Aligning coding sequences with frameshift extension penalties

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Abstract

Background: Frameshift translation is an important phenomenon that contributes to the appearance of novel Coding DNA Sequences (CDS) and functions in gene evolution, by allowing alternative amino acid translations of gene coding regions. Frameshift translations can be identified by aligning two CDS, from a same gene or from homologous genes, while accounting for their codon structure. Two main classes of algorithms have been proposed to solve the problem of aligning CDS, either by amino acid sequence alignment back-translation, or by simultaneously accounting for the nucleotide and amino acid levels. The former does not allow to account for frameshift translations and up to now, the latter exclusively accounts for frameshift translation initiation, not considering the length of the translation disruption caused by a frameshift.

Results: We introduce a new scoring scheme with an algorithm for the pairwise alignment of CDS accounting for frameshift translation initiation and length, while simultaneously considering nucleotide and amino acid sequences. The main specificity of the scoring scheme is the introduction of a penalty cost accounting for frameshift extension length to compute an adequate similarity score for a CDS alignment. The second specificity of the model is that the search space of the problem solved is the set of all feasible alignments between two CDS. Previous approaches have considered restricted search space or additional constraints on the decomposition of an alignment into length-3 sub-alignments. The algorithm described in this paper has the same asymptotic time complexity as the classical Needleman-Wunsch algorithm.

Conclusions: We compare the method to other CDS alignment methods based on an application to the comparison of pairs of CDS from homologous human, mouse and cow genes of ten mammalian gene families from the Ensembl-Compara database. The results show that our method is particularly robust to parameter changes as compared to existing methods. It also appears to be a good compromise, performing well both in the presence and absence of frameshift translations. An implementation of the method is available at https://github.com/UdeS-CoBIUS/FsePSA.

Keywords: Coding DNA sequences; Pairwise alignment; Frameshift; Dynamic programming.

Background

Biological sequence alignment is a cornerstone of bioinformatics and is widely used in such fields as phylogenetic reconstruction, gene finding, genome assembly. The accuracy of the sequence alignments and similarity measures are directly re-
lated to the accuracy of subsequent analysis. CDS alignment methods have many important applications for gene tree and protein tree reconstruction. In fact, they are useful to cluster homologous CDS into groups of orthologous splicing isoforms [1, 2] and combine partial trees on orthology groups into a complete protein tree for a gene family [3, 4]. Aligning and measuring the similarity between homologous CDS requires to account for Frameshift (FS) translations that cannot be detected at the amino acid (AA) level, but lead to a high similarity at the nucleotide level between functionally different sub-sequences.

FS translation consists in alternative AA translations of a coding region of DNA using different translation frames [5]. It is an important phenomenon resulting from different scenarios such as, insertion or deletion of a nucleotide sequence whose length is not a multiple of 3 in a CDS through alternative splicing [6, 7] or evolutionary genomic indels [8, 9], programmed ribosomal frameshifting [10], or sequencing errors [11]. Recent studies have reported the role of FS translations in the appearance of novel CDS and functions in gene evolution [6, 12]. FS translation has also been found to be linked to several diseases such as the Crohn’s Disease [13]. The computational detection of FS translations requires the alignment of CDS while accounting for their codon structure. A classical approach for aligning two CDS used in most alignment tools [14, 15] consists in a three-step method, where the CDS are first translated into AA sequences using their actual coding frame, then AA sequences are aligned, and finally the AA alignment is back-translated to a CDS alignment. This approach does not account for alternative AA translations between two CDS and it leads to incorrect alignment of the coding regions subject to FS translation. The opposite problem of aligning protein sequences while recovering their hypothetical nucleotide CDS sequences and accounting for FS translation was also studied in several papers [16, 17].

Here, we consider the problem of aligning two CDS while accounting for FS translation, by simultaneously accounting for their nucleotide and AA sequences. The problem has recently regained attention due to the increasing evidence for alternative protein production through FS translation by eukaryotic gene families [18, 19].

