# The FASTA program package

# Introduction

This documentation describes the version 36 of the FASTA program package (see W. R. Pearson and D. J. Lipman (1988), "Improved Tools for Biological Sequence Analysis", PNAS 85:2444-2448, [16] W. R. Pearson (1996) "Effective protein sequence comparison" Meth. Enzymol. 266:227-258 [14]; and Pearson et. al. (1997) Genomics 46:24-36 [17]. Version 3 of the FASTA packages contains many programs for searching DNA and protein databases and for evaluating statistical significance from randomly shuffled sequences.

This document is divided into four sections: (1) A summary overview of the programs in the FASTA3 package; (2) A guide to using the FASTA programs; (3) A guide to installing the programs and databases. Section (4) provides answers to some Frequently Asked Questions (FAQs). In addition to this document, the changes\_v36.html, changes\_v35.html and changes\_v34.html files list functional changes to the programs. The readme.v30..v36 files provide a more complete revision history of the programs, including bug fixes.

The programs are easy to use; if you are using them on a machine that is administered by someone else, you can focus on sections (1) and (2) to learn how to use the programs. If you are installing the programs on your own machine, you will need to read section (3) carefully.

*FASTA and BLAST* – FASTA and BLAST have the same goal: to identify statistically significant sequence similarity that can be used to infer homology. The FASTA programs offer several advantages over BLAST:

- 1. Rigorous algorithms unavailable in BLAST (Table I). Smith-Waterman (ssearch36), global: global (ggsearch36), and global:local (glsearch36) programs are available, and these programs can be used with psiblast PSSM profiles.
- 2. Better translated alignments. fastx36, fasty36, tfastx36, and tfastx36 allow frame-shifts in alignments; frame-shifts are treated like gap-penalties, alignments tend to be longer in error-prone reads.
- Better statistics. BLAST calculates very accurate statistics for protein:protein alignments, but its model-based strategy is less robust for translated-DNA:protein and DNA:DNA scores. FASTA uses an empirical estimation strategy, and now provides both search-based, and high-scoring shuffle-based statistics (-z 21).
- 4. More flexible library sequence formats. The FASTA programs can read FASTA, NCBI/ formatdb, and several other sequence formats, and can directly query MySQL and Post-gres databases. The programs offer several strategies for specifying subsets of databases.
- 5. A very efficient threaded implementation. The FASTA programs are fully threaded; both similarity scores and alignments can be calculated in parallel on multi-core hardware. On multi-core machines, FASTA can be faster than BLAST while producing better alignments with more accurate statistical estimates.

In addition, the FASTA programs from fasta-36.3.4 on provide an option to produce very BLAST-like output (-m BB), so that analysis pipelines require minimal modification.

# 1 An overview of the FASTA programs

Although there are a large number of programs in this package, they belong to three groups: (1) Traditional similarity searching programs: fasta36, fastx36, fasty36, tfastx36, tfasty36, ssearch36, ggsearch36, and glsearch36; (2) Programs for searching with short fragments: fasts36, fastf36, tfasts36, tfastf36, and fastm36; (3) A program for finding non-overlapping local alignments: lalign36. Programs that start with fast search protein databases, while tfast programs search translated DNA databases. Table I gives a brief description of the programs.

In addition, there are several programs included. map\_db is used to index FASTA format sequence databases for more efficient scanning. scripts/lav2plt.pl can plot the .lav files produced by lalign -m 11 as postscript (lav2plt.pl --dev ps) or SVG (lav2plt.pl --dev svg) output.

# 2 Using the FASTA Package

### 2.1 Introduction/Overview

All the FASTA sequence comparison programs use similar command line options and arguments. The simplest command line arguments are (in order): the name of a query sequence file, a library file, and (possibly) the *ktup* parameter. If command line options are provided, they *must* precede the standard query-file and library-file arguments. Thus:

fasta36 -s BP62 query.file library.file

will compare the sequences in query.file with those in library.file using the BLOSUM62 scoring matrix with BLASTP gap penalties (-11/-1).

The program can also be run by typing:

fasta36 -I

which presents the "classic" interative mode (this was the default behavior before version 36.3.4). In interactive mode, you will be prompted for: (1) the name of the test sequence file; (2) the name of the library file; (3) whether you want ktup = 1 or 2. (1 – 6 for DNA sequences).

Current versions of the FASTA programs expect a query file and library, if you simply type "fasta36", you will see a short help message:

```
% ssearch36
USAGE
ssearch36 [-options] query_file library_file
ssearch36 -help for a complete option list
DESCRIPTION
SSEARCH performs a Smith-Waterman search
version: 36.3.4 Mar, 2011
COMMON OPTIONS (options must precede query_file library_file)
-s: [BL50] scoring matrix;
-f: [-10] gap-open penalty;
-g: [-2] gap-extension penalty;
```

```
-S filter lowercase (seg) residues;
```

```
-b: high scores reported (limited by -E by default);
```

| FASTA<br>program          | BLAST<br>equiv.   | Description   |
|---------------------------|-------------------|---|
| fasta36                   | blastp/<br>blastn | Compare a protein sequence to a protein sequence database or a DNA sequence to a DNA sequence database using the FASTA algorithm [14, 16]. Search speed and selectivity are controlled with the <i>ktup</i> (wordsize) parameter. For protein comparisons, <i>ktup</i> = 2 by default; <i>ktup</i> =1 is more sensitive but slower. For DNA comparisons, <i>ktup</i> =6 by default; <i>ktup</i> =3 or <i>ktup</i> =4 provides higher sensitivity. |
| ssearch36                 |                   | Compare a protein sequence to a protein sequence database or<br>a DNA sequence to a DNA sequence database using the Smith-<br>Waterman algorithm [20]. ssearch36 uses SSE2 acceleration, and<br>is only 2 - 5X slower than fasta36 [5].   |
| ggsearch36/<br>glsearch36 |                   | Compare a protein sequence to a protein sequence database or<br>a DNA sequence to a DNA sequence database using an optimal<br>global:global (ggsearch36) or global:local (glsearch36) algorithm.  |
| fastx36/<br>fasty36       | blastx            | Compare a DNA sequence to a protein sequence database, by com-<br>paring the translated DNA sequence in three frames and allowing<br>gaps and frameshifts. fastx36 uses a simpler, faster algorithm for<br>alignments that allows frameshifts only between codons; fasty36 is<br>slower but can produce better alignments because frameshifts are<br>allowed within codons [24].  |
| tfastx36/<br>tfasty36     | tblastn           | Compare a protein sequence to a DNA sequence database, calcu-<br>lating similarities with frameshifts to the forward and reverse orienta-<br>tions [24].  |
| fastf36/<br>tfastf36      |                   | Compares an ordered peptide mixture, as would be obtained by Ed-<br>man degradation of a CNBr cleavage of a protein, against a protein<br>(fastf) or DNA (tfastf) database [10].  |
| fasts36/<br>tfasts36      |                   | Compares set of short peptide fragments, as would be obtained from mass-spec. analysis of a protein, against a protein (fasts) or DNA (tfasts) database [10].   |
| lalign36                  |                   | Calculate multiple, non-intersecting alignments using the sim2 imple-<br>mentation of the Waterman-Eggert algorithm [21] developed by Xi-<br>aoqui Huang and Web Miller [7]. Statistical estimates are calculated<br>from Smith-Waterman scores of shuffled sequences.  |

Table 1: Comparison programs in the FASTA36 package

-d: number of alignments shown (limited by -E by default); -I interactive mode;

"fasta36 -help" (or any of the other program names in Table I) provides complete listing of the options available for the program and their default values.

The package includes several test files. To check to make certain that everything is working, you can try:

```
fasta36 ../seq/musplfm.aa ../seq/prot_test.lib
or
tfastx36 ../seq/mgstm1.aa ../seq/gst.nlib
```

#### 2.2 Sequence files

The fasta36 programs can read query and library files in many standard formats (Section 3.5). The default file format for query and library files – the format that will be used if no additional file format information is provided – is FASTA format. Like BLAST, version 36 can compare a query file with multiple query sequences to a sequence database, performing an independent search with each sequence in the query file.

FASTA format files consist of a description line, beginning with a '>' character, followed by the sequence itself:

```
>sequence name and description 1
A F A S Y T .... actual sequence.
F S S .... second line of sequence.
>sequence name and description 2
PMILTYV ... sequence 2
```

All of the characters of the description line are read, and special characters can be used to indicate additional information about the sequence. In general, non-amino-acid/non-nucleotide sequences in the sequence lines are ignored.

FASTA format files from major sequence distributors, like the NCBI and EBI, have specially formatted description lines, e.g.:

>gi|54321|ref|np\_12345| example NCBI refseq sequence

or

>sw:gstm1\_human P01234 glutathione transferase GSTM1 - human

Several sample test files are included with the FASTA distribution: seq/\*.aa and seq/\*.seq, as well as two small sequence libraries, seq/prot\_test.lib and seq/gst.nlib.

You can build your own library by concatenating several sequence files. Just be sure that each sequence is preceded by a line beginning with a '>' followed by a sequence name/description. Sequences entered with word processors should use a "text" mode, e.g. "Save as text" with MS-WORD, with end of line characters and no special formatting characters in the file. The FASTA program cannot read Microsoft Word .DOC files, or rich text (.RTF) files; query and library sequence files should contain only sequence descriptions, sequences, and end-of-line characters.

# 2.3 Running the programs

As mentioned earlier, the FASTA programs can be run either interactively, by typing the name of a FASTA program (and possibly command line options), followed by -I (fasta36 -I) or from the command line, entering command line options, and the query and library file names. For searches of large databases that may take several minutes (or longer), it is more convenient to run searches from the command line, e.g.:

```
fasta36 query.file library.file > output.file
```

The command line shown above could be typed in a Unix or MacOSX terminal window, or from the MS-Windows command line interface (command.exe). The command line syntax shown above works for all the FASTA programs, e.g.:

lalign36 mchu.aa mchu.aa > mchu.laln
fastx36 mgstm1.seq prot\_test.lseg > mgstm1.fx\_out
ssearch36 mgstm1.aa xurtg.aa > mgstm1\_xurtg.ss

Command line options – The FASTA programs provide a variety of command line options that modify the default scoring matrix (-sBL62) and gap penalties (-f -11, -g -1), other algorithm parameters, the output options (-E 0.1, -d 20, -m 9i), and statistical procedures (z -2). A complete list of command line options is shown near the end of this document. Unlike the BLAST programs, all FASTA command line options must precede the query file name and library file name (and there are no command line options available to specify the query and library file names). Thus, you should type:

ssearch36 -s BL62 -f -11 -g -1 query.file library.file > output.file

If you include -I as one of the options, you can provide command line options (e.g. to change the scoring matrix or gap penalties) without a query file or library file, and the program will use the options but prompt for the necessary files.

