HMMER User's Guide

Biological sequence analysis using profile hidden Markov models

http://hmmer.wustl.edu/ Version 2.2; August 2001

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4	4.14.24.3	Expert Options	53 55 56 59 60 61 61 62 63 63 63 64 65
4	4.14.24.3	Expert Options	53 55 56 59 60 61 61 62 63 63 63 63 64 65 65
4	4.14.24.3	Expert Options	 53 55 56 59 60 61 61 62 63 63 63 64 65 66
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Chapter 1

Tutorial

I hate reading documentation. I just want examples of how stuff works, just enough to get me started and doing something productive. So, here's a tutorial walk-through of some small projects with HMMER. If you want the introduction, that's the second chapter. The tutorial should be sufficient to get you started on work of your own. You can read the other chapters later if you want.

1.1 The programs in HMMER

There are currently nine programs supported in the HMMER 2 package:

hmmalign Align sequences to an existing model.

- hmmbuild Build a model from a multiple sequence alignment.
- **hmmcalibrate** Takes an HMM and empirically determines parameters that are used to make searches more sensitive, by calculating more accurate expectation value scores (E-values).
 - hmmconvert Convert a model file into different formats, including a compact HMMER 2 binary format, and "best effort" emulation of GCG profiles.
 - **hmmemit** Emit sequences probabilistically from a profile HMM.
 - hmmfetch Get a single model from an HMM database.
 - hmmindex Index an HMM database.
 - hmmpfam Search an HMM database for matches to a query sequence.
 - hmmsearch Search a sequence database for matches to an HMM.

HMMER also provides a number of utility programs which are not HMM programs, but may be useful. These programs are from the SQUID sequence utility library that HMMER uses:

afetch Retrieve an alignment from an alignment database

alistat Show some simple statistics about a sequence alignment file.

seqstat Show some simple statistics about a sequence file.

sfetch Retrieve a (sub-)sequence from a sequence file.

shuffle Randomize sequences in a sequence file.

sreformat Reformat a sequence file into a different format.

1.2 Files used in the tutorial

The subdirectory /tutorial in the HMMER distribution contains the files used in the tutorial, as well as a number of examples of various file formats that HMMER reads. The important files for the tutorial are:

globins50.msf An MSF format alignment file of 50 aligned globin sequences.

globins630.fa A FASTA format file of 630 unaligned globin sequences.

fn3.slx A SELEX format alignment file of fibronectin type III domains.

rrm.slx A SELEX format alignment file of RNA recognition motif domains.

rrm.hmm An example HMM, built from rrm.slx.

- pkinase.slx A SELEX format alignment file of protein kinase catalytic domains.
- Artemia.fa A FASTA file of brine shrimp globin, which contains nine tandemly repeated globin domains.
- **7LES_DROME** A SWISSPROT file of the *Drosophila* Sevenless sequence, a receptor tyrosine kinase with multiple domains.

Create a new directory that you can work in, and copy all the files in tutorial there. I'll assume for the following examples that you've installed the HMMER programs in your path; if not, you'll need to give a complete path name to the HMMER programs (e.g. something like /usr/people/eddy/hmmer-2.2/binaries/hmmbuild instead of just hmmbuild).

1.3 Searching a sequence database with a single profile HMM

One common use of HMMER is to search a sequence database for homologues of a protein family of interest. You need a multiple sequence alignment of the sequence family you're interested in. (Profile HMMs can be trained from unaligned sequences; however, this functionality is temporarily withdrawn from HMMER. I recommend CLUSTALW as an excellent, freely available multiple sequence alignment program.)

HMM construction with hmmbuild

Let's assume you have a multiple sequence alignment of a protein domain or protein sequence family. To use HMMER to search for additional remote homologues of the family, you want to first build a profile HMM from the alignment. The following command builds a profile HMM from the alignment of 50 globin sequences in globins50.msf:

```
> hmmbuild globin.hmm globins50.msf
```

```
hmmbuild - build a hidden Markov model from an alignment
HMMER 2.2 (August 2001)
Copyright (C) 1992-2001 HHMI/Washington University School of Medicine
Freely distributed under the GNU General Public License (GPL)
globins50.msf
MSF
Alignment file:
File format:
Search algorithm configuration:MSFModel construction strategy:Multiple domain (hmmls)Null model used:(default)
Prior used:
                                (default)
Sequence weighting method: G/S/C tree weights
New HMM file: globin.hmm
Alignment:
                  #1
Number of sequences: 50
Number of columns: 308
Determining effective sequence number ... done. [2]
Weighting sequences heuristically
                                     ... done.
                                      ... done.
Constructing model architecture
                                     ... done.
Converting counts to probabilities
                                      ... done. [globins50]
Setting model name, etc.
Constructed a profile HMM (length 148)
Average score: 194.97 bits
Minimum score: -17.88 bits
Maximum score: 242.22 bits
Std. deviation: 55.12 bits
Finalizing model configuration ... done.
Saving model to file
                                      ... done.
11
```

The process takes a second or two. hmmbuild create a new HMM file called globin.hmm. This is a human and computer readable ASCII text file, but for now you don't care. You also don't care for now what all the stuff in the output means; I'll describe it in detail later. The profile HMM can be treated as a compiled model of your alignment.

HMM calibration with hmmcalibrate

This step is optional, but doing it will increase the sensitivity of your database search.

When you search a sequence database, it is useful to get "E-values" (expectation values) in addition to raw scores. When you see a database hit that scores x, an E-value tells you the number of hits you would've expected to score x or more just by chance in a sequence database of this size.

HMMER will always estimate an E-value for your hits. However, unless you "calibrate" your model before a database search, HMMER uses an analytic upper bound calculation that is extremely conservative. An empirical HMM calibration costs time (about 10% the time of a SWISSPROT search) but it only has to be done once per model, and can greatly increase the sensitivity of a database search. To empirically calibrate the E-value calculations for the globin model, type:

> hmmcalibrate globin.hmm

hmmcalibrate -- calibrate HMM search statistics HMMER 2.2 (August 2001) Copyright (C) 1992-2001 HHMI/Washington University School of Medicine Freely distributed under the GNU General Public License (GPL) HMM file: qlobin.hmm Length distribution mean: 325 Length distribution s.d.: 200 Number of samples: 5000 random seed: 997047045 histogram(s) saved to: [not saved] POSIX threads: 1 HMM : globins50 mu : -38.963402 lambda : 0.256441 max : -11.779000 11

This takes several minutes. Go have a cup of coffee. When it is complete, the relevant parameters are added to the HMM file.

Calibrated HMMER E-values tend to be relatively accurate. E-values of 0.1 or less are, in general, very significant hits. Uncalibrated HMMER E-values are also reliable, erring on the cautious side; uncalibrated models may miss remote homologues.

Sequence database search with hmmsearch

As an example of searching for new homologues using a profile HMM, we'll use the globin model to search for globin domains in the example *Artemia* globin sequence in Artemia.fa:

> hmmsearch globin.hmm Artemia.fa

The output comes in several sections, and unlike building and calibrating the HMM (where we treated the HMM as a black box), now you *do* care about what it's saying.

The first section is the *header* that tells you waht program you ran, on what, and with what options:

```
Query HMM: globins50
Accession: [none]
Description: [none]
[HMM has been calibrated; E-values are empirical estimates]
```

The second section is the *sequence top hits* list. It is a list of ranked top hits (sorted by E-value, most significant hit first), formatted in a BLAST-like style:

Scores for complete sequences (score includes all	domains):		
Sequence Description Score E-value N			
S13421 S13421 GLOBIN - BRINE SHRIMP	496.2	4.3e-150	9

The first field is the name of the target sequence, then followed by the description line for the sequence. The last three fields are the raw score (in units of "bits"), the estimated E-value, and the total number of domains detected in the sequence. By default, every sequence with an E-value less than 10.0 is listed in this output.

The second section is the *domain top hits* list. By default, for every sequence with an E-value less than 10, every domain with a raw score greater than 0 is listed. (Read that carefully. In a later chapter we'll discuss some caveats about how hmmsearch identifies domains, and how to control its output in different ways.) Each domain detected in the search is output in a list ranked by E-value:

Parsed for domains:

Sequence	Domain	seq-f	seq-t		hmm-f	hmm-t		score	E-value
S13421	7/9	927	1075		1	148	[]	82.2	1.8e-25
S13421	2/9	148	293	•••	1	148	[]	66.2	1.2e-20
S13421	3/9	302	450	•••	1	148	[]	63.7	6.8e-20
S13421	8/9	1084	1234	•••	1	148	[]	60.7	5.2e-19
S13421	9/9	1243	1390	••	1	148	[]	55.9	1.5e-17
S13421	4/9	459	607	•••	1	148	[]	52.6	1.5e-16
S13421	6/9	770	918	•••	1	148	[]	46.6	9.6e-15
S13421	1/9	1	143	[.	1	148	[]	44.7	3.5e-14
S13421	5/9	618	762		1	148	[]	23.6	7.8e-08

The first field is the name of the target sequence. The second field is the number of this domain: e.g. "6/9" means the sixth domain of nine total domains detected.

The fields marked "seq-f" and "seq-t" mean "sequence from" and "sequence to": the start and end points of the alignment on the target sequence. After these two fields is a shorthand annotation for whether the alignment is "global" with respect to the sequence or not. A dot (.) means the alignment does not go all the way to the end; a bracket ([or]) means it does. Thus, .. means that the alignment is local within the sequence; [. means that the alignment starts at the beginning of the sequence, but doesn't go all the way to its end; .] means the alignment starts somewhere internally and goes all the way to the end; and [] means the alignment includes the entire sequence.

Analogously, the fields marked "hmm-f" and "hmm-t" indicate the start and end points with respect to the consensus coordinates of the model, and the following field is a shorthand for whether the alignment is global with respect to the *model*. Here, for instance, all the globin domains in the *Artemia* sequence are complete matches to the entire globin model – *because, by default, hmmbuild built the HMM to only look for those kinds of alignments.* We'll discuss later how to modify the profile HMM for other search styles.

The final two fields are the raw score in bits and the estimated E-value, *for the isolated domain*. The scores for the domains sum up to the raw score of the complete sequence.

The next section is the *alignment output*. By default, every domain that appeared in the domain top hits list now appears as a BLAST-like alignment. For example:

```
Alignments of top-scoring domains:
S13421: domain 7 of 9, from 927 to 1075: score 82.2, E = 1.8e-25
                   *->vhlsaeekalvksvwgkveknveevGaeaLerllvvyPetkryFpkF
                       +lsa+e a vk+ w+ v+ ++ vG +++ l++ +P+ +++FpkF
      S13421
              927
                      TGLSAREVAVVKQTWNLVKPDLMGVGMRIFKSLFEAFPAYQAVFPKF 973
                  kdLssadavkgsakvkahgkkVltalgdavkkldd...lkgalakLselH
                                       V t+l++ ++ ld++ +l+
                   d+
                        d+++++ v +h
                                                              ++L+e H
      S13421
               974 SDVPL-DKLEDTPAVGKHSISVTTKLDELIQTLDEpanLALLARQLGEDH 1022
                   aqklrvdpenfkllsevllvvlaeklgkeftpevqaalekllaavataLa
                             fk +++vl+ l++ lq+ f+ ++ +++k++++++ ++
                   + lrv+
      S13421 1023 IV-LRVNKPMFKSFGKVLVRLLENDLGQRFSSFASRSWHKAYDVIVEYIE 1071
                   akYk<-*
                   +
                     +
                           1075
      S13421 1072 EGLQ
```

The top line is the HMM consensus. The amino acid shown for the consensus is the highest probability amino acid at that position according to the HMM (not necessarily the highest *scoring* amino acid, though). Capital letters mean "highly conserved" residues: those with a probability of > 0.5 for protein models, or > 0.9 for DNA models.

The center line shows letters for "exact" matches to the highest probability residue in the HMM, or a "+" when the match has a positive score and is therefore considered to be "conservative" according to the HMM's view of *this particular position in the model* – not the usual definition of conservative changes in general.

The third line shows the sequence itself, of course.

The next section of the output is the *score histogram*. It shows a histogram with raw score increasing along the Y axis, and the number of sequence hits represented as a bar along the X axis. In our example here, since there's only a single sequence, the histogram is very boring:

Notice though that it's a histogram of the whole sequence hits, not the domain hits. You can ignore the rest of the hmmsearch output:

```
Total hits: 1
Satisfying E cutoff: 1
Total memory: 16K
Domain top hits:
tophits_s report:
Total hits: 9
Satisfying E cutoff: 9
Total memory: 21K
```

This is just some trailing internal info about the search that's useful to me sometimes, but probably not to you.

Searching major databases like NR or SWISSPROT

HMMER reads all major database formats and does not need any special database indexing. You can search any large sequence database you have installed locally just by giving the full path to the database file, e.g. something like:

> hmmsearch globin.hmm /nfs/databases/swiss35/sprot35.dat

If you have BLAST installed locally, it's likely that you have a directory (or directories) in which the BLAST databases are kept. These directories are specified in an environment variable called BLASTDB. HMMER will read the same environment variable. For example, if you have BLAST databases in directories called /nfs/databases/blast-db/ and /nfs/databases/golden-path/blast/, and you want to search /nfs/databases/blast-db/swissprot, the following commands will work:

```
> setenv BLASTDB /nfs/databases/blast-db/:/nfs/databases/golden-path/blast/
```

> hmmsearch globin.hmm swissprot

Obviously, you'd tend to have the setenv command as part of the local configuration of your machine, rather than typing it at the command line.

Local alignment searches with hmmsearch

This is extremely important. HMMER does not do local (Smith/Waterman) and global (Needleman/Wunsch) style alignments in the same way that most computational biology analysis programs do it. To HMMER, whether local or global alignments are allowed is part of the *model*, rather than being accomplished by running a different *algorithm.* (This will be discussed in greater detail later; it is part of the "Plan7" architecture of the new HMMER2 models.)

Therefore, you need to choose what kind of alignments you want to allow *when you build the model* with hmmbuild. By default, hmmbuild builds models which allow alignments that are global with respect to the HMM, local with respect to the sequence, and allows multiple domains to hit per sequence. Such models will only find complete domains.

hmmbuild provides some standard options for common alignment styles. The following table shows the four alignment styles supported by hmmbuild, and also shows the equivalent old HMMER 1.x search program style (to orient experienced HMMER users).

Command	w.r.t. sequence	w.r.t. HMM	multidomain	HMMER 1 equivalent
hmmbuild	local	global	yes	hmmls
hmmbuild -f	local	local	yes	hmmfs
hmmbuild -g	local	global	no	hmms
hmmbuild -s	local	local	no	hmmsw

In brief, if you want maximum sensitivity at the expense of only finding complete domains, use the hmmbuild default. If you need to find fragments (local alignments) too, and are willing to give up some sensitivity to see them, use hmmbuild -f. If you want the best of both worlds, build two models and search with both of them.

1.4 Searching a query sequence against a profile HMM database

A second use of HMMER is to look for known domains in a query sequence, by searching a single sequence against a library of HMMs. (Contrast the previous section, in which we searched a single HMM against a sequence database.) To do this, you need a library of profile HMMs. One such library is our PFAM database (Sonnhammer et al., 1997; Sonnhammer et al., 1998), and you can also create your own.

Creating your own profile HMM database

HMM databases are simply concatenated single HMM files. You can build them either by invoking the -A "append" option of hmmbuild, or by concatenating HMM files you've already built. For example, here's two ways to build an HMM database called myhmms that contains models of the rrm RNA recognition motif domain, the fn3 fibronectin type III domain, and the pkinase protein kinase catalytic domain:

```
> hmmbuild rrm.hmm rrm.slx
> hmmbuild fn3.hmm fn3.slx
> hmmbuild pkinase.hmm pkinase.slx
> cat rrm.hmm fn3.hmm pkinase.hmm > myhmms
> hmmcalibrate myhmms
or:
> hmmbuild -A myhmms rrm.slx
> hmmbuild -A myhmms fn3.slx
> hmmbuild -A myhmms pkinase.slx
```

```
> hmmcalibrate myhmms
```

Notice that hmmcalibrate can be run on HMM databases as well as single HMMs.

Parsing the domain structure of a sequence with hmmpfam

Now that you have a small HMM database called myhmms, let's use it to analyze the *Drosophila* Sevenless sequence, 7LES_DROME:

```
> hmmpfam myhmms 7LES_DROME
```

Like hmmsearch, the hmmpfam output comes in several sections. The first section is the header:

The next section is the *sequence family classification* top hits list, ranked by E-value. The scores and E-values here reflect the confidence that this query sequence contains one *or more* domains belonging to a domain family. The fields have the same meaning as in hmmsearch output, except that the name and description are for the HMM that's been hit.

Model	Description	Score	E-value N	1
pkinase		304.1	1.1e-91	1
fn3		176.3	3.5e-53	б
rrm	RNA recognition motif. (aka RRM, RBD, or RNP	-44.5	0.72	1

The next section is the *domain parse* list, ordered by position on the sequence (not by score). Again the fields have the same meaning as in hmmsearch output:

Parsed for domains:									
Model	Domain	seq-f	seq-t		hmm-f	hmm-t		score	E-value
fn3	1/6	437	522		1	84	[]	49.0	7.1e-15
fn3	2/6	825	914		1	84	[]	13.6	4.3e-06
fn3	3/6	1292	1389	••	1	84	[]	16.2	2.4e-06
rrm	1/1	1300	1364	••	1	72	[]	-44.5	0.72
fn3	4/6	1799	1891	••	1	84	[]	63.5	3.2e-19
fn3	5/6	1899	1978	••	1	84	[]	14.6	3.4e-06
fn3	6/6	1993	2107	••	1	84	[]	19.4	1.2e-06
pkinase	1/1	2209	2483	••	1	278	[]	304.1	1.1e-91

Note how it's showing us an "rrm" hit - $7LES_DROME$ doesn't have any RRM domains. You have to notice for yourself that the hit is insignificant (with a negative score, and an E-value of nearly 1). By default, like BLAST, the search programs report well down into the noise. If you want the output to be cleaner, set an E-value threshold; for example hmmpfam $-E \ 0.1$.

The final output section is the *alignment output*, just like hmmsearch:

```
Alignments of top-scoring domains:
fn3: domain 1 of 6, from 437 to 522: score 49.0, E = 7.1e-15
*->P.saPtnltvtdvtstsltlsWsppt.gngpitgYevtyRqpkngge
P saP + +++ ++ 1 ++W p + ngpi+gY+++ ++++g+
7LES_DROME 437 PiSAPVIEHLMGLDDSHLAVHWHPGRfTNGPIEGYRLRL-SSSEGNA 482
```

Downloading the PFAM database

The PFAM database is available from either http://pfam.wustl.edu/ or http://www.sanger.ac.uk/Pfam/. Download instructions are on the Web page. The PFAM HMM library is a single large file, containing several hundred models of known protein domains. Install it in a convenient directory and name it something simple like pfam.

HMMER will look for PFAM and other files in a directory (or directories) specified by the HMMERDB environment variable. For instance, if you install the PFAM HMM library as /nfs/databases/hmmer/pfam, the following commands will search for domains in 7LES_DROME:

```
> setenv HMMERDB /nfs/databases/hmmer/
```

```
> hmmpfam pfam 7LES_DROME
```

1.5 Maintaining multiple alignments with hmmalign

Another use of profile HMMs is to create multiple sequence alignments of large numbers of sequences. A profile HMM can be build of a "seed" alignment of a small number of representative sequences, and this profile HMM can be used to efficiently align any number of additional sequences.

This is in fact how the PFAM database is updated as the main SPTREMBL database increases in size. The PFAM seed alignments are (relatively) stable from release to release; PFAM full alignments are created automatically by searching SPTREMBL with the seed model and aligning all the significant hits into a multiple alignment using hmmalign.

