

learnMSA: learning and aligning large protein families

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Abstract

Background: The alignment of large numbers of protein sequences is a challenging task and its importance grows rapidly along with the size of biological datasets. State-of-the-art algorithms have a tendency to produce less accurate alignments with an increasing number of sequences. This is a fundamental problem since many downstream tasks rely on accurate alignments.

Results: We present learnMSA, a novel statistical learning approach of profile hidden Markov models (pHMMs) based on batch gradient descent. Fundamentally different from popular aligners, we fit a custom recurrent neural network architecture for (p)HMMs to potentially millions of sequences with respect to a maximum *a posteriori* objective and decode an alignment. We rely on automatic differentiation of the log-likelihood, and thus, our approach is different from existing HMM training algorithms like Baum–Welch. Our method does not involve progressive, regressive, or divide-and-conquer heuristics. We use uniform batch sampling to adapt to large datasets in linear time without the requirement of a tree. When tested on ultra-large protein families with up to 3.5 million sequences, learnMSA is both more accurate and faster than state-of-the-art tools. On the established benchmarks HomFam and BaliFam with smaller sequence sets, it matches state-of-the-art performance. All experiments were done on a standard workstation with a GPU.

Conclusions: Our results show that learnMSA does not share the counterintuitive drawback of many popular heuristic aligners, which can substantially lose accuracy when many additional homologs are input. LearnMSA is a future-proof framework for large alignments with many opportunities for further improvements.

Keywords: profile hidden Markov model, multiple sequence alignment, machine learning

Background

Profile hidden Markov models (pHMMs) are probabilistic models for protein families. One of their applications is remote homology search in large databases [1, 2]. Typically, an existing multiple sequence alignment (MSA) is turned into a pHMM, but pHMMs can also be trained on unaligned sequences and a MSA can be decoded from the learned model [3-5]. The training of pHMMs using the Baum-Welch algorithm was originally applied "with handholding" to selected protein families [3], which required a human to decide between specific architectures (e.g., for modeling a domain as opposed to an entire protein). Advantages of the statistical learning approach over traditional aligners are a consistent probabilistic background for position-specific gap penalties and that both training and decoding are linear in the number of sequences. However, profile HMM training has never been popular as a general-purpose alignment method since tabula rasa learning is challenging. Apart from the model architecture being problem dependent, another common issue is that algorithms may get stuck at local optima in the parameter space. Simulated annealing [4] and particle swarm optimization [6, 7] could further improve upon Baum-Welch in this regard but never resulted in applicable tools comparable to modern state-of-the-art aligners. Gradient descent methods like the popular Adam algorithm [8] are a hitherto entirely unexplored class of algorithms for HMM training with increasing relevance in the advent of automatic differentiation [9].

Established tools that construct MSAs are either unfit for large numbers of sequences or their accuracy decreases when the number of aligned sequences grows large [10, 11]. This effect is particularly present for progressive algorithms, which rely on a guide tree

that dictates the order of the sequences to be aligned, by greedily starting with closely related ones. One drawback of this approach is the inability to revert gaps. Early errors accumulate when more and more sequences are added.

One way to revert incorrect gaps is iterative refinement, where intermediate alignments guide the construction of subsequent ones [12]. Although iterative refinement strategies can improve accuracy on moderate sequence numbers, they are unsuitable for large numbers of sequences from a computational perspective. For example, MAFFT G-INS-i produces very accurate alignments but is slow and memory-hungry due to an all-to-all pairwise alignment stage. MAFFT-Sparsecore applies MAFFT G-INS-i to a small set of core sequences and progressively adds the remaining sequences thereafter [13]. This strategy is suitable to scale up iterative refinement to large sequence numbers, but biases in the core sequences have to be avoided by choosing them as diverse as possible.

Divide-and-conquer strategies like PASTA [14] and MAGUS [15] first construct subalignments on relatively small subsets of the sequences and merge them thereafter. MAGUS uses a Graph Clustering Merger for the latter stage. Recently, MAGUS was updated to support recursion for ultra-large datasets [16]. Another technique with improved accuracy is the regressive method, which starts to align sequences containing the most dissimilar ones first and merges subalignments by using an overlapping sequence [10]. Divide-and-conquer strategies have enabled the execution of slow but accurate algorithms like MAFFT G-INS-i on large datasets and improved accuracy compared to progressive strategies [10, 15].

However, they are still heuristics that ignore everything but a subset at first and are prone to errors in their merging steps.

Last, UPP [17] is related to our method by the fact that it also uses a pHMM (or an ensemble of pHMMs) to represent MSAs. However, UPP does not train a model on unaligned sequences. Instead, it first constructs a backbone MSA on a subset of the sequences using tree-guided PASTA in order to estimate the HMM parameters. Afterward, it adds the remaining sequences using the HMM. UPP has shown good performance in the presence of high sequence length heterogeneity.

All mentioned MSA algorithms rely on accurate guide trees, and tree construction often becomes the computational bottleneck. Clustal Omega [11] uses the mbed method to construct a tree. A faster but less accurate alternative is MAFFT-PartTree [18], and another popular algorithm is FastTree [19]. A slow but very accurate tree construction algorithm based on all-to-all pairwise alignments is used in the G-INS-i option of MAFFT [12]. The bottom line is the constant need to balance quality and speed when constructing trees.

To date, deep learning is not commonly used for multiple sequence alignment, and if it is, its function is usually supplementary, for example, by optimizing the order of progressive alignment with reinforcement learning [20] or employing a decision-making model to select from different strategies in a MSA pipeline [21]. While some proof of concepts exist, the respective software is not feasible for large numbers of sequences, generally not optimized (stated by the authors), or not available at all. For pairwise alignment, the traditional dynamic programming framework can be supplemented by reinforcement learning [22] or deep models inspired by recent advances in natural language processing improving accuracy on remote homologs [23]. While deep learning is currently usually not used for their construction, MSAs are, however, a popular input for end-to-end machine learning methods that solve downstream tasks [24-27].

Our proposed aligner learnMSA is based on automated statistical learning of a pHMM with gradient descent. It does not require a tree and has a linear asymptotic runtime in the number of sequences, which is faster than most tree algorithms. No progressive, regressive, or divide-and-conquer heuristic is used. Therefore, we avoid heuristic-based errors when merging subalignments or progressively adding sequences. We provide a more robust framework for (ultra-)large MSAs without the counterintuitive drawback of losing accuracy when many additional homologs are input.

We begin with the description of the underlying model and a batchwise variant of the forward algorithm that plays a central role during parameter training. We empirically show the suitability of learnMSA by testing it on ultra-large protein families from Pfam [28] with up to 3.5 million sequences as well as the established biological benchmarks HomFam and BaliFam.

Methods

Model

Profile hidden Markov models are well-known probabilistic models of sequence consensus. When used to model a protein family, the aim is to define a probability distribution over the space of all possible protein sequences such that member sequences of the family have large probabilities. The resulting statistical model can be used for database searches [1] and MSA construction [3].

In a pHMM, a linear chain of match states represents the consensus sequence of the family in question. Insertions and deletions with respect to the consensus are modeled by positionspecific states and transitions. See Fig. 1A for an illustration of the pHMM.