### 2.4 Interpreting the results

Fig. 1 shows the output from a typical FASTA program (ssearch36). The output file can be viewed as four parts: (a) the initial command line and description of the query sequence used (mgstm1.aa, 218 aa) and library (PIR1, 13,143 entries); (b) a description of the search statistics, algorithm (Smith-Waterman, SSE2 accelerated), and search parameters (BLOSUM50 matrix, gap penalties: -10 to open a gap, -2 for each residue in a gap); (c) a list of high scoring library sequences, descriptions, similarity scores, and statistical significance; (d) the alignments that produced the scores.

### 2.4.1 Identifying homologs

In the description section (which starts: The best scores are:), four numbers after the description of each library sequence are shown: (i) (in parentheses) the length of the library sequence; (ii) the raw Smith-Waterman score for the alignment (s-w; for the fasta36, [t]fast[x,y]36 programs, this column would be labeled opt, for the *opt*imized – banded Smith-Waterman – score), (iii) the *bit* score, and (iv) the expectation (E()), or statistical significance, of the alignment score.

# ../bin/ssearch36 -q -w 80 ../seq/mgstm1.aa a SSEARCH performs a Smith-Waterman search version 36.3.6 June, 2013(preload9) Please cite: T. F. Smith and M. S. Waterman, (1981) J. Mol. Biol. 147:195-197; W.R. Pearson (1991) Genomics 11:635-650 Query: ../seq/mgstm1.aa 1>>>mGSTM1 mouse glutathione transferase M1 - 218 aa Library: PIR1 Annotated (rel. 66) 5121825 residues in 13143 sequences Statistics: Expectation\_n fit: rho(ln(x))= 7.4729+/-0.000484; mu= 2.0282+/- 0.027 mean\_var=56.9651+/-10.957, 0's: 9 Z-trim(119.4): 17 B-trim: 67 in 1/62 Lambda= 0.169930 statistics sampled from 13135 (13143) to 13135 sequences Algorithm: Smith-Waterman (SSE2, Michael Farrar 2006) (7.2 Nov 2010) Parameters: BL50 matrix (15:-5), open/ext: -10/-2 Scan time: 3.820 The best scores are: s-w bits E(13143) sp|P08010|GSTM2\_RAT Glutathione S-transferase Mu 2; GST 4-4; GT (218) 1248 312.0 7.7e-86 sp|P04906|GSTP1\_RAT Glutathione S-transferase P; Chain 7; GST - (210) 344 90.4 3.8e-19 sp|P00502|GSTA1\_RAT Glutathione S-transferase alpha-1; GST 1-1 (222) 237 64.1 3.2e-11 (222) 179 49.9 6.1e-07 sp|P14942|GSTA4\_RAT Glutathione S-transferase alpha-4; GST 8-8 sp|P12653|GSTF1\_MAIZE Glutathione S-transferase 1; GST class-pi (214) 120 35.4 0.013 sp|P04907|GSTF3\_MAIZE Glutathione S-transferase 3; GST class-pi (222) 115 34.2 0.032 sp|P20432|GSTT1\_DROME Glutathione S-transferase 1-1; DDT-dehydr (209) 100 30.5 0.38 sp|P11277|SPTB1\_HUMAN Spectrin beta chain, erythrocytic; Beta-(2137) 108 31.6 1.9 ... (alignments deleted) ... >>sp|P14942|GSTA4\_RAT Glutathione S-transferase alpha-4; GST 8-8; (222 aa) s-w opt: 179 Z-score: 231.0 bits: 49.9 E(13143): 6.1e-07 Smith-Waterman score: 179; 25.6% identity (54.5% similar) in 211 aa overlap (5-207:7-207) 10 20 30 40 50 60 70 MPMILGYWNVRGLTHPIRMLLEYTDSSYDEKRYTMGDAPDFDRSQWLNEKF-KLG-LDFPNLPYL-IDGSHKITQSNA mGSTM : :.. :: . :: :: . ... ..... sp|P14\_MEVKPKLYYFQGRGRMESIRWLLATAGVEFEE-----EFLETREQYEKLQKDGCLLFGQVPLVEIDG-MLLTQTRA 70 10 20 30 40 50 60 80 90 100 110 120 130 140 150 mGSTM ILRYLARKHHLDGETEEERIRADIVENQVMDTRMQLIMLCYNPDFEKQKPEFLKTIPEKMKLYSEF--LGK---RPWFAG \*\* \*\*\* \*\*\*\* \*\* \*\*\*\*\* \*\* \*\*\*\* \*\*\*\* sp|P14 ILSYLAAKYNLYGKDLKERVRIDMYADGTQDLMMMIIGAPFKAPQEKEESLALAVKRAKNRYFPVFEKILKDHGEAFLVG 100 130 80 90 110 120 140 150 160 170 180 190 200 210  ${\tt mGSTM} \quad {\tt DKVTYVDFLAYDILDQYRMFEPKCLDAFPNLRDFLARFEGLKKISAYMKSSRYIATPIFSKMAHWSNK}$ :. :: :. : .:. .. :. ... . ..... sp|P14 NQLSWADIQLLEAILMVEEVSAPVLSDFPLLQAFKTRISNIPTIKKFLQPGSQRKPPPDGHYVDVVRTVLKF 160 170 180 190 200 210 220 ... (alignments deleted) ... 218 residues in 1 query sequences 5121825 residues in 13143 library sequences Tcomplib [36.3.6 May, 2013(preload9)] (4 proc in memory [OG]) start: Thu Jun 6 11:23:28 2013 done: Thu Jun 6 11:23:30 2013 Total Scan time: 3.820 Total Display time: 0.130 Function used was SSEARCH [36.3.6 May, 2013(preload9)]

Figure 1: ssearch36 results

Comparison of seq/mgstm1.aa against a small protein database (pir1.lseg). Some high-scoring sequences and all but one alignment were removed to reduce the output size.

The E()-value depends on the size of the database searched, in this case, 13,143 sequences, so the database size is given at the top of the list.

The bit score is equivalent to a BLAST bit score; together with the length of query and library sequences, it can be used to calculate the significance of the alignment.<sup>1</sup> Bit scores are convenient because they provide a matrix independent score that can be compared with other searches performed with other matrices and gap penalties against other databases. However, the E()-value, or expectation, provides the most direct measure of the statistical significance of the match.

In this example, the GSTP1\_RAT, GSTA1\_RAT, and GSTA4\_RAT proteins share strong significant similarity (better than  $E() < 6.1 \times 10^{-7}$ ), while the GSTF1\_MAIZE, GSTF3\_MAIZE, and GSTT1\_DROME sequences do not share significant similarity (E() < 0.001). However, GSTF1\_MAIZE, GSTF3\_MAIZE, and GSTT1\_DROME are all glutathione transferase homologs, they simply do not share statistically significant similarity with this particular mGSTM1 query. Statistically significant sequence similarity scores *can* be used to infer *homology* (common ancestry), but non-significant scores *cannot* be used to infer *non-homology*.

While percent identity is often used to characterize the quality of an alignment and the likelihood that it reflects homology, the E()-value is a much more reliable value. Often sequences that share less than 30% identity will share very significant similarity (in the example above mgstm1.aa and GSTA4\_RAT, with E() < 6.1E-07 are 25.6% identical). The expectation value captures information about conservative replacements, identities, and alignment length to provide a *single* value that captures the significance of the alignment.

For protein searches, library sequences with E()-values < 0.001 for searches of a 10,000 entry protein database are almost always homologous. Some sequences with E()-values from 1 - 10 may also be related, but unrelated sequences (1–10 per search) will have scores in this range as well.

E()-values < 0.001 can reliably be used to infer homology, assuming that the statistical estimates are accurate. The two most common causes of statistical problems are low-complexity regions and amino-acid composition bias. Low-complexity regions are can be identified using the pseg program [22], and filtered out using the -S option. Composition bias rarely produces highly-significiant E()-values, but can cause unrelated sequences to have E()-values between 0.01 and 0.001. The FASTA programs offer two shuffle-based strategies for evaluating composition bias; calculating similarity scores for random sequences with the same length and amino acid composition (-z 11 .. 16),<sup>2</sup> and calculating statistical estimates derived from shuffles of the high-scoring sequences (-z 21, 22, 24, 25, 26).

When -z 21 ... 26 shuffles are performed, the FASTA36 programs present two E()-values in the list of high scoring sequences and the alignments; the traditional one based on the library search, and a second E2() value, based on the shuffles of the high scoring sequences. -z 21 ... 26 shuffles are most useful for evaluating the significance of translated-DNA:protein searches like fastx36. Out-of-frame translations can produce cryptic low complexity regions, which are most apparent when the high-scoring sequences are shuffled. -z 21 shuffles are more efficient than individual sequence shuffles, because the set of high scoring sequences is shuffled 500 – 1,000 times, rather than 500 shuffles for each of 50 – 100 high scoring library sequences. It is almost as effective, because homologous sequences share similar amino-acid composition.

The statistical routines assume that the library contains a large sample of unrelated sequences.

 $<sup>{}^{1}</sup>E(D) = Dmn2^{-b}$ , where D is the number of sequences in the database, m, n are the lengths of the two sequences, and b is the bit score.

<sup>&</sup>lt;sup>2</sup>Random shuffles are performed for pairwise alignments and lalign36 by default.

>>GST26\_SCHMA Glutathione S-transferase class-mu (218 aa) initn: 422 init1: 359 opt: 407 Z-score: 836.8 bits: 162.0 E(437847): 3.7e-39 Smith-Waterman score: 451; 42.4% identity (73.4% similar) in 203 aa overlap (6-208:6-203) mGSTM1 MPMILGYWNVRGLTHPIRMLLEYTDSSYDEKRYTMGDAPDFDRSQWLNEKFKLGLDFPNLPYLIDGSHKITQSNAILRYL GST26\_ MAPKFGYWKVKGLVQPTR111eh1eetyeeRAY---DRNEIDA--WSNDKFKLGLEFPNLPYYIDGDFKLTQSMAIIRYI mGSTM1 ARKHHLDGETEEERIRADIVENQVMDTRMQLIMLCYNPDFEKQKPEFLKTIPEKMKLYSEFLGKRPWFAGDKVTYVDFLA GST26\_ ADKHNMLGACPKERAEISMLEGAVLDIRMGVLRIAYNKEYETLKVDFLNKLPGRLKMFEDRLSNKTYLNGNCVTHPDFML mGSTM1 YDILDQYRMFEPKCLDAFPNLRDFLARFEGLKKISAYMKSSRYIATPIFSKMAHWSNK :: :: .. .::. ::.: .: .: : .:. :..::::: :. GST26\_ YDALDVVLYMDSQCLNEFPKLVSFKKCIEDLPQIKNYLNSSRYIKWPLQGWDATFGGGDTPPK 

Figure 2: Alignment with -S filtered sequence

If the library contains fewer than 500 sequences (MAX\_RSTATS), then the library sequences are shuffled to produce 500 random scores, from which lambda and K statistical parameters are estimated. If the library contains a large number of *related* sequences, then the statistical parameters should be estimated by using the -z 11–15, options. -z options greater than 10 calculate a shuffled similarity score for each library sequence, in addition to the unshuffled score, and estimate the statistical parameters from the scores of the shuffled sequences.

