.e. different nucleotides) in the pairwise alignment. The default value is -4.
- notmatched** *filename*  
Write query sequences not matching database target sequences to *filename*, in fasta format.
- output\_no\_hits**  
Write both matching and non-matching queries to --alnout, --blast6out, and --userout output files (--uc and --uc\_allhits output files always feature non-matching queries). Non-matching queries are labelled "No hits" in --alnout files.
- qmask** *none/dust/soft*  
Mask simple repeats and low-complexity regions in query sequences using the *dust* or the *soft* algorithms, or do not mask (*none*). Warning, when using *soft* masking search commands become case sensitive. The default is to mask using *dust*.
- query\_cov** *real*  
Reject if the fraction of the query aligned to the target sequence is lower than *real*. The query coverage is computed as (matches + mismatches) / query sequence length. Internal or terminal gaps are not taken into account.
- rightjust**  
Reject the target sequence if the alignment ends with gaps.
- rowlen** *positive integer*  
Width of alignment lines in --alnout output. The default value is 64. Set to 0 to eliminate wrapping.
- self** Reject the alignment if the query and target labels are identical.
- selfid** Reject the alignment if the query and target sequences are strictly identical.
- sizeout**  
Add abundance annotations to the output of the option --dbmatched (using the pattern ";size=integer;").
- strand** *plus/both*  
When searching for similar sequences, check the *plus* strand only (default) or check *both* strands.
- target\_cov** *real*  
Reject if the fraction of the target sequence aligned to the query sequence is lower than *real*. The target coverage is computed as (matches + mismatches) / target sequence length. Internal or terminal gaps are not taken into account.

**--threads** *positive integer*

Number of computation threads to use (1 to 256). The number of threads should be lesser or equal to the number of available CPU cores. The default is to launch one thread per available logical core.

**--top\_hits\_only**

Output only the hits with the highest percentage of identity with the query.

**--uc** *filename*

Output searching results in *filename* using a uclust-like format. See <http://www.drive5.com/usearch/manual/ucout.html> for a description of the format. Output order may vary when using multiple threads.

**--uc\_allhits**

When using the --uc option, show all hits, not just the top hit for each query.

**--usearch\_global** *filename*

Compare target sequences (--db) to the fasta-formatted query sequences contained in *filename*, using global pairwise alignment.

**--userfields** *string*

When using --userout, select and order the fields written to the output file. Fields are separated by "+" (e.g. query+target+id). See the "Userfields" section for a complete list of fields.

**--userout** *filename*

Write user-defined tab-separated output to *filename*. Select the fields with the option --userfields. Output order may vary when using multiple threads. If --userfields is empty or not present, *filename* is empty.

**--weak\_id** *real*

Show hits with percentage of identity of at least *real*, without terminating the search. A normal search stops as soon as enough hits are found (as defined by --maxaccepts, --maxrejects, and --id). As --weak\_id reports weak hits that are not deduced from --maxaccepts, high --id values can be used, hence preserving both speed and sensitivity. Logically, *real* must be smaller than the value indicated by --id.

**--wordlength** *positive integer*

Length of words (i.e. *k*-mers) for database indexing. The range of possible values goes from 3 to 15, but values near 8 are generally recommended. Longer words may reduce the sensitivity for weak similarities, but can increase accuracy. On the other hand, shorter words may increase sensitivity, but can reduce accuracy. Computation time will generally increase with shorter words and decrease with longer words. Memory requirements for a part of the index increase with a factor of 4 each time word length increases by one nucleotide, and this generally becomes significant for long words (12 or more). The default value is 8.

Shuffling options:

**--output** *filename*

Write the shuffled sequences to *filename*, in fasta format.

**--seed** *positive integer*

When shuffling sequence order, use *integer* as seed. A given seed will always produce the same output order (useful for replicability). Set to 0 to use a pseudo-random seed (default behavior).

**--shuffle** *filename*

Pseudo-randomly shuffle the order of sequences contained in *filename*.

**--topn** *positive integer*

Output only the top *integer* sequences.

Sorting options:

Fasta entries are sorted by decreasing abundance (**--sortbysize**) or sequence length (**--sortbylength**). To obtain a stable sorting order, ties are sorted by decreasing abundance and label increasing alpha-numerical order (**--sortbylength**), or just by label increasing alpha-numerical order (**--sortbysize**). Label sorting assumes that all sequences have unique labels. The same applies to the automatic sorting performed during chimera checking (**--uchime\_denovo**), dereplication (**--derep\_fulllength**), and clustering (**--cluster\_fast** and **--cluster\_size**).

