Back-translation for discovering distant protein homologies in the presence of frameshift mutations

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Abstract

Background: Frameshift mutations in protein-coding DNA sequences produce a drastic change in the resulting protein sequence, which prevents classic protein alignment methods from revealing the proteins’ common origin. Moreover, when a large number of substitutions are additionally involved in the divergence, the homology detection becomes difficult even at the DNA level.

Results: We developed a novel method to infer distant homology relations of two proteins, that accounts for frameshift and point mutations that may have affected the coding sequences. We design a dynamic programming alignment algorithm over memory-efficient graph representations of the complete set of putative DNA sequences of each protein, with the goal of determining the two putative DNA sequences which have the best scoring alignment under a powerful scoring system designed to reflect the most probable evolutionary process. Our implementation is freely available at http://bioinfo.lifl.fr/path/.

Conclusions: Our approach allows to uncover evolutionary information that is not captured by traditional alignment methods, which is confirmed by biologically significant examples.

Background

Context and motivation

In protein-coding DNA sequences, frameshift mutations (insertions or deletions of one or more bases) can alter the translation reading frame, affecting all the amino acids encoded from that point forward. Thus, frameshifts produce a drastic change in the resulting protein sequence, preventing any similarity to be visible at the amino acid level. For that reason, classic protein alignment methods, that rely on amino acid comparisons, fail to reveal the proteins’ common origins in the case of divergence by frameshift.

Consequently, it is natural to handle frameshift mutations at the DNA level, by DNA sequence comparisons. Several papers, including [1-4] reported functional frameshifts discovered using classic alignment tools from the BLAST [5,6] family. In all cases, the DNA sequences were relatively well conserved, which allowed the similarity to remain detectable at the DNA level.

However, the divergence may also involve additional base substitutions, that can reduce the similarity of the diverged DNA sequences. It has been shown [7-9] that, in coding DNA, there is a base compositional bias among codon positions, that no longer applies after a reading frame change. A frameshifted coding sequence can be affected by base substitutions leading to a composition that complies with this bias. If, in a long evolutionary time, a large number of codons in one or both sequences undergo such changes, they may be altered to such an extent that the common origin becomes difficult to observe by direct DNA comparison.

In this paper, we address the problem of finding distant protein homologies, in particular when the primary cause of the divergence is a frameshift. We aim at being able to detect the common origins of sequences even if they were affected by an important number of point mutations in addition to the frameshift. Also, when dealing with sequences that have little similarity, we wish to distinguish between sequences that are indeed
(distantly) related, and sequences that resemble by chance. We achieve this by computing the best alignment of DNA sequences that encode the target proteins, with respect to a powerful scoring system that evaluates point mutations in their context, based on codon substitution patterns. Our approach implicitly explores all the pairs of DNA sequences that can be translated into these proteins, which allows a wider vision on the match possibilities at the DNA level.

We designed and implemented an efficient method for aligning putative coding DNA sequences, which builds expressive alignments between hypothetical nucleotide sequences obtained by back-translating the proteins, that can provide some information about the common ancestral sequence, if such a sequence exists. We perform the analysis on memory-efficient graph representations of the complete set of putative DNA sequences of each protein. The proposed method consists of a dynamic programming alignment algorithm that computes the two putative DNA sequences that have the best scoring alignment under an appropriate scoring system.

**Protein back-translation**

Back-translation or reverse translation of a protein usually refers to obtaining one of the DNA sequences that encodes the given protein. Several methods for achieving this exist [10,11], aiming at finding the DNA sequence that is most likely to encode that protein. Several programs use multiple protein alignments to improve the back-translation [12,13]. This can be considered to be the opposite to the “translation way”, where translation is used to improve coding DNA alignments or assess new coding DNA [14-17].

In this paper, we are not interested in just one of the coding sequences, but aim at exploring them exhaustively and aligning them with potential frameshifts. Thus, in the context of our work, the back-translation will refer to all the putative DNA sequences, as explained further in the **Methods** section.

**Similar approaches**

The idea of using knowledge about coding DNA when aligning amino acid sequences has been explored in several papers.

A *non-statistical approach* for analyzing the homology and the “genetic semi-homology” in protein sequences was presented in [18,19]. Instead of using a statistically computed scoring matrix, amino acid similarities are scored according to the complexity of the substitution process at the DNA level, depending on the number and type (transition/transversion) of nucleotide changes that are necessary for replacing one amino acid by the other. This ensures a differentiated treatment of amino acid substitutions at different positions of the protein sequence, thus avoiding possible rough approximations resulting from scoring them equally, based on a classic scoring matrix. The main drawback of this approach is that it was not designed to cope with frameshift mutations.

Regarding *frameshift mutation discovery*, many studies [1-4] preferred the plain BLAST [5,6] alignment approach: BLASTN on DNA and mRNA, or BLASTX on mRNA and proteins, applicable only when the DNA sequences are sufficiently similar. BLASTX programs, although capable of insightful results thanks to the six frame translations, have the limitation of not being able to transparently manage frameshifts that occur inside the sequence, for example by reconstructing an alignment from pieces obtained on different reading frames.

For *handling frameshifts at the protein level*, [20] and [21] propose the use of 5 substitution matrices for aligning amino acids encoded on different reading frames, based on nucleotide pair matches between respective codons and amino acid substitution probabilities. One of the main differences between this scoring scheme and the one we present further in this paper is that our scores target nucleotide symbols explicitly, and are computed by taking into account the changes that occur at the DNA level directly. Also, our alignment method allows more flexibility with respect to frameshift gap placement within the alignment.