#### 2.4.2 Looking at alignments

The description section described above contains the critical information for inferring homology, the E()-value. The alignment section shows the actual alignments that produced the similarity score and statistical estimates. In Fig. 1, the alignment display reports the percent identity, percent similarity (number of aligned residues with BLOSUM50 values  $\geq$  0), and the boundaries of the alignment. Note that for the ssearch36 and fasta36, the alignment shown can include residues that are not part of the best local alignment (e.g. residues 1–5 and 207–218 in mGSTM1 in Fig. 1). The amount of additional sequence context shown is the alignment line length (60 residues, set by -w len) divided by 2 by default, but can be adjusted with the -W context option.

Fig. 2 shows an example of a fasta36 alignment produced using the -S option to filter out lower-case (low complexity) residues. Here, additional scores (initn, init1 are shown, in addition to the opt score which is used to rank the sequences and calculate statistical significance. The init1 score is the highest scoring alignment without gaps; initn is a score that combines consistent (non-overlapping) runs without gaps, and opt is the score of a banded Smith-Waterman of width 16 for *ktup=2* that is applied to sequences with initn scores over the optimization threshold. In Fig. 2, the init1 score is based on the long, un-gapped region from residues 46–208 in mGSTM1, while the initn and opt scores include the other regions joined by gaps. The initn score is higher than the opt score, because it uses a simpler, length-independent, gap penalty.

The init1 and initn scores are shown for historical reasons, and can be used to illustrate the FASTA algorithm. But the opt score is the most reliable and sensitive score for inferring homology; the others can be ignored.

For fasta36 with proteins, the final alignment and score is calculated with the Smith-Waterman algorithm. For DNA sequences, a banded Smith-Waterman is used. (The -A option produces banded Smith-Waterman alignments for proteins, and full Smith-Waterman for DNA.) In Fig. 2, the opt score and Smith-Waterman scores are calculated on exactly the same alignment, but the opt score excludes the contribution from the "low-complexity" region between 19–30 in GST26\_SCHMA.

### 2.4.3 Results without alignments

While sequence alignments are very informative, it is often not practical to examine all the statistically significant alignments in large-scale searches. The -m 9 and -m 8 options present summaries of each alignment (alignment boundaries, percent identity, and other information) in a much more compact form. In addition, -m 9c or -m 9C (see options below) provide a detailed encoding of the alignment, that allows it to be reconstructed. For large-scale searches, we routinely use -m 8 with the -d 0 option, which sets the number of alignments shown to 0 (thus none are shown). Alternatively, the -m 8 and -m 8C ouput options produce BLAST-format tabular results summaries (-m 8C provides commented tabular results). -m BB produces an output that mimics BLAST output (with alignments).

### 2.5 Program Options

Command line options are available to change the scoring parameters and output display. Unlike the NCBI BLAST programs, command line options *must* precede the query file name and library file name arguments. To see the command-line options for a program and their defaults, type program\_name -help, e.g. fasta36 -help or ssearch36 -help. For a quick list of the most common options, just type the program name without any options (e.g. fasta36<ret>).

### 2.5.1 Command line options

- -a (fasta36, ssearch36, glsearch36, fasts36) show both sequences in their entirety.
- -A force Smith-Waterman alignments for fasta36 DNA sequences. By default, only fasta36 protein sequence comparisons use Smith-Waterman alignments. Likewise, for proteins, use band alignments (Smith-Waterman is used by default).
- -b # Number of sequence scores to be shown on output. In the absence of this option, fasta36 (and ssearch36) display all library sequences obtaining similarity scores with expectations less than the expectation (-E) threshold, 10.0 for proteins, and 2.0 for DNA:DNA and protein:translated DNA. The -b # option can limit the display further. There are two "submodes" of -b. -b =100 will force 100 high scores to be displayed, regardless of the expectation (-E) threshold, and -b >1 will show at least 1, but is otherwise limited by -E. Thus, -b 10 will show *no more than* 10 results, limited by -E; -b =10 will always show *exactly* 10 results, and -b >5 will show *at least* 5 results, but could show many more if more results have evalues ≤ -E e\_cut.

-c #,# (fasta36, [t]fast[x,y]36 only) Fraction of alignments optimized (second value is fraction of sequences joined). FASTA36 uses a statistical threshold strategy that joins and optimizes only the fraction of the alignments with an initn score expected -c times. Thus, -c 0.05 should optimize about 5% of sequences. The actual number of sequences optimized (and joined) is displayed in the scoring parameters line. Thus:

```
Parameters: BL50 matrix (15:-5), open/ext: -10/-2
ktup: 2, E-join: 1 (0.687), E-opt: 0.2 (0.294), width: 16
```

reports that 20% of the sequences in the database should have been band-optimized, and 29.4% were. Reducing the -c opt fraction improves performance, but dropping the fraction below 0.02 can reduce the accuracy of the statistical estimates.

-c 0 (letter 'O') sets the joining/optimization thresholds as they were prior to fasta-36.3.3 (original thresholds). Positive values set the thresholds to specific score values, as was the case in older versions of fasta.

- -C length of the sequence name printed at the beginning of alignment lines (default 6 characters).
- -d # Maximum number of alignments to be displayed (must be <= to the number of descriptions, -b #)
- -D Provide some debugging output. Used in conjunction with the -e expand\_script.sh, the link\_acc\_file and link\_lib\_file are not deleted during the run; so that expand\_script.sh scripts can be tested.
- -e expand\_script.sh Expand the set of sequences that are aligned to beyond the set of sequences searched. When the -e expand\_script.sh option is used, the expand\_script.sh script is run after the initial search scan but before the list of high-scoring sequences is displayed. expand\_script.sh is given a single argument, the name of a file that contains a list of accession strings (the text between the > and the first space ('\_') character followed by the E()-value for the sequence (separated by a <tab> character), e.g.:

gi|121719|sp|P08010|GSTM2\_RAT<tab>2.69e-86 gi|121746|sp|P09211|GSTP1\_HUMAN<tab>1.51e-20 gi|121749|sp|P04906|GSTP1\_RAT<tab>1.16e-19 gi|62822551|sp|P00502|GSTA1\_RAT<tab>9.5e-12

The script should produce a fasta-formatted list of additional sequences printed to stdout. The script is run with the command:

expand\_script.sh link\_acc.tmp\_file > link\_lib.tmp\_file

The sequences in link\_lib.tmp\_file (a temporary file name is actually used, and the file is deleted unless the -D option is used) are then compared and, if they are significant, included in the list of high scoring sequences and the alignments. The expanded set of sequences does not change the database size or statisical parameters, it simply expands the set of high-scoring sequences.

The fasta36/misc directory contains expand\_uniref50.pl that uses a mySQL table based on the uniref50 clusters. Using the script and the uniref50 cluster information, one can search uniref50.fasta, but then expand the hits so that uniprot appears to be searched.

- -E e\_cut [e\_cut\_r] Limit the number of scores and alignments shown based on the expected number of scores. Used to override the expectation value of 10.0 (protein:protein; 5.0 translated-DNA:protein; 2.0 DNA:DNA) used by default. -E 2.0 will show all library sequences with scores with an expectation value <= 2.0. With fasta-36, a second value, e\_cut\_r is available to limit the E()-values of additional sequence alignments between the query and library sequences. If not given, the threshold is e\_cut/10.0. If given with a value > 1.0, e\_cut\_r = e\_cut / value; for a value < 1.0, e\_cut\_r = value; If e\_cut\_r < 0, then the additional alignment option is disabled.</p>
- -f # Gap open penalty (-10 by default for proteins, -12 for DNA, -12 for [t]fast[xy]).
- -F # Limit the number of scores and alignments shown based on the expected number of scores.
   -E # sets the highest E()-value shown; -F # sets the lowest E()-value displayed. Thus,
   -F 0.0001 will not show any matches or alignments with E() < 0.0001. This allows one to skip over close relationships to search for more distant relationships.</li>
- -g # Penalty per residue in a gap (-2 by default for proteins, -4 for DNA, -2 for [t]fast[xy]). A single residue gap costs f + g.
- -h Short help message. Help options with ':', e.g. -s:, require an argument (-s BP62). Defaults are shown in square brackets, e.g.: -s: [BL50].
- -help Long help message
- -H Show histogram.
- -i DNA queries search with reverse complement. For tfastx36/y36, search the reverse complement of the library sequence only (complement of -3 option).
- -I Interactive mode (the default for versions older than fasta-36.3.4).
- -j # Penalty for frameshift between codons ([t]fastx36, [t]fasty36) and within a codon (fasty36/ tfasty36 only).
- -J (lalign36 only) show the identity alignment (normally suppressed, -I in versions before fasta-36.3.4).
- -k # number of shuffles for statistical estimates from shuffling.
- -1 file Location of library menu file (FASTLIBS).
- -L Display longer library sequence description.
- -M low-high Range of amino acid sequence lengths to be included in the search.
- -m # Specify alignment type: 0, 1, 2, 3, 4, 5, 6, 8, 9, 10, 11, B, BB, "F# out\_file"

| -m O       | -m 1       | -m 2       | -m 3       | -m 4       |
|------------|------------|------------|------------|------------|
| MWRTCGPPYT | MWRTCGPPYT | MWRTCGPPYT |            | MWRTCGPPYT |
| :::: :::   | xx X       | KSY        | MWKSCGYPYT |            |
| MWKSCGYPYT | MWKSCGYPYT |            |            |            |

If the -V '\*@%' annotation option has been used, annotations can be included in either the coordinate line (the default) or the middle alignment line (-m OM, -m 1M), or both (-m OB, -m 1B). See -V for more details.

-m 5: a combination of -m 4 and -m 0. -m 6 provides -m 5 plus HTML formatting. In addition, independent -m options can be combined. Thus, one can use -m 1 -m 6 -m 9.

- -m 8 provides BLAST tabular format output (a tab delimited line with the query name, library name, percent identity, and other alignment information). "-m 8C" provides some additional information provided by the BLAST tabular format with comment lines.
- -m 9 display alignment coordinates and scores with the best score information. -m 9i provides alignment length, percent identity, and percent similarity only. -m 9 extends the normal best score information:

 The best scores are:
 opt bits E(14548)

 XURTG4 glutathione transferase (EC 2.5.1.18) 4 ( 219) 1248 291.7 1.1e-79

to include the additional information (on the same line, separated by <tab> characters):

%\_id %\_gid sw alen an0 ax0 pn0 px0 an1 ax1 pn1 px1 gapq gapl fs 0.771 0.771 1248 218 1 218 1 218 1 218 1 219 0 0 0

The first two values are fraction identical and fraction similar (score  $\geq 0$ ), followed by the Smith-Waterman alignment score (sw), the alignment length (alen), and the coordinates of the beginning and end of the alignment in the query and target (library) sequences (an0 beginning, ax1 end in query; an1 beginning, ax1 end in target/library), and the coordinate system for the beginning and end of the query and target/library sequence (pn0 is the displayed coordinate of the first residue of the query sequence, px0 is the displayed coordinate of the last residue, pn1,px1 provide the coordinates for the target/library sequence). gapq, gap1 report the number of gaps in the query and library sequence; fs reports the number of frameshifts.