In addition to the standard pHMM architecture, we deploy an augmented model following HMMER's "Plan7" [29, 30] (orange states and transitions in Fig. 1A). The HMM parameters are learned from unaligned protein sequences. In contrast to previous approaches, our method also learns the additional "Plan7" parameters jointly with the pHMM core model. Previously, HMMER used predefined value sets for different alignment modes (local or global, unihit or multihit) [30]. Here, we automatically learn the correct alignment mode jointly with the core pHMM starting tabula rasa. We have special states for the left (L) and right (R) flank of the model. Initialization and regularization of the flanking states differ from ordinary insertion states Ii (see section "Training"). Moreover, the augmented model allows multihit alignments (i.e., sequences may contain repeats of a single domain motif by looping backward). The state C models any unannotated region between 2 domain hits and must be visited to jump from the end state E back to the start state S. The model further handles sequence length heterogeneity (fragmentary sequences) through entry and exit probabilities from S into the consensus and, respectively, from the consensus to E. Note that since version 2, HMMER uses a trick to achieve a uniform distribution over all possible pairs of entry and exit points into the core model [30]. Here, we follow the older construction with explicit entry and exit probabilities, but they are now data dependent instead of ad hoc.

The set of all transition and emission parameters is learned from data with careful initialization and under the use of Dirichlet priors (see section "Training"). In general, we have 1 trainable parameter for each possible state transition and, in case of the emissions, 1 parameter per match state and amino acid. There are exceptions: insertion and flanking states use a fixed background emission distribution that is not optimized. The self-loop (and respectively exit) probabilities for the flanking states L, R, and C are tied to prevent a bias toward one of the sides. Delete states (as well as the domain start and end states S and E) are silent (i.e., they have no emission distribution and do not not advance the position in the observed sequence).

A pHMM can be parameterized by 2 probability matrices for transitions and emissions and an initial state distribution. Let Q be the set of all states and A be the stochastic $|Q| \times |Q|$ matrix of state transitions. Observe that for pHMMs, this matrix is very sparse. We call the number of match states in a model its length l. Let Q': = $Q \setminus \{D_1, ..., D_l, S, E\}$ denote the set of all emitting states. Let B be the $|Q'| \times 25$ emission matrix, which is constructed by concatenating I learnable emission distributions of the match states with background distributions for all insertions and the flanks. The second dimension of B corresponds to the 20 standard amino acids, plus selenocysteine, pyrrolysine, and the ambiguous codes X, B, and Z. The terminal symbol (26th letter) has an implicit probability of 0 at all states except T.

In order to apply gradient descent, we parameterize the model by unconstrained kernels θ^{A} and θ^{B} and enforce the probabilistic constraints that the rows of A and B sum up to 1 with a softmax function defined on a real vector: $softmax(x)_i = \frac{e^{x_i}}{\sum_i e^{x_i}}$.

As seen in Fig. 1A, the emission distribution of, for example, M_i is computed by $softmax(\theta_i^B)$, where $\theta_i^B = (\theta_A^{B_i}, \theta_R^{B_i}, \theta_N^{B_i}, \theta_D^{B_i}, \dots)$ is the ith row of θ^{B} . The matrix B is constructed from the kernel θ^{B}

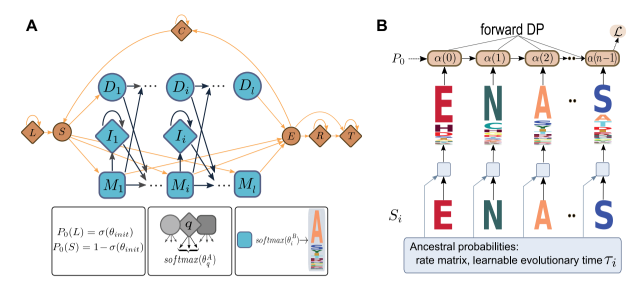


Figure 1: (A) LearnMSAs underlying pHMM based on HMMER's "Plan7" model. For the transition (emission) distributions, unconstrained learnable parameter matrices θ^{A} (θ^{B}) are transformed by softmaxes over the outgoing edges of a state or the amino acid alphabet, respectively. Squares indicate match states, diamonds are insertions, and circles are silent states (either delete states or the start and end state). In contrast to previous approaches, we also learn transition probabilities augmenting the core model (orange). (B) Sketch of a recurrent neural network architecture with a HMM-Cell that implements the forward recursion. The first layer at the bottom computes ancestral distributions of amino acids for a sequence S_i using a rate matrix and an evolutionary time τ_i that is learned jointly with the HMM parameters.

by using softmaxes to compute the match distributions over the amino acid alphabet.

The kernel θ^{A} is a collection of parameter vectors corresponding to different transition types that share the same initialization and prior. For example, we have l-1 parameters for the matchto-match transitions. The total number of allowed transitions in the model as shown in Fig. 1A is linear in l. The probability distribution of transitioning from, for example, match M_i to one of the 4 adjacent states M_{i+1} , I_i , D_{i+1} , or E is calculated by constructing the vector $\theta_{M_i}^A=(\theta_{M_i,M_{i+1}}^A,\theta_{M_i,I_i}^A,\theta_{M_i,D_{i+1}}^A,\theta_{M_i,E}^A)$ and computing softmax(θ_{M}^{A}). We store A (or, in fact, a matrix closely related to A as described in section "Implicit model") in sparse matrix representation where illegal transitions are implicitly zero.

For the initial state distribution P_0 , we use a simple parametrization by introducing a scalar θ_{init} that controls the probability of starting in the left flank. To this end, we define p_{init} $= \sigma(\theta_{init})$, where σ is the sigmoid function. The initial distribution is $P_0(L) = p_{init}$ and $P_0(S) = 1 - p_{init}$ and $P_0(q) = 0$ for $q \neq L$, S.

In the following, let $\theta = (\theta_{init}, \theta^A, \theta^B)$ denote the complete set of learnable parameters for the (augmented) pHMM.

Batchwise forward algorithm

Assume for now that no silent states exist. For the pHMM as introduced in section "Model," we will describe an equivalent implicit model without the silent states D₁, ..., D_l, S, and E in section "Im-

An unaligned protein sequence S can be described by a path π of hidden states in the pHMM. Under our assumption, π = π_0 , ..., π_{n-1} and S = s_0 , ..., s_{n-1} have the same length n. The joint probability of observed and hidden sequence is $P(S, \pi) = P_0(\pi_0)P(S_0 \mid \pi_0) \prod_{i>0} P(\pi_i \mid \pi_{i-1})P(S_i \mid \pi_i)$, where the transition and emission probabilities are computed as described in section "Model" above.

The likelihood of a sequence is the sum of the joint probabilities over all possible hidden paths: $P(S) = \sum_{\pi} P(S, \pi)$, which is related to HMMER's forward score [30]. Intuitively, it describes how well a sequence fits to the consensus when considering all possible alignments. The likelihood can be efficiently computed with dynamic

programming using either the forward or the backward algorithm [31]. We present a batchwise variant of the forward algorithm that plays a central role during parameter training of learnMSA.

The forward probabilities are $\alpha(i)_q$: = $P(\pi_i = q, s_0, ..., s_i)$. The well-known dynamic programming recursion to compute $\alpha(1), \ldots$ $\alpha(n-1)$ is

$$\alpha(i)_{q} = P(s_{i} \mid q) \sum_{q' \in Q} P(q \mid q') \alpha(i-1)_{q'}$$

$$\tag{1}$$

with $\alpha(0)_q = P(s_0|q)P_0(q)$.