On the subject of *aligning coding DNA in presence of frameshift errors*, some related ideas were presented in [22,23]. The author proposed to search for protein homologies by aligning their *sequence graphs* (data structures similar to the ones we use in our method). The algorithm tries to align pairs of codons, possibly incomplete since gaps of size 1 or 2 can be inserted at arbitrary positions. The score for aligning two such codons is computed as the maximum substitution score of two amino acids that can be obtained by translating them. This results in a complex, time costly dynamic programming method that basically explores all the possible translations. Our algorithm addresses the same problem, by employing an approach that is more efficient, since it aligns nucleotides instead of codons, and works with simpler data structures thanks to the IUPAC ambiguity code, without any loss of information, as we will show further in the paper. Also, our alignment algorithm is more generic and is not restricted to a certain scoring function. Additionally, the scoring scheme we propose relies on codon evolution patterns, since we believe that, in frameshift mutation scenarios, the information provided by DNA sequence dynamics provides valuable information in addition to amino acid similarities.

**Methods**

The problem of inferring homologies between distantly related proteins, whose divergence is the result of
frameshifts and point mutations, is approached in this paper by determining the best pairwise alignment between two DNA sequences that encode the proteins.

Given two proteins \( P_A \) and \( P_B \), the objective is to find a pair of DNA sequences, \( D_A \) and \( D_B \), such that translation \( (D_A) = P_A \) and translation \( (D_B) = P_B \) which produce the best pairwise alignment under a given scoring system.

The alignment algorithm incorporates a gap penalty that limits the number of frameshifts allowed in an alignment, to comply with the observed frequency of frameshifts in a coding sequence’s evolution. The scoring system is based on possible mutational patterns of the sequences. This leads to reducing the false positive rate and focusing on alignments that are more likely to be biologically significant.

**Data structures: back-translation graphs**

An explicit enumeration and pairwise alignment of all the putative DNA sequences is not an option, since their number increases exponentially with the protein’s length, as all amino acids are encoded by 2, 3, 4 or 6 codons, with the exception of \( M \) and \( W \), which have a single corresponding codon. Therefore, we represent the protein’s “back-translation” (set of possible source DNAs) as a directed acyclic graph, whose size depends linearly on the length of the protein, and where a path represents one putative sequence. As illustrated in Figure 1(a), the graph is organized as a sequence of length \( 3n \) where \( n \) is the length of the protein sequence. At each position \( i \) in the graph, there is a group of nodes, each node representing a possible nucleotide that can appear at position \( i \) in at least one of the putative coding sequences.

For identical nucleotides that appear at the same position of different codons for the same amino acid, and are preceded by different nucleotides within their respective codon, (as it is the case for bases \( C \) and \( T \) at the second position of the codons corresponding to amino acids \( S \) and \( L \) respectively), different nodes are introduced into the graph in order to avoid the creation of paths that do not correspond to actual putative DNA sequences for the given protein. Also, as the scoring system we propose in this paper requires to differentiate identical symbols by their context, identical nucleotides appearing at the third position of different codons for amino acids \( L, S \) and \( R \) will have different corresponding nodes in the back-translation graph.

Basicly, we can consider that each nucleotide symbol \( \alpha \) from a putative coding DNA sequence, belonging to some codon \( c \), is labeled with a word \( l \) which is its prefix in the codon \( c \). Depending on the position of \( \alpha \) in \( c, l \) will consist of 0, 1 or 2 letters. Here we denote such a labeled symbol by \( \alpha l \). Further in the paper we will drop the \( l \) for notation simplicity, and consider this differentiation implicit. Two symbols that appear at the same position of two putative DNA sequences encoding the same protein are identical (and are represented by the same node) if and only if they represent the same nucleotide and their labels are identical.

Two nodes at consecutive positions are linked by an arc if and only if they are either consecutive nucleotides of the same codon, or they are respectively the third and the first base of two consecutive codons. No other arcs exist in the graph.

The construction of a simple back-translation graph for the amino acid \( R \) is illustrated in Figure 2.

More formally, a back-translation graph of an amino acid sequence \( R \) of length \( n \) is a directed acyclic graph \( G_P = (V_P, E_P) \) where:

\[
V_P = \bigcup_{i=1}^{3n} \{\alpha_i^1| \exists D: P = \text{translation}(D) \land \alpha_i \in \text{suffix}(D[1..i])\} 
\]

where \( \{\alpha_i^1\} \) are the nucleotide symbols that appear at position \( i \) in at least one of the protein’s putative coding sequences, and

\[
E_P = \{(\alpha_i^1, \alpha_{i+1}^1) | \exists D: P = \text{translation}(D) \land \alpha_i \in \text{suffix}(D[1..i]) \land (\alpha_{i+1} \in \text{suffix}(D[1..i+1]))\} 
\]

are arcs between nodes corresponding to symbols that are consecutive in one of the protein’s putative coding sequences.

![Figure 1](http://www.almob.org/content/5/1/6)

**Figure 1** Back-translation graph examples. A fully represented (a) and condensed (b) back-translation graph for the amino acid sequence YSH.

![Figure 2](http://www.almob.org/content/5/1/6)

**Figure 2** Obtaining a simple back-translation graph for the amino acid \( R \). The construction of a simple back-translation graph, for the amino acid \( R \), encoded by 6 codons, is illustrated here. Note that identical nucleotides are associated to different nodes if they have different prefixes in the codons where they appear.
Note that, in the implementation, the number of nodes is reduced by using the IUPAC nucleotide codes [24]. For back-translating an amino acid, only 4 extra nucleotide symbols - R, Y, H and N, representing the sets \{A, G\}, \{C, T\}, \{A, C, T\} and \{A, C, G, T\} respectively - are necessary. In this condensed representation, the number of ramifications in the graph is substantially reduced, as illustrated by Figure 1. More precisely, the only amino acids with ramifications in their back-translation are amino acids R, L and S, each encoded by 6 codons with different prefixes, while the back-translations of all other amino acids are simple sequences of 3 symbols. As we will show below, there is no information loss regarding the actual pair of non-ambiguous symbols aligned.

The reverse complementary of a back-translation graph can be obtained in a classic manner, by reversing the arcs and complementing the nucleotide symbols that label the nodes, as illustrated in Figure 3.

### Alignment algorithm

When aligning two back-translated protein sequences, we are interested in finding the two putative DNA sequences (one for each protein) that are most similar. To achieve this, we use a dynamic programming method, similar to the Smith-Waterman algorithm [25], extended to back-translation graphs, and equipped with gap related restrictions.