 $-{\tt m}$   $_{\rm 9c}$  provides additional information: an encoded alignment string. For example, the alignment:

10 20 30 40 50 60 70 GT8.7 NVRGLTHPIRMLLEYTDSSYDEKRYTMGDAPDFDRSQWLNEKFKL--GLDFPNLPYL-IDGSHKITQ .... . .. . . .... : .: · · . · XURTG NARGRMECIRWLLAAAGVEFDEK-----FIQSPEDLEKLKKDGNLMFDQVPMVEIDG-MKLAQ 20 30 40 50 60

would be encoded: =23+9=13-2=10-1=3+1=5. The numbers in the alignment encoding is with repect to the beginning of the alignment, not the sequences. The beginning coordinate of the alignment is given earlier in the -m 9c line. -m 9C provides the alignment encoding in CIGAR format: 28M9D13M2I10M1I3M1D5M.

(June, 2014) The -m 9c/C option has been extended to -m 9d/D, which encodes the positions of mismatches as well as insertions and deletions. For the example above, the -m 9d encoding would be: =1x1=2x4=2x1=2x7=3-9=1x2=1x4=2x1=1x1=1x1=1x3=1x2+1=3-1x1=1x2=1 while -m 9D would be: 1M1X2M4X2M1X2M7X3M9D1M2X1M4X2M1X1M1X211X1M1X1M3X1M2X1I3M1D1X1M2X1M

-m 10 a parseable format for use with other programs.

-m 11 Provide lav-like output (used by lalign) for graphical output.

lalign36 -m 11 mchu.aa mchu.aa | lav2plt.pl --dev ps > mchu\_laln.ps

Produces a postscript plot of the local alignments. Likewise, lav2plt.pl --dev svg produces SVG output.

- -m BB Format output to mimic BLAST format. -m B formats alignments to look like BLAST alignments (Query/Sbjct), but is FASTA output otherwise. -mBB imitates BLAST as much as possible, and cannot be used with other -m options.
- -m "F# out.file" Send an alternate result format to out.file. Normally, the -m out\_fmt option applies to the default output file, which is either stdout, or specified with -0 out\_file (or within the program in interactive mode). With -m F, an output format can be associated with a separate output file, which will contain a complete FASTA program output. Thus,

```
ssearch36 -m 9c -m "FBB blast.out" -m "F9c,10 m9c_10.out" query library
```

Sends the -m 9c output to stdout, but will also send -m BB output to the blast.out file, and -m 9c -m 10 output to m9\_c10.out. Consistent -m out\_fmt commands can be set to the same file by separating them with ','. Producing alternative format alignments in different files has little additional computational cost.

Because a space ( $_{-}$ ) is used to separate the output format (-m) values from the file name, the -m F argument must typically be surrounded by quotation marks (").

One of the shortcomings of this approach is that it affects only the output format, not the other options that modify the amount of output. Thus, if you specify -E 0.001; that expect threshold will be used for all the output files. When a -m option does modify the output (e.g. -m 8 sets -d 0), that modification is specific to the output file.

- -M low-high Include library sequences with lengths between low and high.
- -n Force the query sequence to be treated as a DNA sequence. Useful when query sequences contain a large number of ambiguous residues, e.g. transcription factor binding sites.
- -N # break long library sequences into blocks of # residues. Useful for bacterial genomes, which have only one sequence entry. -N 2000 works well for well for bacterial genomes. (This option was required when FASTA only provided one alignment between the query and library sequence. It is not as useful, now that multiple alignments are available.)
- -o off1,off2 (Previously -X.) Specifies offsets for the beginning of the query and library sequence. For example, if you are comparing upstream regions for two genes, and the first sequence contains 500 nt of upstream sequence while the second contains 300 nt of upstream sequence, you might try:

```
fasta -o "-500 -300" seq1.nt seq2.nt
```

If the -0 option is not used, FASTA assumes numbering starts with 1. (You should double check to be certain the negative numbering works properly.)

- -0 Send a copy of results to filename. Helpful for environments without STDOUT, but should be avoided (use > filename instead).
- -p Force query to be treated as protein sequence.

-P PSSM\_file Specify a PSI-BLAST format PSSM (Position Specific Scoring Matrix) file. ssearch36, ggsearch36, and glsearch36 can use a PSSM file to improve the sensitivity of a search. The FASTA programs accept two PSSM file formats:

| format | blastpgp  | option        |
|--------|---|---------------|
| 0<br>2 | blastpgp -C pssm.chk -u 0<br>blastpgp -C pssm.asnb -u 2 | •             |
| 2      | prastpgp -t pssm.asnb -u 2                              | Dinary ASN. I |

which can be specified after the file name, e.g.:

ssearch36 -P 'pssm.asnb 2' pssm\_query.aa +sp+

Searches with a PSI-BLAST PSSM must still require a query sequence file, and the query sequence file must match the PSSM seed sequence. The format 0 byte-encoded PSSM is machine dependent; it must be created by blastpgp on the same architecture as ssearch36. In general, you should use the binary ASN.1 (format 2) file.

With the release of NCBI-BLAST+, psiblast replaces blastpgp, and psiblast does not produce the binary ASN.1 PSSM checkpoint data. However, the text ASN.1 PSSM checkpoint file (produced with the psiblast option -out\_pssm) can be converted to a binary ASN.1 format that ssearch36 can read using the NCBI datatool program (available from ftp: //ftp.ncbi.nlm.nih.gov/toolbox/ncbi\_tools++/BIN/CURRENT/datatool) together with http://www.ncbi.nlm.nih.gov/data\_specs/asn/NCBI\_all.asn. More information about datatool is available from http://www.ncbi.nlm.nih.gov/data\_specs/NCBI\_data\_conversion. html. The NCBI BLASTP/PSI-BLAST website provides the same PSSM text ASN.1 file with the downloads link. A text ASN.1 PSSM file can be converted to a binary ASN.1 file using the command:

datatool -m NCBI\_all.asn -v pssm.asn\_txt -e pssm.asnb

The pssm.asnb can then be used with ssearch36 with the -P 'pssm.asnb 2' option shown above.

- -Q,-q Quiet does not prompt for any input. Writes scores and alignments to the terminal or standard output file (on by default, turned off with -I).
- -r +n/-m Specify match/mismatch scores for DNA comparisons. The default is +5/-4. +3/-2 can perform better in some cases.
- -R file Save a results summary line for every sequence in the sequence library. The summary line includes the sequence identifier, superfamily number (if available) position in the library, and the similarity scores calculated. This option can be used to evaluate the sensitivity and selectivity of different search strategies [13, 15].
- -s file Specify the scoring matrix file. fasta36 uses the same scoring matrice format as Blast. Several scoring matrix files are included in the standard distribution in the data/ directory. For protein sequences: codaa.mat - based on minimum mutation matrix; idnaa.mat - identity matrix; pam250.mat - the PAM250 matrix; [4], (-s P250), and pam120.mat - a PAM120 matrix (-s P120). The default scoring matrix is BLOSUM50 (-s BL50). Other matrices include a series of modern PAM-based matrices [8]: MDM40/-s MD40, MDM20/-s MD20, and MDM10/-s MD10, and a selection from the BLOSUM series [6] BLOSUM50, 62, and 80/-s

BL50, -s BL62, -s BL80. -s BP62 sets the scoring matrix to BLOSUM62 and the gap penalties to -11/-1, identical to BLASTP. In addition, the VTML160 matrix (-s VT160) [12] and OPTIMA\_5 (-s OPT5) [9] are available.

If the scoring matrix is prefaced by a question mark, e.g. ?BP62, then the scoring matrix is adjusted for each query to ensure that a 100% identical match can produce a score of at least 40 bits. This is designed for fastx36 searches with potentially short DNA queries; A 120 nt DNA query can only produce a 40 amino-acid alignment, which, with BLOSUM62 -11/-1, cannot produce more than 23 bits of score. A scoring matrix with a higher information content is required; in the set available by default, MD40, with 2.22 bits/position, would be used. For more information about alignment length and information content, see [1].

-S Filter out lower-case characters in the query or library sequences for the initial score calculation (used to filter low-complexity – seg-ed – residues). The pseg program [22] can be used to lower-case mask low complexity regions in protein sequences. With the –S option, lower case characters in the query or database sequences are treated as X's during the initial scan, but are treated as normal residues during the final alignment display. Since statistical significance is calculated from the similarity score calculated during the library search, the lower case residues do not contribute to the score. However, if a significant alignment contains low complexity regions, the residues are shown (as lower case characters, Fig. 2).

The pseg program can be used to produce databases (or query sequences) with lower case residues indicating low complexity regions using the command:

pseg ./swissprot.fasta -z 1 -q > swissprot.lseg

The -S option should always be used with FASTX/Y and TFASTX/Y because out-of-frame translations often generate low-complexity protein sequences. However, only lower case characters in the protein sequence (or protein database) are masked; lower case DNA sequences are translated into upper case protein sequences, and not treated as low complexity by the translated alignment programs. (There is an option in the Makefile, -DDNALIB\_LC, to enable preserving case in DNA sequences.)

-t # Translation table - fastx36, tfastx36, fasty36, and tfasty3 now support the BLAST translation tables. See http://www.ncbi.nih.gov/Taxonomy/Utils/wprintgc.cgi.

-t t or -t t# enables the addition of an implicit termination codon to a protein:translated DNA match. That is, each protein sequence implicitly ends with \*, which matches the termination codes for the appropriate genetic code. -t t# sets implicit termination and a different genetic code.

- -T # set number of threads/workers. Normally on a multi-core machine, the maximum number of processors/cores is used.
- -U Treat the query sequence an RNA sequence. In addition to selecting a DNA/RNA alphabet, this option causes changes to the scoring matrix so that G:A, T:C or U:C are scored as G:G -3.
- -v # Do window shuffles with the window size specified.
- -V str Specify annotation characters that can be included (and will be ignored), in the query sequence file, but are displayed in the alignments. If a query file contains "ACVS\*ITRLFT?",

where "\*" and "?" are used to indicate phosphorylation, giving the option -V '\*?', the annotated characters in the query will (S\*, F?) will be highlighted in the alignment (on the number line). A fasts36 alignment of seq/ngts.aa compared to seq/mgstm1.aa with -V '\*?' produces:

|       | *               | 1         | .0??       |            |              |       |
|-------|-----------------|-----------|------------|------------|--------------|-------|
| GT8.7 | ILGYWN          | ЕҮТ       | DSSYDEKR   |            |              |       |
|       | :::::           |           | :::::::    |            |              |       |
| GT8.7 | MPMILGYWNVRGLTH | PIRMLLEYT | DSSYDEKRYT | MGDAPDFDRS | SQWLNEKFKLGI | DFPNL |
|       | 10              | 20        | 30         | 40         | 50           | 60    |

In addition to showing the alignments of post-translationally modified sites, the -V option can be used to highlight active sites in library sequences. In the -m 9c output, the state of the annotated sites is summarized when -V is used.