Equation (1) lends itself to an efficient implementation for a batch of sequences of size b. Let the $b \times 25$ matrix $S^{(i)}$ denote the tuple of all ith sequence positions in the batch, that is, $S_i^{(i)}$ is an one-hot representation of the ith residue of sequence j. We omit the implementation detail that for variable-length sequences, some positions might be terminal symbols here. In the following, we factor out a partial likelihood term in each forward step to allow an underflow-safe computation of the likelihood. The batchwise forward recursion is

$$\alpha'(i) = \begin{cases} S^{(i)}B^T \circ \frac{\alpha'(i-1)}{Z(i-1)}A, & i > 0 \\ S^{(0)}B^T \circ P_0, & i = 0 \end{cases}$$

$$\mathcal{Z}(i) = \sum_{q \in Q} \alpha'(i)_q$$

$$\mathcal{L}(i) = \ln \mathcal{Z}(i). \tag{2}$$

where i is a sequence index, $\alpha'(i)$ are $b \times |Q|$ batches of scaled forward variables, o denotes element-wise multiplication (with shape broadcasting, where required), and the matrix multiplication that involves A uses an efficient implementation that exploits the sparse representation. Observe that $\alpha(i) = \alpha'(i) \circ \prod_{i'=0}^{i-1} \mathcal{Z}(i')$.

The likelihood (for a single sequence) can eventually be computed as $P(S) = \sum_{q} \alpha (n-1)_q$. However, we prevent numerical underflow by equivalently using the partial log-likelihood values in Equation (2):

$$ln P(S) = \sum_{i=0}^{n-1} \mathcal{L}(i).$$
(3)

Viterbi decoding

When we decode an alignment, we are interested in the hidden path of a sequence with maximum probability (i.e., $\operatorname{argmax}_{\pi} P(S, \pi)$). This can be computed efficiently using the Viterbi algorithm [31], which is closely related to the forward algorithm.

A Viterbi MSA can be constructed by aligning the most likely hidden sequences of all input sequences [3]. Currently, we leave insertions unaligned and left-adjusted except for the left flank, which is right-adjusted. Moreover, if domain repeats occur, the ith occurrences of the domain in multiple sequences respectively are currently aligned with each other. With both simplifications, we accept that we are in a slight disadvantage compared to state-of-the-art aligners, which will align all residues globally.

Implicit model

Conventionally, the forward recursion for pHMMs is implemented in linear time per step by explicitly handling silent states (the delete states D_i , the starting state S, and the ending state E) [32]. This requires a long-winded sequential computation of the forward variable for the delete states where $\alpha(i)_{D_j}$ depends on $\alpha(i)_{D_{j-1}}$. Here, we treat all silent states as implicit states, that is, internally we use an equivalent model that has only emitting states, by folding all transitions entering and leaving a silent state. That means all possible partial state paths that start and end in an emitting state and consist only of silent states else are replaced by single transitions that have probability equal to the probability of the respective partial path. In detail, each partial path $M_i \rightarrow D_{i+1} \rightarrow ...D_{j-1} \rightarrow M_j$ for j > i+1 is replaced by an edge with probability

$$P(M_j \mid M_i) = P(D_{i+1} \mid M_i) \left(\prod_{i'=i+1}^{j-2} P(D_{i'+1} \mid D_{i'}) \right) P(M_j \mid D_{j-1}).$$
 (4)

This changes the asymptotic runtime of the forward algorithm, because the number of possible transitions from each match state is not constant anymore. However, we can now implement Equation (2) by taking full advantage of modern (GPU-accelerated) computing frameworks. We found that given the typical length of a protein (our benchmarks contain sequences of length up to 800), the asymptotic downgrade is acceptable in the light of parallelism: we can compute all values of $\alpha(i)$ in parallel given $\alpha(i-1)$. In the batchwise forward algorithm, the bottleneck is the matrix multiplication with the transition matrix, which should use an efficient implementation that exploits sparseness.

Folding all edges adjacent to silent states is referred to as the implicit model, represented by a transition matrix A_{impl} replacing A from section "Model." Note that A_{impl} is still very sparse. Transitions over the start state S and the end state E (i.e., deletions of initial or terminal parts) are handled analogously. Also note that empty, infinite silent loops through the model are not possible, because the unannotated segment state C is an insertion that emits at least 1 amino acid and cannot be skipped.

Training

During training, learnMSA uses a recurrent neural network architecture with a pHMM cell that scans a batch of sequences column-of-residues-wise and successively applies Equation (2). This architecture is visualized in Fig. 1B with the addition of "Ancestral probabilities" as described later. Given θ , the parameters of the model,

the log-likelihood of a random batch of b sequences is

$$\mathcal{L}(\theta; S_1, \dots, S_b) = \sum_{i=1}^b \ln P(S_i \mid \theta).$$
 (5)

The general goal while successively observing random batches is to adjust θ such that $\mathcal L$ increases over time. In practice, we minimize a loss function related to $\mathcal L$ that also incorporates prior knowledge about proteins.

Existing optimization algorithms like Baum-Welch [3] or simulated annealing [4] avoid using gradients of \mathcal{L} and use the forwardbackward algorithm for parameter updates instead. An advantage of learnMSA is the possibility to optimize the HMM jointly with other layers. Currently, we demonstrate this as described in section "Ancestral probabilities," but a broader field opens up in this direction as discussed later. Gradient-based optimization can also be applied to objectives that are not based on likelihood, for instance, the discrimination or classification of (sub)families [33]. Traditional HMM learning algorithms are not used for online learning, although such variants exist [5]. Typically, they require more technical work to include priors than our gradient-based approach. None of the methods can guarantee globally optimal results. However, learnMSA can make use of the advancing optimization toolbox for machine learning problems based on automatic differentiation [9].

Maximum a posteriori loss

Models found by maximizing \mathcal{L} might generalize weakly. This is especially true if the number of training sequences m is low. Our experiments will mainly focus on cases where m is large (i.e., 10,000 to millions of sequences). However, we can still have overfitting problems. Domain motifs of subfamilies might be underrepresented in the sequence set, leading to a skewed model. Moreover, we might end up with a result that fits the data well but is not biologically plausible (e.g., a model that allows very long insertions or many gap openings). A maximum a posteriori estimate attempts to fit the data while at the same time penalizing implausible models [3]. In this sense, we define our loss function as

$$\ell(\theta; S_1, \dots, S_b) = -\frac{1}{h} \mathcal{L}(\theta; S_1, \dots, S_b) - \frac{1}{m} \ln(\rho(\theta)). \tag{6}$$

The loss ℓ has a foundation in Bayesian statistics. The first term is the log-likelihood per sequence averaged over a batch of sequences. Usually, we choose b < m and consequently perform stochastic gradient descent. This allows us to rapidly train models even on millions of homologous sequences. We use random uniform batch sampling. The second term is the prior density (i.e., ρ is a function that rewards plausible models). We normalize by $\frac{1}{m}$ to make the estimate consistent. The effect of the prior is reduced proportional to the number of training sequences. This is particularly important because we use a general (i.e., family-agnostic) prior that should work over the full range of dataset sizes. Following conventional standards [3], we use Dirichlet densities [34, 35] over the different types of transition distributions and the match emissions.