![Figure 3 Example of reverse complementary back-translation graphs for the amino acid sequence YSH](image)

#### Figure 3 Example of reverse complementary back-translation graphs for the amino acid sequence YSH. The reverse complementary of a back-translation graph can be obtained in a classic manner, by reversing the arcs and complementing the nucleotide symbols that label the nodes.

Given input graphs \(G_A\) and \(G_B\) obtained by back-translating proteins \(P_A\) and \(P_B\), the algorithm finds the best scoring local alignment between two DNA sequences comprised in the back-translation graphs (illustrated in Figure 4). The alignment is built by filling each entry \(M[i, j, (\alpha, \beta)]\) of a dynamic programming matrix \(M\), where \(i\) and \(j\) are positions of \(G_A\) and \(G_B\) respectively, and \((\alpha, \beta)\) enumerates the possible pairs of nodes that can be found in \(G_A\) at position \(i\), and in \(G_B\) at position \(j\) respectively. An example of matrix \(M\) is given in Figure 5.

The dynamic programming algorithm begins with a classic local alignment initialization (0 at the top and left borders), followed by the recursion step described in relation (3). The partial alignment score of each matrix entry \(M[i, j, (\alpha, \beta)]\) is computed as the maximum of 6 types of values:

1. (a) 0 (similarly to the classic Smith-Waterman algorithm, only non-negative scores are considered for local alignments).
2. (b) the substitution score of symbols \((\alpha, \beta)\), denoted \(\text{score}(\alpha, \beta)\), added to the score of the best partial alignment ending in \(M[i-1, j-1]\), provided that the partially aligned paths contain \(\alpha_i\) at position \(i\) and \(\beta_j\) on position \(j\) respectively; this condition is ensured by restricting the entries of \(M[i-1, j-1]\) to those labeled with symbols that precede \(\alpha\) and \(\beta\) in the graphs, and is expressed in (3) by \(\alpha_{i-1} \in \text{pred}_{C_A}(\alpha), \beta_{j-1} \in \text{pred}_{C_B}(\beta)\).
3. (c) the cost \(\text{singleGapPenalty}\) of a frameshift (gap of size 1 or extension of a gap of size 1) in the first sequence, added to the score of the best partial alignment that ends in a cell \(M[i, j-1, (\alpha, \beta_{j-1})]\), provided that \(\beta_{j-1}\) precedes \(\beta_j\) in the second graph \(\beta_{j-1} \in \text{pred}_{C_B}(\beta)\); this case is considered only if the number of allowed frameshifts on the current path is not exceeded, or a gap of size 1 is extended.
4. (d) the cost of a frameshift in the second sequence, added to a partial alignment score defined as above.

![Figure 4 Alignment example](image)

#### Figure 4 Alignment example. A path (corresponding to a putative DNA sequence) was chosen from each graph so that the match/mismatch ratio is maximized.
The cost of removing an entire codon from the first sequence, added to the score of the best partial alignment ending in a cell \(M[i, j - 3, (\alpha, \beta)]\).

(f) the cost of removing an entire codon from the second sequence, added to the score of the best partial alignment ending in a cell \(M[i - 3, j, (\alpha, \beta)]\)

We adopted a non-monotonic gap penalty function, where insertions and deletions of full codons are less penalized than reading frame disruptive gaps. Additionally, since frameshifts are considered to be very rare events, their number in an alignment is restricted. More precisely, as can be seen in equation (3), two particular kinds of gaps are considered: i) **frameshifts** - gaps of size 1 or 2, with high penalty, whose number in a local alignment is limited, and ii) **codon skips** - gaps of size 3 which correspond to the insertion or deletion of a whole codon.

\[
M[i, j, (\alpha, \beta)] = \begin{cases} 
0 & \text{if } i = 0 \text{ or } j = 0, \\
M[i-1, j-1, (\alpha_{i-1}, \beta_{j-1})] + \text{score}(\alpha_{i}, \beta_{j}), & \alpha_{i-1} \in \text{prod}_{\alpha_{i-1}}(\alpha_{i}), \beta_{j-1} \in \text{prod}_{\beta_{j}}(\beta_{j}), \\
\max_{\beta_{j-1} \in \text{prod}_{\beta_{j}}(\beta_{j})} \left( M[i-1, j-1, (\alpha_{i-1}, \beta_{j-1})] + \text{singleGapPenalty} \right) & \beta_{j-1} \in \text{prod}_{\beta_{j}}(\beta_{j}), \\
M[i-1, j, (\alpha_{i-1}, \beta_{j})] + \text{singleGapPenalty} \quad \alpha_{i-1} \in \text{prod}_{\alpha_{i-1}}(\alpha_{i}), & j \geq 3, \\
M[i, j-3, (\alpha_{i}, \beta_{j})] + \text{tripleGapPenalty} \quad \beta_{j} \in \text{prod}_{\beta_{j}}(\beta_{j}), & j \geq 3, \\
M[i-3, j, (\alpha_{i}, \beta_{j})] + \text{tripleGapPenalty} \quad \alpha_{i} \in \text{prod}_{\alpha_{i}}(\alpha_{i}), & j \geq 3.
\end{cases}
\tag{3}
\]

Although the algorithm is defined for back-translated protein alignment, it can also be used for aligning two DNA sequences or a DNA sequence to a protein. The graph corresponding to a DNA sequence has only one node at each position. Thus, the method can be used for aligning proteins to longer DNA sequences containing coding regions. However, when long sequences are aligned by dynamic programming methods, time and space complexity issues need to be addressed.

**Complexity and improvements**

In this section we discuss the time and space complexity of our method and show how we can improve the latter using an approach inspired by [26].