(fasta-36.3.6 June 2012) The -V option has been extended to: (1) allow feature descriptions to be specified in a file, e.g. -V =annot.defs where annot.defs contains:

```
*:phosphorylation
@:active site
^:binding site
```

>accession1

The annotation character is left of the '.', the definition is on the right. The annot.defs file can also be specified by setting the FA\_ANNOT\_DEF environment variable to the file name;

(2) to include optional annotation file, e.g. -V '<features.annot', or script, e.g. -V '!features.pl' for library annotations and -V 'q!features.pl' for query annotations. (Some shells require \!features.pl.) Similar to the library expansion script, the features.pl script is run against a temporary file containing the list of high scoring sequence accessions (the text before the first space), e.g.

gi|121735|sp|P09488.3|GSTM1\_HUMAN gi|1170096|sp|Q03013.3|GSTM4\_HUMAN gi|67461004|sp|Q5R8E8.3|GSTM2\_PONAB ...

The features.pl script then produces a file of annotations on those sequences, in the format:

```
position label value
>accession2
...
For example:
>gi|121735|sp|P09488.3|GSTM1_HUMAN
23 *
33 *
34 *
116 ^
173 V N
210 V T
>gi|1170096|sp|Q03013.3|GSTM4_HUMAN
2 V P
116 ^
```

160 V V 208 V V 209 V F 211 V K 212 V M >gi|67461004|sp|Q5R8E8.3|GSTM2\_PONAB ....

The same format is used for the -V '<feature.annot' file.

The V label is special; it indicates that the feature is a variant residue and specifies the alternative residue in the label field. Thus,  $GSTM4\_HUMAN$  can have a M at position 2. Unlike modification or active site annotations, variant residues can change the sequence of the library sequence if replacing the canonical library residue with the variant residue improves the score. Thus, without the -V '!feature.pl' script, the human GSTM1B variant with db-SNP:rs449856 would align to GSTM1\_HUMAN (P04988) like this (the -m 1 format option was used to highlight differences) :

The best scores are: opt bits E(1) sp|P09488.3|GSTM1\_HUMAN Glutathione S-transferase Mu 1; GST HB subuni (218) 1490 335.9 3.7e-97 >>sp|P09488.3|GSTM1\_HUMAN Glutathione S-transferase Mu 1; GST HB s (218 aa) initn: 1490 init1: 1490 opt: 1490 Z-score: 1776.8 bits: 335.9 E(1): 3.7e-97 Smith-Waterman score: 1490; 99.1% identity (100.0% similar) in 218 aa overlap (1-218:1-218) . . . 170 180 190 200 210 gtm1\_h YDVLDLHRIFEPNCLDAFPNLKDFISRFEGLEKISAYMKSSRFLPRPVFTKMAVWGNK х sp|P09 YDVLDLHRIFEPKCLDAFPNLKDFISRFEGLEKISAYMKSSRFLPRPVFSKMAVWGNK 180 190 200 170 210 With a -V 'feature.pl' script to annotate the variants, the alignment becomes: The best scores are: opt bits E(1) sp|P09488.3|GSTM1\_HUMAN Glutathione S-transferase Mu 1; GST HB s (218) 1500 338.1 8e-98 >>sp|P09488.3|GSTM1\_HUMAN Glutathione S-transferase Mu 1; GST HB s (218 aa) Variant: K173N;S210T; initn: 1500 init1: 1500 opt: 1500 Z-score: 1788.7 bits: 338.1 E(1): 8e-98 Smith-Waterman score: 1500; 100.0% identity (100.0% similar) in 218 aa overlap (1-218:1-218) . . . 170 180 190 200 210 gtm1\_h YDVLDLHRIFEPNCLDAFPNLKDFISRFEGLEKISAYMKSSRFLPRPVFTKMAVWGNK

sp|P09 YDVLDLHRIFEPNCLDAFPNLKDFISRFEGLEKISAYMKSSRFLPRPVFTKMAVWGNK 170 V 180 190 200 21V

In addition to removing the two differences as residues 173 and 210, which produces a 100% identical alignment, inclusion of the variant library sequence also improves the raw similarity score and E()-value. An example script (misc/up\_feats.pl) that extracts annotations from a mysql database of Uniprot features is provided.

If the annotation script produces lines beginning with '=', then these lines are taken as annotation definitions, similar to the annot.defs file described above. Thus:

=\*:phosphorylation =@:active site

```
=^:binding site
>gi|121735|sp|P09488.3|GSTM1_HUMAN
23 *
33 *
34 *
116 ^
173 V N
210 V T
```

will produce the same annotation descriptions as the annot.defs file.

Scripts to produce annotations are available in the ../scripts/directory as ../scripts/ann\_feats\*.pl. These scripts require a MySQL database that associates features or domains with sequence identifiers (accessions).

- -w # Display width value (<200). Sets the approximate width of the high-score descriptions and the length of residue alignments. -w 60 by default.
- -W # context length (default is 1/2 of line width -w) for alignment, for programs like fasta36 and ssearch36, that provide additional sequence context.
- -X extended\_option A number of rarely used options are now only available as extended options:
  - X1 sort output by init1 score (for compatibility with FASTP; obsolete).
  - XB (Previously -B.) Show the z-score, rather than the bit-score in the list of best scores (rarely used, provided for backward compatibility).
  - XI Modify rounding used in percent identity/percent similarity display to ensure that sequences that have a mismatch are not shown as 100.0% identical. Without this option, a single mismatch in a 10,000 residue alignment would be shown as 100.0% identical; with this option, it would be shown as 99.9% identical.
  - Xo (fasta36, [t]fast[x/y]36 only) (Previously -o.) Turn off the default opt score calculation and sort results by initn scores (reduces sensitivity and statistical accuracy, obsolete).
  - XM The maximum amount of memory available for storing the library in multi-sequence searches. The value is specified in MBytes (-XL16) or GBytes (-XL4G) and can also be set using the LIB\_MEMK environment variable (LIB\_MEMK=4G). Negative values remove the memory restriction. By default (set as a compile-time option, -DMAX\_MEMK=2), set to 2 GBytes in 32-bit environments and 12 GBytes in 64-bit environments.
  - XN/XX Alter the treatment of N:N (DNA) or X:X (protein) alignments for counts of identities and similarities. By default the FASTA programs count N:N or X:X as identical, but not similar, because their alignment scores are typically negative. -XNS, -XN+, -XXS, and -XX+ treat N:N and X:X alignments as "similar", even though their alignment scores are negative, when calculating percent similarity. -XND, -XN-, -XXD, and -XX- treat N:N and X:X alignments as non-identical for calculating percent identity.
  - Xx (Previously -x) Specify the penalty for a match to an X, and mismatch to X, independently of the PAM matrix. Particularly useful for fastx3/fasty36, where termination codons are encoded as X. For example, -Xx=0,-1 scores an X:X match as 0, and X:not-X as -1.

- Xy (Previously -y.) Set the width of the band used for calculating "optimized" scores. For proteins and ktup=2, the width is 16. For proteins with ktup=1, the width is 32 by default. For DNA the width is 16.
- -z -1,0,1,2,3,4,5,6

-z -1 turns off statistical calculations. z 0 estimates the significance of the match from the mean and standard deviation of the library scores, without correcting for library sequence length. -z 1 (the default) uses a weighted regression of average score vs library sequence length; -z 2 uses maximum likelihood estimates of  $\lambda$  and K; -z 3 uses Altschul-Gish parameters [2]; -z 4 - 5 uses two variations on the -z 1 strategy. -z 1 and -z 2 are the best methods, in general.

-z 11,12,14,15,16

estimate the statistical parameters from shuffled copies of each library sequence. This allows accurate statistics to be estimated for libraries comprised of a single protein family.

-z 21,22,24,25,26

estimate the statistical parameters from shuffled copies of the highest scoring sequences reported in the search. library sequence. This shuffling strategy is much more like prss, since the sequences shuffled share compositional similarity to the query.

-Z db\_size sets the apparent size of the database to be used when calculating expectation E()-values. If you searched a database with 1,000 sequences, but would like to have the E()-values calculated in the context of a 100,000 sequence database, use -Z 100000.

-3 translate only three forward frames or search with only the forward strand (complement of -i).

Thus, to tell fasta36 to align seq1.aa with seq2.aa showing the entirety of both sequences, with 80 characters per line, one would type:

fasta36 -w 80 -s BP62 -a seq1.aa seq2.aa

The -w 80 and -a options must precede the file names. If you just enter the options on the command line followed by -I, the program will prompt for the file names.

In addition, the FASTA programs can accept query sequence data from STDIN. To specify that stdin be used as the query or library file, the file name should be specified as @. Thus:

cat query.aa | fasta36 @:25-75 /slib/swissprot

would take residues 25-75 from query.aa and search the /slib/swissprot.

#### 2.5.2 Environment variables

FASTA allows virtually every option to be set on the command line (except the *ktup*, which must be set as the third command line argument), but it is often convenient to set the FASTLIBS environment variable to specify the location of the fastlibs database description file.

FASTLIBS – FASTLIBS specifies the location of the file that contains the list of library descriptions, locations, and library types (see section on finding library files). LIB\_MEMK – Set the maximum amount of memory (MBytes) to be available for library buffering (equivalent to -XM#, see above). By default, 2GB is available on 32-bit systems (LIB\_MEMK=2G); 8GB on 64-bit systems.

REF\_URL, SRCH\_URL and SRCH\_URL1 — These environment variables are used in HTML mode (-m 6) to provide links from the sequence alignment (see the links at http://fasta.bioch. virginia.edu/fasta\_www2/). REF\_URL is associated with the Entrez Lookup link; SRCH\_URL with the Re-search database link, and SRCH\_URL1 with the General re-search link. In each case, the text corresponds to a HTML URL, but with positions containing the %s or %ld (for numbers) part of a 'C' sprintf() call for specific variables. REF\_URL uses the database (protein or nucleotide), together with a query term (typically the gi number). SRCH\_URL and SRCH\_URL1 use db, query (gi, pgm (fa, ss, fx, etc.), and start, stop, and n1 (library sequence length), where start and stop are the boundaries of the alignment, for sub-sequence searches. The values of these environment variables are used with sprintf to build a new URL that is linked in the output.