To reduce the total number of hyperparameters that have to be set by hand, we salvaged as much general-purpose information as possible from Pfam HMMs. For the core model probabilities, we took over 3 million example transition distributions and maximized the likelihoods of 3 Dirichlets: one for matches, insertions, and deletions, respectively (see Table 1).

For the emissions, we tested Dirichlet mixtures with different component counts (1, 9, 32, 64, 128, 512), which we trained on the

Table 1. Dirichlet parameters for the core pHMM transition distributions estimated from Pfam HMMs

α	Match	Insert	Delete
Match	40.59	0.96	0.68
Insert	26.75	23.32	_
Delete	37.79	_	25.15

match emission distributions of Pfam HMMs, but found that for large sequence counts, a single Dirichlet density (i.e., a mixture with 1 component) is enough. The expectation of this Dirichlet distribution is also used to initialize the match emissions as well as the (fixed) insertion emissions and the flanks.

As described earlier, we optimize the transition probabilities for flanking states, domain multihits, and the entry and exit probabilities jointly with the core model. We found that these transitions require strict regularization. We defined a simplified set of hyperparameters α_{flank} , α_{single} , and α_{global} and (currently only roughly) searched for suitable values based on the quality of the produced alignments. These hyperparameters have a probabilistic foundation as parameters of Dirichlet priors over specific Bernoulli distributions that were defined to favor the probability p = 1 for particular, carefully defined events. That means the prior can be maximized by maximizing p, but this choice has to be balanced with the likelihood. For each possible choice of p and α , the logarithmic prior densities are $(\alpha - 1) \ln p + (\alpha' - 1) \ln (1 - p)$, where we set α' = 1. The motivation behind this ad hoc choice was to keep the set of hyperparameters for the method simple while maintaining the probabilistic interpretation of the regularization term.

In particular, α_{flank} controls the pressure to align to the core model (rather than using the flanking states), that is, increasing α_{flank} will result in longer insertions at the flanks and between repeated domain segments. The parameter α_{flank} regularizes the self-loop probabilities of all flanking states, as well as $P_0(L)$ and P(R|E). Furthermore, we introduce α_{single} to penalize core model repeats favoring large values for the probability 1 - P(C|E) = P(R|E)+ P(T|E). Last, α_{global} penalizes local alignments that use entry and exit transitions other than $S \to M_1$ and $M_l \to E.$ The probabilities regularized by α_{alohal} were chosen such that all choices of start and end points into the consensus $S \to M_i \to ... \to M_i \to E$ for $1 \le i \le M_i \to ... \to M_i \to E$ $j \le l$, $(i, j) \ne (1, l)$ are penalized uniformly. More precisely, we favor large probabilities $1 - P(M_i|S)P(E|M_j)$ for $1 \le i \le j \le l$, $(i, j) \ne (1, l)$. The values used for this article are $\alpha_{flank} = 7,000$, $\alpha_{single} = 1e9$, and $\alpha_{alohal} = 1e4.$

Initialization

First, we guess an initial model length l by taking the median of the sequence lengths and scaling it by a constant c. We found that c = 0.8 works well. It is easier to find a rough initial consensus if the number of match states is limited, which forces the model to restrict itself to the more relevant parts of the sequences. The median is more robust against fragmentary sequences than the average.

The initialization of θ could in principle use prior knowledge about the protein family at hand. However, we are interested in tabula rasa training with an universal initial parameter set independent of the input sequences. We chose an ad hoc position independent initialization that reflects the prior distributions. Intuitively, we want the initial model to focus its probability mass on paths that use all match states. We do this by having larger probability for the initial match-match transitions. We took care

to initialize the entry probabilities dependent on the model length such that $P(M_1|S)$ is always roughly $\frac{1}{2}$. Moreover, we initialize the repeat transition $E \rightarrow C$ with a very small probability, and for the flanking states L, R, and C, we initialize such that the self-loops are more likely than the exits.

Model surgery

After training, we might observe rarely used match states or overused insertion states. We can discard or expand those positions and adapt the model length, which is known as model surgery [3].

Given a trained model, we discard match positions that are used by less than 50% of the sequences. Likewise, we expand positions where more than 50% of all sequences have an insertion by a number of new match states equal to the average insertion length. If a match position is discarded, all incident edges are removed and new edges with default initialization are carefully inserted to close the holes (there is a hole for each consecutive segment of discarded positions). If an insertion is expanded, edges at the position of interest that connect left and right model parts are removed. Eventually, all edges incident to a new match state are default initialized. After each surgery iteration, the flanking states, θ_{init} , the kernel for the transition distribution of the end state E, and the evolutionary times τ of the ancestral probability layer (for details, see section "Ancestral probabilities") are reset to default and the model is trained again. This is repeated at most 4 times, which we found is a good compromise between speed and accuracy. Per default, we train 5 independent models and optimize them with model surgery. Eventually, we choose the model with parameters θ that maximizes $\frac{1}{m}(\mathcal{L}(\theta; S_1, \dots, S_m) + \ln(\rho(\theta)))$ to decode the final alignment.

If the number of surgery iterations is >1, we found it beneficial (both performance- and accuracy-wise) to restrict training in all but the last iterations to sequences with lengths above the qth quantile while keeping a minimum of k sequences. Therefore, initial parameter updates are always on sequences that have roughly full length. Short fragmentary sequences may disturb early training epochs. It is easier to incorporate them, if a rough consensus is established and the matter simplifies to fine-tuning the entry, exit, and repeat probabilities. We found that q = 50% and k =10,000 work well. This is in line with other large-scale MSA methods, where a common denominator is a strong preliminary focus on putative full-length sequences (i.e., sequences with lengths from the upper quantiles). For example, MAFFT-Sparsecore only considers sequences with lengths above the median for its core alignment and the regressive strategy favors the longest sequence as representatives of subtrees (i.e., longer sequences are aligned first).

Ancestral probabilities

We naturally assume the existence of a single whole-protein consensus sequence C that represents the sequence set we wish to align. Homologous sequences Si may be closely or distantly related to C (i.e., we assume they have independent expected mutations per site with respect to the consensus). Model-wise, we introduce evolutionary times τ_i to estimate the distance of S_i to C. The process is conventionally described by the general timereversible substitution model parameterized by a 20 \times 20 matrix Q of instantaneous substitution rates from 1 amino acid to any other [36, 37]. Like the scoring matrices used by traditional alignment algorithms, Q models prior biological knowledge on the relative expected frequencies of amino acid substitutions. From Q, the amino acid mutation probabilities after time τ given an initial amino acid can be derived as follows:

$$P(\tau) = \exp(\tau Q), \tag{7}$$

where exp denotes the matrix exponential. The ath row of this matrix, $P(\tau)_a$, corresponds to the expected amino acid distribution after time τ when starting with amino acid a. As the model is timereversible, it is also the distribution of amino acids τ time units ago at a site where amino acid a is observed now.

We initialize τ with zeros and optimize it under the constraints $0 \le \tau_S \le 2.5$, where the maximal value of 2.5 corresponds to the PAM250 matrix and zero is the identity. The vector τ is learned jointly with the HMM parameters θ . Put differently, we learn the branch lengths of a starlike tree jointly with the sequence model. For each batch of sequences, the correct subset of τ is gathered. The ancestral probabilities with the final values τ are also used during Viterbi decoding of the alignment. More precisely, we replace all likelihoods $P(S_i | \theta)$ with $P(S_i | \theta, \tau_i)$.