For example, to align the 630 globin sequences in globins630.fa to our globin model globin.hmm, and create a new alignment file called globins630.ali, we'd do:

```
> hmmalign -o globins630.ali globin.hmm globins630.fa
hmmalign - align sequences to an HMM profile
HMMER 2.2 (August 2001)
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HMM file: globin.hmm
Sequence file: globins630.fa
```

Alignment saved in file globins630.ali

Using the -0 option to specify a save file for the final alignment is a good idea; else, the alignment will be displayed on the screen as output (and an alignment of several hundred sequences will give a fairly voluminous output).

Chapter 2

Introduction

You discover a new protein sequence family, and carefully construct a multiple sequence alignment. Your family, like most protein families, has a number of strongly (but not absolutely) conserved key residues, separated by characteristic spacing. You wonder if there are more members of your family in the sequence databases, but the family is so evolutionarily diverse, a BLAST search with any individual sequence doesn't even find the rest of the sequences you already know about; you're sure there are some distantly related sequences in the "noise". You spend many pleasant evenings scanning weak BLAST alignments by eye to find ones with the right key residues are in the right places. You sure wish there was a computer program that did this better.

You get the sequence of another favorite protein and eagerly BLAST it against the NCBI server, and nothing significant shows up. It looks like you'll get no easy information from the sequence.

You BLAST another favorite sequence against the NCBI server; a thousand hits come back, and the top hundred are hypothetical sequences from genome projects. It looks like you'll have to wade through a lot of BLAST output to find the informative hits.

Another sequence comes back a slew of hits to receptor tyrosine kinases, but before you decide to call your sequence an RTK homologue, you recall that RTK's are, like many proteins, composed of multiple functional domains. Is your sequence really an RTK? Or is it a novel sequence that also happens to have a protein kinase catalytic domain, fibronectin type III domain, or immunoglobulin superfamily domain?

These (and other problems in sequence analysis) are the kinds of problems that HMMER tries to address.

2.1 Profile HMMs

Profile hidden Markov models (profile HMMs) are statistical models of the primary structure consensus of a sequence family. Anders Krogh, David Haussler, and co-workers at UC Santa Cruz introduced profile HMMs (Krogh et al., 1994), adopting HMM techniques which have been used for years in speech recognition. HMMs had been used in biology before the Krogh/Haussler work, but the Krogh paper had a particularly dramatic impact, because HMM technology was so well-suited to the popular "profile" methods for searching databases using multiple sequence alignments instead of single query sequences. Since then, several computational biology groups (including ours) have rapidly adopted HMMs as the underlying formalism for sequence profile analysis.

"Profiles" were introduced by Gribskov and colleagues (Gribskov et al., 1987; Gribskov et al., 1990) at about the same time that other groups introduced similar approaches, such as "flexible patterns" (Barton, 1990), and "templates" (Bashford et al., 1987; Taylor, 1986). The term "profile" has stuck.¹ All of these are

¹There has been some agitation to call all such models "position specific scoring matrices", PSSMs, but certain small nocturnal

more or less statistical descriptions of the consensus of a multiple sequence alignment. They use positionspecific scores for amino acids (or nucleotides) and position specific scores for opening and extending an insertion or deletion. Traditional pairwise alignment (for example, BLAST (Altschul et al., 1990), FASTA (Pearson and Lipman, 1988), or the Smith/Waterman algorithm (Smith and Waterman, 1981)) uses position*independent* scoring parameters. This property of profiles captures important information about the degree of conservation at various positions in the multiple alignment, and the varying degree to which gaps and insertions are permitted.

The advantage of using HMMs is that HMMs have a formal probabilistic basis. We can use Bayesian probability theory to guide how all the probability (scoring) parameters should be set. Though this might sound like a purely academic issue, this probabilistic basis lets us do things that the more heuristic methods cannot do easily. For example, an HMM can be trained from unaligned sequences, if a trusted alignment isn't yet known. Another consequence is that HMMs have a consistent theory behind gap and insertion scores.² In most details, HMMs are a slight improvement over a carefully constructed profile – but far less skill and manual intervention is necessary to train a good HMM and use it. This allows us to make libraries of hundreds of profile HMMs and apply them on a very large scale to whole-genome or EST sequence analysis. One such database of protein domain models is Pfam (Sonnhammer et al., 1997); the construction and use of Pfam is tightly tied to the HMMER software package.

HMMs do have important limitations. One is that HMMs do not capture any higher-order correlations. An HMM assumes that the identity of a particular position is independent of the identity of all other positions.³ HMMs make poor models of RNAs, for instance, because an HMM cannot describe base pairs. Also, compare protein "threading" methods, which include scoring terms for nearby amino acids in a three-dimensional protein structure.

A general definition of HMMs and an excellent tutorial introduction to their use has been written by Rabiner (Rabiner, 1989). Throughout, I will often use "HMM" to refer to the specific case of profile HMMs as described by Krogh et al. (Krogh et al., 1994). This shorthand usage is for convenience only. For a review of profile HMMs, see (Eddy, 1996), and for a complete book on the subject of probabilistic modeling in computational biology, see (Durbin et al., 1998).

2.2 Primary changes from HMMER 1.x

HMMER 2 is an almost complete rewrite of the original 1992-1996 HMMER code. A list of the major changes follows.

Plan7 The model architecture is changed. The "Plan 7" architecture⁴ allows detailed control over parameters of local, global, and multidomain alignments. In Plan 7, you choose the alignment style when you build the model with hmmbuild, not when you search. A single database search program hmmsearch replaces the four search programs in HMMER 1.x.

Pfam support There is much better support for the Pfam HMM database. HMMs have names; the HMM file format allows libraries of multiple HMMs per file; and there is a search

North American marsupials have a prior claim on the mnemonic.

²Surprisingly, this is still controversial. People have been saying "there is no theory for setting gap scores" for so long that many people believe it.

³This is not strictly true. There is a subtle difference between an HMM's state path (a first order Markov chain) and the sequence described by an HMM (generated from the state path by independent emissions of symbols at each state).

⁴The origin of the codename is obscure, involving both details of the state transition probability distributions in profile HMM architecture and an inordinate fondness for bad science fiction movies. Frighteningly, David Haussler understood the codename immediately.

program hmmpfam for searching a single query sequence against an HMM library. HMM databases like Pfam can be indexed for rapid access using hmmindex, and single HMMs can be retrieved from an HMM database using hmmfetch.

- E-values HMMER now reports both log-odds scores and a E-value. E-values can detect significant matches even when the log-odds score is negative. A new program hmmcalibrate empirically determines the necessary parameters for estimating E-values as best as possible.
 - **Output** Database searches now sort and postprocess their results for prettier and more useful output. Output styles are deliberately similar to BLAST, to try to simplify the design of postprocessors.
- **Save file format** HMMER save files are now in an ASCII format which is easy for humans to read, portable across all architectures without byteswapping issues, and compact.
- **Decent defaults** The "best" average behavior of HMMER is now its default behavior. For example, mixture Dirichlet priors are now the default for protein model building. In HMMER 1.x, you had to invoke a number of "experimental" options to get better performance this led to a bunch of really frustrating benchmarking in the literature, in which people compared fully optimized program X to the default behavior of HMMER 1, which was the 1994 basic Krogh/Haussler behavior rather than its actual power. (However, there's still a lot of bad benchmarking, because people don't understand how to benchmark profile software yet.)

2.3 Plan 7

The Plan 7 architecture

The Plan 7 architecture is substantially different from the original HMMER model architecture. The abbreviations for the states are as follows:

- \mathbf{M}_x Match state x. Has K emission probabilities.
- \mathbf{D}_x Delete state x. Non-emitter.
- I_x Insert state x. Has K emission probabilities.
- S Start state. Non-emitter.
- **N** N-terminal unaligned sequence state. Emits *on transition* with K emission probabilities.
- **B** Begin state (for entering main model). Non-emitter.
- E End state (for exiting main model). Non-emitter.
- **C** C-terminal unaligned sequence state. Emits *on transition* with *K* emission probabilities.
- **J** Joining segment unaligned sequence state. Emits *on transition* with K emission probabilities.

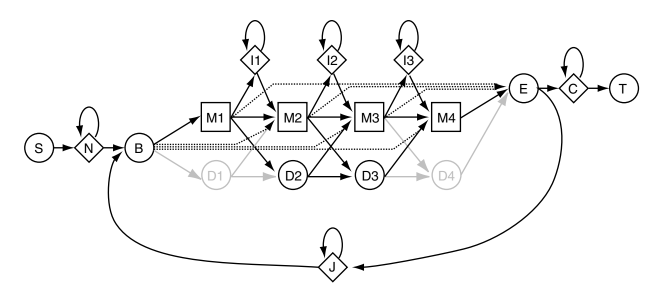


Figure 2.1: The Plan7 architecture. Squares indicate match states (modeling consensus positions in the alignment). Diamonds indicate insert states (modeling insertions relative to consensus) and special random sequence emitting states. Circles indicate delete states (modeling deletions relative to consensus) and special begin/end states. Arrows indicate state transitions. See text for more details.

The section of the model composed of M, D, and I states, and the B and E states, is essentially a Krogh/Haussler profile HMM. I refer to this as the "main model". A group of three states M/D/I at the same consensus position in the alignment is called a "node". The main model controls the *data dependent* features of the model. The probability parameters in the main model are generally learned from data in a multiple sequence/structure alignment.

Unlike the original Krogh/Haussler and HMMER model architecture, Plan 7 has no $D \rightarrow I$ or $I \rightarrow D$ transitions. This reduction from 9 to 7 transitions per node in the main model is the origin of the codename Plan 7. (The original HMMER architecture is called Plan 9 in parts of the code.)

The other states (S,N,C,T,J) are called "special states". They (combined with special entry probabilities from B and exit probabilities to E) control the *algorithm dependent* features of the model: how likely the model is to generate various sorts of local or multihit alignments. The algorithm dependent parameters are typically not learned from data, but rather set externally by choosing a desired alignment style.

Local alignments in Plan 7

The Plan 7 architecture models a complete sequence, regardless of how much of that sequence matches the main model. *All alignments to a Plan 7 model are "global" alignments*, but some of the sequence may be assigned to Plan 7 states (N,C,J) that generate "random" sequence that is not aligned to the main model. Thus, the algorithm dependent parts of the model control the *apparent* locality of the alignments.

Local alignments with respect to the sequence (i.e., allowing a match to the main model anywhere internal to a longer sequence) are controlled by the N and C states. If the $N \rightarrow N$ transition is set to 0, alignments are constrained to start in the main model at the very first residue. Similarly, if the $C \rightarrow C$ transition is set to 0, alignments are constrained to match the main model at the very last residue in the sequence.

Local alignments with respect to the model (i.e., allowing fragments of the model to match the sequence) are controlled by $B \rightarrow M$ "entry" transitions and $M \rightarrow E$ "exit" transitions, shown as dotted lines in the Plan

7 figure. Setting all entries but the $B \to M_1$ transition to 0 forces a partially "global" alignment in which all alignments to the main model must start at the first match or delete state. Setting all exits to 0 but the final $M \to E$ transition (which is always 1.0) forces a partially global alignment in which all alignments to the main model must end at the final match or delete state.

Multiple hit alignments are controlled by the $E \rightarrow J$ transition and the J state. If the $E \rightarrow J$ transition is set to 0, a sequence may only contain one domain (one alignment to the main model). If it is nonzero, more than one domain per sequence can be aligned to the main model. The $J \rightarrow J$ transition controls the expected length of the intervening sequence between domains; the lower this probability, the more clustered the domains are expected to be.

The original HMMER1 search programs are encoded in Plan 7 models as follows:

- **hmms** Simple global alignment. t_{NN} and t_{CC} set to t_{GG} from the null model (see below). t_{EJ} set to zero. Internal entries and exits set to zero.
- **hmm1s** Akin to "profile" alignment (as Waterman and others call it): global with respect to main model, local with respect to sequence, multiple nonoverlapping domains allowed. t_{NN} and t_{CC} set to t_{GG} from the null model. t_{EJ} set to 0.5. Internal entries and exits set to zero.
- **hmmsw** Smith/Waterman alignment: fully local with respect to both the main model and the sequence; single domain. t_{NN} and t_{CC} set to t_{GG} from the null model. t_{EJ} set to zero. Internal entries are set to 0.5/(M-1) for a model with M match states. Exit probabilities are set such that the overall chance of exiting from an internal match state is 0.5, and the posterior probability distribution over which match state is exited is equal (I'm trying to word this carefully; the $M \rightarrow E$ probabilities are not equal, because there's a small compensation for the fact that if you can leave from M_5 , then the chance that you'll even get to M_6 is reduced.)

hmmfs Multihit Smith/Waterman alignment. Same as the above, but t_{EJ} is set to 0.5.

One advantage of Plan 7 is great flexibility in choosing an alignment style. Complicated alignment styles are easily encoded in the model parameters without changing the alignment algorithm. For example, say you wanted to model human L1 retrotransposon elements. Because of the way L1 elements are inserted by reverse transcriptase (RVT), L1 elements tend to have a defined 3' end (RVT starts replication at the same place in each new L1) but a ragged 5' end (RVT prematurely falls off a new L1 in an unpredictable fashion). A specialized L1 model could define non-zero internal entry probabilities and zero internal exit probabilities to model this biological situation.

One disadvantage of Plan 7 is that if you decide you want to do both local and global alignments, you need two different models (or you need to do one search, then change the model). This wouldn't be a terrible burden except for the fact that the algorithm-dependent parameters strongly affect the values of the μ and λ parameters that E-value statistics depend on. If the algorithm dependent parameters are changed, these parameters are lost and the model should be recalibrated with hmmcalibrate – and hmmcalibrate is relatively slow.

The Plan 7 null model

When HMM alignments are scored, they are scored by a log-odds score relative to a "null model" of random sequence composition (Barrett et al., 1997). In Plan 7, this model is now specified as a full probabilistic model too:



The G state has a symbol emission probability distribution for K symbols in the alphabet. By default, this distribution is set either to the average amino acid composition of SWISSPROT 34, or to 0.25 for each nucleotide. The $G \rightarrow G$ transition controls the expected length of observed random sequences; in practice, this transition probability is so close to 1 that it has very little effect. The F state is just a dummy end state like the T state in the Plan 7 architecture.

Wing retraction in Plan 7 dynamic programming

In the figure of the Plan 7 architecture, you may have noticed that the first and last delete states are greyed out. Internally in HMMER, these delete states exist in the "probability form" of the model (when the model is being worked with in every way except alignments) but they are carefully removed in the "search form" of the model (when the model is converted to log-odds scores and used for alignments). This process is called "wing retraction" in the code, by analogy to a swept-wing fighter changing from a wings-out takeoff and landing configuration to a wings-back configuration for high speed flight.

The problem is that the Plan 7 model allows cycles through the J state. If a continuous nonemitting "mute cycle" were possible (J, B, D states, E, and back to J), dynamic programming recursions would fail. This is why special mute states like delete states must be handled carefully in HMM dynamic programming algorithms; see (Durbin et al., 1998) for further discussion. The easiest way to prevent a mute cycle is to make sure that the model must pass through at least one match state per path through the main model.

Wing retraction involves folding the probabilities of the terminal delete paths into the Plan 7 entry and exit probabilities. For example, in wing retraction the "algorithm dependent" $B \rightarrow M_3$ entry probability is incremented by the probability of the "data dependent" path $B \rightarrow D_1 \rightarrow D_2 \rightarrow M_3$.

Having the wing retraction step, rather than *always* folding these probabilities together, is a design decision, preserving a distinction between the "algorithm dependent" and "data dependent" parts of the model.

2.4 Sequence file formats

For all the programs, unaligned sequence files can be in FASTA, Genbank, EMBL, or SWISS-PROT format, as well as a few other common file formats. The programs automatically detect what format the file is in and whether the sequences are DNA, RNA, or protein.

Aligned sequence files can be in ClustalW, GCG MSF, or SELEX format. SELEX format is a simple format of one line per sequence, containing the name first, followed by the aligned sequence. ClustalW, MSF and SELEX alignment files can also be used where unaligned format files are required; the sequences will be read in and their gaps removed.

Full specifications of these file formats and the other formats recognized by the HMM package are in the file formats chapter near the end of the guide.

The programs work on RNA, DNA, and protein sequence. They automatically detect what your sequences are. The behavior of the programs when a nucleic acid model is used to analyze protein sequences, or vice versa, is undefined. Certain other situations may arise (trying to search the "complementary strand" of a protein database, for example) that are nonsensical in certain contexts. Be forewarned. If you're lucky, the software will issue a snide warning to you if you try to do something nonsensical, but usually it will assume you know what you're doing.

2.5 Command line options

If you forget the command-line syntax or available options of any of the programs, you can type the name of the program with no other arguments and get a short help message, including summaries of the common options.

Commonly used options are generally small letters, like -a. More infrequently used options are generally large letters, like -A. Expert or experimental options are generally in the GNU long form, like -- null2.

If you call any program with an option -h, you get an augmented help message, including version info (the software version number is helpful if you report bugs or other problems to me) and a complete summary of all the available options, including rarely used and expert/experimental ones.

2.6 Environment variables

HMMER is built to coexist peacefully with the BLAST suite of database search programs (Altschul, 1991). HMMER reads the following environment variables (the examples given use UNIX csh syntax):

- **BLASTDB** Location of sequence databases that hmmsearch will look in, in addition to the current working directory. Multiple directories are allowed, separated by colons. A trailing slash on each path is important to BLAST, but not to HMMER. Examples:
 - > setenv BLASTDB /nfs/databases/
 - > setenv BLASTDB /nfs/databases/:/nfs/moredatabases/
- **BLASTMAT** Location of substitution matrices that hmmbuild --pam (the PAM prior option) can read. Although HMMER can parse a colon-separated list, BLAST must have a single directory path here. Example:
 - > setenv BLASTMAT /nfs/databases/matrix/
 - **HMMERDB** Location of HMMs, PFAM, or other HMMER specific data files. Any program that reads an HMM file looks in both HMMERDB and the current working directory. Multiple directories are allowed, colon-separated. Examples:
 - > setenv HMMERDB /usr/local/lib/hmmer/
 - > setenv HMMERDB /usr/local/lib/hmmer/:/nfs/databases/pfam/
- HMMER_NCPU On multiprocessors that support POSIX threads (this includes almost all modern UNIX multiprocessors; for example, SGI Origin servers), the programs hmmcalibrate, hmmpfam, and hmmsearch run as parallelized, multithreaded applications. Normally they will take over all available CPUs in the machine. HM-MER_NCPU sets a maximum number of CPUs to utilize, so HMMER searches are "good citizens", leaving some CPU power for other jobs. An example of configuring HMMER to use only 16 processors on a 32-processor Origin: > setenv HMMER_NCPU 16

2.7 Other profile HMM implementations

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Several implementations of profile HMM methods and related PSSM methods are available. Some are listed in the table below.

Software	URL
HMMER	http://hmmer.wustl.edu/
SAM	http://www.cse.ucsc.edu/research/compbio/sam.html
PFTOOLS	http://ulrec3.unil.ch:80/profile/
HMMpro	http://www.netid.com/html/hmmpro.html
GENEWISE	http://www.sanger.ac.uk/Software/Wise2/
PROBE	ftp://ncbi.nlm.nih.gov/pub/neuwald/probe1.0/
META-MEME	http://www.cse.ucsd.edu/users/bgrundy/metameme.1.0.html
BLOCKS	http://www.blocks.fhcrc.org/
PSI-BLAST	http://www.ncbi.nlm.nih.gov/BLAST/newblast.html

HMMER, SAM, PFTOOLS, and HMMpro are the most closely related to the profile HMM methods introduced by Krogh et al. HMMpro is commercial, not free software.