TMP\_DIR - Location (if defined) of the temporary files used by the -e expand\_script.sh option. In addition, environment variables can be used inside both the fastlibs file and in the @db.nam files of file names. The fasta36/conf/fast\_libs\_e.www file, included with the distribution, shows an example, as do the descriptions of file of file names files shown below. Whenever a word of the form \${WORD} is found in fastlibs or a file of file names, the \${WORD} environment variable is expanded and inserted in the string. Thus, if <\${SLIB}/blast\_dbs/ describes where a list of files will be found and \${SLIB} is "/seqdata", then the resulting substitution yields: </seqdata/blast\_dbs/.

# 3 Installing FASTA and the sequence databases

#### 3.1 Obtaining/preparing the sequence libraries

The FASTA program package does not include any protein or DNA sequence libraries. Protein and DNA sequence databases are available via anonymous FTP from the NCBI (ftp://ftp. ncbi.nih.gov/blast/db), ftp://ftp.ncbi.nih.gov/blast/db), UniProt (ftp://ftp.uniprot. org/pub/databases/uniprot), and the EBI (ftp.ebi.ac.uk/pub/databases).

Protein Sequence Databases – Protein sequence databases are available from the NCBI, UniProt, and the EBI. The NCBI provides a "raw" database, nr, and a well-curated, less redundant database, refseq\_protein, and a copy of the very well annotated swissprot database. Protein sequence databases can also be downloaded from UniProt and the EBI; both sites provide the same UniProt [3] database.

Protein libraries, particularly those used for translated-DNA:protein comparisons with fastx36 or fasty36, show be scanned to remove low-complexity regions. Matches between low complexity regions can violate the composition assumptions used by the FASTA statistical estimates. The pseg program ([22], ftp://ftp.ncbi.nih.gov/pub/seg/pseg) can be used to lower-case low complexity regions, which then can be ignored during the initial database search by using the -S option. To lower-case low complexity regions, run the pseg program against the protein sequence database:

```
pseg /seqdata/swissprot.fa -z 1 -q > /seqdata/swissprot.lseg
```

And then you can run most FASTA programs with -S:

ssearch36 -S mgstm1.aa /seqdata/swissprot.lseg

Fig. 2 shows the effect of including the -S option with lower-cased low-complexity sequences. The opt score (407), which is used to sort the results and calculate statistics, is lower than the Smith-Waterman score (451), even though exactly the same residues are aligned for each score. The opt score excludes residues 19-30, because they were marked as low-complexity by pseg; thus they are shown as lower-case. The Smith-Waterman score includes the contribution from that part of the alignment.

Out-of-frame translated DNA sequences often produce low-complexity regions [17], so it is particularly important to avoid low-complexity alignments when using fastx36 and fasty36

### 3.2 Searching taxonomic subsets

Because increasing database size reduces search sensitivity (an alignment with an E()-value of 0.001 in a search of a 100,000 entry database will have an E()-value of 0.1, not significant, if found in a database of 10,000,000 sequences), it is much more effective to search smaller, less redundant databases (you can always search the larger database later). Thus, the refseq\_protein database from the NCBI is preferred over nr; even better are databases that reflect a limited phylogenetic range (e.g. refseq\_human for vertebrate sequences).

While the NCBI provides organism-specific refseq subsets on their FTP site, they can be difficult to find. Alternatively, you can use the NCBI Entrez web site to download a list of gi numbers specific to a particular organism or taxonomic range. The FASTA programs can search a subset of a large sequence database that is specified by a list of gi numbers by using library format 10. For example, given a list of gi numbers for the human proteins in swissprot.lseg, the file sp\_human.db, with the content:

```
<${SLIB}/swissprot.lseg 0:2 4|
3121763
51701705
7404340
205831112
74735515
```

could be used to search the human subset of swissprot.lseg. The gi numbers for the Swiss-Prot entries begin with the second line. The first line specifies the location of the file where the sequences containing the gi numbers can be found (\${SLIB}/swissprot.lseg, the libtype of that file (0:fasta), the character offset to the beginning of the sequence identifier in that file (2), the identifier type (4), and the character that separates the fields in the FASTA descriptor (1). The identifier type can take four formats:

- 1 ordered accession strings (letters or numbers)
- 2 ordered numbers (digits only)
- 3 un-ordered accession strings
- 4 un-ordered numbers

(Ordered accession strings/numbers are ordered in both the library and the subset file.)

Thus, given the 0:2 4| specification above, the line:

```
>gi|3121763|sp|015143.3|ARC1B_HUMAN Actin-related protein 2/3 ...
```

would be parsed, looking for an number starting at column 4 (the first column is numbered 0), and ending with |. The order of sequences in the library do not have to correspond to the order in the sp\_human.db file (un-ordered). Given a the sp\_human.db file, a file swissprot.lseg in the directory specified by the environment variable \${SLIB}, and a command of the form:

fasta36 -S mgstm1.aa 'sp\_human.db 10'

Would use the sp\_human.db file to search the subset of swissprot.lseg that contained the specified gi numbers.

#### 3.3 DNA sequence libraries

Because of the large size of DNA databases, you will probably want to keep DNA databases in only one format. The FASTA3 programs that search DNA databases — fasta36, fastm36, and tfastx/y36 — can read DNA databases in Genbank flatfile (not ASN.1), FASTA, and BLAST2.0 (formatdb) formats, as well as EMBL format. BLAST2.0 format is preferred for DNA sequence libraries, because the files are considerably more compact than GenBank format. The NCBI does not provide software for converting from Genbank flat files to Blast2.0 DNA databases, but you can use the Blast formatdb program to convert ASN.1 formatted Genbank files, which are available from the NCBI ftp site.

The NCBI also provides the comprehensive nt DNA database, and several EST databases in Blast2.0/formatdb format from ftp://ncbi.nih.gov/blast/db.

#### 3.4 Finding the library files

All the FASTA programs comparison programs have the command line syntax:

```
fasta36 query.file /seqdata/library
```

However, in addition to simply specifying the location of the database to be searched (/seqdata/library), the FASTA programs provide several methods for referring to sequence databases without specifying a specific file. These methods can be used to provide abbreviations for sequence libraries, e.g.:

```
fasta36 query.file s Or fasta36 query.file +sp+
```

To use abbreviations like 's' or '+sp+' to reference a sequence database, a FASTLIBS file must be used, see section 3.5.

Large DNA and protein databases are often distributed across several files. For example, the NCBI nr protein database is found in 5 files, nr.00 ... nr.04. To search databases in multiple files, the names of the files are specified in a file of filenames, nr.nam:

<\${SLIB}/blast\_dbs/ nr.00 12 nr.01 12 nr.02 12 nr.03 12 nr.04 12 In this file, the first line <\${SLIB2}/blast\_dbs/, beginning with <, specifies the location and format (Blast2.0 formatdb) the data files. Text of the form \${SLIB} refers to Unix/MacOSX/Windows environment variables; the value of \${SLIB} is set by a Unix/MacOSX shell environment command. Thus, if the value of \${SLIB} is /seqdata, then the first sequence library file to be read will be /seqdata/blast\_dbs/nr.00, in format 12 (Blast2.0 formatdb).

To refer to the nr.nam file as a file of file names, it must be prefixed by a @ character, e.g.

fasta36 query.file @nr.nam

Files of file names can contain references to other files of file names:

<\${SLIB}/fasta\_dbs/ @pdb.nam @swissprot.nam

The FASTA file of file names is similar to the NCBI prot\_db.pal and dna\_db.nal, files, but unfortunately they are different, and currently FASTA cannot read NCBI .pal or .nal files that contain a DBLIST line. FASTA can read NCBI .pal or .nal files that do not contain a DBLIST line.

FASTA version fasta-36.3.6 provides an alternative way to generate a database to be searched: the !script.sh file. Like the -e expand\_file.sh script, a shell script or program can be used to produce a database to a temporary file, which is then seached. For example, if the file cat\_db.sh contains the command echo /seqdb/swissprot.lseg, the command:

fasta36 query.aa \!@cat\_db.sh

will cause cat\_db.sh to produce a temporary file with the line swissprot.lseg, which is interpreted as an indirect file of filenames; thus, because of the @, the file will be interpreted as an indirect file, and the swissprot.lseg file will be searched. Note that on Unix systems, the '!' must be preceeded by a '\' so that it is not interpreted by the shell, as shown above.

#### **3.5** FASTLIBS

All the search programs in the FASTA3 package can use the environment variable FASTLIBS to find the protein and DNA sequence libraries. (Alternatively, you can specify the FASTLIBS file with the -l fastlibs.file option.) The FASTLIBS variable contains the name of a file that has the actual filenames of the libraries. The fastlibs file included with the distribution is an example of a file that can be referred to by FASTLIBS. To use the fastlibs file, type:

```
setenv FASTLIBS /seqdata/info/fastgbs (csh/tcsh)
or
export FASTLIBS=/seqdata/info/fastgbs (bash/ksh)
```

Then edit the fastlibs file to indicate the location of the protein and DNA sequence libraries. If the protein sequence library is kept in the file /seqdata/aa/swissprot.lseg and your Genbank DNA sequence library is kept in the directory: /seqdata/genbank, then the fastlibs file might contain:

```
SwissProt$OP/seqdata/aa/swissprot.lseg 0
UniProt$O+uniprot+@/seqdata/aa/uniprot.nam
```

The first line of this file says that there is a copy of the SwissProt sequence database (a protein database) that can be selected by typing "P" on the command line or when the database menu is presented in interactive mode.

Note that there are 4 (or 5) fields in the lines in the fastlibs file. The first field describes library and is displayed by FASTA program; it ends with the '\$'. The second field (1 character), is a 0 if the library is a protein library and 1 if it is a DNA library. The third field can either be a single character (P) or a word surrounded by the + symbol (+uniprot+), and can be used to specify the library on the command line or in interactive mode.

The fourth field is the name of the library file. In the example above, the /seqdata/aa/swissprot.lseg file contains the entire protein sequence library. Alternatively, /seqdata/aa/uniprot.nam is a file of file names, which contains a list of one or more library files. Likewise, the DNA library files are files of file names.