The τ_i are related to sequence weights, but they are learned from data and do not require a tree or any other pairwise sequence comparison. Assume that for some suitable distance metric, one sequence S_i has a large total distance to all other sequences. In a sequence weighting scheme, Si would typically have a larger weight than sequences with many close relatives to account for the underrepresentation. Choosing a large τ_i can increase $P(S_i|\theta)$ τ_i) by smearing S_i toward the consensus. But this increase is independent of all other sequences and involves no change of θ .

Technical background

We use TensorFlow [38] to automatically compute the gradients of ℓ with respect to θ and τ . We use the Adam optimizer [8] with a learning rate of 0.1 to minimize ℓ . Note that automatic differentiation allows low-effort changes to the HMM architecture and the prior. Moreover, the addition of any type of preliminary deep learning layer (e.g., ancestral probabilities) is possible. Using a machine learning back end provides access to GPU acceleration and other computational benefits out of the box. Our method does not strictly require a GPU, but it is highly recommended to use one to train models beyond length 100. The training automatically scales to multiple GPUs by splitting the batches.

Data description

We tested learnMSA on HomFam [11], BaliFam [39], and the 10 largest Pfam [28] families. The former two are benchmark collections based on reference alignments from HOMSTRAD [40] and BAliBase [41], respectively. Each reference set is embedded into a large set of putative homologs gathered from Pfam. BaliFam has 2 variants where the references are embedded into 100 and 10,000 homologs, respectively. Low sequence numbers were not our target of interest, but we included the small BaliFam variant specifically to test the upscaling ability of our model. See Table 2 for further details. We did not modify, extend, or reduce HomFam or BaliFam other than the embedding step as just described.

To test the ability of our method to align under high sequence length heterogeneity, we constructed a fragmentary variant of BaliFam10000 by following the procedure that was used to test UPP before [17]. We chose BaliFam10000, because the homologs had lengths comparable to the references, whereas HomFam homologs in many cases appear to be not full-length. We constructed a high-fragmentation collection BaliFrag by randomly selecting 40% of the sequences per dataset in BaliFam10000. For

each of these sequences, we sampled a fragment length from a normal distribution with mean equal to 33% of the mean length of the full-length sequences and a standard deviation of 15. We sampled uniformly from all valid starting positions of the fragment in the whole sequence.

Finally, we experimented with 10 ultra-large datasets that were acquired from Pfam by selecting the largest families (based on the number of sequences in the full alignments) and downloading the respective UniProt datasets that were generated by searching the UniProtKB database using the Pfam family HMM. We also downloaded the corresponding seed alignments to use them as a reference. For the training datasets, we added the seed sequences to the UniProt datasets if not already present and removed all gaps. The families are Zinc finger C2H2 type (PF00096), WD domain G-beta repeat (PF00400), ABC transporter (PF00005), Protein kinase domain (PF00069), Ankyrin repeats (PF12796), Major Facilitator Superfamily (PF07690), Leucine rich repeat (PF13855), Fibronectin type III domain (PF00041), Response regulator receiver domain (PF00072), and Immunoglobulin I-set domain (PF07679). All have known 3-dimensional structure. ABC transporter is the largest dataset with about 3.5 million sequences. See Table 3 for details.

Analysis

We compared learnMSA to the following aligners: Clustal Omega (version 1.2.4), regressive T-Coffee (version 13.45.0.4846264), MA-GUS (git hash f9a3676 from 2022-01-21), UPP (version 4.5.2), and MAFFT-Sparsecore (MAFFT Version 7.490). To the best of our knowledge, there is no mature deep learning-based tool for large multiple alignment of proteins available for comparison.

The command lines to align HomFam and BaliFam were (input/output and CPU arguments omitted):

```
MsaHmm.py
clustalo -t protein --outfmt=fa
mafft-sparsecore.rb
run_upp.py -M -1 -m amino
magus.py -t clustal --recursive false
t_coffee -reg -nseq 100 -tree mbed
         -method mafftginsi msa
```

and for the ultra-large datasets (commands equal to the Hom-Fam/BaliFam case omitted):

```
mafft --parttree
magus.py -t random --recurse True
         --recurseguidetree clustal
t_coffee -reg -nseq 1000 -tree parttree
         -method mafftfftnsi msa
```

We ran learnMSA as well as UPP on all datasets (including ultralarge) in default mode without manual parameter adjustments. We did not attempt to align the ultra-large files with Clustal Omega, because we already observed a severe drop in accuracy on sequences in the thousands. MAFFT-Sparsecore refused to align the ultra-large datasets. We used MAFFT with the parttree option instead. For MAGUS, we enabled recursion for the ultra-large datasets, set the guide tree for the highest recursion level to "random" due to very long runtimes with other choices, and used clustal trees for all other recursion levels. To use T-Coffee regressive on the ultra-large datasets, we increased the maximum number of sequences in the subalignments to 1,000 in the hope that we could avoid very long MSAs due to concatenated independent gaps during the merging steps. For a speedup, we also ran T-Coffee with parttree and MAFFT FFT-NS-i. All parameter changes in order to align the ultra-large datasets were done reactively after testing

Table 2. Dataset properties

Collection	Number of families	Number of sequences			Sequence length		
		Min	Max	Avg	Min	Max	Avg
HomFam (refs.)	94	5	41	8	14	854	215
HomFam	94	93	93,681	8,007	12	854	148
BaliFam (refs.)	59	4	142	27	22	471	158
BaliFam100	59	104	242	127	20	764	161
BaliFam10000	36	10,004	10,142	10,031	7	607	175
BaliFrag	36	10,004	10,142	10,031	7	607	129

Table 3. Ultra-large dataset properties

Family	No. of sequences %id			Sequence length			
	Combined	Seed		Min	Max	Avg	
PF00005		55	26	18	683	146	
	3,489,586						
PF07690		192	13	37	577	284	
	1,861,106						
PF00096		159	41	12	34	23	
	1,783,511						
PF00072		52	25	28	156	110	
	1,767,045						
PF00400		1,465	24	12	101	35	
	1,594,257						
PF00069		38	21	24	511	227	
	1,154,714						
PF12796	945,198	184	24	27	153	78	
PF13855	766,271	62	28	26	73	57	
PF00041	666,310	98	20	27	139	81	
PF07679	579,519	48	21	25	149	83	

Sequence identity is based on full alignment. Sequence lengths are given for

the slower and more accurate settings used for HomFam and BaliFam first.

Our method was run using 8 CPU cores, 100 GB of RAM, and a NVIDIA GeForce RTX 3090 GPU for all datasets, including the ultra-large ones. All other aligners did not utilize a GPU and were run using 8 CPU cores and 100 GB of RAM for HomFam and Bali-Fam and 16 cores and 500 GB of RAM for the ultra-large datasets. We chose all memory numbers as a safe upper limit and did no further experiments to evaluate tight requirements. We used a wall clock limit of 3 days for each individual ultra-large alignment.

Sum-of-pairs (SP) score and total column (TC) score were computed by comparing the subalignments induced by the reference sequences to a structure-based alignment (in case of HomFam and BaliFam) or the Pfam seed alignment (in case of the ultralarge datasets). We used T-Coffee with the aln_compare option. The reference sequences are not known to the aligning method.