**Space complexity of the back-translation graphs**

The space necessary for storing the back-translation graph of a protein sequence \(P\) of size \(n\) depends linearly on \(n\). Basically, as mentioned in the section dedicated to data structures, the back-translation graph \(G_P = (V_P, E_P)\) consists of \(3n\) groups of nodes \(\{\alpha_i\}\) (as each of the \(n\) amino-acids are encoded by sequences of 3 nucleotides). Every group \(i\) contains the nodes corresponding to the nucleotides that can appear at position \(i\) in at least one of the putative coding sequences (see (1)).

The number of nodes in a group is limited by the number of codons that encode an amino acid, that we denote \(C\) (6 in the
worse case scenario for non-ambiguous symbols) and thus does not depend on the protein's length.

Arcs exist only between nodes in consecutive groups (equation (2)), therefore each node can have a limited number of neighbors. Consequently, the overall memory consumption for storing the back-translation graph of a protein sequence $P$ of size $n$ is $\mathcal{O}(n)$. The worst case scenario is a protein sequence composed only of the amino acids $L$, $S$, $R$, which are encoded by 6 codons each, and hence have the most complex back-translation. For each such amino acid, 10 nodes and 20 arcs are necessary, yielding a maximum memory size of $30n$ for the entire graph. For the ambiguous nucleotide symbol encoding though, 6 nodes and 6 arcs are necessary in the worst case for each amino acid, while most amino acids only require 3 nodes and 3 arcs for their back-translated representation.

**Complexity of the alignment algorithm**

Let $G_A$ and $G_B$ be graphs obtained by back-translating proteins $P_A$ and $P_B$, of lengths $n_A$ and $n_B$ respectively. The dynamic programming matrix $M$ computed by the alignment algorithm will have $3n_A + 1$ rows and $3n_B + 1$ columns. Each cell of the matrix $M[i,j]$ has several entries corresponding to the possible pairs of nodes from each sequence. The number of entries is bounded by the square of the number of nodes that can appear on each position in the graph ($C^2$). Consequently, the total number of entries in the matrix is at most $C^2(3n_A + 1)(3n_B + 1)$, hence $\mathcal{O}(n_An_B)$.

Each entry holds the score of the partial alignment ending at the corresponding positions, as well as the number of frameshifts that occurred on the path so far (to ensure the established limit in the complete alignment) and a reference to the previous matrix entry of the alignment path, to facilitate the traceback. The storage space requirements for this supplementary information are bounded by a constant.

For computing each score in the matrix, the expressions that need to be evaluated are given by equation (3), by querying some of the entries from 5 other cells in the matrix. Since the number of entries in each cell is bounded by $C^2$, this operation is considered to be performed in constant time. Consequently, the overall time complexity of the algorithm is $\mathcal{O}(n_An_B)$. To recover the best alignment and the two actual sequences that contribute to the computation of the score, and computed, the best scoring cell will provide all the necessary information for reconstructing the alignment: the start of the alignment, the intersection with each cut-point, and its end, which is the cell itself. According to these coordinates, subgraphs of the two back-translation graphs are extracted and aligned globally (ensuring that the start and end node pair of each fragment are preserved). The obtained global alignments, combined, will give the best local alignment of the two large sequences.

**Translation-dependent scoring function**

In this section, we present a new translation-dependent scoring system suitable for our alignment algorithm. Our scoring scheme incorporates information about possible mutational patterns for coding sequences, based on a codon substitution model, with the aim of filtering out alignments between sequences that are unlikely to have common origins.
Mutation rates have been shown to vary within genomes, under the influence of different factors, including neighbor bases [27]. Consequently, a model where all base mismatches are equally penalized is oversimplified, and ignores possibly precious information about the context of the substitution.

With the aim of retracing the sequence’s evolution and revealing which base substitutions are more likely to occur within a given codon, our scoring system targets pairs of triplets \((\alpha, p, a)\), were \(\alpha\) is a nucleotide, \(p\) is its position in the codon, and \(a\) is the amino acid encoded by that codon, thus differentiating various contexts of a substitution. There are 99 valid triplets out of the total of 240 hypothetical combinations. Pairwise alignment scores are computed for all possible pairs of valid triplets \((t_i, t_j) = ((\alpha_i, p_i, a_i), (\alpha_j, p_j, a_j))\) as a classic log-odds ratio:

\[
score(t_i, t_j) = \lambda \log \frac{f_{t_i,t_j}}{b_{t_i,t_j}}
\]

where \(f_{t_i,t_j}\) is the frequency of the \(t_i \rightarrow t_j\) substitution in related sequences, and \(b_{t_i,t_j} = p(t_i)p(t_j)\) is the background probability. This scoring function is used in the algorithm as shown by equation (3)(b), where we refer to it as \(score(\alpha_a, \alpha_b)\), without explicitly mentioning the context - amino acid and position in the corresponding codon - of the paired nucleotides. These details were omitted in equation (3) for generality (other scoring functions, that do not depend on the translation, can be used by the algorithm too) and for notation simplicity.

In order to obtain the foreground probabilities \(f_{t_i,t_j}\) we consider the following scenario, depicted in Figure 6: two proteins are encoded on the same DNA sequence, on different reading frames; at some point, the sequence was duplicated and the two copies diverged independently; we assume that the two coding sequences undergo, in their independent evolution, synonymous and non-synonymous point mutations, or full codon insertions and removals.

The insignificant amount of available real data that fits our hypothesis does not allow classical, statistical computation of the foreground and background probabilities. Therefore, instead of doing statistics on real data directly, we will rely on codon frequency tables and codon substitution models, either mechanistic or empirically constructed.