In addition, an optional fifth field can be used to specify the format of the library file. Alternatively, you can specify the library format in a file of file names. This field must be separated from the file name by a space character (' ') from the filename. FASTA can read the libraries in the following formats:

- 0 FASTA (>SEQID comment/sequence)
- 1 Uncompressed Genbank (LOCUS/DEFINITION/ORIGIN)
- 2 NBRF CODATA (ENTRY/SEQUENCE) (obsolete)
- 3 EMBL/SWISS-PROT (ID/DE/SQ)
- 4 Intelligenetics (;comment/SEQID/sequence) (obsolete)
- 5 NBRF/PIR VMS (>P1; SEQID/comment/sequence) (obsolete)
- 6 GCG (version 8.0) Unix Protein and DNA (compressed)
- 7 FASTQ (sequence only, quality ignored)
- 10 subset format (j/slib2/swissprot.lseg 0:2 4---)
- 11 NCBI Blast1.3.2 format (unix only) (obsolete)
- 12 NCBI Blast2.0 format
- 16 MySQL (requires special compilation)
- 17 Postgres (requires special compilation)

Today, the most popular formats are FASTA, type '0', the default, and the NCBI Blast2.0 formatdb formats (type '12'). The FASTA programs cannot read NCBI ASN.1 formatted databases. If a library format is not specified, for example, because you are just comparing two sequences, FASTA (format 0) is used by default. To specify a library type on the command line, add it to the library filename and surround the filename and library type in quotes:

```
fasta36 query.file "/seqdb/genbank/gbmam 12"
```

NCBI formatdb databases are built from multiple files, e.g. gbmam.nsq, gbmam.nhr, gbmam.nin; to refer to the complete set of files, simply use name before the suffixes, e.g. gbmam. When NCBI databases distributed across several files, e.g. gbbct.00, gbbct.01, etc, those files must be included in a gbbct.nam file of file names.

FASTA subset format (10) allows users to search a subset of a sequence database, by specifying a list of gi numbers or accessions in a larger database. The format begins with a line naming the file sequence file followed by information about how to extract the gi number or accession. Thus, the line.

<library\_file lib\_fmt:id\_fmt id\_loc</pre>

where lib\_fmt is the library format (0), id\_fmt is the format of the sequence identifier (:1, :2 - ordered strings or numbers; :3, :4 - unordered strings or numbers), and id\_loc is the location of the sequence identifier. For example,

```
</slib2/blast/swissprot.lseg 0:2 4|
3121763
51701705
7404340
74735515
...
```

specifies the file containing all the sequences and the file is in FASTA format ('0:'), the sequence identifier is a number (':2'), and the identifier starts at character 4 and ends with the '|' symbol.

The major problem that most new users of the FASTA package have is in setting up the program to find the databases and their library type. In general, if you cannot get fasta36 to read a sequence database, there is probably something wrong with the FASTLIBS file. A common problem is that the database file is found, but either no sequences are read, or an incorrect number of entries is read. This is almost always because the library format (libtype) is incorrect.

Test the setup by running FASTA. Enter the sequence file 'mgstm1.aa' when the program requests it (this file is included with the programs). The program should then ask you to select a protein sequence library. Alternatively, if you run the tfastx36 -I program and use the mgstm1.aa query sequence, the program should show you a selection of DNA sequence libraries. Once the fastlibs file has been set up correctly, you can set FASTLIBS=fastgbs in your AUTOEXEC.BAT file, and you will not need to remember where the libraries are kept or how they are named.

# 4 Frequently Asked Questions (FAQs)

Where can I get FASTA? - http://faculty.virginia.edu/wrpearson/fasta has the latest versions of the FASTA programs. This document describes fasta-36.3.6, which is available from http://faculty.virginia.edu/wrpearson/fasta/fasta3.tar.gz. In addition, pre-compiled versions of the programs are available for MacOSX and Windows.

| QueryLibraryFASTA pgm.BLAST pgm.Prot.Prot.fasta36<br>ssearch36<br>ggearch36<br>ggearch36<br>ggearch36<br>blastnblastpheuristic local similarity<br>optimal local sim.<br>global:global sim.<br>global:local sim.<br>global:local sim.DNADNAfasta36*blastnProt.Prot.lalign36multiple non-intersecting<br>alignmentsDNAProt.fastx36<br>fasty36blastxProt.DNAfastx36<br>fasty36blastxProt.DNAtfastx36<br>tfasty36blastxProt.DNAtfastx36<br>tfasty36blastnProt.Prot.fasts36<br>unordered peptides<br>Unordered peptides<br>Unordered peptidesProt.Prot.fasts36<br>fasts36<br>   |       |         |                        |            |  |
|---|-------|---------|------------------------|------------|--|
| NomProtectionProt | Query | Library | FASTA pgm.             | BLAST pgm. |  |
| DNADNAfasta36*blastnProt.Prot.lalign36multiple non-intersecting<br>alignmentsDNADNAlalign36trans.DNAProt.fastx36<br>fasty36blastxDNAProt.fastx36<br>fasty36blastxProt.DNAtfastx36<br>tfasty36blastnProt.DNAtfastx36<br>tfasty36blastnProt.DNAtfastx36<br>tfasts36<br>DNAblastnProt.Prot.fasts36<br>tfasts36<br>DNAUnordered peptides<br>Unordered peptides<br>Unordered peptides<br>Ordered peptides  | Prot. | Prot.   | ssearch36<br>ggearch36 | blastp     | optimal local sim.<br>global:global sim. |
| DNADNAalignmentsDNAProt.fastx36<br>fasty36blastxtrans. DNA:protein sim.Prot.DNAtfastx36<br>tfasty36blastnprotein:trans. DNAProt.Prot.fasts36<br>tfasty36blastnprotein:trans. DNAProt.Prot.fasts36<br>tfasts36<br>DNAUnordered peptides<br>Unordered peptides<br>  | DNA   | DNA     | 00                     | blastn     | g.e.e.ae.a.e                             |
| Prot.DNAtfastx36<br>tfasty36blastnprotein:trans. DNAProt.Prot.fasts36Unordered peptidesProt.DNAtfasts36Unordered peptidesDNADNAfasts36Unordered peptidesProt.Prot.fasts36Ordered peptidesDNADNAfasts36Unordered peptidesProt.Prot.fasts36Ordered peptides   |       |         | lalign36               |            |  |
| tfasty36Prot.Prot.fasts36Unordered peptidesProt.DNAtfasts36Unordered peptidesDNADNAfasts36Unordered oligonucleotidesProt.Prot.fastm36Ordered peptides   | DNA   | Prot.   | 2 40 0110 0            | blastx     | trans. DNA:protein sim.                  |
| Prot.DNAtfasts36Unordered peptidesDNADNAfasts36Unordered oligonucleotidesProt.Prot.fastm36Ordered peptides  | Prot. | DNA     |                        | blastn     | protein:trans. DNA                       |
| DNADNAfasts36Unordered oligonucleotidesProt.Prot.fastm36Ordered peptides  | Prot. | Prot.   | fasts36                |            | Unordered peptides                       |
| Prot. Prot. fastm36 Ordered peptides  | Prot. | DNA     | tfasts36               |            | Unordered peptides                       |
|   | DNA   | DNA     | fasts36                |            | Unordered oligonucleotides               |
| DNA DNA fastm36 Ordered oligos  | Prot. | Prot.   | fastm36                |            | Ordered peptides                         |
|   | DNA   | DNA     | fastm36                |            | Ordered oligos                           |

#### Which program should I use? – See Table I, also:

\*ssearch36 can also be used for DNA:DNA, but is much slower and no more sensitive.

### How do I make FASTA act/look like BLAST? -

fasta36 -s BP62 -m BB query.file library.file

-s BP62 sets the same scoring matrix (BLOSUM62) and gap-penalties (-11/-1) as BLAST (FASTA uses BLOSUM50 by default). -m BB produces very BLAST-like output.

When I search Genbank - the program reports: 0 residues in 0 sequences? This typically happens because the program does not know that you are searching a Genbank flatfile database and is looking for a FASTA format database. Be certain to specify the library type ("1" for Genbank flatfile) with the database name.

The search seemed to work, but I do not see any results. – In command line mode (the default), all the FASTA programs limit the number of high scoring sequences shown using an expectation value cutoff (E() < 10 for proteins; E() < 2 for DNA). Sometimes, a search will complete successfully (you see the message XXXX residues in YYY sequences) but the message: !! No sequences with E() < 10 instead of The best scores are:. Typically, this happens because of a problem with the statistical estimation process; in particular, if the library contains only related sequences and -z 11 was not used, none of the hits may be "significant". To trouble shoot this problem, you can search with -z -1, which turns off all the statistical estimation procedures, and will show the 20 highest scoring sequences (-b # sets the default number of sequences shown).

What is the difference between fastx3 and fasty3? (or tfastx3 and tfasty3)? - [t]fastx3 uses a simpler codon based model for alignments that does not allow frameshifts in some codon positions (see ref. [24]). fastx3 is about 30% faster, but fasty3 can produce higher quality alignments in some cases.

**What is ktup**? – All of the programs with fast in their name use a computer science method called a lookup table to speed the search. For proteins with ktup=2, this means that the program does not look at any sequence alignment that does not involve matching two identical residues in both sequences. Likewise with DNA and ktup = 6, the initial alignment of the sequences looks for 6 identical adjacent nucleotides in both sequences. Because it is less likely that two identical amino-acids will line up by chance in two unrelated proteins, this speeds up the comparison. But very distantly related sequences may never have two identical residues in a row but will have single aligned identities. In this case, ktup = 1 may find alignments that ktup=2 misses.

**How do I turn off statistics**? – The FASTA programs are designed to identify homologs based on statistically significant similarity; to infer homology you need accurate statistical estimates. Sometimes, however, you know the sequences are related, and searching against libraries of related sequences can confuse FASTA if you do not use -z 11. If all you want are scores and alignments, use -z -1 to turn off statistical estimates.

**Where are** prss **and** prfx? – Earlier FASTA3 releases included prss3 and prfx3. With FASTA version 35 and 36, these programs have been incorporated into ssearch36 and fastx36. FASTA version 35 and 36 programs now automatically estimate statistical parameters by shuffling - the function of prss and prfx, when searching for libraries with fewer than 500 members.

**Where is** tfasta? – Although it is possible to make tfasta36, it is not compiled by default. tfastx36 and tfasty36 allow frame-shifts to be joined into a single alignment; tfasta did not. tfastx36 produces better alignments with better statistics.

**Can I run the FASTA programs on a cluster**? – With version 36.3.4, almost all of the FASTA programs can be run on clusters of computers using MPI (Message Passaging Interface). The programs can be compiled using make <code>-f .../make/Makefile.mpi\_sse2</code> from the <code>fasta36/src</code> directory. Except for <code>lalign36</code>, all the programs in Table I are available as <code>fasta36\_mpi</code>, <code>ssearch36\_mpi</code>, etc.

Unfortunately, the current MPI implementation involves substantially more communications overhead than the threaded versions. The FASTA programs are very efficient on threaded machines; if the preload option is used (edit make/Makefile36m.common to use comp\_lib8.c), the FASTA programs can obtain more than 40-fold speedup on a 48-core machine (the largest I have tested).

Sometimes, in the list of best scores, the same sequence is shown twice with exactly the same score. Sometimes, the sequence is there twice, but the scores are slightly different? – When any of the FASTA programs searches a long sequence, it breaks the sequence up into *overlapping* pieces. If the highest scoring alignment is at the end of one piece, it will be scored again at the beginning of the next piece. If the alignment is not be completely included in the overlap region, one of the pieces will give a higher score than the other. These duplications can be detected by looking at the coordinates of the alignment. If either the beginning or end coordinate is identical in two alignments, the alignments are at least partially duplicates.