**--maxsize** *positive integer*

When using **--sortbysize**, discard sequences with an abundance value greater than *integer*.

**--minsize** *positive integer*

When using **--sortbysize**, discard sequences with an abundance value smaller than *integer*.

**--output** *filename*

Write the sorted sequences to *filename*, in fasta format.

**--relabel** *string*

Relabel sequence using the prefix *string* and a ticker (1, 2, 3, etc.) to construct the new headers. Use **--sizeout** to conserve the abundance annotations.

**--sizeout** When using **--relabel**, report abundance annotations to the output fasta file (using the pattern "**size=integer**").

**--sortbylength** *filename*

Sort by decreasing length the sequences contained in *filename*. See the general options **--minseqlength** and **--maxseqlength** to eliminate short and long sequences.

**--sortbysize** *filename*

Sort by decreasing abundance the sequences contained in *filename* (the pattern "**[>]size=integer**:" has to be present). See the options **--minsize** and **--maxsize** to eliminate rare and dominant sequences.

**--topn** *positive integer*

Output only the top *integer* sequences (i.e. the longest or the most abundant).

Userfields (fields accepted by the **--userfields** option):

<b>aln</b>	Print a string of M (match), D (delete, i.e. a gap in the query) and I (insert, i.e. a gap in the target) representing the pairwise alignment. Empty field if there is no alignment.
<b>alnlen</b>	Print the length of the query-target alignment (number of columns). The field is set to 0 if there is no alignment.
<b>bits</b>	Bit score (not computed for nucleotide alignments). Always set to 0.
<b>caln</b>	Compact representation of the pairwise alignment using the CIGAR format (Compact Idiosyncratic Gapped Alignment Report): M (match), D (deletion) and I (insertion). Empty field if there is no alignment.
<b>evaluate</b>	E-value (not computed for nucleotide alignments). Always set to -1.
<b>exts</b>	Number of columns containing a gap extension (zero or positive integer value).
<b>gaps</b>	Number of columns containing a gap (zero or positive integer value).
<b>id</b>	Percentage of identity (real value ranging from 0.0 to 100.0). The percentage identity is defined as $100 * (\text{matching columns}) / (\text{alignment length} - \text{terminal gaps})$ .
<b>id0</b>	CD-HIT definition of the percentage of identity, using the shortest sequence in the pairwise alignment as numerator (real value ranging from 0.0 to 100.0).

<b>id1</b>	The percentage of identity (real value ranging from 0.0 to 100.0) is defined as the edit distance: $100 * (\text{matching columns}) / (\text{alignment length})$ .
<b>id2</b>	The percentage of identity (real value ranging from 0.0 to 100.0) is defined as the edit distance, excluding terminal gaps. The field id2 is an alias for the field id.
<b>id3</b>	Marine Biological Lab definition of the percentage of identity (real value ranging from 0.0 to 100.0), counting each extended gap as a single difference.
<b>id4</b>	BLAST definition of the percentage of identity (real value ranging from 0.0 to 100.0), equivalent to --iddef 2 in a context of global pairwise alignment.
<b>ids</b>	Number of matches in the alignment (zero or positive integer value).
<b>mism</b>	Number of mismatches in the alignment (zero or positive integer value).
<b>opens</b>	Number of columns containing a gap opening (zero or positive integer value).
<b>pairs</b>	Number of columns containing only nucleotides. That value corresponds to the length of the alignment minus the gap-containing columns (zero or positive integer value).
<b>pctgaps</b>	Number of columns containing gaps expressed as a percentage of the alignment length (real value ranging from 0.0 to 100.0).
<b>pctpv</b>	Percentage of positive columns. When working with nucleotide sequences, this is equivalent to the percentage of matches (real value ranging from 0.0 to 100.0).
<b>pv</b>	Number of positive columns. When working with nucleotide sequences, this is equivalent to the number of matches (zero or positive integer value).
<b>qcov</b>	Fraction of the query sequence that is aligned with the target sequence (real value ranging from 0.0 to 100.0). The query coverage is computed as $100.0 * (\text{matches} + \text{mismatches}) / \text{query sequence length}$ . Internal or terminal gaps are not taken into account. The field is set to 0.0 if there is no alignment.
<b>qframe</b>	Query frame (-3 to +3). That field only concerns coding sequences and is not computed by <b>vsearch</b> . Always set to +0.
<b>qhi</b>	Last nucleotide of the query aligned with the target. Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
<b>qihi</b>	Last nucleotide of the query aligned with the target (ignoring terminal gaps). Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
<b>qilo</b>	First nucleotide of the query aligned with the target (ignoring initial gaps). Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
<b>ql</b>	Query sequence length (positive integer value). The field is set to 0 if there is no alignment.
<b>qlo</b>	First nucleotide of the query aligned with the target. Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
<b>qrow</b>	Print the sequence of the query segment as seen in the pairwise alignment (i.e. with gap insertions if need be). Empty field if there is no alignment.
<b>qs</b>	Query segment length. Always equal to query sequence length.
<b>qstrand</b>	Query strand orientation (+ or - for nucleotide sequences). Empty field if there is no alignment.
<b>query</b>	Query label.
<b>raw</b>	Raw alignment score (negative, null or positive integer value). The score is the sum of match rewards minus mismatch penalties, gap openings and gap extensions. The field is set to 0 if there is no alignment.