**Codon substitution models**

Mechanistic codon substitution models

We can assume that codon substitutions in our scenarios are modeled by a Markov model presented in [28] that specifies the relative instantaneous substitution rate from codon \(i\) to codon \(j\) as:

\[
Q_{ij} = \begin{cases} 
0 & \text{if } i \text{ or } j \text{ is a stop codon, or} \\
\pi_j & \text{if } i \rightarrow j \text{ requires more than 1 nucleotide substitution,} \\
\pi_k & \text{if } i \rightarrow j \text{ is a synonymous transversion,} \\
\pi_o & \text{if } i \rightarrow j \text{ is a synonymous transition,} \\
\pi_p & \text{if } i \rightarrow j \text{ is a non synonymous transversion,} \\
\pi_n & \text{if } i \rightarrow j \text{ is a non synonymous transition.} 
\end{cases}
\]

for all \(i \neq j\). Here, the parameter \(\omega\) represents the non-synonymous-synonymous rate ratio, \(\kappa\) the transition-transversion rate ratio, and \(\pi_j\) the equilibrium frequency of codon \(j\). As in all Markov models of sequence evolution, absolute rates are found by normalizing the relative rates to a mean rate of 1 at equilibrium, that is, by enforcing \(\sum_i \sum_{j \neq i} \pi_i Q_{ij} = 1\) and completing the instantaneous rate matrix \(Q\) by defining \(Q_{ii} = -\sum_{j \neq i} Q_{ij}\) to give a form in which the transition probability matrix is calculated as \(P(\theta) = e^{\theta Q}\) [29]. Evolutionary times \(\theta\) are measured in expected number of nucleotide substitutions per codon.

Note that there exist some more advanced codon substitution models, targeting sequences with overlapping reading frames [30]. However, such models do not fit our scenario, because they are designed for overlapping reading frames, where a mutation affects both translated sequences, while in our case the sequences become at one point independent and undergo mutations independently.

**Empirical codon substitution model**

The mechanistic codon substitution model presented above simulates substitutions with accurate parameters, but does not take into account the selective pressure and the resulting effects on the final codon conservation.

One of these effects, most commonly known and most observable in alignments of coding sequences, is the “third base mutation”: in most cases, the encoded amino acid is not changed by a transition mutation of the codon third base; this is true in some cases of transversion mutations as well.

There are several other specific conservation families for groups of amino acids, as the *aliphatic* conservation (amino acids \(L, I, V\) where corresponding amino acid codons share \(T\) at their second base. The last base is, within this group, almost a free choice, while the first has a large degree of freedom. It is thus expected to frequently observe the second \(T\) conserved on such codons when aligned with the *aliphatic* group. A similar phenomenon (however with a weaker frequency) appears for the subset \((A, S, T)\) of the “small” amino acids, where the codons have in common the second base \(C\).

In other chemically related amino acid groups, the succession of nucleotide substitutions at the codon level
follows more complex paths, as it is the case for positively charged amino acids \((R, K)\), aromatic amino acids \((F, Y, W)\), etc.

Such different and complex conservation patterns are difficult to express and model with simple rules. As most of the matrices built for proteins, an empirical estimation gives a very good global approximation. In \([31]\), the first empirical codon substitution matrix entirely built from alignments of coding sequences from vertebrate DNA is presented. A set of 17,502 alignments of orthologous sequences from five vertebrate genomes yielded 8.3 million aligned codons from which the number of substitutions between codons were counted.

From this data, \(64 \times 64\) probability matrices and similarity score matrices ("1-codon PAM") were computed. One can use these probability matrices as an alternative to the ones obtained using the mechanistic model.

**Foreground probabilities**

Once the codon substitution probabilities are obtained, \(f_{i,j}\) can be deduced in several steps. Basically, we first need to identify all pairs of codons with a common subsequence, that have a perfect semi-global alignment (for instance, codons CAT and ATG satisfy this condition, having the common subsequence AT; this example is further explained below). Then we assume that the codons from each pair undergo independent evolution, according to the codon substitution model. For the resulting codons, we compute, based on all possible original codon pairs, \(p((\alpha_o, p_o, c_o), (\alpha_p, p_p, c_p))\) - the probability that nucleotide \(\alpha_o\), located at position \(p_o\) of codon \(c_o\) and nucleotide \(\alpha_p\), situated on position \(p_p\) of codon \(c_p\) have a common origin (equation (7)). From these, we can immediately compute, as shown by equation (8) below, \(p((\alpha_o, p_o, a_o), (\alpha_p, p_p, a_p))\), corresponding to the foreground probabilities \(f_{i,j}\), where \(t_i = (\alpha_o, p_o, a_o)\) and \(t_j = (\alpha_p, p_p, a_p)\).

In the following, \(p(c_j \rightarrow c_i, \theta)\) stands for the probability of the event **codon** \(c_i\) **mutates into codon** \(c_j\) **in evolutionary time** \(\theta\), and is given by a codon substitution probability matrix \(P_{c_i,c_j}(\theta)\).

The notation \(c_i[\text{interval}] = c_j[\text{interval}]\) states that codon \(c_i\) restricted to the positions given by \(\text{interval}_i\) is a sequence identical to \(c_j\) restricted to \(\text{interval}_j\). This is equivalent to having a word \(w\) obtained by "merging" the two codons. For instance, if \(c_i = CAT\) and \(c_j = ATG\), with their common substring being placed in \(\text{interval}_i = [2..3]\) and \(\text{interval}_j = [1..2]\) respectively, \(w\) is CATG.

We denote by \(p(c, [\text{interval}]_i \equiv [\text{interval}]_j)\) the probability to have \(c_i\) and \(c_p\), in the relation described above, and we compute it as the probability of the word \(w\) obtained by "merging" the two codons. This function should be symmetric, it should depend on the codon distribution, and the probabilities of all the words \(w\) of a given length should sum to 1. However, since we consider the case where the same DNA sequence is translated on two different reading frames, one of the two translated sequences would have an atypical composition. Consequently, the probability of a word \(w\) is computed as if the sequence had the known codon composition when translated on the reading frame imposed by the first codon, or on the one imposed by the second. This hypothesis can be formalized as:

\[
p(w) = p(w \text{ on } rf_1 \text{ OR } w \text{ on } rf_2) = p^{rf_1}(w) + p^{rf_2}(w) - p^{rf_1}(w) \cdot p^{rf_2}(w)
\]  