On the ultra-large datasets, learnMSA is most accurate and fastest in almost all cases (see Table 4). All other methods except UPP required manual adjustment of the default parameters to get them to work. In the end, not all tested aligners were able to align all datasets indicating technical limitations of state-of-theart tools. In addition to timeout and memory issues, we observed a tendency of the divide-and-conquer methods (T-Coffee, MAGUS) to construct MSAs with much larger column counts than the reference (see the expansion column in Table 4), sometimes to the extent that the output file was too large for further usage. This is most likely due to their merging of subalignments in which independent gaps are stacked rather than aligned. LearnMSAs alignments do not grow in length with increasing number of sequences. Fig. 2 shows representatively that ultra-large MSAs computed by learnMSA tend to be tighter than those of comparable tools and do not suffer from underalignment. In the case of PF00096, learn-MSA has no clear advantage, but this family has relatively high sequence identity and very short sequences and is therefore easier to align than the others. Below 1 million sequences, learnMSA loses its runtime advantage and is about as fast as MAFFT and T-Coffee but at the same time much more accurate.

Fig. 3 shows the distribution of SP and TC scores for HomFam and BaliFam. We were able to match state-of-the-art performance on HomFam. If restricted to the 20 sequence sets with at least 10,000 sequences, the benefit of using pHMM-based alignment increases. Note that the number of sequences in the HomFam collection varies significantly (see Table 2). Likewise, HMM matches state-of-the-art performance on BaliFam10000 but falls behind on BaliFam100.

LearnMSA aligned HomFam and BaliFam10000 in a total of 40 hours (sequential training of 5 independent models on the same machine). For the same, Clustal Omega took 3.5 hours, MAFFT-Sparsecore 24 hours, UPP 19 hours, T-Coffee regressive 9 hours, and MAGUS 48 hours.

We also evaluated how increasing the number of homologs that are aligned together with the reference sequences affects alignment accuracy. To create a biologically realistic test setting, we took the 10 Pfam datasets from Table 3 and aligned the combination of the respective seed sequences (called references in the following) with random subsets of the remaining homologs. We started by aligning only the references. Note that the reference set sizes vary between 38 and 1,465 (Table 3). Homologs were drawn randomly without replacement from the UniProtKB datasets to fill up the aligned datasets to monotonically increasing sizes, such that the resulting sets are nested (in a series of MSAs, homologs are only added, never removed). We repeated this serial sampling procedure 10 times and averaged the results over equal-sized alignments. We compared learnMSA with T-Coffee regressive and MAFFT using the commands above from previous experiments, both the accurate and fast variants.

As seen in Fig. 4, the accurate variants of T-Coffee regressive, MAFFT-Sparsecore, and learnMSA are similar in SP score when only aligning the references. Further, all alignment methods lose accuracy after adding homologs at all. However, the asymptotic accuracy of learnMSA is barely affected by the number of added homologs, whereas we observe clearly decreasing trends for the other methods. The relative performance of the methods is dataset dependent and indicates that learnMSA has advantages for the global alignment of protein families. Starting at 200,000 sequences, we observed that regressive T-Coffee and MAFFT-Sparsecore failed for some MSA tasks (we allowed 200 GB

Table 4. Results for the ultra-large datasets

Family	Method	SP	TC	Hours	Expansion
PF00005	learnMSA	74.9	22.2	10.0	1.89
	UPP	73.5	10.2	52.5	1.98
	MAFFT		e	rror	
	MAGUS		tim	neout	
	Regressive T-Coffee		e	rror	
PF07690	learnMSA	56.1	0.0	30.2	1.82
	UPP	51.6	0.0	35.5	2.48
	MAFFT		e	rror	
	MAGUS		tim	neout	
	Regressive T-Coffee			rror	
PF00096	learnMSA	92.9	6.5	0.9	1.16
	UPP	86.3	0.0	1.7	2.23
	MAFFT	84.1	16.1	0.3	2.74
	MAGUS	94.8	3.2	3.6	4.68
	Regressive T-Coffee	69.9	0.0	0.9	6.55
PF00072	learnMSA	92.4	39.2	2.9	1.1
	UPP	91.4	34.6	6.7	1.32
	MAFFT	64.9	4.6	7.6	3.69
	MAGUS	85.8	33.1	24.8	2.41
	Regressive T-Coffee		output	too large	
PF00400	learnMSA	18.0	0.0	1.1	1.29
	UPP	3.6	0.0	2.0	2.62
	MAFFT	0.0	0.0	2.3	7.71
	MAGUS	6.9	0.0	12.6	17.32
	Regressive T-Coffee	0.0	0.0	2.0	51.28
PF00069	learnMSA	83.4	24.9	11.3	1.37
	UPP	83.3	20.2	19.5	1.6
	MAFFT	54.9	5.4	53.0	3.52
	MAGUS	65.4	18.1	29.1	4.77
	Regressive T-Coffee			rror	
PF12796	learnMSA	72.4	0.0	1.3	0.85
	UPP	40.8	0.0	4.3	3.18
	MAFFT	40.4	0.4	7.5	6.36
	MAGUS	58.9	0.0	67.2	5.62
	Regressive T-Coffee			too large	
PF13855	learnMSA	94.7	26.2	0.8	1.05
	UPP	91.0	21.5	2.5	1.71
	MAFFT	80.6	3.1	1.2	3.05
	MAGUS	94.7	38.5	54.1	1.47
	Regressive T-Coffee	49.2	0.0	0.8	7.21
PF00041	learnMSA	79.1	16.5	1.0	1.34
	UPP	74.9	22.0	2.3	2.18
	MAFFT	43.2	0.0	2.0	7.83
	MAGUS	72.6	10.1	53.8	6.4
	Regressive T-Coffee	37.0	0.0	0.8	15.16
PF07679	learnMSA	94.1	50.0	0.9	1.11
	UPP	88.7	46.0	2.9	1.43
	MAFFT	68.1	13.0	1.1	3.36
	MAGUS	84.0	42.0	4.3	2.12
	Regressive T-Coffee	44.2	2.0	0.6	8.55

Expansion denotes the ratio of the length of the predicted alignment (induced by the reference sequences) to the reference alignment length. Values greater than 1 indicate underalignment (i.e., the estimated alignment is longer than the reference). Timeout: the alignment could not be completed by the method within a wall clock limit of 3 days. Error: the alignment failed with an error (either out of memory or another unknown reason). Output too large: the alignment was successful, but the output file was impractically large to be properly postprocessed (e.g., PF12796: T-Coffee 445 GB, learnMSA 1.2 GB). For each cell and column, the best value is in boldface.

of RAM per MSA). The only methods able to align all datasets were learnMSA and MAFFT with the partree option. For our evaluation, we decided to replace each failed MSA with the largest successful alignment in the respective series of nested sets, assuming, in favor of the aligning method, that the failed MSA is in theory unaffected by an increase in sequence numbers. Despite that, as seen in Fig. 4, typically a further increase in the number of homologs

still leads to a decrease in accuracy of the established algorithmic aligners.