<b>target</b>	Target label. The field is set to "*" if there is no alignment.
<b>tcov</b>	Fraction of the target sequence that is aligned with the query sequence (real value ranging from 0.0 to 100.0). The target coverage is computed as $100.0 * (\text{matches} + \text{mismatches}) / \text{target sequence length}$ . Internal or terminal gaps are not taken into account. The field is set to 0.0 if there is no alignment.
<b>tframe</b>	Target frame (-3 to +3). That field only concerns coding sequences and is not computed by <b>vsearch</b> . Always set to +0.
<b>thi</b>	Last nucleotide of the target aligned with the query. Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
<b>tihi</b>	Last nucleotide of the target aligned with the query (ignoring terminal gaps). Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
<b>tilo</b>	First nucleotide of the target aligned with the query (ignoring initial gaps). Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
<b>tl</b>	Target sequence length (positive integer value). The field is set to 0 if there is no alignment.
<b>tlo</b>	First nucleotide of the target aligned with the query. Nucleotide numbering starts from 1. The field is set to 0 if there is no alignment.
<b>trow</b>	Print the sequence of the target segment as seen in the pairwise alignment (i.e. with gap insertions if need be). Empty field if there is no alignment.
<b>ts</b>	Target segment length. Always equal to target sequence length. The field is set to 0 if there is no alignment.
<b>tstrand</b>	Target strand orientation (+ or - for nucleotide sequences). Always set to "+", so reverse strand matches have tstrand "+" and qstrand "-". Empty field if there is no alignment.

## DELIBERATE CHANGES

If you are a usearch user, our objective is to make you feel at home. That's why **vsearch** was designed to behave like usearch, to some extent. Like any complex software, usearch is not free from quirks and inconsistencies. We decided not to reproduce some of them, and for complete transparency, to document here the deliberate changes we made.

During a search with usearch, when using the options `--blast6out` and `--output_no_hits`, for queries with no match the number of fields reported is 13, where it should be 12. This is corrected in **vsearch**.

The field `raw` of the `--userfields` option is not informative in usearch. This is corrected in **vsearch**.

The fields `qlo`, `qhi`, `tlo`, `thi` now have counterparts (`qilo`, `qihi`, `tilo`, `tihi`) reporting alignment coordinates ignoring terminal gaps.

In usearch, when using the option `--output_no_hits`, queries that receive no match are reported in `blast6out` file, but not in the alignment output file. This is corrected in **vsearch**.

**vsearch** introduces a new `--cluster_size` command that sorts sequences by decreasing abundance before clustering.

**vsearch** reintroduces `--iddef` alternative pairwise identity definitions that were removed from usearch.

**vsearch** extends the `--topn` option to sorting commands.

**vsearch** extends the `--sizein` option to dereplication (`--derep_fulllength`) and clustering (`--cluster_fast`).

**vsearch** treats T and U as identical nucleotides for dereplication.

**vsearch** sorting is stabilized by using sequence abundances or sequences labels as secondary or tertiary keys.

## NOVELTIES

**vsearch** introduces new options not present in usearch. They are described in the "Options" section of this manual. Here is a short list:

- alignwidth
- cluster\_size
- fasta\_width
- iddef
- maxuniquesize
- shuffle

## EXAMPLES

Cluster with a 97% similarity threshold, collect cluster centroids, and write cluster descriptions using a uclust-like format:

```
vsearch --cluster_fast queries.fas --id 0.97 --centroids centroids.fas --uc clusters.uc
```

Dereplicate the sequences contained in *queries.fas*, take into account the abundance information already present, write unwrapped sequences to output with the new abundance information, discard all sequences with an abundance of 1:

```
vsearch --derep_fulllength queries.fas --output queries_masked.fas --sizein --sizeout --fasta_width 0 --minuniquesize 2
```