The problem was first addressed by Hein et al. [20, 21] who proposed a DNA/Protein model such that the score of an alignment between two CDS of length \(n\) and \(m\) is a combination of its score at the nucleotide level and its score at the AA level. They described a \(O(n^2m^2)\) algorithm in [20], later improved to a \(O(nm)\) algorithm in [21] for computing an optimal score alignment, under the constraint that the search space of the problem is restricted. Arvestad [22] later proposed a CDS alignment scoring model with a \(O(nm)\) alignment algorithm accounting for codon structures and FS translations based on the concept of generalized substitutions introduced in [23]. In this model, the score of a CDS alignment depends on its decomposition into a concatenation of codon fragment alignments, such that a codon fragment of a CDS is defined as a substring of length \(w\), \(0 \leq w \leq 5\). This decomposition into codon fragment alignments allows to define a score of the CDS alignment at the AA level. More recently, Ranwez et al. [18] proposed a simplification of the model of Arvestad limiting the maximum length of a codon fragment to 3. Under this model, a \(O(nm)\) CDS alignment algorithm was described
and extended in the context of multiple sequence alignment [18]. In the models of Arvestad [22] and Ranwez et al. [18], several scores may be computed for the same alignment based on different decompositions into codon fragment alignments. The corresponding algorithms for aligning two CDS then consist in computing an optimal score decomposition of an alignment between the two CDS. This optimal score exclusively accounts for FS translation initiations, i.e. a FS translation in an alignment is penalized by adding a constant FS cost, which only penalizes the initiation of the FS, not accounting for the length of this FS translation. However, taking account of FS translation lengths is important in order to increase the precision of CDS alignment scores, as these lengths induce more or less disruptions between the protein sequences.

In this paper, we propose the first alignment algorithm that accounts for both the initiation and the length of FS translations in order to compute the similarity scores of CDS alignments. The remaining of the paper is organized as follows. In the "Motivation" section, we illustrate the importance of accounting for FS translation length when aligning CDS. In the "Preliminaries" section, we give some preliminary definitions and we introduce a new CDS alignment scoring model with a self-contained definition of the score of an alignment penalizing both the initiation and the extension of FS translations. In the "Method" section, a dynamic programming algorithm for computing an optimal score alignment between two CDS is described. Finally, in the "Results" section, we present and discuss the results of a comparison of our method with other CDS alignment methods for a pairwise comparison of CDS from homologous genes of ten mammalian gene families.

**Motivation: Importance of accounting for FS translation length**

The two main goals of aligning biological sequences are to evaluate the similarity and to identify similar regions between the sequences, used thereafter to realize molecular analyses such as evolutionary, functional and structural predictions. In practice, CDS alignment can be used to exhaustively identify the conserved features of a set of proteins. Thus, the definition of CDS similarity must account for sequence conservation and disruptions at both the nucleotide and the protein levels.

Figure 1 illustrates the importance of accounting for AA translations and FS translation length in order to compute an adequate similarity score for a CDS alignment. It describes an example of three CDS Seq1, Seq2 and Seq3. Seq1 has a length of 45. The CDS Seq2 has length 60 and is obtained from Seq1 by deleting the nucleotide ‘G’ at position 30 and adding 16 nucleotides at the end. The CDS Seq3 has length 60 and is obtained from Seq1 by deleting the nucleotide ‘G’ at position 15 and adding 16 nucleotides at the end.

When looking at the AA translations of Seq1, Seq2 and Seq3, we observe that the similarity between Seq2 and Seq1 is higher than the similarity between Seq3 and Seq1 at the protein level, because Seq1 and Seq2 share a longer AA prefix "M T E S K Q P W H" (amino acids in black characters in the alignments). However, the pairwise CDS alignment algorithms that do not account for the length of FS translations would return the same score for the two following optimal alignments of Seq1 with Seq2 and Seq1 with Seq3, penalizing only the initiation of one
FS translation in both cases (positions marked with a "!" symbol in the alignments), and not penalizing the sequence disruptions at the protein level.

From an evolutionary point of view, a good scoring model for evaluating the similarity between two CDS in the presence of FS translations should then penalize not only the initiation of FS but also the length of FS translations extension (amino acids in gray characters in the alignments). The alignment of Seq1 with Seq2 would then have a higher similarity score than the alignment of Seq1 with Seq3.

**Preliminaries: Score of CDS alignment**

In this section, we formally describe a new definition of the score of a CDS alignment that penalizes both the initiation and the extension of FS translations.