Chapter 3

Manual pages

3.1 HMMER - profile hidden Markov model software

Synopsis

hmmalign	Align multiple sequences to a profile HMM.
hmmbuild	Build a profile HMM from a given multiple sequence alignment.
hmmcalibrate	Determine appropriate statistical significance parameters for a profile HMM prior to doing database searches.
hmmconvert	Convert HMMER profile HMMs to other formats, such as GCG profiles.
hmmemit	Generate sequences probabilistically from a profile HMM.
hmmfetch	Retrieve an HMM from an HMM database
hmmindex	Create a binary SSI index for an HMM database
hmmpfam	Search a profile HMM database with a sequence (i.e., annotate various kinds of domains in the query sequence).
hmmsearch	Search a sequence database with a profile HMM (i.e., find additional homologues of a modeled family).

Description

These programs use profile hidden Markov models (profile HMMs) to model the primary structure consensus of a family of protein or nucleic acid sequences.

Options

All **HMMER** programs give a brief summary of their command-line syntax and options if invoked without any arguments. When invoked with the single argument, **-h** (i.e., help), a program will report more verbose

command-line usage information, including rarely used, experimental, and expert options. **-h** will report version numbers which are useful if you need to report a bug or problem to me.

Each **HMMER** program has its own man page briefly summarizing command line usage. There is also a user's guide that came with the software distribution, which includes a tutorial introduction and more detailed descriptions of the programs. See http://hmmer.wustl.edu/ for on-line documentation and the current HMMER release.

In general, no command line options should be needed by beginning users. The defaults are set up for optimum performance in most situations. Options that are single lowercase letters (e.g. -a) are "common" options that are expected to be frequently used and will be important in many applications. Options that are single uppercase letters (e.g. -B) are usually less common options, but also may be important in some applications. Options that are full words (e.g. -verbose) are either rarely used, experimental, or expert options. Some experimental options are only there for my own ongoing experiments with HMMER, and may not be supported or documented adequately.

Sequence File Formats

In general, **HMMER** attempts to read most common biological sequence file formats. It autodetects the format of the file. It also autodetects whether the sequences are protein or nucleic acid. Standard IUPAC degeneracy codes are allowed in addition to the usual 4-letter or 20-letter codes.

Unaligned sequences	Unaligned sequence files may be in FASTA, Swissprot, EMBL, GenBank, PIR, In- telligenetics, Strider, or GCG format. These formats are documented in the User's Guide.
Sequence alignments	Multiple sequence alignments may be in CLUSTALW, SELEX, or GCG MSF for- mat. These formats are documented in the User's Guide.

Environment Variables

For ease of using large stable sequence and HMM databases, **HMMER** looks for sequence files and HMM files in the current working directory as well as in system directories specified by environment variables.

- **BLASTDB** Specifies the directory location of sequence databases. Example: /seqlibs/blastdb/. In installations that use BLAST software, this environment variable is likely to already be set.
- HMMERDB Specifies the directory location of HMM databases. Example: /seqlibs/pfam/.

3.2 hmmalign - align sequences to an HMM profile

Synopsis

hmmalign [options] hmmfile seqfile

Description

hmmalign reads an HMM file from *hmmfile* and a set of sequences from *seqfile*, aligns the sequences to the profile HMM, and outputs a multiple sequence alignment.

seqfile may be in any unaligned or aligned file format accepted by HMMER. If it is in a multiple alignment format (e.g. Stockholm, MSF, SELEX, ClustalW), the existing alignment is ignored (i.e., the sequences are read as if they were unaligned - hmmalign will align them the way it wants).

Options

- -h Print brief help; includes version number and summary of all options, including expert options.
- -m Include in the alignment only those symbols aligned to match states. Do not show symbols assigned to insert states.
- -o <f> Save alignment to file <f> instead of to standard output.
 - -q quiet; suppress all output except the alignment itself. Useful for piping or redirecting the output.

Expert Options

- --informat <*s*> Assert that the input *seqfile* is in format <*s*>; do not run Babelfish format autodection. This increases the reliability of the program somewhat, because the Babelfish can make mistakes; particularly recommended for unattended, high-throughput runs of HMMER. Valid format strings include FASTA, GENBANK, EMBL, GCG, PIR, STOCKHOLM, SELEX, MSF, CLUSTAL, and PHYLIP. See the User's Guide for a complete list.
 - --mapali < f > Reads an alignment from file < f > and aligns it as a single object to the HMM; e.g. the alignment in < f > is held fixed. This allows you to align sequences to a model with **hmmalign** and view them in the context of an existing trusted multiple alignment. The alignment to the alignment is defined by a "map" kept in the HMM, and so is fast and guaranteed to be consistent with the way the HMM was constructed from the alignment. The alignment in the file < f > must be exactly the alignment that the HMM was built from. Compare the --withali option.

--withali $\langle f \rangle$ Reads an alignment from file $\langle f \rangle$ and aligns it as a single object to the HMM; e.g. the alignment in $\langle f \rangle$ is held fixed. This allows you to align sequences to a model with **hmmalign** and view them in the context of an existing trusted multiple alignment. The alignment to the alignment is done with a heuristic (nonoptimal) dynamic programming procedure, which may be somewhat slow and is not guaranteed to be completely consistent with the way the HMM was constructed (though it should be quite close). However, any alignment can be used, not just the alignment that the HMM was built from. Compare the --mapali option.

3.3 hmmbuild - build a profile HMM from an alignment

Synopsis

hmmbuild [options] hmmfile alignfile

Description

hmmbuild reads a multiple sequence alignment file *alignfile*, builds a new profile HMM, and saves the HMM in *hmmfile*.

alignfile may be in ClustalW, GCG MSF, SELEX, Stockholm, or aligned FASTA alignment format. The format is automatically detected.

By default, the model is configured to find one or more nonoverlapping alignments to the complete model: multiple global alignments with respect to the model, and local with respect to the sequence. This is analogous to the behavior of the **hmmls** program of HMMER 1. To configure the model for multiple *local* alignments with respect to the model and local with respect to the sequence, a la the old program **hmmfs**, use the **-f** (fragment) option. More rarely, you may want to configure the model for a single global alignment (global with respect to both model and sequence), using the **-g** option; or to configure the model for a single local/local alignment (a la standard Smith/Waterman, or the old **hmmsw** program), use the **-s** option.

Options

- -f Configure the model for finding multiple domains per sequence, where each domain can be a local (fragmentary) alignment. This is analogous to the old **hmmfs** program of HMMER 1.
- -g Configure the model for finding a single global alignment to a target sequence, analogous to the old **hmms** program of HMMER 1.
- -h Print brief help; includes version number and summary of all options, including expert options.
- **-n** $\langle s \rangle$ Name this HMM $\langle s \rangle$. $\langle s \rangle$ can be any string of non-whitespace characters (e.g. one "word"). There is no length limit (at least not one imposed by HMMER; your shell will complain about command line lengths first).
- -o $\langle f \rangle$ Re-save the starting alignment to $\langle f \rangle$, in Stockholm format. The columns which were assigned to match states will be marked with x's in an #=RF annotation line. If either the --hand or --fast construction options were chosen, the alignment may have been slightly altered to be compatible with Plan 7 transitions, so saving the final alignment and comparing to the starting alignment can let you view these alterations. See the User's Guide for more information on this arcane side effect.
 - -s Configure the model for finding a single local alignment per target sequence. This is analogous to the standard Smith/Waterman algorithm or the **hmmsw** program of HMMER 1.

- -A Append this model to an existing *hmmfile* rather than creating *hmmfile*. Useful for building HMM libraries (like Pfam).
- **-F** Force overwriting of an existing *hmmfile*. Otherwise HMMER will refuse to clobber your existing HMM files, for safety's sake.

Expert Options

- --amino Force the sequence alignment to be interpreted as amino acid sequences. Normally HMMER autodetects whether the alignment is protein or DNA, but sometimes alignments are so small that autodetection is ambiguous. See --nucleic.
- --archpri $\langle x \rangle$ Set the "architecture prior" used by MAP architecture construction to $\langle x \rangle$, where $\langle x \rangle$ is a probability between 0 and 1. This parameter governs a geometric prior distribution over model lengths. As $\langle x \rangle$ increases, longer models are favored a priori. As $\langle x \rangle$ decreases, it takes more residue conservation in a column to make a column a "consensus" match column in the model architecture. The 0.85 default has been chosen empirically as a reasonable setting.
 - --binary Write the HMM to *hmmfile* in HMMER binary format instead of readable ASCII text.
 - --cfile <f> Save the observed emission and transition counts to <f> after the architecture has been determined (e.g. after residues/gaps have been assigned to match, delete, and insert states). This option is used in HMMER development for generating data files useful for training new Dirichlet priors. The format of count files is documented in the User's Guide.
 - --fast Quickly and heuristically determine the architecture of the model by assigning all columns will more than a certain fraction of gap characters to insert states. By default this fraction is 0.5, and it can be changed using the --gapmax option. The default construction algorithm is a maximum a posteriori (MAP) algorithm, which is slower.
- --gapmax $\langle x \rangle$ Controls the --*fast* model construction algorithm, but if --*fast* is not being used, has no effect. If a column has more than a fraction $\langle x \rangle$ of gap symbols in it, it gets assigned to an insert column. $\langle x \rangle$ is a frequency from 0 to 1, and by default is set to 0.5. Higher values of $\langle x \rangle$ mean more columns get assigned to consensus, and models get longer; smaller values of $\langle x \rangle$ mean fewer columns get assigned to consensus, and models get smaller. $\langle x \rangle$
 - --hand Specify the architecture of the model by hand: the alignment file must be in SELEX or Stockholm format, and the reference annotation line (#=RF in SELEX, #=GC RF in Stockholm) is used to specify the architecture. Any column marked with a non-gap symbol (such as an 'x', for instance) is assigned as a consensus (match) column in the model.
 - --idlevel <*x*> Controls both the determination of effective sequence number and the behavior of the --*wblosum* weighting option. The sequence alignment is clustered by percent

identity, and the number of clusters at a cutoff threshold of $\langle x \rangle$ is used to determine the effective sequence number. Higher values of $\langle x \rangle$ give more clusters and higher effective sequence numbers; lower values of $\langle x \rangle$ give fewer clusters and lower effective sequence numbers. $\langle x \rangle$ is a fraction from 0 to 1, and by default is set to 0.62 (corresponding to the clustering level used in constructing the BLOSUM62 substitution matrix).

- --informat <*s*> Assert that the input *seqfile* is in format <*s*>; do not run Babelfish format autodection. This increases the reliability of the program somewhat, because the Babelfish can make mistakes; particularly recommended for unattended, high-throughput runs of HMMER. Valid format strings include FASTA, GENBANK, EMBL, GCG, PIR, STOCKHOLM, SELEX, MSF, CLUSTAL, and PHYLIP. See the User's Guide for a complete list.
 - --noeff Turn off the effective sequence number calculation, and use the true number of sequences instead. This will usually reduce the sensitivity of the final model (so don't do it without good reason!)
 - --nucleic Force the alignment to be interpreted as nucleic acid sequence, either RNA or DNA. Normally HMMER autodetects whether the alignment is protein or DNA, but sometimes alignments are so small that autodetection is ambiguous. See -- amino.
 - --null <f> Read a null model from <f>. The default for protein is to use average amino acid frequencies from Swissprot 34 and p1 = 350/351; for nucleic acid, the default is to use 0.25 for each base and p1 = 1000/1001. For documentation of the format of the null model file and further explanation of how the null model is used, see the User's Guide.
 - --pam <f> Apply a heuristic PAM- (substitution matrix-) based prior on match emission probabilities instead of the default mixture Dirichlet. The substitution matrix is read from <f>. See --pamwgt. The default Dirichlet state transition prior and insert emission prior are unaffected. Therefore in principle you could combine --prior with --pam but this isn't recommended, as it hasn't been tested. (--pam itself hasn't been tested much!)
- --pamwgt $\langle x \rangle$ Controls the weight on a PAM-based prior. Only has effect if --pam option is also in use. $\langle x \rangle$ is a positive real number, 20.0 by default. $\langle x \rangle$ is the number of "pseudocounts" contributed by the heuristic prior. Very high values of $\langle x \rangle$ can force a scoring system that is entirely driven by the substitution matrix, making HMMER somewhat approximate Gribskov profiles.
- --pbswitch $\langle n \rangle$ For alignments with a very large number of sequences, the GSC, BLOSUM, and Voronoi weighting schemes are slow; they're O(N²) for N sequences. Henikoff position-based weights (PB weights) are more efficient. At or above a certain threshold sequence number $\langle n \rangle$ hmmbuild will switch from GSC, BLOSUM, or Voronoi weights to PB weights. To disable this switching behavior (at the cost of compute time, set $\langle n \rangle$ to be something larger than the number of sequences in your alignment. $\langle n \rangle$ is a positive integer; the default is 1000.

- --prior <*f*> Read a Dirichlet prior from <*f*>, replacing the default mixture Dirichlet. The format of prior files is documented in the User's Guide, and an example is given in the Demos directory of the HMMER distribution.
- --swentry $\langle x \rangle$ Controls the total probability that is distributed to local entries into the model, versus starting at the beginning of the model as in a global alignment. $\langle x \rangle$ is a probability from 0 to 1, and by default is set to 0.5. Higher values of $\langle x \rangle$ mean that hits that are fragments on their left (N or 5'-terminal) side will be penalized less, but complete global alignments will be penalized more. Lower values of $\langle x \rangle$ mean that fragments on the left will be penalized more, and global alignments on this side will be favored. This option only affects the configurations that allow local alignments, e.g. -s and -f; unless one of these options is also activated, this option has no effect. You have independent control over local/global alignment behavior for the N/C (5'/3') termini of your target sequences using --swentry and --swexit.
 - --swexit $\langle x \rangle$ Controls the total probability that is distributed to local exits from the model, versus ending an alignment at the end of the model as in a global alignment. $\langle x \rangle$ is a probability from 0 to 1, and by default is set to 0.5. Higher values of $\langle x \rangle$ mean that hits that are fragments on their right (C or 3'-terminal) side will be penalized less, but complete global alignments will be penalized more. Lower values of $\langle x \rangle$ mean that fragments on the right will be penalized more, and global alignments on this side will be favored. This option only affects the configurations that allow local alignments, e.g. -s and -f; unless one of these options is also activated, this option has no effect. You have independent control over local/global alignment behavior for the N/C (5'/3') termini of your target sequences using --swentry and --swexit.
 - --verbose Print more possibly useful stuff, such as the individual scores for each sequence in the alignment.
 - --wblosum Use the BLOSUM filtering algorithm to weight the sequences, instead of the default. Cluster the sequences at a given percentage identity (see --idlevel); assign each cluster a total weight of 1.0, distributed equally amongst the members of that cluster.
 - --wgsc Use the Gerstein/Sonnhammer/Chothia ad hoc sequence weighting algorithm. This is already the default, so this option has no effect (unless it follows another option in the --w family, in which case it overrides it).
 - --wme Use the Krogh/Mitchison maximum entropy algorithm to "weight" the sequences. This supercedes the Eddy/Mitchison/Durbin maximum discrimination algorithm, which gives almost identical weights but is less robust. ME weighting seems to give a marginal increase in sensitivity over the default GSC weights, but takes a fair amount of time.
 - --wnone Turn off all sequence weighting.
 - --wpb Use the Henikoff position-based weighting scheme.
 - --wvoronoi Use the Sibbald/Argos Voronoi sequence weighting algorithm in place of the default GSC weighting.

3.4 hmmcalibrate - calibrate HMM search statistics

Synopsis

hmmcalibrate [options] hmmfile

Description

hmmcalibrate reads an HMM file from *hmmfile*, scores a large number of synthesized random sequences with it, fits an extreme value distribution (EVD) to the histogram of those scores, and re-saves *hmmfile* now including the EVD parameters.

hmmcalibrate may take several minutes (or longer) to run. While it is running, a temporary file called *hmmfile.xxx* is generated in your working directory. If you abort **hmmcalibrate** prematurely (ctrl-C, for instance), your original *hmmfile* will be untouched, and you should delete the *hmmfile.xxx* temporary file.

Options

-h Print brief help; includes version number and summary of all options, including expert options.

Expert Options

- --cpu <*n*> Sets the maximum number of CPUs that the program will run on. The default is to use all CPUs in the machine. Overrides the HMMER_NCPU environment variable. Only affects threaded versions of HMMER (the default on most systems).
- --fixed $\langle n \rangle$ Fix the length of the random sequences to $\langle n \rangle$, where $\langle n \rangle$ is a positive (and reasonably sized) integer. The default is instead to generate sequences with a variety of different lengths, controlled by a Gaussian (normal) distribution.
- --histfile $\langle f \rangle$ Save a histogram of the scores and the fitted theoretical curve to file $\langle f \rangle$.
- --mean $\langle x \rangle$ Set the mean length of the synthetic sequences to $\langle x \rangle$, where $\langle x \rangle$ is a positive real number. The default is 350.
- --num $\langle n \rangle$ Set the number of synthetic sequences to $\langle n \rangle$, where $\langle n \rangle$ is a positive integer. If $\langle n \rangle$ is less than about 1000, the fit to the EVD may fail. Higher numbers of $\langle n \rangle$ will give better determined EVD parameters. The default is 5000; it was empirically chosen as a tradeoff between accuracy and computation time.
 - --pvm Run on a Parallel Virtual Machine (PVM). The PVM must already be running. The client program **hmmcalibrate-pvm** must be installed on all the PVM nodes. Optional PVM support must have been compiled into HMMER.
 - --sd $\langle x \rangle$ Set the standard deviation of the synthetic sequence length distribution to $\langle x \rangle$, where $\langle x \rangle$ is a positive real number. The default is 350. Note that the Gaussian is left-truncated so that no sequences have lengths $\langle = 0$.

--seed <n> Set the random seed to <n>, where <n> is a positive integer. The default is to use time() to generate a different seed for each run, which means that two different runs of hmmcalibrate on the same HMM will give slightly different results. You can use this option to generate reproducible results for different hmmcalibrate runs on the same HMM.

3.5 hmmconvert - convert between profile HMM file formats

Synopsis

hmmconvert [options] oldhmmfile newhmmfile

Description

hmmconvert reads an HMM file from *oldhmmfile* in any HMMER format, and writes it to a new file *newhmmfile* in a new format. *oldhmmfile* and *newhmmfile* must be different files; you can't reliably overwrite the old file. By default, the new HMM file is written in HMMER 2 ASCII format. Available formats are HMMER 2 ASCII (default), HMMER 2 binary (*-b*) GCG profile (*-p*), and Compugen XSW extended profile (*-P*).

Options

- -a Convert to HMMER 2 ASCII file. This is the default, so this option is unnecessary.
- -b Convert to HMMER 2 binary file.
- -h Print brief help; includes version number and summary of all options, including expert options.
- -p Convert to GCG profile .prf format.
- -A Append mode; append to *newhmmfile* rather than creating a new file.
- -F Force; if *newhmmfile* already exists, and -A is not being used to append to the file, hmmconvert will refuse to clobber the existing file unless -F is used.
- -P Convert the HMM to Compugen XSW extended profile format, which is similar to GCG profile format but has two extra columns for delete-open and delete-extend costs. (I do not believe that Compugen publicly supports this format; it may be undocumented.)

3.6 hmmemit - generate sequences from a profile HMM

Synopsis

hmmemit [options] hmmfile

Description

hmmemit reads an HMM file from *hmmfile* containing one or more HMMs, and generates a number of sequences from each HMM; or, if the **-c** option is selected, generate a single majority-rule consensus. This can be useful for various applications in which one needs a simulation of sequences consistent with a sequence family consensus.