As always, please inform me of bugs as soon as possible.

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# Appendix

## A FASTA Makefile compile time options

The fasta-36/make directory includes Makefiles appropriate for a broad range of environments, including Linux/Unix, BSD, MacOSX, and Windows. Makefiles are regularly tested against MacOSX, Linux, and Windows. Table 2 summarizes the Makefile options that can be modified.

As distributed, the Makefiles in fasta36/make, build a version of the FASTA programs that is optimized for single searches against arbitrary sized databases, using bit scores, efficient sampled statistics, and gap-open/extend penalties. The default compilation configuration can be changed either by changing the compile time defines (Table 2) in the main Makefile, e.g. make/Makefile.linux64\_sse2, or by editing make/Makefile36m.common.

High-performance searches with many queries – By default, the comp\_lib5e.c program specified in Makefile36m.common builds FASTA programs that re-read the library sequence database for every query sequence. This has the advantage that sequence comparison begins almost

| #define          | Status* | Target file(s)   | Function   |
|------------------|---------|--|--|
| ALLOCNO          | obs     | dropnfa.c,<br>dropfx.c,<br>dropfz2.c                     | allows FASTA algorithm to use memory $\sim$ query length (n0), not query + library (n0+n1).                  |
| DNALIB_LC        | undef   | initfa.c   | enable lower case masking for DNA libraries  |
| HTML_HEAD        | undef   | comp_lib5e.c,<br>comp_lib8.c                             | <pre>wrap -m 6 HTML output with <html> <body> </body> </html></pre>  |
| M10_CONS         | def     | c_dispn.c  | show consensus line (:. ) with $-m$ 10 output.   |
| OLD_FASTA_GAP    | undef   | drop*.c  | use first-residue/additional residue penalties, not open/extend.   |
| PGM_DOC          | def     | comp_lib5e.c,<br>comp_lib8.c                             | provide #pgm_name -opt1 -opt2 query<br>file copy of command line   |
| PROGRESS         | def     | comp_lib5e.c,<br>comp_lib8.c                             | provide progress symbols in interactive mode   |
| SAMP_STATS       | def     | comp_lib5e.c,<br>comp_lib8.c                             | scores are sampled for statistical estimates   |
| SAMP_STATS_LESS  | def     | compacc.c  | a slower sampling strategy is used   |
| SHOW_ALIGN_SCORE | undef   | wm_align.c   | print score, cummulative score, during align-<br>ment (for teaching)   |
| SHOW_HELP        | def     | <pre>comp_lib5e.c, comp_lib8.c, initfa.c, doinit.c</pre> | print out help information with '-help', or no arguments given. Undef SHOW_HELP reverts to pre-fasta-35.4.4. |
| SHOW_HIST        | undef   | doinit.c   | inverts current meaning of -H (shows by de-<br>fault for non-PCOMPLIB (MPI) programs).                       |
| SHOWSIM          | def     | mshowbest.c<br>mshowalign2.c                             | display percent similarity   |
| USE_LNSTATS      | obs     | scaleswn.c   | use $ln()$ -scaling for scores, removed in fasta2.0.   |

#### Table 2: FASTA Makefile compile time #defines

\*Status: def: #defined in standard Makefiles; undef: undefined; obs: obsolete, provided backwards compatibility with FASTA2.0 or earlier.

immediately, but if thousands of searches are being performed, the database is re-read thousands of times. Makefile36m.common can be edited to use comp\_lib8.c in place of comp\_lib5e.c and the database is read only once, then held in memory for additional searches. Of course, if comp\_lib8.c is used, the computer must have enough memory to store the complete database. Keeping the database in memory allows the FASTA programs to very efficiently used large, multicore computers.

Parallel searches with MPI – By default under Unix/Linux/MacOSX, the FASTA programs are threaded; they will spawn as many threads as CPU cores are available (this can be limited with the -t n-threads option). Using comp\_lib8.c, we see almost 48-fold speedup on a 48-core machine. The FASTA programs can also be run in parallel in the MPI environment on clusters of computers. To build the MPI versions of the programs, use make .../make/Makefile.mpi\_sse2 ssearch36\_mpi, fastx36\_mpi, etc. The MPI programs currently substantially more communications overhead than the threaded versions, so they may not scale as well to large clusters.

# **B FASTA version history**

|           | FASTA v33, Oct, 1999 – Dec, 2000   |
|-----------|--|
| Oct 1999  | Add support for NCBI Blast2.0 formatted libraries, and memory mapped databases. FASTA now reads both BLAST1.4 and BLAST2.0 formatted databases. (version 3.2t08)   |
|           | Include Maximum Likelihood Estimates for Lambda and K (-z 2)<br>Include a new strategy for searching with low complexity regions. The pseg<br>program can produce libraries with low complexity regions as lower case char-<br>acters, which can be ignored during the initial FASTA/SSEARCH scan, but are<br>considered when producing the final alignments. (3.3t01) |
|           | Change output to report bit scores, which are also used by BLAST.  |
| Mar 2000  | Another new statistics option, -z 6, uses Mott's approach [11] for calculating a composition dependent Lambda for each sequence. (3.3t05)  |
| Dec 2000  | Automatically change the gap penalties when alternate (known) scoring matrices are used using Reese and Pearson gap penalties [18]. First implementation to read from MySQL databases.   |
| May 2001  | change all FASTA gap penalties from first-residue, additional residue to the gap-<br>open, gap-extend values used by BLAST.  |
|           | FASTA v34, Jan, 2001 – Jan, 2007   |
| Jun 2002  | Modify statistical estimation strategy to sample all the sequences in the database, not just the first 60,000. (3.4t11)  |
| Jan 2003  | Implementation of vector-accelerated (Altivec) code for Smith-Waterman (SSEARCH) and banded Smith-Waterman (FASTA) using the Rognes and Seebug [19] algorithm. This code was removed in Sept, 2003, because of possible conflict with a patent application, but was restored using a different algorithm in Nov. 2004.   |
| Jun 2003  | Provide PSI-SEARCH — an implementation of SSEARCH that can search with PSI-BLAST PSSM profile files. PSI-SEARCH estimates statistical significance from the distribution of actual alignment scores; thus the estimates are much more reliable than PSI-BLAST estimates. Also, change the similarity display to work with profiles. (3.4t22)                           |
| July 2003 | Provide ASN.1 definition line parsing for BLAST formatdb v.4 libraries. Restruc-<br>ture the programs to use a table-driven approach to parameter setting. Two<br>tables now define the algorithm, query sequence type, library type, scoring ma-<br>trix, and gap penalties for all programs.   |
| Sept 2003 | A new option -V for annotating alignments provided. Designed for highlight-<br>ing post-translational modifications with fasts, it can also be used to highlight   |
| Dec 2003  | active sites and other conserved residues. (3.4t23)<br>Addition of -U option for RNA sequence comparison. G:A matches score like<br>G:G matches to account for G:U basepairs. Change default <i>ktup</i> for short query<br>sequences. Increase band-width for DNA banded final alignments.  |

### FASTA version history (cont.)

- July 2004Allow searching of Postgres, as well as MySQL database queries.Nov 2004(fa34t24) Incorporation of Erik Lindahl "anti-diagonal" Altivec implementation<br/>of [23] for Smith-Waterman only. Altivec ssearch34 is now faster than fasta34<br/>for query sequences < 250 amino acids.</th>Jan 2005Change FASTS to accommodate very large numbers of peptides (>100) for full
- Jun. 2006 (fa34t26) Incorporation of Smith-Waterman algorithm for the SSE2 vector in-
- structions written by Michael Farrar [5]. The SSE code speeds up Smith-Waterman 8 – 16-fold.

FASTA v35, March, 2007 – March, 2010

Mar. 2007 fasta v35 – Accurate shuffle-based E()-values for all searches and alignments; statistics from searches against small libraries are supplemented with shuffled alignments.

More efficient threading strategies on multi-core computers, for 12X speedup on 16-core machines.

Inclusion of lalign (SIM) local domain alignments. lalign alignments now have accurate shuffle-based E()-values.

- Apr. 2007 Introduction of ggsearch, for global alignment searches, and glsearch, for searches with scores that are global in the query and local in the library. ggsearch and glsearch calculate E()-values using the normal distribution. Both programs can search with PSI-BLAST PSSMs.
- Dec. 2007 Efficient strategy for searching subsets of databases (lists of GI or accession numbers)
- Feb. 2008 Annotations in either query or library sequences can be highlighted in the alignment, and the state of annotated residues is compactly summarized with -m 9c.
- Oct. 2008 Modification lsim4.c (lalign35) provided by Xiaoqui Huang to ensure that selfalignments do not cross the identity diagonal.

FASTA v36, March, 2010 -

Mar. 2010 FASTA v36 displays all significant alignments between query and library sequence. BLAST has always displayed multiple high-scoring alignments (HSPs) between the query and library sequence; previous versions of the FASTA programs displayed only the best alignment, even when other high-scoring alignments were present.

New statistical options, -z 21, 22, 26, provide a second E2()-value estimate based on shuffles of the highest scoring sequences.

#### FASTA version history (cont.)

Improved performance using statistics-based thresholds for gap-joining and band-optimization in the heuristic FASTA local alignment programs, increasing speed 2 - 3X.

Greater flexibility in specifying combinations of library files and subsets of libraries. FASTA v36 programs can include indirect files of library names inside of indirect files of library names.

FASTA 36 programs are fully threaded, both for searches, and for alignments. The programs routinely run 12 - 15X faster on 8-core machines with "hyper-threading" (effectively 16 cores).

-z 21 .. 26 E2() statistical estimates from shuffled best scores.

- Sep. 2010 -m 8, -m 8C BLAST tabular output.
- Nov, 2010 Variable scoring matrices (-m ?BP62).
- Dec, 2010 (fasta-36.3.1) SSE2 vectorized ggsearch36, glsearch36 (Michael Farrar).
- Jan, 2011 (fasta-36.3.2) MPI versions implemented and tested.
- Feb, 2011 Introduce -m B, -m BB BLAST-like output.
- Mar, 2011 (fasta-36.3.4) Program is no longer interactive by default. fasta36 -h and fasta36 -help provide common/complete options, with many defaults. doc/fasta\_guide.pdf available.
- May, 2011 (fasta-36.3.5) Introduce (1) -e expand.sh scripts to extend the effective size of the database searched, based on significant hits; (2) -m "F# output.file" to send different output formats to different files; and (3) -X expanded options, -o replaces the old -X and -Xo replaces -o.
- Jan, 2012 Include .fastq files as library type 7
- May, 2012 allow reverse-complement alignments with ggsearch and glsearch
- Jun, 2012 Introduce -V !script.pl driven alignments, and variant scoring.
- Aug, 2012 Introduce -V !ann\_feats.pl sub-alignment (region-based) scoring.
- Apr, 2013 Extend ENV options to introduce a domain-plotting option for FASTA web sites.