(6)

where \(p^{rf_1}(w)\) and \(p^{rf_2}(w)\) are the probabilities of the word \(w\) in the reading frame imposed by the position of the first and second codon, respectively. This is computed as the products of the probabilities of the codons and codon pieces that compose the word \(w\) in the established reading frame. In the previous example, the probabilities of \(w = CATG\) in the first and second reading frame are:

\[
p^{rf_1}(\text{CATG}) = p(\text{CAT}) \cdot p(\text{G} \cdot \ast) = p(\text{CAT}) \cdot \sum_{c \text{ starts with G}} p(c)
\]

\[
p^{rf_2}(\text{CATG}) = p(\ast \cdot C) \cdot p(\text{ATG}) = \sum_{c \text{ ends with C}} p(c) \cdot p(\text{ATG})
\]
The values of $p((\alpha_i, p_i, c_i), (\alpha_j, p_j, c_j))$ are computed as:

$$
\sum_{c' \in \text{internal}_c|\text{internal}_c} p(c'|\text{internal}_c) \cdot p(c|\text{internal}_c) = c'|\text{internal}_c \cdot p(c'|\text{internal}_c) \cdot p(c|\text{internal}_c)
$$  \hspace{1cm} (7)

from which obtaining the foreground probabilities is straightforward:

$$
f_{t_{ij}} = p((\alpha_i, p_i, a_i), (\alpha_j, p_j, a_j)) = \sum_{c_i \in \text{internal}_c} \sum_{c_j \in \text{internal}_c} p(c_i|\text{internal}_c) \cdot p(c_j|\text{internal}_c)
$$  \hspace{1cm} (8)

Background probabilities

The background probabilities of $(t_i, t_j)$, $b_{t_{ij}}$, can be simply expressed as the probability of the two symbols appearing independently in the sequences:

$$
b_{t_{ij}} = b((\alpha_i, p_i, a_i), (\alpha_j, p_j, a_j)) = \sum_{c_i \in \text{internal}_c} \sum_{c_j \in \text{internal}_c} \pi_{c_i} \pi_{c_j}
$$  \hspace{1cm} (9)

Substitution matrix for ambiguous symbols

Earlier we have shown how to compute the translation dependent scores for non-ambiguous nucleotide symbols. However, as mentioned in the section concerning data structures, we work with ambiguous nucleotide symbols, because their usage improves time and memory consumption while providing the same final results. The scores for ambiguous nucleotide symbol pairs are easily obtained as follows:

$$
score((\overline{\alpha_i}, p_i, a_i), (\overline{\alpha_j}, p_j, a_j)) = \max_{\alpha_i, \alpha_j \in \text{internal}_c|\text{internal}_c} score((\alpha_i, p_i, a_i), (\alpha_j, p_j, a_j))
$$  \hspace{1cm} (10)

where $\overline{\alpha_i}$ is an ambiguous nucleotide symbol representing the possible nucleotides that can appear on position $p_i$ of the codons that encode the amino acid $a_i$, and set $\text{set}_{\overline{\alpha_i}}$ denotes the set of non-ambiguous nucleotide symbols represented by $\overline{\alpha_i}$. Basically, the score of pairing two ambiguous symbols is the maximum over all substitution scores for all pairs of nucleotides from the respective sets.

By using ambiguous symbols, less triplets are formed for each amino acid when compared with the non-ambiguous symbol case. 17 amino acids can be anti-translated as tri-mers with just one ambiguous symbol per position, while the others have two alternatives each of the three positions. Therefore, there are 69 different triplets with ambiguity codes to be paired (as opposed to 99), which means more than twice less storage space necessary for the score matrix.

For the reconstruction of the non-ambiguous putative DNA sequences at traceback, the actual pair of nucleotides that have the highest substitution score from the sets corresponding to two paired ambiguous symbols is required. These are easily obtained for each pair of ambiguous symbols as

$$
sym((\overline{\alpha_i}, p_i, a_i), (\overline{\alpha_j}, p_j, a_j)) = \max_{\alpha_i, \alpha_j \in \text{internal}_c|\text{internal}_c} \text{score}((\alpha_i, p_i, a_i), (\alpha_j, p_j, a_j))
$$  \hspace{1cm} (11)

Parametrization

In this section we have presented a general framework that helps to compute a translation dependent scoring function for DNA sequence pairs, parametrized by a codon substitution model and an evolutionary time measured in expected number of mutations per codon. We consider that the sequences evolve independently, and the distance is relative to the original sequence.

Score evaluation

The score significance is estimated according to the Gumbel distribution, where the parameters $\lambda$ and $K$ are computed with the method described in [32,33]. In the future, we aim at improving our estimation by using a computation method more suited for gapped alignments, such as [34].

We use two different score evaluation parameter sets for the forward alignment (where the two back-translated graphs that are aligned have the same translation sense) and the reverse complementary alignment (where one of the graphs is aligned with the reverse complementary of the other), because these are two independent cases with different score distributions.

In order to obtain a more refined evaluation of the alignments, we introduce $(\lambda, K)$ parameters for estimating the score significance of alignment fragments inside which the reading frame difference is preserved. Therefore, there are eight $(\lambda, K)$ parameters that help to evaluate the alignments (four for the forward alignment sense and four for the reverse complementary alignment sense):

- $(\overline{\lambda}_\text{FW}, K_{\text{FW}})$ for the forward sense and $(\overline{\lambda}_\text{RC}, K_{\text{RC}})$ for the reverse complementary sense respectively, that are used for evaluating the score of the whole alignment.

- $(\lambda_{\alpha_i}, K_{\alpha_i})$ for the forward sense and $(\lambda_{\alpha_j}, K_{\alpha_j})$ for the reverse complementary sense respectively, with $i \in \{0, 1, 2\}$ that are used for evaluating the scores of each alignment fragment within which the reading frame difference is preserved. This second evaluation aims at providing a measure of the actual contribution of each such fragment to the score of the alignment.

The parameters $(\lambda_{\alpha_i}, K_{\alpha_i})$ are estimated on alignments restricted to the respective reading frame difference, where further frameshifts are not allowed, while $(\overline{\lambda}_\text{FW}, K_{\text{FW}})$ and $(\overline{\lambda}_\text{RC}, K_{\text{RC}})$ are computed in a more flexible setup, where a limited number of frameshifts is accepted.
**Behavior in the non-frameshifted case**

In this section we discuss the behavior of the proposed scoring system when aligning protein sequences without a frameshift. Given their construction method, we expect the scores to reflect the amino acid similarities, but also to be influenced by similarities at the DNA level.