**Definition 1 (Coding DNA sequence (CDS))**

A coding DNA sequence (CDS) is a DNA sequence on the alphabet of nucleotides \( \Sigma_N = \{A, C, G, T\} \) whose length \( n \) is a multiple of 3. A coding sequence is composed of a concatenation of \( \frac{n}{3} \) codons that are the words of length 3 in the sequence ending at positions \( 3i, 1 \leq i \leq \frac{n}{3} \). The AA translation of a CDS is a protein sequence of length \( \frac{n}{3} \) on the alphabet \( \Sigma_A \) of AA such that each codon of the CDS is translated into an AA symbol in the protein sequence.

Note that, in practice an entire CDS begins with a start codon "ATG" and ends with a stop codon "TAA", "TGA" or "TAG".

**Definition 2 (Alignment between DNA sequences)**

An alignment between two DNA sequences \( A \) and \( B \) is a pair \( (A', B') \) where \( A' \) and \( B' \) are two sequences of same length \( L \) derived by inserting gap symbols \(-1\) in \( A \) and \( B \), such
Given an alignment \((A', B')\) of length \(L\) between two CDS \(A\) and \(B\), let \(S\) be the sequence \(A'\) or \(B'\). We denote by \(S[k .. l]\), \(1 \leq k \leq l \leq L\), the substring of \(S\) going from position \(k\) to position \(l\). \(|S[k .. l]|\) denotes the number of letters in \(S[k .. l]\) that are different from the gap symbol ‘\(\_\)’. For example, if \(A' = \text{ACCAT}\text{--GTAG}\) and \(B' = \text{AC}\text{--TACGTAG}\), \(|A'[4 .. 8]| = |\text{AT}\text{--G}| = 3\). A codon of \(A\) or \(B\) is grouped in the alignment \((A', B')\) if its three nucleotides appear in three consecutive columns of the alignment. For example, the first codon \(\text{ACC}\) of \(A\) is grouped, while the first codon \(\text{ACT}\) of \(B\) is not grouped.

In the following, we give our definition of the score of an alignment \((A', B')\) between two CDS \(A\) and \(B\). It is based on a partition of the codons of \(A\) (resp. \(B\)) into four sets depending on the alignment of codons (see Figure 2 for an illustration):

1. The set of In-frame Matching codons (IM) contains the codons that are grouped in the alignment and aligned with a codon of the other CDS.
2. The set of Frameshift extension codons (FSext) contains the codons that are grouped in the alignment and aligned with a concatenation of three nucleotides that overlaps two codons of the other CDS.
3. The set of Deleted/Inserted codons (InDel) contains the codons that are grouped in the alignment and aligned with a concatenation of 3 gap symbols.
4. All other codons constitutes Frameshift initiation codons (FSinit).

The set of Matching nucleotides in FSinit codons (MFS) contains all the nucleotides belonging to FSinit codons and aligned with a nucleotide of the other CDS.

\[
\begin{align*}
\text{pos} & \quad 0000000001111111112222222233333334444444444 \\
& \quad 123456789012345678901234567890123456789012345678 \\
& \quad M \quad T \quad E \quad S \quad K \quad Q \quad P \quad W \quad P \quad D \quad Q \quad G \quad * \\
A' & \quad \text{ATGACCGAATCCAAG---CAGCCCTGCGCAG---AT---CAAGG--TCTG} \\
B' & \quad \text{ATG---GAGTCGAATATCGAC---TGG-CAGGCCATTTGGCAATGAC} \\
& \quad M \quad E \quad S \quad N \quad I \quad S \quad W \quad Q \quad A \quad I \quad G \quad N \quad D \quad * \\
\end{align*}
\]

Figure 2: An alignment \((A', B')\) of length 48 between two CDS, \(A\) (13 codons) and \(B\) (14 codons). The number arrays indicate the positions of the consecutive alignment columns. Codons of \(A\) and \(B\) are colored according to the set to which they belong: IM codons in blue color, FSext codons in red color, InDel codons in green color and FSinit codons in black color. MFS nucleotides contained in FSinit codons are underlined.