For the high-fragmentation collection BaliFrag, learnMSA can compete with MAFFT-Sparsecore, UPP, and MAGUS (Fig. 3). All rely on robust ways to exclude putative fragmentary sequences in early alignment stages by restricting initial backbone alignments to sequences from the upper quantiles [13, 15, 17]. Clustal Omega

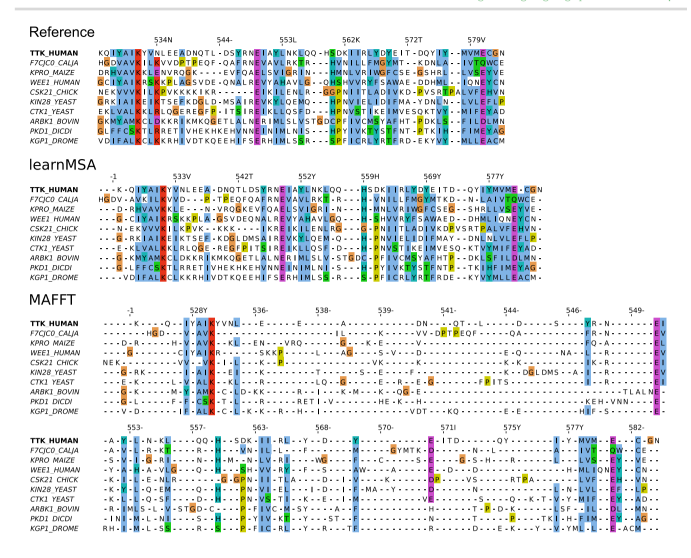


Figure 2: Vertical MSA slices for the ultra-large family PF00069 with more than a million sequences. The 10 most informative sequences (i.e., the most dissimilar ones based on the reference MSA) were extracted using T-Coffee. We took a random vertical slice ranging from columns 25 to 90 in the reference MSA and computed vertical slices for the predicted MSAs as induced by the sequence fragments. We used Jalview 2.11.2.2 with clustalx coloring for visualization. For better comparability, TTK_HUMAN was selected as reference sequence.

and T-Coffee regressive fall behind in this benchmark. This analysis confirms that learnMSA can accurately adapt to fragmentary sequences by first training a pHMM on sequences that are deemed full-length and fitting to the complete sequence set thereafter. Partial-domain hits correctly use the entry and exit transitions as seen in Fig. 5. The difference of learnMSA to the competing methods is that we do not restrict the initial stages to a constant-sized subset of the sequences and that the final alignment is, in principle, able to correct incorrect decisions from earlier iterations. A suitable number of full-length examples is required to find a correct initial model length and to build a consensus. However, UPP teaches us that it is easy to add fragmentary sequences with pH-MMs once a full-length consensus is established [17].

Discussion

We have proposed learnMSA, a novel unsupervised learning approach for the alignment of large protein families. In contrast to state-of-the-art aligners, learnMSA does not require a tree, which eliminates a crucial performance bottleneck and makes learn-MSA asymptotically fast—linear in the number of sequences. It is interesting to see that state-of-the-art performance on large sequence numbers can be reached without a tree by uniform batch sampling. Our method does not rely on progressive, regressive, or divide-and-conquer heuristics. We showed empirically that learn-MSA, when aligning millions of sequences, is both more accurate and faster (even though the measured time was for 5 independent, sequentially trained models). Moreover, when aligning Pfam families, additional homologs decrease the accuracy of traditional, heuristic methods (if they are feasible for large sequence numbers at all), whereas learnMSA is more robust. Whether this statement also applies to established benchmarks like HomFam remains an open question that can be answered if more homologs are gathered for these datasets in the future. A similar scaling experiment, which was done for T-Coffee regressive [10] based on HomFam, suffers from limited data coverage for large sequence numbers (i.e., the number of available families decreases when the MSA depth increases). This is not the case in our study as enough homologs were available from the UniProtKB datasets.

LearnMSA generalizes and automatizes earlier pHMM training approaches for protein families. It does this by taking HMMER's "Plan7" model but avoids the manual adjustment of the "alignment mode" (local versus glocal or unihit versus multihit). Instead, the extra states and transitions (orange in Fig. 1A) are opti-

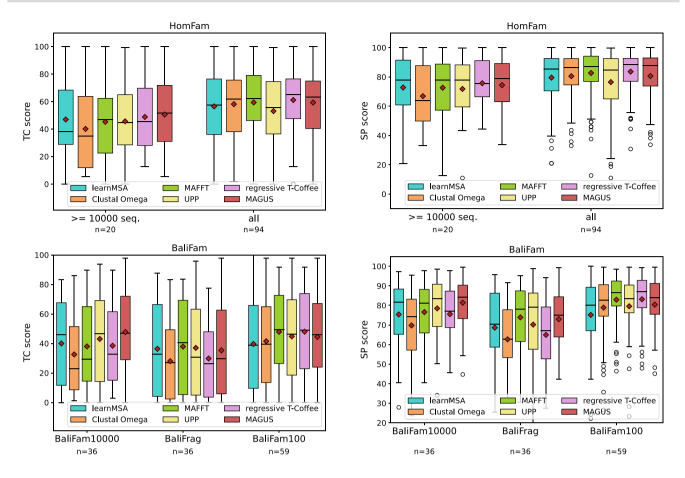


Figure 3: Total column (TC, left) and sum-of-pairs (SP, right) scores for the HomFam (top) and BaliFam (bottom) collections.

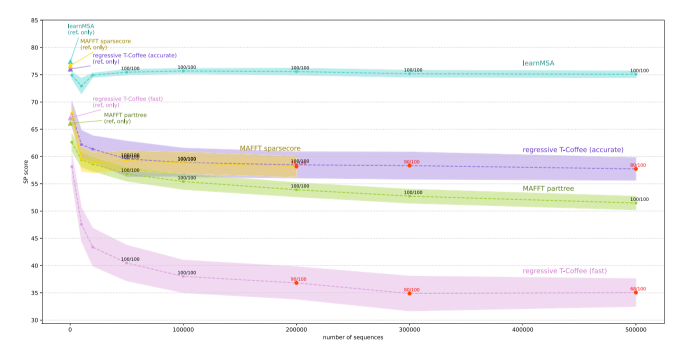


Figure 4: Alignment accuracy as a function of family size. For evaluation purposes, increasing numbers of further homologs are added to a static set of reference sequences. The data points are labeled with the fractions of alignment tasks that produced an usable MSA at all. Missing data points indicate that the aligning method failed for the entirety of the datasets. In case of a failed alignment (due to hardware constraints), we inserted the score of the largest successful alignment in the respective series of nested sets, in favor of the aligning method. Therefore, the plot shows the behavior of the accuracy of the remaining MSAs under the (obliging) assumption that the failed MSAs are in theory unaffected by an increase in sequence numbers. Such incomplete data points are colored red. The shaded area is the standard deviations over the 10 samples, averaged over the families.