By default, hmmemit generates 10 sequences and outputs them in FASTA (unaligned) format.

Options

- -a Write the generated sequences in an aligned format (SELEX) rather than FASTA.
- -c Predict a single majority-rule consensus sequence instead of sampling sequences from the HMM's probability distribution. Highly conserved residues ($p \ge 0.9$ for DNA, $p \ge 0.5$ for protein) are shown in upper case; others are shown in lower case. Some insert states may become part of the majority rule consensus, because they are used in $\ge 50\%$ of generated sequences; when this happens, insert-generated residues are simply shown as "x".
- -h Print brief help; includes version number and summary of all options, including expert options.
- -**n** <*n*> Generate <*n*> sequences. Default is 10.
- -o <*f*> Save the synthetic sequences to file <*f*> rather than writing them to stdout.
 - -q Quiet; suppress all output except for the sequences themselves. Useful for piping or directing the output.

Expert Options

--seed $\langle n \rangle$ Set the random seed to $\langle n \rangle$, where $\langle n \rangle$ is a positive integer. The default is to use **time**() to generate a different seed for each run, which means that two different runs of **hmmemit** on the same HMM will give slightly different results. You can use this option to generate reproducible results.

3.7 hmmfetch - retrieve an HMM from an HMM database

Synopsis

hmmfetch [options] database name

Description

hmmfetch is a small utility that retrieves an HMM called *name* from a HMMER model database called *database*. in a new format, and prints that model to standard output. For example, *hmmfetch* Pfam rrm retrieves the RRM (RNA recognition motif) model from Pfam, if the environment variable HMMERDB is set to the location of the Pfam database. The retrieved HMM file is written in HMMER 2 ASCII format.

The database must have an associated GSI index file. To index an HMM database, use the program **hmmindex.**

Options

-h Print brief help; includes version number and summary of all options, including expert options.

3.8 hmmindex - create a binary SSI index for an HMM database

Synopsis

hmmindex [options] database

Description

hmmindex is a utility that creates a binary SSI ("squid sequence index" format) index for an HMM database file called *database*. The new index file is named *database.ssi*. An SSI index file is required for **hmmfetch** to work, and also for the PVM implementation of **hmmpfam**.

Options

-h Print brief help; includes version number and summary of all options, including expert options.

3.9 hmmpfam - search one or more sequences against an HMM database

Synopsis

hmmpfam [options] hmmfile seqfile

Description

hmmpfam reads a sequence file *seqfile* and compares each sequence in it, one at a time, against all the HMMs in *hmmfile* looking for significantly similar sequence matches.

hmmfile will be looked for first in the current working directory, then in a directory named by the environment variable *HMMERDB*. This lets administrators install HMM library(s) such as Pfam in a common location.

There is a separate output report for each sequence in *seqfile*. This report consists of three sections: a ranked list of the best scoring HMMs, a list of the best scoring domains in order of their occurrence in the sequence, and alignments for all the best scoring domains. A sequence score may be higher than a domain score for the same sequence if there is more than one domain in the sequence; the sequence score takes into account all the domains. All sequences scoring above the -E and -T cutoffs are shown in the first list, then *every* domain found in this list is shown in the second list of domain hits. If desired, E-value and bit score thresholds may also be applied to the domain list using the *--domE* and *--domT* options.

Options

- -h Print brief help; includes version number and summary of all options, including expert options.
- -n Specify that models and sequence are nucleic acid, not protein. Other HMMER programs autodetect this; but because of the order in which **hmmpfam** accesses data, it can't reliably determine the correct "alphabet" by itself.
- -A < *n*> Limits the alignment output to the < *n*> best scoring domains. -A0 shuts off the alignment output and can be used to reduce the size of output files.
- -**E** <*x*> Set the E-value cutoff for the per-sequence ranked hit list to <*x*>, where <*x*> is a positive real number. The default is 10.0. Hits with E-values better than (less than) this threshold will be shown.
- **-T** <*x*> Set the bit score cutoff for the per-sequence ranked hit list to <*x*>, where <*x*> is a real number. The default is negative infinity; by default, the threshold is controlled by E-value and not by bit score. Hits with bit scores better than (greater than) this threshold will be shown.
- -Z < *n*> Calculate the E-value scores as if we had seen a sequence database of < *n*> sequences. The default is arbitrarily set to 59021, the size of Swissprot 34.

Expert Options

- --acc Report HMM accessions instead of names in the output reports. Useful for high-throughput annotation, where the data are being parsed for storage in a relational database.
- --compat Use the output format of HMMER 2.1.1, the 1998-2001 public release; provided so 2.1.1 parsers don't have to be rewritten.
- --cpu <*n*> Sets the maximum number of CPUs that the program will run on. The default is to use all CPUs in the machine. Overrides the HMMER_NCPU environment variable. Only affects threaded versions of HMMER (the default on most systems).
 - --cut_ga Use Pfam GA (gathering threshold) score cutoffs. Equivalent to --globT <GA1> -domT <GA2>, but the GA1 and GA2 cutoffs are read from each HMM in *hmmfile* individually. hmmbuild puts these cutoffs there if the alignment file was annotated in a Pfam-friendly alignment format (extended SELEX or Stockholm format) and the optional GA annotation line was present. If these cutoffs are not set in the HMM file, --cut_ga doesn't work.
 - --cut_tc Use Pfam TC (trusted cutoff) score cutoffs. Equivalent to --globT <TC1> --domT <TC2>, but the TC1 and TC2 cutoffs are read from each HMM in *hmmfile* individually. hmmbuild puts these cutoffs there if the alignment file was annotated in a Pfam-friendly alignment format (extended SELEX or Stockholm format) and the optional TC annotation line was present. If these cutoffs are not set in the HMM file, --cut_tc doesn't work.
 - --cut_nc Use Pfam NC (noise cutoff) score cutoffs. Equivalent to --globT <NC1> --domT <NC2>, but the NC1 and NC2 cutoffs are read from each HMM in *hmmfile* individually. hmmbuild puts these cutoffs there if the alignment file was annotated in a Pfam-friendly alignment format (extended SELEX or Stockholm format) and the optional NC annotation line was present. If these cutoffs are not set in the HMM file, --cut_nc doesn't work.
- --domE $\langle x \rangle$ Set the E-value cutoff for the per-domain ranked hit list to $\langle x \rangle$, where $\langle x \rangle$ is a positive real number. The default is infinity; by default, all domains in the sequences that passed the first threshold will be reported in the second list, so that the number of domains reported in the per-sequence list is consistent with the number that appear in the per-domain list.
- --domT $\langle x \rangle$ Set the bit score cutoff for the per-domain ranked hit list to $\langle x \rangle$, where $\langle x \rangle$ is a real number. The default is negative infinity; by default, all domains in the sequences that passed the first threshold will be reported in the second list, so that the number of domains reported in the per-sequence list is consistent with the number that appear in the per-domain list. *Important* note: only one domain in a sequence is absolutely controlled by this parameter, or by --domT. The second and subsequent domains in a sequence have a de facto bit score threshold of 0 because of the details of how HMMER works. HMMER requires at least one pass through the main model per sequence; to do more than one pass (more than one domain) the multidomain alignment must have a better score than the single domain alignment,

and hence the extra domains must contribute positive score. See the Users' Guide for more detail.

- --forward Use the Forward algorithm instead of the Viterbi algorithm to determine the persequence scores. Per-domain scores are still determined by the Viterbi algorithm. Some have argued that Forward is a more sensitive algorithm for detecting remote sequence homologues; my experiments with HMMER have not confirmed this, however.
- --informat <*s*> Assert that the input *seqfile* is in format <*s*>; do not run Babelfish format autodection. This increases the reliability of the program somewhat, because the Babelfish can make mistakes; particularly recommended for unattended, high-throughput runs of HMMER. Valid format strings include FASTA, GENBANK, EMBL, GCG, PIR, STOCKHOLM, SELEX, MSF, CLUSTAL, and PHYLIP. See the User's Guide for a complete list.
 - --null2 Turn off the post hoc second null model. By default, each alignment is rescored by a postprocessing step that takes into account possible biased composition in either the HMM or the target sequence. This is almost essential in database searches, especially with local alignment models. There is a very small chance that this postprocessing might remove real matches, and in these cases --null2 may improve sensitivity at the expense of reducing specificity by letting biased composition hits through.
 - --pvm Run on a Parallel Virtual Machine (PVM). The PVM must already be running. The client program hmmpfam-pvm must be installed on all the PVM nodes. The HMM database *hmmfile* and an associated GSI index file *hmmfile*.gsi must also be installed on all the PVM nodes. (The GSI index is produced by the program hmmindex.) Because the PVM implementation is I/O bound, it is highly recommended that each node have a local copy of *hmmfile* rather than NFS mounting a shared copy. Optional PVM support must have been compiled into HMMER for --pvm to function.
 - --xnu Turn on XNU filtering of target protein sequences. Has no effect on nucleic acid sequences. In trial experiments, --xnu appears to perform less well than the default post hoc null2 model.

3.10 hmmsearch - search a sequence database with a profile HMM

Synopsis

hmmsearch [options] hmmfile seqfile

Description

hmmsearch reads an HMM from *hmmfile* and searches *seqfile* for significantly similar sequence matches.

seqfile will be looked for first in the current working directory, then in a directory named by the environment variable *BLASTDB*. This lets users use existing BLAST databases, if BLAST has been configured for the site.

hmmsearch may take minutes or even hours to run, depending on the size of the sequence database. It is a good idea to redirect the output to a file.

The output consists of four sections: a ranked list of the best scoring sequences, a ranked list of the best scoring domains, alignments for all the best scoring domains, and a histogram of the scores. A sequence score may be higher than a domain score for the same sequence if there is more than one domain in the sequence; the sequence score takes into account all the domains. All sequences scoring above the -E and -T cutoffs are shown in the first list, then *every* domain found in this list is shown in the second list of domain hits. If desired, E-value and bit score thresholds may also be applied to the domain list using the --domE and --domT options.

Options

- -h Print brief help; includes version number and summary of all options, including expert options.
- -A < n > Limits the alignment output to the < n > best scoring domains. -A0 shuts off the alignment output and can be used to reduce the size of output files.
- -**E** <*x*> Set the E-value cutoff for the per-sequence ranked hit list to <*x*>, where <*x*> is a positive real number. The default is 10.0. Hits with E-values better than (less than) this threshold will be shown.
- **-T** <*x*> Set the bit score cutoff for the per-sequence ranked hit list to <*x*>, where <*x*> is a real number. The default is negative infinity; by default, the threshold is controlled by E-value and not by bit score. Hits with bit scores better than (greater than) this threshold will be shown.
- -Z < *n*> Calculate the E-value scores as if we had seen a sequence database of < *n*> sequences. The default is the number of sequences seen in your database file < *seqfile*>.

Expert Options

- --compat Use the output format of HMMER 2.1.1, the 1998-2001 public release; provided so 2.1.1 parsers don't have to be rewritten.
- --cpu <*n*> Sets the maximum number of CPUs that the program will run on. The default is to use all CPUs in the machine. Overrides the HMMER_NCPU environment variable. Only affects threaded versions of HMMER (the default on most systems).
 - --cut_ga Use Pfam GA (gathering threshold) score cutoffs. Equivalent to --globT <GA1> --domT <GA2>, but the GA1 and GA2 cutoffs are read from the HMM file. hmmbuild puts these cutoffs there if the alignment file was annotated in a Pfam-friendly alignment format (extended SELEX or Stockholm format) and the optional GA annotation line was present. If these cutoffs are not set in the HMM file, --cut_ga doesn't work.
 - --cut_tc Use Pfam TC (trusted cutoff) score cutoffs. Equivalent to --globT <TC1> --domT <TC2>, but the TC1 and TC2 cutoffs are read from the HMM file. hmmbuild puts these cutoffs there if the alignment file was annotated in a Pfam-friendly alignment format (extended SELEX or Stockholm format) and the optional TC annotation line was present. If these cutoffs are not set in the HMM file, --cut_tc doesn't work.
 - --cut_nc Use Pfam NC (noise cutoff) score cutoffs. Equivalent to --globT <NC1> --domT <NC2>, but the NC1 and NC2 cutoffs are read from the HMM file. hmmbuild puts these cutoffs there if the alignment file was annotated in a Pfam-friendly alignment format (extended SELEX or Stockholm format) and the optional NC annotation line was present. If these cutoffs are not set in the HMM file, --cut_nc doesn't work.
- --domE $\langle x \rangle$ Set the E-value cutoff for the per-domain ranked hit list to $\langle x \rangle$, where $\langle x \rangle$ is a positive real number. The default is infinity; by default, all domains in the sequences that passed the first threshold will be reported in the second list, so that the number of domains reported in the per-sequence list is consistent with the number that appear in the per-domain list.
- --domT $\langle x \rangle$ Set the bit score cutoff for the per-domain ranked hit list to $\langle x \rangle$, where $\langle x \rangle$ is a real number. The default is negative infinity; by default, all domains in the sequences that passed the first threshold will be reported in the second list, so that the number of domains reported in the per-sequence list is consistent with the number that appear in the per-domain list. *Important* note: only one domain in a sequence is absolutely controlled by this parameter, or by --domT. The second and subsequent domains in a sequence have a de facto bit score threshold of 0 because of the details of how HMMER works. HMMER requires at least one pass through the main model per sequence; to do more than one pass (more than one domain) the multidomain alignment must have a better score than the single domain alignment, and hence the extra domains must contribute positive score. See the Users' Guide for more detail.
 - --forward Use the Forward algorithm instead of the Viterbi algorithm to determine the persequence scores. Per-domain scores are still determined by the Viterbi algorithm.

Some have argued that Forward is a more sensitive algorithm for detecting remote sequence homologues; my experiments with HMMER have not confirmed this, however.

- --informat <*s*> Assert that the input *seqfile* is in format <*s*>; do not run Babelfish format autodection. This increases the reliability of the program somewhat, because the Babelfish can make mistakes; particularly recommended for unattended, high-throughput runs of HMMER. Valid format strings include FASTA, GENBANK, EMBL, GCG, PIR, STOCKHOLM, SELEX, MSF, CLUSTAL, and PHYLIP. See the User's Guide for a complete list.
 - --null2 Turn off the post hoc second null model. By default, each alignment is rescored by a postprocessing step that takes into account possible biased composition in either the HMM or the target sequence. This is almost essential in database searches, especially with local alignment models. There is a very small chance that this postprocessing might remove real matches, and in these cases --null2 may improve sensitivity at the expense of reducing specificity by letting biased composition hits through.
 - --pvm Run on a Parallel Virtual Machine (PVM). The PVM must already be running. The client program **hmmsearch-pvm** must be installed on all the PVM nodes. Optional PVM support must have been compiled into HMMER.
 - --xnu Turn on XNU filtering of target protein sequences. Has no effect on nucleic acid sequences. In trial experiments, --xnu appears to perform less well than the default post hoc null2 model.

3.11 afetch - retrieve an alignment from an alignment database

Synopsis

afetch [options] alignmentdb key afetch --index alignmentdb

Description

afetch retrieves the alignment named key from an alignment database in file alignmentdb.

alignmentdb is a "multiple multiple alignment" file in Stockholm (e.g. native Pfam) format.

key is either the name (ID) of the alignment, or its accession number (AC).

The *alignmentdb* file should first be SSI indexed with **afetch --index** for efficient retrieval. An SSI index is not required, but alignment retrieval without one may be painfully slow.

Options

-h Print brief help; includes version number and summary of all options, including expert options.

Expert Options

--index Instead of retrieving a *key*, the special command **afetch** --index *alignmentdb* produces an SSI index of the names and accessions of the alignments in the file *alignmentdb*. This should be run once on the *alignmentdb* file to prepare it for all future afetch's.

3.12 alistat - show statistics for a multiple alignment file

Synopsis

alistat [options] alignfile

Description

alistat reads a multiple sequence alignment from the file *alignfile* in any supported format (including SE-LEX, GCG MSF, and CLUSTAL), and shows a number of simple statistics about it. These statistics include the name of the format, the number of sequences, the total number of residues, the average and range of the sequence lengths, the alignment length (e.g. including gap characters).

Also shown are some percent identities. A percent pairwise alignment identity is defined as (*idents* / MIN(len1, len2)) where *idents* is the number of exact identities and *len1*, len2 are the unaligned lengths of the two sequences. The "average percent identity", "most related pair", and "most unrelated pair" of the alignment are the average, maximum, and minimum of all (N)(N-1)/2 pairs, respectively. The "most distant seq" is calculated by finding the maximum pairwise identity (best relative) for all N sequences, then finding the minimum of these N numbers (hence, the most outlying sequence).

Options

- -a Show additional verbose information: a table with one line per sequence showing name, length, and its highest and lowest pairwise identity. These lines are prefixed with a * character to enable easily grep'ing them out and sorting them. For example, *alistat* -a foo.slx | grep "*" | sort -n +3 gives a ranked list of the most distant sequences in the alignment. Incompatible with the -f option.
- -f Fast; use a sampling method to estimate the average %id. When this option is chosen, alistat doesn't show the other three pairwise identity numbers. This option is useful for very large alignments, for which the full (N)(N-1) calculation of all pairs would be prohibitive (e.g. Pfam's GP120 alignment, with over 10,000 sequences). Incompatible with the -a option.
- -h Print brief help; includes version number and summary of all options, including expert options.
- -q be quiet suppress the verbose header (program name, release number and date, the parameters and options in effect).
- -B (Babelfish). Autodetect and read a sequence file format other than the default (FASTA). Almost any common sequence file format is recognized (including Genbank, EMBL, SWISS-PROT, PIR, and GCG unaligned sequence formats, and Stockholm, GCG MSF, and Clustal alignment formats). See the printed documentation for a complete list of supported formats.

Expert Options

--informat <*s*> Specify that the sequence file is in format <*s*>, rather than the default FASTA format. Common examples include Genbank, EMBL, GCG, PIR, Stockholm, Clustal, MSF, or PHYLIP; see the printed documentation for a complete list of accepted format names. This option overrides the default format (FASTA) and the -*B* Babelfish autodetection option.

3.13 seqstat - show statistics and format for a sequence file

Synopsis

seqstat [options] seqfile

Description

seqstat reads a sequence file *seqfile* and shows a number of simple statistics about it. The printed statistics include the name of the format, the residue type of the first sequence (protein, RNA, or DNA), the number of sequences, the total number of residues, and the average and range of the sequence lengths.

Options

- -a Show additional verbose information: a table with one line per sequence showing name, length, and description line. These lines are prefixed with a * character to enable easily **grep**'ing them out and sorting them.
- -h Print brief help; includes version number and summary of all options, including expert options.
- -B (Babelfish). Autodetect and read a sequence file format other than the default (FASTA). Almost any common sequence file format is recognized (including Genbank, EMBL, SWISS-PROT, PIR, and GCG unaligned sequence formats, and Stockholm, GCG MSF, and Clustal alignment formats). See the printed documentation for a complete list of supported formats.

Expert Options

- --informat <*s*> Specify that the sequence file is in format <*s*>, rather than the default FASTA format. Common examples include Genbank, EMBL, GCG, PIR, Stockholm, Clustal, MSF, or PHYLIP; see the printed documentation for a complete list of accepted format names. This option overrides the default expected format (FASTA) and the -*B* Babelfish autodetection option.
 - --quiet Suppress the verbose header (program name, release number and date, the parameters and options in effect).

3.14 sfetch - get a sequence from a flatfile database.

Synopsis

sfetch [options] seqname

Description

sfetch retrieves the sequence named *seqname* from a sequence database.