The following notations and conventions are used in Definition 3 to denote the different sets of codons and nucleotides in \(A\) and \(B\). The set of IM codons in \(A\) (resp. \(B\)) is denoted by \(\text{IM}_{A \rightarrow B}\) (resp. \(\text{IM}_{B \rightarrow A}\)). The set of FSext codons in \(A\) (resp. \(B\)) is denoted by \(\text{FSext}_{A \rightarrow B}\) (resp. \(\text{FSext}_{B \rightarrow A}\)). The set of InDel codons in \(A\) (resp. \(B\)) is denoted by \(\text{InDel}_{A \rightarrow B}\) (resp. \(\text{InDel}_{B \rightarrow A}\)). The set of MFS nucleotides in \(A\) (resp. \(B\)) is denoted by \(\text{MFS}_{A \rightarrow B}\) (resp. \(\text{MFS}_{B \rightarrow A}\)). In these sets, the codons of \(A\) and \(B\) are simply identified by the position (column) of their last nucleotide in the alignment. In this case, we always have \(\text{IM}_{A \rightarrow B} = \text{IM}_{B \rightarrow A}\) as in the example below. The MFS nucleotides are also identified by their positions in the alignment.
For example, for the alignment depicted in Figure 2, the composition of the different sets are: $\text{IM}_{A \to B} = \text{IM}_{B \to A} = \{3, 9, 12, 15, 26, 48\}; \text{FSext}_{A \to B} = \{20, 41\}; \text{InDel}_{A \to B} = \{6\}; \text{FSext}_{A \to B} = \{21, 28, 29, 30, 34, 35, 42, 43, 45\}; \text{FSext}_{B \to A} = \{21, 30, 42\}; \text{InDel}_{B \to A} = \{33\};$ and $\text{FSext}_{B \to A} = \{18, 34, 35, 39, 43, 45\}$.

In the alignment scoring model described in Definition 3, the substitutions of $\text{IM}$ and $\text{FSext}$ codons are scored using an AA scoring function $s_{\text{aa}}$ such that aligned codons with silent nucleotide mutations get the same score as identity. A fixed FS extension cost denoted by $fs_{\text{extend} \text{ cost}}$ is added for each $\text{FSext}$ codon. The insertions/deletions of $\text{InDel}$ codons are scored by adding a fixed gap cost denoted by $\text{gap \ cost}$ for each $\text{InDel}$ codon. The alignment of $\text{FS}$ nucleotides are scored independently from each other, using a nucleotide scoring function $s_{\text{aa}}$. The insertions or deletions of nucleotides in $\text{FSinit}$ codons are responsible for the initiation of FS translations. They are then scored by adding a fixed FS opening cost denoted by $fs_{\text{open} \text{ cost}}$ for each $\text{FSinit}$ codon. Note that, by convention, the values of all penalty costs for gap and FS ($\text{gap \ cost}, fs_{\text{open} \text{ cost}}, fs_{\text{extend} \text{ cost}}$) are negative. Note also that the scoring scheme assumes that the AA and the nucleotide scoring functions, $s_{\text{aa}}$ and $s_{\text{aa}}$, are symmetric.

**Definition 3 (Score of an alignment)**

Let $(A', B')$ be an alignment of length $L$ between two CDS $A$ and $B$. The score of the alignment $(A', B')$ is defined by:

\[
\text{score}(A', B') = \sum_{k \in \text{IM}_{A \to B}} s_{\text{aa}}(A'[k-2..k], B'[k-2..k]) + \\
\sum_{k \in \text{FSext}_{A \to B}} \left( s_{\text{aa}}(A'[k-2..k], B'[k-2..k]) + fs_{\text{extend} \text{ cost}} \right) + \\
\sum_{k \in \text{FSext}_{B \to A}} \left( s_{\text{aa}}(A'[k-2..k], B'[k-2..k]) + fs_{\text{extend} \text{ cost}} \right) + \\
\sum_{k \in \text{FSext}_{B \to A}} \left( s_{\text{aa}}(B'[k-2..k], A'[k-2..k]) + fs_{\text{extend} \text{ cost}} \right) + \\
\sum_{k \in \text{FSinit}_{A \to B}} \left( s_{\text{aa}}(A'[k-2..k], B'[k-2..k]) \right) + \\
\sum_{k \in \text{FSinit}_{B \to A}} \left( s_{\text{aa}}(B'[k-2..k], A'[k-2..k]) \right).
\]

**Method**

In this section, we describe a $O(nm)$ time and space complexity algorithm that solves the problem of finding a maximum score alignment between two CDS $A$ and $B$ of lengths $n$ and $m$. Similarly to other classical sequence alignment algorithms [24], we use dynamic programming tables that are indexed by the pairs of prefixes of the two CDS. The table $D$ stores the maximum scores of the alignments between prefixes of $A$ and $B$. The table $D_F$ is used to account for potential cases of FS extensions that are counted subsequently.