Mask simple repeats and low complexity regions in the input fasta file (masked regions are lowercased), and write the results to the output file:

```
vsearch --maskfasta queries.fas --output queries_masked.fas --qmask dust
```

Sort by decreasing abundance the sequences contained in *queries.fas* (using the "size=*integer*" information), relabel the sequences while preserving the abundance information (with --sizeout), keep only sequences with an abundance equal to or greater than 2:

```
vsearch --sortbysize queries.fas --output queries_sorted.fas --relabel sampleA_ --sizeout --min-size 2
```

Align all sequences in a database with each other and output all pairwise alignments:

```
vsearch --allpairs_global database.fas --alnout results.aln --acceptall
```

Search queries in a reference database, with a 80%-similarity threshold, take terminal gaps into account when calculating pairwise similarities:

```
vsearch --usearch_global queries.fas --db references.fas --alnout results.aln --id 0.8 --iddef 1
```

Search a sequence dataset against itself (ignore self hits), get all matches with at least 60% identity, and collect results in a blast-like tab-separated format:

```
vsearch --usearch_global queries.fas --db queries.fas --id 0.6 --self --blast6out results.blast6 --maxaccepts 0 --maxrejects 0
```

Shuffle the input fasta file (change the order of sequences) in a repeatable fashion (fixed seed), and write unwrapped fasta sequences to the output file:

```
vsearch --shuffle queries.fas --output queries_shuffled.fas --seed 13 --fasta_width 0
```

## AUTHORS

Implementation by Torbjørn Rognes and Tomas Flouri, documentation by Frédéric Mahé.

## REPORTING BUGS

Submit suggestions and bug-reports at <<https://github.com/torognes/vsearch/issues>>, send a pull request on <<https://github.com/torognes/vsearch>>, or compose a friendly or curmudgeont e-mail to Torbjørn Rognes <[torognes@ifi.uio.no](mailto:torognes@ifi.uio.no)>.



## AVAILABILITY

Source code and binaries are available at <<https://github.com/torognes/vsearch>>.

## COPYRIGHT

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**vsearch** includes code from Google's CityHash project by Geoff Pike and Jyrki Alakuijala, providing some excellent hash functions available under a MIT license.

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## SEE ALSO

**swipe**, an extremely fast pairwise local (Smith-Waterman) database search tool by Torbjørn Rognes, available at <<https://github.com/torognes/swipe>>.

**swarm**, a fast and accurate amplicon clustering method by Frédéric Mahé and Torbjørn Rognes, available at <<https://github.com/torognes/swarm>>.

## VERSION HISTORY

New features and important modifications of **vsearch** (short lived or minor bug releases may not be mentioned):

**v1.0.0** released November 28th, 2014

First public release.

**v1.0.1** released December 1st, 2014

Bug fixes (sortbysize, semicolon after size annotation in headers) and minor changes (labels as secondary sort key for most sorts, treat T and U as identical for dereplication, only output size in dbmatched file if sizeout specified).

**v1.0.2** released December 6th, 2014

Bug fixes (ssse3/sse41 requirement, memory leak).

**v1.0.3** released December 6th, 2014

Bug fix (now writes help to stdout instead of stderr).

**v1.0.4** released December 8th, 2014

Added --allpairs\_global option. Reduced memory requirements slightly. Removed memory leaks.

**v1.0.5** released December 9th, 2014

Fixes a minor bug with --allpairs\_global and --acceptall options.

**v1.0.6** released December 14th, 2014

Fixes a memory allocation bug in chimera detection (--uchime\_ref option).

**v1.0.7** released December 19th, 2014

Fixes a bug in the output from chimera detection with the --uchimeout option.

**v1.0.8** released January 22nd, 2015

Introduces several changes and bug fixes:

- a new linear memory aligner for alignment of sequences longer than 5,000 nucleotides,
- a new `--cluster_size` command that sorts sequences by decreasing abundance before clustering,
- meaning of userfields `qlo`, `qhi`, `tlo`, `thi` changed for compatibility with `usearch`,
- new userfields `qilo`, `qihi`, `tilo`, `tihi` gives coordinates ignoring terminal gaps,
- in `--uc` output files, a perfect alignment is indicated with a "=" sign,
- `--cluster_fast` will now sort sequences by decreasing length, then by decreasing abundance and finally by sequence identifier,
- default `--maxseqlength` value set to 50,000 nucleotides,
- fix for bug in alignment in rare cases,
- fix for lack of detection of under- or overflow in SIMD aligner.

**v1.0.9** released January 22nd, 2015

Fixes a bug in the function sorting sequences by decreasing abundance (`--sortbysize`).