Which database is used is controlled by the **-d** and **-D** options, or "little databases" and "big databases". The directory location of "big databases" can be specified by environment variables, such as \$SWDIR for Swissprot, and \$GBDIR for Genbank (see **-D** for complete list). A complete file path must be specified for "little databases". By default, if neither option is specified and the name looks like a Swissprot identifier (e.g. it has a _ character), the \$SWDIR environment variable is used to attempt to retrieve the sequence *seqname* from Swissprot.

A variety of other options are available which allow retrieval of subsequences (-f,-t); retrieval by accession number instead of by name (-a); reformatting the extracted sequence into a variety of other formats (-F); etc.

If the database has been GSI indexed, sequence retrieval will be extremely efficient; else, retrieval may be painfully slow (the entire database may have to be read into memory to find *seqname*). GSI indexing is recommended for all large or permanent databases. This program was originally named **getseq**, and was renamed because it clashed with a GCG program of the same name.

Options

- -a Interpret *seqname* as an accession number, not an identifier.
- -d <*seqfile*> Retrieve the sequence from a sequence file named <*seqfile*>. If a GSI index <*seqfile*>.*gsi* exists, it is used to speed up the retrieval.
 - -f <*from*> Extract a subsequence starting from position <*from*>, rather than from 1. See -t. If <*from*> is greater than <*to*> (as specified by the -t option), then the sequence is extracted as its reverse complement (it is assumed to be nucleic acid sequence).
 - -h Print brief help; includes version number and summary of all options, including expert options.
- -o *<outfile>* Direct the output to a file named *<outfile>*. By default, output would go to stdout.
- -r <*newname*> Rename the sequence <*newname*> in the output after extraction. By default, the original sequence identifier would be retained. Useful, for instance, if retrieving a sequence fragment; the coordinates of the fragment might be added to the name (this is what Pfam does).

- -t < to> Extract a subsequence that ends at position < to>, rather than at the end of the sequence. See -f. If < to> is less than < from> (as specified by the -f option), then the sequence is extracted as its reverse complement (it is assumed to be nucleic acid sequence)
 - -B (Babelfish). Autodetect and read a sequence file format other than the default (FASTA). Almost any common sequence file format is recognized (including Genbank, EMBL, SWISS-PROT, PIR, and GCG unaligned sequence formats, and Stockholm, GCG MSF, and Clustal alignment formats). See the printed documentation for a complete list of supported formats.
- -D < database> Retrieve the sequence from the main sequence database coded < database>. For each code, there is an environment variable that specifies the directory path to that database. Recognized codes and their corresponding environment variables are Dsw (Swissprot, \$SWDIR); -Dpir (PIR, \$PIRDIR); -Dem (EMBL, \$EMBLDIR); -Dgb (Genbank, \$GBDIR); -Dwp (Wormpep, \$WORMDIR); and -Dowl (OWL, \$OWLDIR). Each database is read in its native flatfile format.
 - -F <*format*> Reformat the extracted sequence into a different format. (By default, the sequence is extracted from the database in the same format as the database.) Available formats are **embl, fasta, genbank, gcg, strider, zuker, ig, pir, squid,** and **raw.**

Expert Options

--informat <*s*> Specify that the sequence file is in format <*s*>, rather than the default FASTA format. Common examples include Genbank, EMBL, GCG, PIR, Stockholm, Clustal, MSF, or PHYLIP; see the printed documentation for a complete list of accepted format names. This option overrides the default format (FASTA) and the -*B* Babelfish autodetection option.

3.15 shuffle - randomize the sequences in a sequence file

Synopsis

shuffle [options] seqfile

Description

shuffle reads a sequence file *seqfile*, randomizes each sequence, and prints the randomized sequences in FASTA format on standard output. The sequence names are unchanged; this allows you to track down the source of each randomized sequence if necessary.

The default is to simply shuffle each input sequence, preserving monosymbol composition exactly. To shuffle each sequence while preserving both its monosymbol and disymbol composition exactly, use the -d option.

The -0 and -1 options allow you to generate sequences with the same Markov properties as each input sequence. With -0, for each input sequence, 0th order Markov statistics are collected (e.g. symbol composition), and a new sequence is generated with the same composition. With -1, the generated sequence has the same 1st order Markov properties as the input sequence (e.g. the same disymbol frequencies). Note that the default and -0, or -d and -1, are similar; the shuffling algorithms preserve composition exactly, while the Markov algorithms only expect to generate a sequence of similar composition on average. Other shuffling algorithms are also available, as documented below in the options.

Options

- -0 Calculate 0th order Markov frequencies of each input sequence (e.g. residue composition); generate output sequence using the same 0th order Markov frequencies.
- -1 Calculate 1st order Markov frequencies for each input sequence (e.g. diresidue composition); generate output sequence using the same 1st order Markov frequencies. The first residue of the output sequence is always the same as the first residue of the input sequence.
- -d Shuffle the input sequence while preserving both monosymbol and disymbol composition exactly. Uses an algorithm published by S.F. Altschul and B.W. Erickson, Mol. Biol. Evol. 2:526-538, 1985.
- -h Print brief help; includes version number and summary of all options, including expert options.
- -I Look only at the length of each input sequence; generate an i.i.d. output protein sequence of that length, using monoresidue frequencies typical of proteins (taken from Swissprot 35).
- -n < *n*> Make < *n*> different randomizations of each input sequence in *seqfile*, rather than the default of one.

- -r Generate the output sequence by reversing the input sequence. (Therefore only one "randomization" per input sequence is possible, so it's not worth using *-n* if you use reversal.)
- -t <*n*> Truncate each input sequence to a fixed length of exactly <*n*> residues. If the input sequence is shorter than <*n*> it is discarded (therefore the output file may contain fewer sequences than the input file). If the input sequence is longer than <*n*> a contiguous subsequence is randomly chosen.
- -w <n> Regionally shuffle each input sequence in window sizes of <n>, preserving local residue composition in each window. Probably a better shuffling algorithm for biosequences with nonstationary residue composition (e.g. composition that is varying along the sequence, such as between different isochores in human genome sequence).
 - -B (Babelfish). Autodetect and read a sequence file format other than the default (FASTA). Almost any common sequence file format is recognized (including Genbank, EMBL, SWISS-PROT, PIR, and GCG unaligned sequence formats, and Stockholm, GCG MSF, and Clustal alignment formats). See the printed documentation for a complete list of supported formats.

Expert Options

- --informat <*s*> Specify that the sequence file is in format <*s*>, rather than the default FASTA format. Common examples include Genbank, EMBL, GCG, PIR, Stockholm, Clustal, MSF, or PHYLIP; see the printed documentation for a complete list of accepted format names. This option overrides the default expected format (FASTA) and the -*B* Babelfish autodetection option.
 - --nodesc Do not output any sequence description in the output file, only the sequence names.
 - --seed $\langle s \rangle$ Set the random number seed to $\langle s \rangle$. If you want reproducible results, use the same seed each time. By default, **shuffle** uses a different seed each time, so does not generate the same output in subsequent runs with the same input.

3.16 sreformat - convert sequence file to different format

Synopsis

sreformat [options] format seqfile

Description

sreformat reads the sequence file *seqfile* in any supported format, reformats it into a new format specified by *format*, then prints the reformatted text.

Supported input formats include (but are not limited to) the unaligned formats FASTA, Genbank, EMBL, SWISS-PROT, PIR, and GCG, and the aligned formats Stockholm, Clustal, GCG MSF, and Phylip.

Available unaligned output file format codes include *fasta* (FASTA format); *embl* (EMBL/SWISSPROT format); *genbank* (Genbank format); *gcg* (GCG single sequence format); *gcgdata* (GCG flatfile database format); *strider* (MacStrider format); *zuker* (Zuker MFOLD format); *ig* (Intelligenetics format); *pir* (PIR/CODATA flatfile format); *squid* (an undocumented St. Louis format); *raw* (raw sequence, no other information).

The available aligned output file format codes include *stockholm* (PFAM/Stockholm format); *msf* (GCG MSF format); *a2m* (aligned FASTA format, called A2M by the UC Santa Cruz HMM group); *PHYLIP* (Felsenstein's PHYLIP format); and *selex* (old SELEX/HMMER/Pfam annotated alignment format);

All thee codes are interpreted case-insensitively (e.g. MSF, Msf, or msf all work).

Unaligned format files cannot be reformatted to aligned formats. However, aligned formats can be reformatted to unaligned formats -- gap characters are simply stripped out.

This program was originally named **reformat**, but that name clashes with a GCG program of the same name.

Options

- -a Enable alignment reformatting. By default, sreformat expects that the input file should be handled as an unaligned input file (even if it is an alignment), and it will not allow you to convert an unaligned file to an alignment (for obvious reasons). This may seem silly; surely if sreformat can autodetect and parse alignment file formats as input, it can figure out when it's got an alignment! There are two reasons. One is just the historical structure of the code. The other is that FASTA unaligned format and A2M aligned format (aligned FASTA) are impossible to tell apart with 100% confidence.
- -d DNA; convert U's to T's, to make sure a nucleic acid sequence is shown as DNA not RNA. See -r.
- -h Print brief help; includes version number and summary of all options, including expert options.
- -l Lowercase; convert all sequence residues to lower case. See -u.

- -r RNA; convert T's to U's, to make sure a nucleic acid sequence is shown as RNA not DNA. See -d.
- -u Uppercase; convert all sequence residues to upper case. See -l.
- -x For DNA sequences, convert non-IUPAC characters (such as X's) to N's. This is for compatibility with benighted people who insist on using X instead of the IUPAC ambiguity character N. (X is for ambiguity in an amino acid residue). Warning: the code doesn't check that you are actually giving it DNA. It simply literally just converts non-IUPAC DNA symbols to N. So if you accidentally give it protein sequence, it will happily convert most every amino acid residue to an N.
- -B (Babelfish). Autodetect and read a sequence file format other than the default (FASTA). Almost any common sequence file format is recognized (including Genbank, EMBL, SWISS-PROT, PIR, and GCG unaligned sequence formats, and Stockholm, GCG MSF, and Clustal alignment formats). See the printed documentation for a complete list of supported formats.

Expert Options

- --informat <*s*> Specify that the sequence file is in format <*s*>, rather than the default FASTA format. Common examples include Genbank, EMBL, GCG, PIR, Stockholm, Clustal, MSF, or PHYLIP; see the printed documentation for a complete list of accepted format names. This option overrides the default format (FASTA) and the -*B* Babelfish autodetection option.
 - --mingap If *seqfile* is an alignment, remove any columns that contain 100% gap characters, minimizing the overall length of the alignment. (Often useful if you've extracted a subset of aligned sequences from a larger alignment.)
 - --pfam For SELEX alignment output format only, put the entire alignment in one block (don't wrap into multiple blocks). This is close to the format used internally by Pfam in Stockholm and Cambridge.
 - --sam Try to convert gap characters to UC Santa Cruz SAM style, where a . means a gap in an insert column, and a means a deletion in a consensus/match column. This only works for converting aligned file formats, and only if the alignment already adheres to the SAM convention of upper case for residues in consensus/match columns, and lower case for residues in insert columns. This is true, for instance, of all alignments produced by old versions of HMMER. (HMMER2 produces alignments that adhere to SAM's conventions even in gap character choice.) This option was added to allow Pfam alignments to be reformatted into something more suitable for profile HMM construction using the UCSC SAM software.
- --samfrac <x> Try to convert the alignment gap characters and residue cases to UC Santa Cruz SAM style, where a . means a gap in an insert column and a means a deletion in a consensus/match column, and upper case means match/consensus residues and lower case means inserted residues. This will only work for converting aligned file formats, but unlike the --sam option, it will work regardless of whether the file

adheres to the upper/lower case residue convention. Instead, any column containing more than a fraction $\langle x \rangle$ of gap characters is interpreted as an insert column, and all other columns are interpreted as match columns. This option was added to allow Pfam alignments to be reformatted into something more suitable for profile HMM construction using the UCSC SAM software.

Chapter 4

File formats

4.1 HMMER save files

The file Demos/rrm.hmm gives an example of a HMMER ASCII save file. An abridged version is shown here, where (...) mark deletions made for clarity and space:

HMMER2.0									
NAME rrm									
ACC PF00076									
DESC RNA recognition motif. (aka RRM, RBD, or RNP domain)									
LENG 72									
ALPH Amino									
RF no									
CS no									
MAP yes									
COM/src/hmmbuild -F rrm.hmm rrm.slx									
COM/src/hmmcalibrate rrm.hmm									
NSEQ 70									
DATE Wed Jul 8 08:13:25 1998									
CKSUM 2768									
GA 13.3 0.0									
TC 13.40 0.60									
NC 13.20 13.20									
XT -8455 -4 -1000 -1000 -8455 -4 -8455	-4								
NULT -4 -8455									
NULE 595 -1558 85 338 -294 453 -1158 ()								
EVD -49.999123 0.271164									
HMM A C D E F G H	I ()								
m->m m->i m->d i->m i->i d->m d->d b-	->m m->e								
-21 * -6129									
1 -1234 -371 -8214 -7849 -5304 -8003 -7706 23	384 () 1								
149 -500 233 43 -381 399 106 -6	626 ()								
11 -11284 -12326 -894 -1115 -701 -1378 -	-21 *								
2 -3634 -3460 -5973 -5340 3521 -2129 -4036 -8	831 () 2								
149 -500 233 43 -381 399 106 -6	626 ()								
11 -11284 -12326 -894 -1115 -701 -1378	* *								
()									
71 -1165 -4790 -240 -275 -5105 -4306 1035 -20	009 () 90								
149 -500 233 43 -381 398 106 -6	626 ()								

	-	-43	-6001	-12336	-150	-3342	-701	-1378	*	*	
	72	-1929	1218	-1535	-1647	-3990	-4677	-3410	1725	() 92	
	-	*	*	*	*	*	*	*	*	()	
	-	*	*	*	*	*	*	*	*	0	
11											

HMMER2 profile HMM save files have a very different format compared to the previous HMMER1 ASCII formats. The HMMER2 format provides all the necessary parameters to compare a protein sequence to a HMM, including the search mode of the HMM (hmmls, hmmfs, hmmsw, and hmms in the old HMMER1 package), the null (background) model, and the statistics to evaluate the match on the basis of a previously fitted extreme value distribution.

The format consists of one or more HMMs. Each HMM starts with the identifier "HMMER2.0" on a line by itself and ends with // on a line by itself. The identifier allows backward compatibility as the HMMer software evolves. The closing // allows multiple HMMs to be concatenated into a single file to provide a database of HMMs.

The format for an HMM is divided into two regions. The first region contains text information and miscalleneous parameters in a (roughly) tag-value scheme, akin to EMBL formats. This section is ended by a line beginning with the keyword HMM. The second region is of a more fixed format and contains the main model parameters. It is ended by the // that ends the entire definition for a single profile-HMM.

Both regions contain probabilities that are used parameterize the HMM. These are stored as integers which are related to the probability via a log-odds calculation. The log-odds score calculation is defined in mathsupport.c and is:

$$score = (\texttt{int})\texttt{floor}(0.5 + (\texttt{INTSCALE} * \log_2(\texttt{prob}/\texttt{null-prob})))$$

so conversely, to get a probability from the scores in an HMM save file:

$$prob = null-prob * 2^{score/INTSCALE}$$

INTSCALE is defined in config.h as 1000.

Notice that you must know a null model probability to convert scores back to HMM probabilities.

The special case of prob = 0 is translated to "*", so a score of * is read as a probability of 0. Null model probabilities are not allowed to be 0.

This log-odds format has been chosen because it has a better dynamic range than storing probabilities as ASCII text, and because the numbers are more meaningful to a human reader to a certain extent: positive values means a better than expected probability, and negative values a worse than expected probability. However, because of the conversion from probabilities, it should be noted that *you should not edit the numbers in a HMMER save file directly*. The HMM is a probabilistic model and expects state transition and symbol emission probability distributions to sum to one. If you want to edit the HMM, you must understand the underlying Plan7 probabilistic model, and ensure the correct summations yourself.

A more detailed description of the format follows.

Header section

In the header section, each line after the initial identifier has a unique tag of five characters or less. For shorter tags, the remainder of the five characters is padded with spaces. Therefore the first six characters of these lines are reserved for the tag and a space. The remainder of the line starts at the seventh character. The parser does require this.

- HMMER2.0 File format version; a unique identifier for this save file format. Used for backwards compatibility. *Not* necessarily the version number of the HMMER software that generated it; rather, the version number of the last HMMER that changed the format so much that a whole new function had to be introduced to do the parsing. (i.e., HMMER 2.8 might still be writing save files that are headed HMMER2.0). Mandatory.
- NAME <s> Model name; <s> is a single word name for the HMM. No spaces or tabs may occur in the name. hmmbuild will use the #=ID line from a SELEX alignment file to set the name. If this is not present, or the alignment is not in SELEX format, hmmbuild sets the HMM name using the name of the alignment file, after removing any file type suffix. For example, an HMM built from the alignment file rrm.slx would be named rrm by default. Mandatory.
 - ACC **<s>** Accession number; **<s>** is a one-word accession number for an HMM. Used in Pfam maintenance. Accessions are stable identifiers for Pfam models, whereas names may change from release to release. Added in v2.1.1. **Optional.**
- DESC <s> Description line; <s> is a one-line description of the HMM. hmmbuild will use the #=DE line from a SELEX alignment file to set the description line. If this is not present, or the alignment is not in SELEX format, the description line is left blank; one can be added manually (or by Perl script) if you wish. Optional.
- **LENG** <**d>** Model length; <**d>**, a positive nonzero integer, is the number of match states in the model. **Mandatory.**
- ALPH <s> Symbol alphabet; <s> must be either Amino or Nucleic. This determines the symbol alphabet and the size of the symbol emission probability distributions. If Amino, the alphabet size is set to 20 and the symbol alphabet to "ACDEFGHIKLM-NPQRSTVWY" (alphabetic order). If Nucleic, the alphabet size is set to 4 and the symbol alphabet to "ACGT". Case insensitive. Mandatory.
 - **RF <s>** Reference annotation flag; **<s>** must be either no or yes (case insensitive). If set to yes, a character of reference annotation is read for each match state/consensus column in the main section of the file (see below); else this data field will be ignored. Reference annotation lines are currently somewhat inconsistently used. The only major use in HMMER is to specify which columns of an alignment get turned into match states when using the hmmbuild --hand manual model construction option. Reference annotation can only be picked up from SELEX format alignments. See description of SELEX format for more details on reference annotation lines. **Optional**; assumed to be no if not present.
 - **CS <s>** Consensus structure annotation flag; **<s>** must be either no or yes (case insensitive). If set to yes, a character of consensus structure annotation is read for each match state/consensus column in the main section of the file (see below); else this data field will be ignored. Consensus structure annotation lines are currently somewhat inconsistently used. Consensus structure annotation can only be picked up from SELEX format alignments. See description of SELEX format for more details on consensus structure annotation lines. **Optional**; assumed to be no if not present.