To evaluate how our scores, used in non-frameshifted alignments, would relate to the classic scoring systems used by biological sequence comparison methods, we first compute, for each scoring matrix $T$ corresponding to an evolutionary distance $\theta$, the expected score for each amino acid pair, as:

\[
T_{AA}(a_i, a_j) = \sum_{p_i=1}^{3} \sum_{t=t' \in \{a_i, p_i, a_j\}, t'=(a_i, p_i, a_j)} p(t,t')T(t,t')
\]  

(12)

Where

\[
p(t,t') = \sum_{c \in \text{encodes } a_i} p(c) \cdot \sum_{c' \in \text{encodes } a_j} p(c')
\]  

(13)

Then, considering each amino acid pair as an observation, we compute the correlation coefficient of these expected scores and the BLOSUM matrices as given by [35].

We also evaluate the correlation with the expected amino acid pair scores obtained when the sequences are aligned using a classic nucleotide match/mismatch system. The latter expected amino acid pair scores are also obtained as weighted sums of scores, in a manner similar to the one described by equations (12) and (13), where the score for aligning two symbols has one of the three established values for match, transition mutation or transversion mutation. For these classic scores, we used the values +5, -3, -4 in the examples reported below, although we have not noticed any drastic changes when different sets of values are used. The obtained correlation coefficients are reported in Table 1.

They suggest that the obtained translation dependent score matrices, either obtained from mechanistic or empirical codon substitution models, are a compromise between the “fully selective” BLOSUM matrices and the non-selective DNA scores.

On the one hand, the scores obtained using the mechanistic model do not make use of the selective pressure, and for this reason are more likely to be correlated with the classic DNA scores. On the other hand, the scores based on empirical codon substitution models reflect the constraints imposed by the similarity of the amino acids encoded by the codons. Hence, they show a strong correlation with the BLOSUM matrices when used without a frameshift.

**Results and Discussion**

We have proposed a method for aligning protein sequences with frameshifts, by back-translating the proteins into graphs that implicitly contain all the putative DNA sequences, and aligning them with a dynamic programming algorithm that uses a scoring system designed for this particular purpose.

**Implementation and availability**

A Java implementation of our method is available at http://bioinfo.lifl.fr/path/. The files containing translation dependent score matrices computed for several evolutionary distances can be downloaded at the same address.

**Experimental results**

We will further discuss several significant frameshifted alignments obtained with our method. The experimental results presented here were obtained in the following experimental setup: a search for frameshifted forward alignments was launched on samples from the full NCBI protein databases for several species, using a 00.50 base per codon divergence scoring matrix; we selected only the alignments with an E-value < 10^-9, presenting at least one significant frameshift.

**Yersinia pestis: Frameshifted transposases**

Figure 7 displays the alignment of two transposase variants from *Yersinia pestis*. Both proteins are widely present on the NCBI nr database. The mechanism involved is (most probably) a programmed translational frameshifting since such mechanism has been quite frequently observed in several other transposases from related species, e.g. as in E. coli [36].

**Xylella fastidiosa: Frameshifted β-glucosidases**

Two β-glucosidase variants from *Xylella fastidiosa* are aligned on Figure 8 with both variants widely present on the NCBI nr database. *Xylella fastidiosa* is a plant
pathogen transmitted by Cicadellidae insects (Homalodisca vitripennis, Homalodisca liturata) and responsible for phony disease on the peach tree, leaf scorch on the oleander, and Pierce’s disease on grape. The β-glucosidase is usually required by several organisms to consume cellulose.

Interestingly, β-glucosidase frameshifts have already been studied in [37] on several bacteria including such γ-proteobacteria as Erwinia herbicola and Escherichia coli of Enterobacteriales but not directly observed in Xanthomonadales.

Figure 7 Yersinia pestis: transposases
The alignment of two transposase variants from Yersinia pestis: [GenBank:167423046] - subsequence 4-167 of the back-translation, and [GenBank:EDR63673.1] - subsequence 225-389 of the back-translation. The frameshift mutation at position 115/336 corrects the reading frame. The frameshifted alignment fragment has an E-value of 10^{-7}.

Figure 8 Xylella fastidiosa: glucosidases
Two β-glucosidase variants from Xylella fastidiosa: [GenBank:AAO29662.1] - subsequence 202-2645 of the back-translation, and [GenBank:EAO32640.1] - subsequence 2-2444 of the back-translation. We only show in this image a fragment of the full alignment (the first 239 base pairs). The second part is not particularly interesting in our context because the sequences are aligned on the same reading frame, with a very small number of mismatches. In the first part, the sequences are aligned with a reading frame difference that is corrected starting with positions 304/104. The frameshifted alignment fragment has an E-value of 10^{-8}.

Figure 9 Elapidae: neurotoxins (1)
Two presynaptic neurotoxins from two higher snakes of the Elapidae family (Bungarus candidus and Naja kaouthia): [Swiss-Prot:Q8AY47.1] - subsequence 64-407 of the back-translation, and [PIR:PSNJ2K] - subsequence 13-354 of the back-translation. The sequences are aligned on the same reading frame up to position 186/135, and on a +1 reading frame from that point forward. The frameshifted fragment has an E-value of 10^{-9}.
Venom neurotoxins

Diversification of venom toxins has been studied in [38] for advanced snakes: frameshifts were one of the most significant mechanisms involved in the "evolution of the arsenal" and its diversification toward specialized prey capture, sometime with a loss of neurotoxicity [39]. We thus studied neurotoxins from several higher snakes. In Figure 9, we show the alignment of two presynaptic neurotoxins from two higher snakes of the Elapidae family (Bungarus candidus and Naja kaouthia). Most of the sites are conserved: the primary metal binding site and the putative hydrophobic channel remain before the frameshift, and only the fourth (and last) part of the catalytic network seems changed. We also noticed that, in the original second sequence, the Cysteine regions are more conserved at the DNA level than other amino acids, even after the frameshift, which is a strong hint of the non randomness of this part of the alignment.