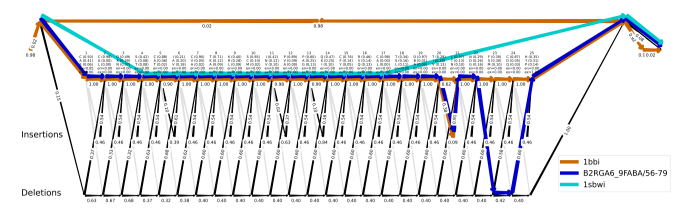


Figure 5: A learned pHMM for the Bowman-Birk serine protease inhibitor family in the HomFam collection with Viterbi paths for 3 different sequences: A single domain hit (blue), a multihit (brown) and, a partial hit (cyan). Numbers on edges are transition probabilities, numbers on nodes are self-loop probabilities. For each match states, the top 3 amino acids and their probabilities are printed, along with the probability of entering and exiting at the respective match.

mized jointly with the core model starting with a tabula rasa configuration, which greatly reduces the required hand-holding. This is also beneficial, if a suitable alignment mode for a dataset is unknown. LearnMSA is designed in a way that minimizes the assumptions a user has to make. Note that for all tested datasets, including dramatically varying sequence numbers and levels of fragmentation, we used learnMSA's default configuration of hyperparameters. It should be pointed out that learnMSA is particularly accurate compared to other methods when aligning families that contain multihits. This is clearly visible in Fig. 6, for example, in the cases of Beta gamma crystallin ("cryst," PF00030), Bowman-Birk protease inhibitor ("bowman," PF00228), or Annexin ("annexin," PF00191).

On HomFam and BaliFam, we match state-of-the-art performance but observe reduced relative accuracy for low sequence numbers. This indicates that there is a lower limit on the sequence numbers below which learnMSA's performance decreases relatively to other methods, but this is not surprising for a statistical learning approach and can currently be solved by falling back to a traditional aligner. There is a slight disadvantage of HMM in average scores for HomFam over all 94 datasets compared to only the largest 20. HomFam contains datasets with a few as 93 sequences. Further evaluation revealed that the disadvantage is not fully explained by low sequence numbers alone, however. Instead, we observed problems if the reference sequences are significantly longer than the homologs (for instance, rhv references are on average 5 times as long as the homologs). Fig. 6 (left) indicates a negative correlation between relative reference length with respect to homologs and score difference. The low-score cases frequently map to "multidomain" secondary structures. In those cases, the references are full-length proteins and the homologs pruned to a specific domain (i.e., information is cut away). This effect is present for all comparison tools except UPP, which is shown in Fig. 6 on the right. For statistical learning, the choice of homologs in HomFam constitutes a problem. The number of reference sequences is very low (8 on average for HomFam), and they can contain information that the homologs miss, which means that potentially important motifs are underrepresented in the dataset. In such situations, it is both hard to guess a suitable initial model length and train a full-length model from scratch. Moreover, this reveals a potential weak spot of the HomFam collection: a method that aligns the longest sequences in a dataset first will most likely catch the references early. The score, which is estimated on the

references only, might therefore overestimate the true score on the complete dataset.

Note that in principle, learnMSA could also align DNA/RNA sequences, but this feature is not implemented yet. Machine learning methods can likely play out their advantages more for proteins due to the relative complexity of parameter space and priors. Further, learnMSA is currently best suited for short- or mediumlength sequences.

Conclusion

Our proposed approach constitutes a probabilistically grounded framework for a large MSA that has potential for further improvements in several directions. Further development might be straightforward because of the extensible nature of our method.

A natural extension of the work presented here are ensembles of pHMMs. They are used in UPP, where a subset of the sequences is aligned and subsequently represented by an ensemble. Recently, MAGUS combined with an HMM ensemble has shown improved accuracy as well [42]. On the HomFam collection, UPP's performance decreased slightly when replacing the ensemble with a single HMM [17]. The latter is related to our approach, with the difference that for learnMSA, the HMM parameters depend on all input sequences instead of a randomly selected backbone set. This might explain why learnMSA aligns HomFam slightly more accurately than UPP, as seen in Fig. 3, even though learnMSA does not currently use an ensemble.

When benchmarking learnMSA, we observed decreasing relative performance when reducing the number of sequences to align. The behavior of state-of-the-art tools is usually complementary: they are more accurate for lower sequence numbers. Moreover, Fig. 6 shows that the relative performance of learnMSA greatly depends on the particular (reference) dataset. This suggests the idea of a combined approach to multiple sequence alignment, where a prior (e.g., the number of sequences) or posterior (e.g., the likelihood) criterion is used to decide between the MSA of either learnMSA or of an established heuristic aligner.

In contrast to traditional learning algorithms for HMMs, gradient-based learning can, in principle, be a module of a larger machine learning model that is trained end-to-end. By design, learnMSA can incorporate any type of sequence context encoded into the HMM alphabet. For instance, single-sequence secondary structure predictions can be incorporated. Secondary structure is

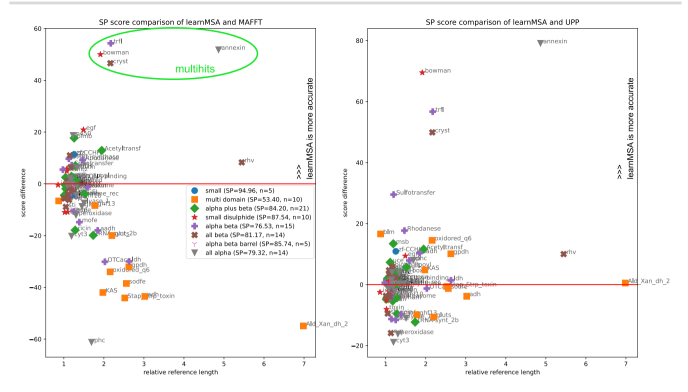


Figure 6: A detailed comparison of the performance of learnMSA relative to MAFFT-Sparsecore (left) and UPP (right) for all 94 HomFam families grouped by secondary structure. Score difference is defined as SP(learnMSA) — SP(other). Relative reference length is defined as the ratio of the average reference length and the average length of the combined dataset including the homologs. For example, "Ald_Xan_dh_2" references are on average about 7 times as long as the respective homologs. The legend contains the average SP score of learnMSA per structure group.

more conserved than primary sequence, and this approach has been shown to increase accuracy in the presence of low sequence identity [43]. There are many kinds of interactions in proteins that are not easily modeled by our current approach (e.g., pairwise correlations between amino acid distributions in positions that are widely separated in the primary sequence but close in the 3-dimensional structure). The field of protein language modeling where parameter-rich sequence models are learned semisupervised [44, 45] based on Attention [46, 47] or LSTMs [48] is also compatible and complementary to our approach. Currently, we use very limited prior knowledge about proteins in the form of parameters as we simply one-hot encode amino acids and only use a rate matrix to compute ancestral probabilities. Using instead semantically rich [44] residual-level embedding vectors from pretrained language models may benefit the predictions.

Availability of Source Code and Requirements

- Project name: learnMSA
- Project homepage: https://github.com/Gaius-Augustus/lear nMSA
- Operating system(s): Platform independent
- Programming language: Python3
- Other requirements: Python packages tensorflow, optional for visualization: networkx, logomaker
- · License: MIT
- RRID: SCR_022572
- biotoolsID: learnMSA

Data Availability

The datasets supporting the results of this article are available in the repository https://github.com/felbecker/MSA-HMM-Analysis. An archival copy of code and data is also available via the *Giga-Science* database GigaDB [49].

Abbreviations

GPU: graphics processing unit; MSA: multiple sequence alignment; (p)HMM: (profile) hidden Markov model.

Competing Interests

The authors declare that they have no competing interests.

Authors' Contributions

F.B. designed and implemented learnMSA, prepared the data, ran all software, and wrote the manuscript. M.S. conceived the idea and designed and implemented an initial version of a recurrent machine learning layer for HMMs and provided prototype code for the usage of ancestral probabilities. Both authors approved the final manuscript.

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