- MAP <s> Map annotation flag; <s> must be either no or yes (case insensitive). If set to yes, each line of data for the match state/consensus column in the main section of the file is followed by an extra number. This number gives the index of the alignment column that the match state was made from. This information provides a "map" of the match states (1...M) onto the columns of the alignment (1...alen). It is used for quickly aligning the model back to the original alignment, e.g. when using hmmalign --mapali. Added in v2.0.1. Optional; assumed to be no if not present.
- **COM <s>** Command line log; **<s>** is a one-line command. There may be more than one COM line per save file. These lines record the command line for every HMMER command that modifies the save file. This helps us automatically log Pfam construction strategies, for example. **Optional.**
- **CKSUM** <d> Training alignment checksum; <d> is a nonzero positive integer. This number is calculated from the training alignment and stored when hmmbuild is used. It is used in conjunction with the alignment map information to to verify that some alignment is indeed the alignment that the map is for. Added in v2.0.1. **Optional.**
- **GA** <**f**> <**f**> Pfam gathering thresholds GA1 and GA2. This is a feature in progress. See Pfam documentation of GA lines. Added in v2.1.1. **Optional.**
- **TC** <**f**> <**f**> Pfam trusted cutoffs TC1 and TC2. This is a feature in progress. See Pfam documentation of TC lines. Added in v2.1.1. **Optional.**
- NC <f> <f> Pfam noise cutoffs NC1 and NC2. This is a feature in progress. See Pfam documentation of NC lines. Added in v2.1.1. **Optional.**
 - **NSEQ** <**d>** Sequence number; <**d>** is a nonzero positive integer, the number of sequences that the HMM was trained on. This field is only used for logging purposes. **Optional.**
 - **DATE <s>** Creation date; **<s>** is a date string. This field is only used for logging purposes. **Optional.**
 - **XT** <d>*8 Eight "special" transitions for controlling parts of the algorithm-specific parts of the Plan7 model. The null probability used to convert these back to model probabilities is 1.0. The order of the eight fields is $N \rightarrow B$, $N \rightarrow N$, $E \rightarrow C$, $E \rightarrow J$, $C \rightarrow T$, $C \rightarrow C$, $J \rightarrow B$, $J \rightarrow J$. (Another way to view the order is as four transition probability distributions for N,E,C,J; each distribution has two probabilities, the first one for "moving" and the second one for "looping".) For an explanation of these special transitions (and definition of the state names), read the Plan7 architecture documentation. **Mandatory.**
- **NULT** <d> <d> The transition probability distribution for the null model (single G state). The null probability used to convert these back to model probabilities is 1.0. The order is G \rightarrow G, G \rightarrow F. Mandatory.
 - **NULE** $\langle d \rangle * \kappa$ The symbol emission probability distribution for the null model (G state); consists of K (e.g. 4 or 20) integers. The null probability used to convert these back to model probabilities is 1/K. (Yes, it's a little weird to have a "null probability"

for the null model symbol emission probabilities; this is strictly an aesthetic decision, so one can look at the null model and easily tell which amino acids are more common than chance expectation in the background distribution.) **Mandatory.**

- **EVD** $\langle \mathbf{f} \rangle \langle \mathbf{f} \rangle$ The extreme value distribution parameters μ and λ , respectively; both floating point values. λ is positive and nonzero. These values are set when the model is calibrated with hmmcalibrate. They are used to determine E-values of bit scores. If this line is not present, E-values are calculated using a conservative analytic upper bound. **Optional.**
 - **HMM** HMM flag line; flags the end of the header section. Otherwised not parsed. Strictly for human readability, the symbol alphabet is also shown on this line, aligned to the NULE fields and the fields of the match and insert symbol emission distributions in the main model. The immediately next line is also an unparsed human annotation line: column headers for the state transition probability fields in the main model section that follows. Both lines are **mandatory.**

Main model section

All the remaining fields are **mandatory**, except for the alignment map.

The first line in the main model section is atypical; it contains three fields, for transitions from the B state into the first node of the model. The only purpose of this line is to set the $B \to D$ transition probability. The first field is the score for $1 - t(B \to D)$. The second field is always "*" (there is no $B \to I$ transition). The third field is the score for $t(B \to D)$. The null probability used for converting these scores back to probabilities is 1.0. In principle, only the third number is needed to obtain $t(B \to D)$. In practice, HMMER reads both the first and the third number, converts them to probabilities, and renormalizes the distribution to obtain $t(B \to D)$. *

The remainder of the model has three lines per node, for M nodes (where M is the number of match states, as given by the LENG line). These three lines are:

Match emission line The first field is the node number (1..M). The HMMER parser verifies this number as a consistency check (it expects the nodes to come in order). The next K numbers for match emission scores, one per symbol. The null probability used to convert them to probabilities is the relevant null model emission probability calculated from the NULE line.

If MAP was yes, then there is one more number on this line, representing the alignment column index for this match state. See MAP above for more information about the alignment map, and also see the man pages for hmmalign --mapali. Added in v2.0.1. This field is optional, for backwards compatibility with 2.0.

Insert emission line The first field is a character of reference annotation (RF), or "-" if there is no reference annotation. The remaining fields are K numbers for insert emission scores, one per symbol, in alphabetic order. The null probability used to convert them to probabilities is the relevant null model emission probability calculated from the NULE line.

^{*}OK, it's more complicated than it has to be. Sometimes you only discover your temporary insanity when you're trying to document it.

State transition line The first field is a character of consensus structure annotation (CS), or "-" if there is no consensus structure annotation. The remaining 9 fields are state transition scores. The null probability used to convert them back from log odds scores to probabilities is 1.0. The order of these scores is given by the annotation line at the top of the main section: it is $M \to M, M \to I, M \to D; I \to M, I \to D; D \to M, D \to D; B \to M; M \to E.$

The insert emission and state transition lines for the final node M are special. Node M has no insert state, so all the insert emissions are given as "*". (In fact, this line is skipped by the parser, except for its RF annotation.) There is also no next node, so only the $B \rightarrow M$ and $M \rightarrow E$ transitions are valid; the first seven transitions are always "*". (Incidentally, the $M \rightarrow E$ transition score for the last node is always 0, because this probability has to be 1.0.)

Finally, the last line of the format is the "//" record separator.

Renormalization

After the parser reads the file and converts the scores back to probabilities, it renormalizes the probability distributions to sum to 1.0 to eliminate minor rounding/conversion/numerical imprecision errors. If you're trying to emulate HMMER save files, it might be useful to know what HMMER considers to be a probability distribution. See Plan7Renormalize() in plan7.c for the relevant function.

null emissions The K symbol emissions given on the NULE line.

null transitions The two null model transitions given on the NULT line.

- **N,E,C,J specials** Each of the four special states N,E,C,J have two state transition probabilities (move and loop). All four distributions are specified on the XT line.
 - **B** transitions $M \to M$ entry probabilities are given by the 9th field in the state transition line of each of the M nodes. The $B \to D$ transition (from the atypical first line of the main model section) is also part of this state transition distribution.
- **match transitions** One distribution of 4 numbers per node; $M \to M, M \to I, M \to D$, and $M \to E$ (fields 2, 3, 4, and 10 in the state transition line of each node). Note the asymmetry between B \to M and M \to E; entries are a probability distribution of their own, while exits are not.
- insert transitions One distribution of 2 numbers per node; $I \rightarrow M$, $I \rightarrow I$ (fields 5 and 6 of the state transition line of each node).
- **delete transitions** One distribution of 2 numbers per node; $D \rightarrow M$, $D \rightarrow D$ (fields 7 and 8 of the state transition line of each node).
- **match emissions** One distribution of K numbers per node; the K match symbol emissions given on the first line of each node in the main section.
- insert emissions One distribution of K numbers per node; the K insert symbol emissions given on the second line of each node in the main section.

Note to developers

Though I make an effort to keep this documentation up to date, it may lag behind the code. For definitive answers, please check the parsing code in hmmio.c. The relevant function to see what's being written is WriteAscHMM(). The relevant function to see how it's being parsed is read_asc20hmm().

4.2 HMMER null model files

A "null model" is used to calculate HMMER log odds scores. The null model states the expected background occurrence frequencies of the 20 amino acids or the 4 nucleotide bases. The null model also contains a parameter called p1, which is the $G \rightarrow G$ transition probability in the Plan7 null model (see the figure in the Introduction).

For protein models, by default, the 20 residue frequencies are set to the amino acid composition of SWISS-PROT 34, and p1 is set to 350/351 (which, because the Plan7 null model implies a geometric length distribution, states that the mean length of a protein is about 350 residues). For DNA/RNA models, by default, the 4 residue frequencies are set to 0.25 each, and p1 is set to 1000/1001. [In the code, see prior.c:p7DefaultNullModel(), and the amino acid frequencies are set in iupac.c:aafg.]

Each HMM carries its own null model (see above, HMM file format). The null model is determined when the model is built using hmmbuild. The default null model can be overridden using the --null < f > option to hmmbuild, where < f > is the name of a null model file.

Two example null model files, amino.null and nucleic.null, are provided in the Demos subdirectory of the HMMER distribution. (They are copies of the internal default HMMER null model settings.) nucleic.null looks like this:

```
# nucleic.null
#
# Example of a null model file for DNA/RNA sequences.
# The values in this file are the HMMER 2 default
# settings.
Nucleic
0.25  # A
0.25  # C
0.25  # G
0.25  # T
0.999001 # p1
```

Anything on a line following a # is a comment, and is ignored by the software. Blank lines are also ignored. Valid fields are separated by blanks or new lines. Only the order that the fields occur in the file is important, not how they're put on lines; for example, 20 required fields can all occur on one line separated by blanks, or on 20 separate lines.

There must be 6 or 22 non-comment fields in a null model file, occurring in the following order:

- Alphabet type The first (non-comment) word in the file must be Nucleic or Amino, specifying what kind of sequence the null model is for.
- **Emission probabilities** 4 or 20 background frequencies for the amino acids or nucleotides. These *must* come in alphabetical order (the A, C, G, T comments in the example above are only for easier human viewing, and aren't parsed by the software).

p1 probability The $G \to G$ transition probability in the null model. Basically, if the expected mean length of target sequences is x, p1 should be $\frac{x}{x+1}$.

Null model files are parsed in prior.c:P7ReadNullModel().

4.3 HMMER prior files

Observed counts of emissions (residues) and transitions (insertions and deletions) in a multiple alignment are combined with *Dirichlet priors* to convert them to probabilities in an HMM.

For protein models, by default, HMMER uses a nine-component mixture Dirichlet prior for match emissions, and single component Dirichlet priors for insert emissions and transitions. The nine-component match emission mixture Dirichlet comes from the work of Kimmen Sjölander (Sjölander et al., 1996).

For DNA/RNA models, by default, HMMER uses single component Dirichlets.

Two example null model files, amino.pri and nucleic.pri, are provided in the Demos subdirectory of the HMMER distribution. (They are copies of the internal default HMMER prior settings.)

The way the format of these files is parsed is identical to null models: everything after a # on a line is a comment, the order of occurrence of the fields is important, and fields must be separated by either blanks or newlines.

A prior file consists of the following fields:

Strategy Must be the keyword Dirichlet. Currently this is the only available prior strategy in the public HMMER release.

Alphabet type Must be either Amino or Nucleic.

- **Transition priors** 1 + 8a fields, where a is the number of transition mixture components. The first field is the number of transition prior components, a (often just 1). Then, for each component, eight fields follow: the prior probability of that mixture component (1.0 if there is only one component), then the Dirichlet alpha parameters for the seven transitions, in order of $M \to M$, $M \to I$, $M \to D$, $I \to M$, $I \to I$, $D \to M$, $D \to I$.
- Match emission priors 1 + (5 or 21)b fields, where b is the number of match emission mixture components. The first field is the number of match emission mixture components, b. Then, for each component, 5 or 21 fields follows: the prior probability of that mixture component (1.0 if there is only one component), then the Dirichlet alpha parameters for the 4 or 20 residue types, in alphabetical order.
- **Insert emission priors** 1 + (5 or 21)c fields, where c is the number of insert emission mixture components. The first field is the number of insert emission mixture components, c. Then, for each component, 5 or 21 fields follows: the prior probability of that mixture component (1.0 if there is only one component), then the Dirichlet alpha parameters for the 4 or 20 residue types, in alphabetical order.

In the code, prior files are parsed by prior.c:P7ReadPrior().

4.4 Sequence files

Supported file formats

HMMER can automatically recognize and parse a number of common file formats. The best supported of these formats are listed below. If you know your sequence database is in one of these formats, you can use the file. If you are formatting sequences yourself, see the section of FASTA format below for unaligned sequences, and the section on SELEX format for alignments; these are the simplest formats to hand type

Unaligned sequence formats Pearson FASTA format; BLAST databases FASTA SWISS-PROT SWISS-PROT protein sequence database PIR protein sequence database PIR EMBL DNA sequence database EMBL GenBank GenBank DNA database flat files Wisconsin GCG sequence database format GCGdata GCG Wisconsin GCG single sequence format Multiple sequence alignment formats Stockholm The preferred format for HMMER/Pfam (see below) **SELEX** The old format for HMMER/Pfam MSF GCG alignment format CLUSTALW and CLUSTALV format CLUSTAL A2M "Aligned FASTA"

For programs that do sequential, database-style access (i.e. where you'd usually use an unaligned flat file), the alignment formats are read as if they were multiple unaligned sequences.

There is no provision for enforcing that single unaligned sequence formats (i.e. GCG unaligned sequence format) are single sequence only. HMMER will happily try to read more than one sequence if your file contains more than one. However, this may not give the results you expected.

Staden "experiment file" format is parsed using the EMBL file parser, but this functionality is relatively unsupported. There is one wrinkle in this. Staden experiment files use '-' characters to indicate 'N' – i.e., that a base is present in a sequence read, but its identity is unknown. Therefore, the software replaces any '-' in an EMBL sequence with an 'N'. Sometimes people use the unaligned formats to distribute aligned sequences by including gap characters. If EMBL files are used i this way for aligned strings, they must use a different character than '-' to indicate gaps.

FASTA unaligned sequence format

An example of a simple FASTA file:

FASTA is probably the simplest of formats for unaligned sequences. FASTA files are easily created in a text editor. Each sequence is preceded by a line starting with >. The first word on this line is the name of the sequence. The rest of the line is a description of the sequence (free format). The remaining lines contain the sequence itself. You can put as many letters on a sequence line as you want.

Blank lines in a FASTA file are ignored, and so are spaces or other gap symbols (dashes, underscores, periods) in a sequence. Any other non-amino or non-nucleic acid symbols in the sequence should produce an appropriately strident string of warnings on yur terminal screen when you try to use the file.

HMMER currently has a limit of 4095 characters on lines in files. You may see this limitation in NCBI NR databases, where some description lines will be truncated by HMMER.

Stockholm, the recommended multiple sequence alignment format

While we recommend a community standard format (FASTA) for unaligned sequence files, the recommended multiple alignment file format is not a community standard. The Pfam Consortium developed a format (based on extended SELEX) called "Stockholm format". The reasons for this are two-fold. First, there really is no standard accepted format for multiple sequence alignment files, so we don't feel guilty about inventing a new one. Second, the formats of popular multiple alignment software (e.g. CLUSTAL, GCG MSF, PHYLIP) do not support rich documentation and markup of the alignment. Stockholm format was developed to support extensible markup of multiple sequence alignments, and we use this capability extensively in both RNA work (with structural markup) and the Pfam database (with extensive use of both annotation and markup).

A minimal Stockholm file

STOCKHOLM 1.0
seq1 ACDEF...GHIKL
seq2 ACDEF...GHIKL
seq3 ...EFMNRGHIKL
seq1 MNPQTVWY
seq2 MNPQTVWY
seq3 MNPQT...

The simplest Stockholm file is pretty intuitive, easily generated in a text editor. It is usually easy to convert alignment formats into a "least common denominator" Stockholm format. For instance, SELEX, GCG's MSF format, and the output of the CLUSTALV multiple alignment program are all similar interleaved formats.

The first line in the file must be # STOCKHOLM 1.x, where x is a minor version number for the format specification (and which currently has no effect on my parsers). This line allows a parser to instantly identify the file format.

In the alignment, each line contains a name, followed by the aligned sequence. A dash or period denotes a gap. If the alignment is too long to fit on one line, the alignment may be split into multiple blocks, with blocks separated by blank lines. The number of sequences, their order, and their names must be the same in every block. Within a given block, each (sub)sequence (and any associated #=GR and #=GC markup, see below) is of equal length, called the *block length*. Block lengths may differ from block to block; the block length must be at least one residue, and there is no maximum.

Other blank lines are ignored. You can add comments to the file on lines starting with a #.

All other annotation is added using a tag/value comment style. The tag/value format is inherently extensible, and readily made backwards-compatible; unrecognized tags will simply be ignored. Extra annotation includes consensus and individual RNA or protein secondary structure, sequence weights, a reference coordinate system for the columns, and database source information including name, accession number, and coordinates (for subsequences extracted from a longer source sequence) See below for details.

Syntax of Stockholm markup

There are four types of Stockholm markup annotation, for per-file, per-sequence, per-column, and perresidue annotation:

> **#=GF** <tag> <s> Per-file annotation. <s> is a free format text line of annotation type <tag>. For example, #=GF DATE April 1, 2000. Can occur anywhere in the file, but usually all the #=GF markups occur in a header.

#=GS <seqname> <tag> <s> Per-sequence annotation. **<**s**>** is a free format text line of annotation type tag associated with the sequence named **<**seqname**>**. For example, **#**=GS seq1 SPECIES_SOURCE Caenorhabditis elegans. Can occur anywhere in the file, but in single-block formats (e.g. the Pfam distribution) will typically follow on the line after the sequence itself, and in multi-block formats (e.g. HMMER output), will typically occur in the header preceding the alignment but following the **#**=GF annotation.

#=GC < tag> <...s. Per-column annotation. <...s. is an aligned text line of annotation type < tag>. #=GC lines are associated with a sequence alignment block; <...s. is aligned to the residues in the alignment block, and has the same length as the rest of the block. Typically #=GC lines are placed at the end of each block.

#=GR <seqname> <tag> <....> Per-residue annotation. **<...s...>** is an aligned text line of annotation type **<tag>**, associated with the sequence named **<seqname>**. **#=GR** lines are associated with one sequence in a sequence alignment block; **<...s...>** is aligned to the residues in that sequence, and has the same length as the rest of the block. Typically **#=GR** lines are placed immediately following the aligned sequence they annotate.

Semantics of Stockholm markup

Any Stockholm parser will accept syntactically correct files, but is not obligated to do anything with the markup lines. It is up to the application whether it will attempt to interpret the meaning (the semantics) of the markup in a useful way. At the two extremes are the Belvu alignment viewer and the HMMER profile hidden Markov model software package.

Belvu simply reads Stockholm markup and displays it, without trying to interpret it at all. The tag types (#=GF, etc.) are sufficient to tell Belvu how to display the markup: whether it is attached to the whole file, sequences, columns, or residues.

HMMER uses Stockholm markup to pick up a variety of information from the Pfam multiple alignment database. The Pfam consortium therefore agrees on additional syntax for certain tag types, so HMMER can parse some markups for useful information. This additional syntax is imposed by Pfam, HMMER, and other software of mine, not by Stockholm format per se. You can think of Stockholm as akin to XML, and what my software reads as akin to an XML DTD, if you're into that sort of structured data format lingo.

The Stockholm markup tags that are parsed semantically by my software are as follows:

Recognized #=GF annotations

- **ID <s>** Identifier. **<s>** is a name for the alignment; e.g. "rrm". One word. Unique in file.
- AC <s> Accession. <s> is a unique accession number for the alignment; e.g. "PF00001". Used by the Pfam database, for instance. Often a alphabetical prefix indicating the database (e.g. "PF") followed by a unique numerical accession. One word. Unique in file.
- **DE <s>** Description. **<s>** is a free format line giving a description of the alignment; e.g. "RNA recognition motif proteins". One line. Unique in file.
- AU <s> Author. <s> is a free format line listing the authors responsible for an alignment;e.g. "Bateman A". One line. Unique in file.
- GA <f> <f> Gathering thresholds. Two real numbers giving HMMER bit score per-sequence and per-domain cutoffs used in gathering the members of Pfam full alignments. See Pfam and HMMER documentation for more detail.
- NC <f> <f> Noise cutoffs. Two real numbers giving HMMER bit score per-sequence and perdomain cutoffs, set according to the highest scores seen for unrelated sequences when gathering members of Pfam full alignments. See Pfam and HMMER documentation for more detail.
- **TC** <**f**> <**f**> Trusted cutoffs. Two real numbers giving HMMER bit score per-sequence and per-domain cutoffs, set according to the lowest scores seen for true homologous sequences that were above the GA gathering thresholds, when gathering members of Pfam full alignments. See Pfam and HMMER documentation for more detail.