Following the discovered frameshift of Figure 9, we took into consideration the sequences of Bungarus candidus species that were similar to the non-frameshifted presynaptic neurotoxin of Naja kaouthia. An interesting alignment is presented in Figure 10, showing a protein that aligns to it well but not perfectly (at least 4 non synonymous transitions before the frameshift and 1 transversion after): this lets open the potential "duplicated first then frameshifted" origin of the frameshifted protein. This assumption was strongly supported by the alignment of the two corresponding cDNA [GenBank:AY057881.1] and [GenBank:AY057880.1] of two homologous proteins.

Furthermore, Figure 11 displays an alignment of two presynaptic neurotoxins from two higher snakes of the Elapidae family (Laticauda colubrina and Laticauda laticaudata): It shows that the unidentified peptide is in fact an alternative splicing (or frameshifted) variant of the neurotoxin. The frameshifted fragment has an E-value of $10^{-10}$.

**Figure 10 Elapidae: neurotoxins (2).** Two Bungarus candidus proteins, very similar at the DNA level ([Swiss-Prot:Q8AY47.1] and [Swiss-Prot: Q8AY48.1]). From the first 94 amino acid pairs, only 4 present mismatches (which are transitions at the coding DNA level). A frameshift mutation is visible at position 284 of the back-translated sequences. The fragments following it are almost perfectly aligned with a frameshift, with an E-value of $10^{-9}$.

**Figure 11 Elapidae: neurotoxins (3).** Two presynaptic neurotoxins from two higher snakes of the Elapidae family ([DDBJ:BAA75760.1] of Laticauda colubrina and [DDBJ:BAC78208.1] of Laticauda laticaudata): It shows that the unidentified peptide is in fact an alternative splicing (or frameshifted) variant of the neurotoxin. The frameshifted fragment has an E-value of $10^{-10}$.
the neurotoxin. Note that BLASTP identification of the frameshifted peptide on the NCBI nr database gives high E-values (minimal is of 0.67) and is thus expected to be missed by automatic prediction tools (since most of the features specific to neurotoxin are not present), whereas the non frameshifted peptide fragment, once identified, gives several E-values <10^{-10} on NCBI nr.

Platelet-derived growth factor
Platelet-derived growth factor is a potent mitogen for cells of mesenchymal origin. Binding of this growth factor to its affinity receptor elicits a variety of cellular

Figure 12 Platelet-derived growth factor proteins. The alignment of the back-translated platelet-derived growth factor proteins from Homo sapiens and Ratus sp ([Swiss-Prot:P04085.1] and [DDBJ:BAA00987.1]). The two proteins share high similarity at the amino acid level on the subsequences 1-84 and 113-195. The amino acids 85-112 can be easily aligned with a frameshift, with an E-value of 10^{-6}. Both the "inducing" and "correcting" frameshifts are located on two different exons.

Figure 13 Classic protein alignment of the platelet-derived growth factor proteins. The classic protein alignment of the platelet-derived growth factor proteins from Homo sapiens and Ratus sp ([Swiss-Prot:P04085.1] and [DDBJ:BAA00987.1]) shows very little amino acid similarity between the 85-112 subsequences, that we have successfully aligned on a +1 frameshift.
responses. It is released by platelets upon wounding and plays an important role in stimulating adjacent cells to grow and thereby heals the wound. In Figure 12, we show the alignment of the back-translated human and rat platelet-derived growth factor proteins. The two proteins share high similarity at the amino acid level on the subsequences 1-84 and 113-195. The amino acids 85-112 can be easily aligned with a frameshift, as can be seen in Figure 12, while classic protein alignment reveals little similarity in these areas (Figure 13).

It is interesting to notice that this double frameshift (if confirmed) may have little influence on the protein (only the beginning of the receptor binding interface is modified). It is also interesting to notice that both the “inducing” and “correcting” frameshifts are located on two different exons.

Conclusions
In this paper, we addressed the problem of finding distant protein homologies, in particular affected by frameshift events, from a codon evolution perspective. We search for protein common origins by implicitly aligning all their putative coding DNA sequences, stored in efficient data structures called back-translation graphs. Our approach relies on a dynamic programming alignment algorithm for these graphs, which involves a non-monotonic gap penalty that handles frameshifts and full codon indels differently. We designed a powerful translation-dependent scoring function for nucleotide pairs, based on codon substitution models, whose purpose is to reflect the expected dynamics of coding DNA sequences.

We illustrated our approach with several alignment examples of known and hypothetical frameshifted proteins, some of which are not detectable via classic alignment methods because of the low coding sequence similarity. Such examples support our method’s applicability in the discovery of distant protein homologies and functional frameshifting, without any explicit information about the coding sequences. Future work includes further improvements of the scoring system, for example in order to focus on the detection of short double correcting frameshifts (two frameshifts separated by a small number of bases, where the second corrects the reading frame disrupted by the first). Such cases have been shown to occur frequently [37], but are often highly penalized by sequence comparison methods, that discard the correct alignment in favor of an ungapped one with a higher score.

Some natural extensions of our work include the support for multiple alignments of back-translation graphs. This feature can be useful for confirming frameshifts by similarity of the frameshifted subsequence with the corresponding fragments from several members of a family. Also, for boosting the efficiency, seeding techniques for back-translation graphs could be explored, possibly inspired by the BLASTP score-based seeds.

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Authors’ contributions
GX initiated and guided the project, and proposed the first version of the method. MG refined the method, proposed the initial scoring system, did the implementation and the web interface. LN contributed to defining and improving the scoring system, and did most of the experimentation and the analysis of the results. MG drafted the manuscript, which was completed by LN with the “Experimental results” section, and then finalized and approved by GK.

Competing interests
The authors declare that they have no competing interests.

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