Recognized #=GS annotations

- WT <f> Weight. <f> is a positive real number giving the relative weight for a sequence, usually used to compensate for biased representation by downweighting similar sequences. Usually the weights average 1.0 (e.g. the weights sum to the number of sequences in the alignment) but this is not required. Either every sequence must have a weight annotated, or none of them can.
- **AC <s>** Accession. **<s>** is a database accession number for this sequence. (Compare the #=GF AC markup, which gives an accession for the whole alignment.) One word.
- **DE <s>** Description. **<s>** is one line giving a description for this sequence. (Compare the #=GF DE markup, which gives a description for the whole alignment.)

Recognized #=GC annotations

- **RF** Reference line. Any character is accepted as a markup for a column. The intent is to allow labeling the columns with some sort of mark.
- **SS_cons** Secondary structure consensus. For protein alignments, DSSP codes or gaps are accepted as markup: [HGIEBTSCX.-_], where H is alpha helix, G is 3/10-helix, I is p-helix, E is extended strand, B is a residue in an isolated b-bridge, T is a turn, S is a

bend, C is a random coil or loop, and X is unknown (for instance, a residue that was not resolved in a crystal structure). For RNA alignments the symbols > and < are used for base pairs (pairs point at each other). + indicate definitely single-stranded positions, and any gap symbol indicates unassigned bases or single-stranded positions. This description roughly follows (Konings and Hogeweg, 1989). RNA pseudoknots are represented by alphabetic characters, with upper case letters representing the 5' side of the helix and lower case letters representing the 3' side. Note that this limits the annotation to a maximum of 26 pseudoknots per sequence.

SA_cons Surface accessibility consensus. 0-9, gap symbols, or X are accepted as markup. 0 means ;10% accessible residue surface area, 1 means ;20%, 9 means ;100%, etc. X means unknown structure.

Recognized #=GR annotations

- **ss** Secondary structure consensus. See #=GC SS_cons above.
- **SA** Surface accessibility consensus. See #=GC SA_cons above.

SELEX alignment format

An example of a simple SELEX alignment file:

Example selex file
seq1 ACGACGACGACG.
seq2 ..GGGAAAGG.GA
seq3 UUU..AAAUUU.A
seq1 ..ACG
seq2 AAGGG
seq3 AA...UUU

SELEX is an interleaved multiple alignment format that arose as an intuitive format that was easy to write and manipulate manually with a text editor like emacs. It is usually easy to convert other alignment formats into SELEX format, even with a couple of lines of Perl, but it can be harder to go the other way, since SELEX is more free-format than other alignment formats. For instance, GCG's MSF format and the output of the CLUSTALV multiple alignment program are similar interleaved formats that can be converted to SELEX just by stripping a small number of non-sequence lines out. Because SELEX evolved to accomodate different user input styles, it is very tolerant of various inconsistencies such as different gap symbols, varying line lengths, etc.

Each line contains a name, followed by the aligned sequence. A space, dash, underscore, or period denotes a gap. If the alignment is too long to fit on one line, the alignment is split into multiple blocks, separated by blank lines. The number of sequences, their order, and their names must be the same in every block (even if a sequence has no residues in a given block!) Other blank lines are ignored. You can add comments to the file on lines starting with a #.

SELEX stands for "Systematic Evolution of Ligands by Exponential Enrichment" – it refers to the Tuerk and Gold technology for evolving families of small RNAs for particular functions (Tuerk and Gold, 1990).

SELEX files were what we used to keep track of alignments of these small RNA families. It's an interesting piece of historical baggage – most of my lab works on RNA, not protein.

As the format evolved, more features have been added. To maintain compatibility with past alignment files, the new features are added using a reserved comment style. You don't have to worry about adding these extra information lines. They are generally the province of automated SELEX-generating software, such as my koala sequence alignment editor or the COVE and HMMER sequence analysis packages. This extra information includes consensus and individual RNA or protein secondary structure, sequence weights, a reference coordinate system for the columns, and database source information including name, accession number, and coordinates (for subsequences extracted from a longer source sequence) See below for details.

Detailed specification of a SELEX file

- 1. Any line beginning with a #= as the first two characters is a machine comment. #= comments are reserved for parsed data about the alignment. Often these features are maintained by software such as the koala editor or the HMMER software, not by hand. The format of #= lines is usually quite specific, since they must be parsed by the file-reading software.
- 2. All other lines beginning with a % or # as the first character are user comments. User comments are ignored by all software. Anything may appear on these lines. Any number of comments may be included in a SELEX file, and at any point.
- 3. Lines of data consist of a name followed by a sequence. The total length of the line must be smaller than 4096 characters.
- 4. Names must be a single word. Any non-whitespace characters are accepted. No spaces are tolerated in names: names MUST be a single word. Names must be less than 32 characters long.
- 5. In the sequence, any of the characters –_. or a space are recognized as gaps. Any other characters are interpreted as sequence. Sequence is case-sensitive. There is a common assumption by my software that upper-case symbols are used for consensus (match) positions and lower-case symbols are used for inserts. This language of "match" versus "insert" comes from the hidden Markov model formalism (Krogh et al., 1994). To almost all of my software, this isn't important, and it immediately converts the sequence to all upper-case after it's read.
- 6. Multiple different sequences are grouped in a block of data lines. Blocks are separated by blank lines. No blank lines are tolerated between the sequence lines in a block. Each block in a multi-block file of a long alignment must have its sequences in the same order in each block. The names are checked to verify that this is the case; if not, only a warning is generated. (In manually constructed files, some users may wish to use shorthand names in subsequent blocks after an initial block with full names but this isn't recommended.)

Optional SELEX file annotation

Header

Several pieces of information can be added to the header of the file using the following machine comments:

#=ID <s> Name of the alignment; **<s>** is one word, no spaces (example: "rrm"). If present, HMMER's hmmbuild picks this up and uses it as the name of an HMM.

- **#=AC <s>** Accession number of the alignment; **<s>** is one word, no spaces (example: "PF999999"). If present, HMMER's hmmbuild picks this up and uses it as the accession number of an HMM. (This is fairly specific to the support of the Pfam HMM database.)
- **#=DE <s>** Description line; **<s>** is a one-line description of the alignment. If present, HM-MER's hmmbuild picks this up and uses it as the description line of an HMM.
- **#=AU <s>** Author line; **<s>** is a one-line description of who made the alignment. HMMER doesn't use this line for anything.
- **#=GA <f1> <f2>** Gathering cutoffs; <f1> is the per-sequence GA1 cutoff, <f2> is the per-domain GA2 cutoff. Both numbers are floating point numbers (HMMER bit scores). This is very specific to Pfam support. The GA cutoffs specify the HMMER score cutoffs used in automatically generating the Pfam full alignments.
- **#=TC <f1> <f2>** Trusted cutoffs; **<f1>** is the per-sequence TC1 cutoff, **<f2>** is the per-domain TC2 cutoff. Both numbers are floating point numbers (HMMER bit scores). This is very specific to Pfam support. The TC cutoffs specify the HMMER scores of the lowest-scoring sequence/domain included in a Pfam full alignment.
- **#=NC** <**f1>** <**f2>** Noise cutoffs; <**f1>** is the per-sequence NC1 cutoff, <**f2>** is the per-domain NC2 cutoff. Both numbers are floating point numbers (HMMER bit scores). This is very specific to Pfam support. The NC cutoffs specify the HMMER scores of the highest scoring sequence/domain that were not included in a Pfam full alignment.

Sequence header

Additional per-sequence information can be placed after the main header lines and before any blocks appear. These lines, one per sequence and in exactly the same order as the sequences appear in the alignment, are formatted like

#=SQ <name> <wgt> <source> <accession> <start..stop::orig length> <desc>

This information includes a sequence weight (for compensating for biased representation of subfamilies of sequences in the alignment); source information, if the sequence came from a database, consisting of identifier, accession number, and source coordinates; and a description of the sequence.

If a #=SQ line is present, all the fields must be present on each line and one #=SQ line must be present for each sequence. If no information is available for a field, use '-' for all the fields except the source coordinates, which would be given as '0'.

Secondary structure annotation

Lines beginning with #=SS or #=CS are individual or consensus secondary structure data, respectively. #=SS individual secondary structure lines must immediately follow the sequence they are associated with. There can only be one #=SS per sequence. All, some, or none of the sequences may have #=SS annotation.

#=CS consensus secondary structure predictions precede all the sequences in each block. There can only be one #=CS per file.

I use one-letter codes to indicate secondary structures. Secondary structure strings are aligned to sequence blocks just like additional sequences. For RNA secondary structure, the symbols > and < are used for base pairs (pairs point at each other). + indicate definitely single-stranded positions, and any gap symbol indicates unassigned bases or singlestranded positions. This description roughly follows (Konings and Hogeweg, 1989). For protein secondary structure, I use E to indicate residues in β -sheet, H for those in α -helix, L for those in loops, and any gap symbol for unassigned or unstructured residues.

RNA pseudoknots are represented by alphabetic characters, with upper case letters representing the 5' side of the helix and lower case letters representing the 3' side. Note that this restricts the annotation to a maximum of 26 pseudoknots per sequence.

Reference coordinate system

Alignments are usually numbered by some reference coordinate system, often a canonical molecule. For instance, tRNA positions are numbered by reference to the positions of yeast tRNA-Phe.

A line beginning with #=RF preceding the sequences in a block gives a reference coordinate system. Any non-gap symbol in the #=RF line indicates that sequence positions in its columns are numbered. For instance, the #=RF lines for a tRNA alignment would have 76 non-gap symbols for the canonical numbered columns; they might be the aligned tRNA-Phe sequence itself, or they might be just X's.

The #=RF line also provides a good way to mark conserved key residues in a protein alignment. HM-MER picks up both the #=RF and #=CS lines when it builds an HMM, if they're there, and will display them in the context of any alignments to target sequences.

An example SELEX file with annotation

Here's an example SELEX file with examples of all the optional fields. The sequences are evolved RNA ligands for the phage R17 coat protein (Schneider et al., 1992).

```
#=ID r17
#=AC RNA00001
#=DE RNA ligands evolved to bind phage R17 coat protein
#=GA 0.0 0.0
#=TC 0.0 0.0
#=NC 0.0 0.0
#=AU COVE 2.2.3 Wed Dec 2 08:39:58 1998
#=SQ lig28 1.0 - - 1..29:29 R17 coat protein ligand 28
#=SQ lig1 1.0 - - 1..17:17 R17 coat protein ligand 1
#=SQ lig2 1.0 - - 1..29:29 R17 coat protein ligand 2
            ***A** loop *****
\#=RF
#=CS
      .....>>>+>> ++++ <<<<<.....
liq28 GGAGUAAGAUAGC AUCA GCAUCUUGUUCC
#=SS
      >>>>>>> ++++ <<<<<<<<
liql
           GUUCACC AUCA GGGGAC
#=SS
           >>>+>> ++++ <<<<<<
lig2 AUGGAUGCGCACC AUCA GGGCGUAUCUAU
#=SS >>>>>>>+>> ++++ <<<<<<<<
```

4.5 Count vector files

hmmbuild saves a "count vector file" when the --cfile option is invoked. The count vector file contains observed weighted counts for match emissions, insert emissions, and state transitions. The intended use of the count vector file is for training mixture Dirichlet priors (something we do locally), but since it may be useful for other purposes, the format is documented here.

Each line of the file is an annotated count vector. The format of this line is:

- **Vector type** The first field is a single letter M, I, or T, specifying whether the line is for a match emission, insert emission, or transition vector.
 - Counts The next several fields are the counts themselves.

For match and insert emission vectors, there are either 4 or 20 real numbers, depending on whether the alignment was DNA/RNA or protein, respectively; and the counts are in alphabetical order by residue ("ACGT" for DNA/RNA, "AC..WY" for protein).

For state transition vectors, there are 7 real numbers, in order MM, MI, MD, IM, II, DM, DD.

The counts are real numbers, not integers, because they're weighted. Both relative weights (default: tree weights) and absolute weights (effective sequence number) affect the observed counts. (They are exactly the numbers used by HMMER before addition of Dirichlet prior pseudocounts and renormalization.) Options affecting either relative or absolute weighting will therefore affect the count vectors. To see unweighted raw count vectors, add --wnone --idlevel 1.0 to the hmmbuild command line.

- Alignment name If the alignment was in Stockholm format (for example, a Pfam distribution), the name of the alignment is recorded in this field. (A count vector file may contain vectors from a large number of alignments.) If this information is unavailable, "-" appears instead.
- Alignment column An integer indicating which alignment column this count vector corresponds to. In combination with the alignment name, this allows you to retrieve other annotation from the vicinity of this count vector in the original alignment file if you need to.
 - **HMM node** An integer indicating which HMM node this vector corresponds to. This number is always \leq the alignment column index, because some alignment columns are skipped in constructing an HMM.

Note that count vectors are only recorded for the alignment columns that became HMM match states. If you want to collect a count vector from *every* alignment column, you need to make hmmbuild assign every column to a match state. You can do this by invoking the --fast --gapmax 1.0 options to hmmbuild.

Structure annotation The next 2 fields are single characters representing structural annotation for this column in the alignment, if available. If not, "-" characters appear.

The first field is Stockholm format's #=GC SS_cons secondary structure consensus annotation for the column. This is a single letter DSSP code.

The second field is Stockholm format's $\#=GC \ SA_cons$ surface accessibility annotation for the column. This is a single character 0-9 (representing <10% to <100% accessibility), or X for unknown, or - or . for columns that are gaps in all the known structures in the alignment.

More structure annotation Transition vectors (only) have another two fields of structure annotation for the *next* (k+1'th) HMM node. Transition vectors are thought of as being between the k'th and the k+1'th HMM node, so most applications would probably want to see the annotation for both. Note that this does not necessarily correspond to the annotation on the next alignment column, because that column is not necessarily assigned to an HMM node.

Chapter 5

Installation

5.1 System requirements and portability

HMMER is designed to run on UNIX platforms. The code is POSIX-compliant ANSI C. You need a UNIX machine and an ANSI C compiler to build and use it.

Some optional tests and utilities use Perl. make check will only work if you have Perl installed. However, Perl isn't necessary to make HMMER work.

HMMER includes support for two kinds of parallelization: POSIX threads and PVM (Parallel Virtual Machine). Most modern UNIX OS's support POSIX threads. HMMER's configure script attempts to automatically detect your threads library, if you have one, and will build in multithreading by default – so you don't need to do anything special to get multithreading. PVM, in contrast, is a separate, third-party software application, and you will have to install and configure it before building HMMER with PVM support. For this reason, PVM support in HMMER is not enabled by default. See the PVM section of this chapter for more details.

HMMER 2 should be easy to port to non-UNIX operating systems, provided they support ANSI C and some reasonable level of POSIX compliance. A WinNT distribution is available by anonymous FTP from Time Logic, Inc. HMMER 1 was ported by other people to Digital VAX/VMS, Apple MacOS, Win95, and WinNT with relatively little difficulty, and I've made efforts to improve the portability of the HMMER 2 code since then. I would greatly appreciate receiving diffs for any ports of HMMER to any platform.

5.2 Installing a precompiled distribution

Thanks to generous hardware support from Compaq, Hewlett-Packard, IBM, Intel, Silicon Graphics, and Sun Microsystems, precompiled binary distributions of HMMER are available for at least the following platforms:

- Intel PC/Linux
- Intel PC/FreeBSD
- Intel PC/OpenBSD
- Intel PC/Sun Solaris
- Silicon Graphics/IRIX
- Sun Sparc/Solaris

- Compaq Alpha/Tru64
- Hewlett Packard P/HP-UX
- IBM/AIX

To install a precompiled distribution, do the following:

- 1. Download the distribution from http://hmmer.wustl.edu/.
- 2. Uncompress and un-tar it; for example,
 - > uncompress hmmer.bin.intel-linux.tar.gz
 - > tar xf hmmer.bin.intel-linux.tar

```
A new directory hmmer-xx is created, where "xx" is the current HMMER version number.
```

- 3. Go to the top level directory and run the configure script.
 - > cd hmmer-xx
 - > ./configure
- 4. Edit the Makefile in the top level directory, if you want.

The executables are in the binaries directory. The man pages are in the documentation/man directory. The PDF copy of the Userguide is in the top level HMMER directory (Userguide.pdf).

To run HMMER in the distribution directory, you don't need to edit the Makefile at all. But to permanently install HMMER on your system, set the make variables BINDIR and MANDIR to be the directories where you want HMMER executables and man pages to be installed. If you are installing in directories /usr/local/bin/ and /usr/local/man/man1/, you don't need to change anything.

- 5. Type make install to install the programs and man pages. You may have to become root, depending on where you're installing.
- 6. Type make clean to clean up.

5.3 Compiling from a source-only distribution

- 1. Download the distribution from http://hmmer.wustl.edu/.
- 2. Uncompress and un-tar it:

```
> uncompress hmmer.tar.gz
```

```
> tar xf hmmer.tar
```

- A new directory hmmer-xx is created, where "xx" is the current HMMER version number.
- 3. Go to the top level directory and run the configure script.

```
> cd hmmer-xx
```

> ./configure

If you want to include the optional PVM support, do:

> ./configure --with-pvm

If you include PVM, the system that you're compiling on must already be set up for PVM; specifically,

the environment variables PVM_ROOT and PVM_ARCH must be set, so HMMER can find the PVM headers and library.

If you want to disable POSIX threads support, do:

> ./configure --disable-threads

Sometimes this is necessary, if your system has a screwy Pthreads installation. HMMER can run fine without threads, it just won't utilize multiple processors.

If you want to specify a different default installation directory, such as /usr/share, do:

> ./configure -prefix=/usr/share

In a multiplatform environment, you might have your binaries in one platform-specific hierarchy (say, /usr/share/Linux/bin and your man pages and other platform independent stuff somewhere else (say /usr/share/man. You can install in separate binary and data directories:

> ./configure -prefix=/usr/share -exec_prefix=/usr/share/Linux

If you want to change the choice of compilation flags (CFLAGS) or the compiler (CC), set these as environment variables:

> env CC=gcc CFLAGS="-06" ./configure

For other generic configuration options, see the documentation for GNU autoconf 2.12. – but it may be easier to just edit the Makefile (see below).

4. Edit the top of the Makefile, if needed.

To build and test HMMER in its source directory, you don't need to edit the Makefile at all. If the configure script did a good job with the installation directories, you also don't need to edit the Makefile, but you might check to be sure.

To permanently install HMMER on your system, be sure that the make variables BINDIR and MANDIR are set to be the directories where you want HMMER executables and man pages to be installed. If you are installing the programs in /usr/local/bin and the man pages in /usr/local/man/man1, you don't need to change anything.

The default Makefile gets configured into a reliable but not necessarily optimal choice of compiler and compilation flags. The package is known to build "out of the box" on SGI IRIX, Sun Solaris, Intel/Linux, Intel/FreeBSD, Intel/OpenBSD, HP/HP-UX, IBM/AIX, or Compaq Tru64 Unix platforms without any special modifications. However, you may want to change the CC or CFLAGS variables to suit your system. In particular, you can play with the compiler options in CFLAGS to try to get more speed, if you're compiler-fluent. The Makefile may include some hints for various platforms.

On SunOS 4.1.x systems (God help you), you will have to use the GNU gcc compiler, because ancient SunOS cc is not ANSI-compliant.

- 5. Type make to build the programs.
- 6. (Optional) Type make check to compile and run a test suite. Some of the tests require that you have Perl installed. None of these tests should fail. Ever.
- 7. Type make install to install the programs and man pages. You may have to become root, depending on where you're installing.
- 8. Type make clean to clean up.

5.4 Environment variable configuration

These steps are optional, and will only apply (I think) to sufficiently POSIX-compliant operating systems like UNIX, Linux, or WinNT.

HMMER reads four environment variables, and is designed to coexist peacefully with an installation of WUBLAST or NCBI BLAST:

BLASTDB - directory location of FASTA-formatted sequence databases

BLASTMAT - directory location of BLAST substitution matrices

HMMERDB - directory location of HMM databases (e.g. PFAM)

HMMER_NCPU - maximum number of CPUs to utilize in a multiprocessor

If you have installed BLAST, you probably already have the two BLAST environment variables set in system-wide or user-specific .cshrc files.

All four variables are optional. If they are set up, you can simplify command lines to:

```
> hmmpfam pfam my.query
```

```
> hmmsearch my.hmm swiss35
```

instead of

> hmmpfam /some/long/path/to/databases/pfam my.query

> hmmsearch my.hmm /some/long/path/to/databases/swiss35

5.5 Parallelization using threads

HMMER includes support for two kinds of parallelization: POSIX threads and PVM (Parallel Virtual Machine). Threads support is built in by default; PVM is not.

If you have a multi-processor UNIX machine, odds are that you have a POSIX threads library. This kind of parallelization is easy from the standpoint of a HMMER user. HMMER will happily use all your processors (fewer, if you wish). The binary distributions of HMMER all come with multithreading capability built in.

To *disable* threads support, add --disable-threads to the command line for ./configure before you compile a source distribution. Multithreading is completely optional, and the software will work fine without it.

Like multithreaded BLAST, a multithreaded HMMER search will use all the available CPUs. Sometimes this is not desired; you may want HMMER searches to leave some spare processing power for other work. The environment variable HMMER_NCPU sets the maximum number of CPUs that any HMMER job will use. You can also set a limit on an individual process using the -cpu < n > option.

5.6 Parallelization using PVM

Parallelization across multiple machines (as opposed to multithreading on a single multiprocessor machine) can be done with PVM, the Parallel Virtual Machine software from Oak Ridge National Labs.

PVM is freely available. You can obtain it from http://www.epm.ornl.gov/. You must install and configure PVM before compiling PVM support into HMMER. During compilation, HMMER needs to

see the environment variables PVM_ROOT and PVM_ARCH, and the PVM header files and libraries must be found in the appropriate place under \$PVM_ROOT.

To enable PVM support in HMMER, add --with-pvm to the command line for ./configure before you compile a source distribution. PVM is completely optional, and the software will work fine without it.

Whereas multithreading requires no special configuration once support is compiled into HMMER, configuring a PVM cluster and using HMMER on it is a little more involved.

Configuring a PVM cluster for HMMER

Here, I will assume you're already familiar with PVM.

Designate one machine as the "master", and the other machines as "slaves". You will start your HMMER process on the master, and the master will spawn jobs on the slaves using PVM.

Install PVM on the master and all the slaves. On the master, make sure the environment variables PVM_ROOT and PVM_ARCH are set properly (ideally, in a system-wide .cshrc file).

Add the master's name to your .rhosts or /etc/hosts.equiv file on the slaves, so the slaves accept rsh connections from the master.

Put copies of HMMER executables in a directory on the master and all the slaves. For each PVMcapable program (hmmcalibrate, hmmpfam, and hmmsearch, there is a corresponding slave PVM program (hmmcalibrate-pvm, hmmpfam-pvm, and hmmsearch-pvm). The master machine needs copies of all the HMMER programs, including the slave PVM programs. The slaves only need copies of the three slave PVM programs. (You never need to start the slave programs yourself; PVM does that. You just need to make sure they're installed where PVM can see them.)

The PVM implementation of hmmpfam needs a copy of any HMM databases you may search to be installed on the master and every slave. All HMM databases must be indexed with hmmindex. The reason is that hmmpfam is I/O bound; the PVM implementation can't distribute an HMM database fast enough over a typical cluster's Ethernet. Instead, each PVM node accesses its own copy of the HMM database, distributing the I/O load across the nodes. hmmcalibrate and hmmsearch, in contrast, are freestanding. Only the master node needs to be able to access any HMM and/or sequence files.

Write a PVM hostfile for the cluster. Specify the location of the HMMER executables using the ep= directive. Specify the location of pvmd on the slaves using the dx= directive (alternatively, you can make sure PVM_ROOT and PVM_ARCH get set properly on the slaves). For the slaves, use the wd= directive to specify the location of the HMM databases for hmmpfam (alternatively, you can make sure HMMERDB gets set properly on the slaves). Use the sp= directive to tell HMMER how many processors each node has (and hence, how many independent PVM processes it should start); sp=1000 means 1 CPU, sp=2000 means 2 CPUs, sp=4000 means 4 CPUS, etc.

Start the PVM by typing

> pvm hostfile

(where "hostfile" is the name of your hostfile) on the master. Make sure all the nodes started properly by typing > conf

at the PVM console prompt. Type > quit

to exit from the PVM console, which leaves the PVM running in the background. You should only need to start PVM once. (We have a PVM running continuously on our network right now, waiting for HMMER jobs.)

Once PVM is running, at any time you can run HMMER programs on the master and exploit your PVM, just by adding the option – -pvm; for instance,

> hmmpfam --pvm Pfam my.query

parallelizes a search of a query sequence in the file my.query against the Pfam database.

Once PVM is properly configured and your slave nodes have the required slave programs (and databases, in the case of hmmpfam), the only difference you will notice between the serial and the PVM version is a (potentially massive) increase in search speed. Aside from the addition of the --pvm option on the command line, all other options and input/output formats remain identical.

Example of a PVM cluster

The St. Louis Pfam server runs its searches using HMMER on a PVM cluster called Wulfpack. I'll use it as a specific example of configuring a PVM cluster. It's a little more intricate than you'd usually need for personal use, just because of the details of running PVM jobs in a standalone way from CGI scripts on a Web server.

The master node is the Web server, fisher. The slave nodes are eight rack-mounted dual processor Intel/Linux boxes called wulf01 through wulf08. Collectively, we refer to this cluster as Wulfpack; it is a Beowulf-class Linux computing cluster.

PVM 3.3.11 is installed in /usr/local/pvm3 on the master and the slaves.

On fisher, all HMMER executables are installed in /usr/local/bin. On the wulf slave nodes, the three PVM slave executables are installed in /usr/local/wulfpack.

Pfam and PfamFrag, two Pfam databases, are installed on the wulf slave nodes in /usr/local/wulfpack. They are converted to binary format using hmmconvert -b, then indexed using hmmindex. (Using binary format databases is a big performance win for hmmpfam searches, because hmmpfam is I/O bound and binary HMM databases are smaller.)

An ls of /usr/local/wulfpack on any wulf node looks like:

[eddy@wulf01 /home]\$ ls /usr/local/wulfpack/			
Pfam	PfamFrag	hmmcalibrate-pvm	hmmsearch-pvm
Pfam.gsi	PfamFrag.gsi	hmmpfam-pvm	

The PVM hostfile for the cluster looks like:

```
# Config file for Pfam Web server PVM
#
* ep=/usr/local/bin sp=1000
fisher.wustl.edu
* lo=pfam dx=/usr/local/pvm3/lib/pvmd ep=/usr/local/wulfpack sp=2000
wulf01
wulf02
wulf03
wulf04
wulf05
wulf06
wulf07
wulf08
```

Note one wrinkle specific to configuring Web servers: the web server is running HMMER as user "nobody" because it's calling HMMER from a CGI script. We can't configure a shell for "nobody" on the slaves, so we create a dummy user called "pfam" on each wulf node. The lo= directive in the PVM hostfile tells the master to connect to the slaves as user "pfam". On each slave, there is a user "pfam" with a .rhosts that looks like:

fisher nobody fisher.wustl.edu nobody

which tells the wulf node to accept rsh connections from fisher's user "nobody".

Also note how we use the sp= directive to tell HMMER (via PVM) that the wulf nodes are dual processors. fisher is actually a dual processor too, but by setting sp=1000, HMMER will only start one PVM process on it (leaving the other CPU free to do all the things that keep Web servers happy).

The trickiest thing is making sure PVM_ROOT and PVM_ARCH get set properly. For my own private PVM use, my .cshrc contains the lines:

```
setenv PVM_ROOT /usr/local/pvm3
setenv PVM_ARCH `$PVM_ROOT/lib/pvmgetarch`
```

But for the web server PVM, it's a little trickier. We start the Web server PVM as user "nobody" on fisher using a local init script, /etc/rc.d/init.d/pvm_init. With its error checking deleted for clarity, this script basically looks like:

```
#!/bin/sh
wulfpack_conf=/home/www/pfam/pfam-3.1/wulfpack.conf
. /usr/local/pvm3/.pvmprofile
$PVM_ROOT/lib/pvmd $wulfpack_conf >/dev/null &
```

We call this at boot time by adding the line su nobody -c "sh /etc/rc.d/init.d/pvm_init to our rc.local file. .pvmprofile is a little PVM-supplied script that properly sets PVM_ROOT and PVM_ARCH, and wulfpack.conf is our PVM hostfile.

The relevant lines of the CGI Perl script that runs HMMER jobs from the Web server (again, heavily edited for clarity) are:

```
# Configure environment for PVM
$ENV{'HMMERDB'} = "/usr/local/wulfpack:/home/www/pfam/data/"
$ENV{'PVM_EXPORT'} = "HMMERDB";
$output = `/usr/local/bin/hmmpfam --pvm Pfam /tmp/query`;
```

The trick here is that we export the HMMERDB environment variable via PVM, so the PVM processes on wulf nodes will know where to find their copy of Pfam.

PVM is relatively complex, but with luck, this and the PVM documentation give you enough information to get HMMER running on a cluster. It's well worth it. Wulfpack was simple to assemble; besides the eight rack-mounted machines, there's a terminal switch, a single console, a 10baseT Ethernet switch, and a UPS. Each machine runs stock Red Hat Linux (we don't need no steenking Extreme Linux hype). The whole thing cost us \$20K, but it runs HMMER searches as fast as a 16-processor SGI Origin – and it's bigger and has more blinking lights than an Origin, so it's more impressive to look at.

5.7 **Recommended systems**

HMMER is currently developed and maintained on Intel/GNU/Linux systems. Since these are the development systems, similar systems are the least likely to show portability problems.

At each release, HMMER is also tested on a compile farm that includes most major commercial and open source UNIX platforms (see start of the chapter for a list). These systems are also unlikely to show portability problems.

5.8 Make targets

There are a variety of make targets in the toplevel Makefile. For completeness, they are summarized here. *You shouldn't need to know this stuff.* Targets marked (Private) are targets which are not supported in the public HMMER release. They only run in my local development environment.

- all Compiles the source code; puts the compiled programs in the binaries/ subdirectory.
- check Compiles and runs the testsuite.
- install Installs the programs in BINDIR and the man pages in MANDIR.
- clean Cleans up the directory, leaving the distribution files and binaries.
- **distclean** Like 'make clean', but removes everything that isn't part of a pristine source distribution.
 - verify Runs consistency checks on the package. (Private)
 - doc Builds the Userguide from source. (Private)
 - dist Checks out a complete distribution from RCS and assembles the .tar.Z file. (Private)
 - ftpdist Installs a new release on the FTP site. (Private)
 - stable Symlinks a new FTP release to hmmer.tar.Z. (Private)

Chapter 6

License terms

If you're just going to use HMMER to analyze sequence, it's free to you, and you can stop reading the legalese. If you have a specially licensed (non-GPL) copy of HMMER from Washington University, you have your own legalese, in the form of a separate written WashU licensing contract; this chapter does not apply to you. But if you have a public copy of HMMER and you're interested in modifying or borrowing the source code or distributing the software yourself, keep reading.

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 - if you're a proprietary software developer, contact Washington University to arrange for non-GPL licensing terms. For commercial licenses, we will typically ask for a fee and/or royalties.

6.1 "Manifesto"

Because HMMER has commercial value, I spend an inordinate amount of time explaining the choice of the GPL. Open source licenses are widely misunderstood.

Much of my funding comes from the National Human Genome Research Insitute at the National Institutes of Health (NIH). NIH grants policy states that the results of NIH-funded research must be made available to qualified scientists upon request. The free exchange of information is also a fundamental tenet of publicly funded science. The simplest means of disseminating information about how HMMER works is to distribute documented source code. I take this responsibility seriously. While I am in academia, my software will be freely available in source code form.

At the same time, I'm not going to put HMMER in the public domain. If I did, someone could incorporate HMMER into a proprietary software package without my permission or knowledge. I couldn't see the improvements they were making to my work. They might sell the proprietary software and not contribute a fair share of the profits to Washington University or my research lab. So I need a distribution license that lets me distribute it freely, while at the same time trying to make sure that my university and I don't get taken advantage of.

Some developers license their software under restricted "academic only" licenses, such that "commercial" users have to pay license fees. Having worked in a biotech startup, I strongly oppose this sort of license. Most industry scientists contribute to basic research, and they consider themselves no different from their counterparts in academia. The line I want to draw is between open and proprietary, not academic and commercial. Commercial scientists can be open, and academic scientists can be proprietary. (And NIH's free resource distribution policy does not distinguish between academic and industrial peers.)

Open Source licenses (aka "free" software licenses) like the BSD license, the Perl Artistic License, the Netscape Public License, and the GNU Public License do what I need. The GNU Public License is the best known of these licenses. So long as you play the game – that you are working in a nonproprietary manner, accepting that any of your modifications to my code must also be GPL'ed and made freely available as open source code – you get free, open access to HMMER source code. But if you want to use HMMER in a proprietary way, the GPL does not grant you that right, so you must contact the Washington University tech transfer people to arrange a special proprietary license. And, unsurprisingly, we'll happily charge you fair license fees. You can feel warm and fuzzy about this - I opt to take all licensing funds into a research fund for the lab, not into personal income. I still quite contently drive a Civic; I'm too busy coding to own a Boxster.

The GPL is often misinterpreted as a socialist license, and I sometimes field questions about why I'm so determined to oppose commercial development, or even undermine it (by distributing free software). This is a misunderstanding. American universities have a responsibility, under the 1980 Bayh/Dole act, to transfer technology to the private sector. I respect the economic goals of Bayh/Dole. In my view, there is a productive tension between Bayh/Dole and scientific ethics. The purpose of the HMMER licensing policy is not to impede commercial development, but rather to maintain this tension. As an academic, my primary responsibility is to make my work freely available. As an American research university, the job of Washington University and our technology transfer office is to make HMMER for proprietary use. We're just not going to *give it away* for proprietary use, that's all. If you want to make a profit from HMMER, we want a fair share so we can put it towards the research that develops HMMER.

Whew. You think that was long-winded? That's nothing; the full text of the GNU GPL follows:

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Version 2, June 1991

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END OF TERMS AND CONDITIONS

Chapter 7

Acknowledgements and history

It must be remembered that there is nothing more difficult to plan, more doubtful of success, nor more dangerous to manage, than the creation of a new system. For the initiator has the emnity of all who would profit by the preservation of the old institutions and merely lukewarm defenders in those who would gain by the new ones.

- Niccolo Machiavelli

HMMER 1 was developed at the MRC Laboratory of Molecular Biology, Cambridge UK, while I was a postdoc with Dr. Richard Durbin. I thank the Human Frontier Science Program and the National Institutes of Health for their support.

HMMER 1.8 (and subsequent minor releases) was the first public release of HMMER in April 1995. A number of modifications and improvements went into HMMER 1.9 code, but 1.9 was never released. Some versions of HMMER 1.9 did inadvertently escape St. Louis and make it to other sites, but it was never documented or supported. HMMER 1.9 collapsed under its own weight in 1996, when the number of ugly hacks increased to a critical mass.

HMMER 2 is a nearly complete rewrite, based on a new model architecture dubbed "Plan 7". Implementation was begun in November 1996 at Washington University in St. Louis. I thank the Washington University Dept. of Genetics, the NIH National Human Genome Research Institute, and Monsanto for their support during this (extremely stressful) time. There is nothing like throwing four years of work away and starting fresh to make you question your sanity. Working on a book (Durbin et al., 1998) and starting up a lab at the same time made it all doubly "exciting". If you are so bored as to actually read the code, you will run across inexplicable comments that note where I was when I wrote various parts, making up a sort of disjointed diary of this period; amongst other places, parts of HMMER 2 were written in airport lounges, on TWA flights 720 and 721 to and from London, in Graeme Mitchison's kitchen in Cambridge, and on vacations while my (ex-)wife wasn't watching (which probably explains the divorce, come to think of it). I therefore owe special thanks (I think) to the Biochemistry Academic Contacts Committee at Eli Lilly & Co. for a gift that paid for the trusty Linux laptop on which much of HMMER 2 was written.

HMMER 2.1.1 was the stable release for three years, from 1998-2001. After a long developmental gestation period, during which my research lab had mostly switched to RNA work, and I'd mostly switched to being a PI instead of a researcher, HMMER 2.2 was finally released in summer 2001. HMMER has now settled into a comfortable middle age, like its author - still actively maintained, but no dramatic new changes are planned.

The MRC-LMB computational molecular biology discussion group has contributed many ideas to HM-MER. In particular, I thank Richard Durbin, Graeme Mitchison, Erik Sonnhammer, Alex Bateman, Ewan Birney, Gos Micklem, Tim Hubbard, Roger Sewall, David MacKay, and Cyrus Chothia. Any errors in the code, though, are my fault alone, of course.

Sequence format parsing (sqio.c) in HMMER is derived from an early release of the READSEQ package by Don Gilbert, Indiana University. Thanks to Don for an excellent piece of software; and apologies for the mangling I've put it through since. The file hsregex.c is a derivative of Henry Spencer's regular expression library; thanks, Henry. Several miscellaneous functions in sre_math.c are taken from public domain sources and are credited in the code's comments. masks.c includes a modified copy of the XNU source code from David States and Jean-Michel Claverie.

In many other places, I've reimplemented algorithms described in the literature. These are too numerous to credit and thank here. The original references are given in the comments of the code. However, I've borrowed (stolen?) more than once from the following folks that I'd like to be sure to thank: Steve Altschul, Pierre Baldi, Phillip Bucher, Warren Gish, David Haussler, Steve and Jorja Henikoff, Richard Hughey, Kevin Karplus, Anders Krogh, Bill Pearson, and Kimmen Sjölander.

HMMER is developed on Silicon Graphics and Linux machines in my lab. I thank Compaq, IBM, Intel, Sun Microsystems, Silicon Graphics, Hewlett-Packard, and Paracel for generous hardware support. Dave Cortesi at SGI contributed much useful advice on the POSIX threads implementation. I am proud to acknowledge a tremendous debt to the development tools that I use from the free software community: an incomplete list includes GNU gcc, gdb, emacs, and autoconf; Cygnus' and others' egcs compiler; Conor Cahill's dbmalloc library; Bruce Perens' ElectricFence; the cast of thousands that develops CVS, the Concurrent Versioning System; Larry Wall's perl; LaTeX and TeX from Leslie Lamport and Don Knuth; Nikos Drakos' latex2html; Thomas Phelps' PolyglotMan; Linus Torvalds' Linux operating system; and the folks at Red Hat Linux and Mandrake Linux.

Finally, I'd like to cryptically thank Dave "Mr. Frog" Pare and Tom "Chainsaw" Ruschak for a totally unrelated free software product that was historically instrumental in HMMER's development, for reasons that are best not discussed while sober.

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