**Definition 4 (Dynamic programming tables)**

Given two CDS $A$ and $B$ as input, the algorithm uses two dynamic programming tables $D$ and $D_F$ of size $(n+1) \times (m+1)$. The cell $D(i, j)$ contains the maximum score of an alignment between the prefixes $A[1..i]$ and $B[1..j]$. The table $D_F$ is filled only for values
of \(i\) and \(j\) such that \(i \mod 3 = 0\) or \(j \mod 3 = 0\). If \(i \mod 3 \neq 0\) (resp. \(j \mod 3 \neq 0\)), the cell \(D_F(i, j)\) contains the score of an alignment between the prefixes \(A[1..i+a]\) and \(B[1..j+a]\) where \(a = (3-i)(\mod 3)\) (resp. \(a = (3-j)(\mod 3)\)). The table \(D_F\) is filled as follows:

- If \(i \mod 3 = 0\) and \(j \mod 3 = 0\), \(D_F(i, j) = D(i, j)\).
- If \(i \mod 3 = 0\) and \(j \mod 3 = 2\), or \(i \mod 3 = 2\) and \(j \mod 3 = 0\), \(D_F(i, j)\) contains the maximum score of an alignment between \(A[1..i+1]\) and \(B[1..j+1]\) such that \(A[i+1]\) and \(B[j+1]\) are aligned together and half of the score for aligning \(A[i+1]\) with \(B[j+1]\) is subtracted.
- If \(i \mod 3 = 0\) and \(j \mod 3 = 1\), or \(i \mod 3 = 1\) and \(j \mod 3 = 0\), \(D_F(i, j)\) contains the maximum score of an alignment between \(A[1..i+2]\) and \(B[1..j+2]\) such that \(A[i+1]\), \(B[j+1]\) and \(A[i+2]\), \(B[j+2]\) are aligned together and half of the scores of aligning \(A[i+2]\) with \(B[j+2]\) and \(A[i+1]\) with \(B[j+1]\) is subtracted.

**Lemma 1 (Filling up table D)**

1. If \(i \mod 3 = 0\) and \(j \mod 3 = 0\)

   \[
   D(i, j) = \max \begin{cases} 
   1. \quad s_{mv}(A[i-2..i], B[j-2..j]) + D(i-3, j-3) \\
   2. \quad s_{mv}(A[i, B[j]]) + s_{mv}(A[i-1, B[j-1]]) + D(i-3, j-2) + 2 \cdot fs_{open cost} \\
   3. \quad s_{mv}(A[i, B[j]]) + s_{mv}(A[i-2, B[j-1]]) + D(i-3, j-2) + 2 \cdot fs_{open cost} \\
   4. \quad s_{mv}(A[i, B[j]]) + D(i-3, j-1) + 2 \cdot fs_{open cost} \\
   5. \quad s_{mv}(A[i, B[j]]) + s_{mv}(A[i-1, B[j-1]]) + D(i-2, j-3) + 2 \cdot fs_{open cost} \\
   6. \quad s_{mv}(A[i, B[j]]) + s_{mv}(A[i-2, B[j-1]]) + D(i-2, j-3) + 2 \cdot fs_{open cost} \\
   7. \quad s_{mv}(A[i, B[j]]) + D(i-1, j-3) + 2 \cdot fs_{open cost} \\
   8. \quad s_{mv}(A[i, B[j]]) + D(i-1, j-1) + 2 \cdot fs_{open cost} \\
   9. \quad s_{mv}(A[i, B[j]]) + s_{mv}(A[i-2, B[j]]) + D(i-3, j-2) + fs_{open cost} \\
   10. \quad s_{mv}(A[i-1, B[j]]) + D(i-3, j-1) + 2 \cdot fs_{open cost} \\
   11. \quad s_{mv}(A[i-2, B[j]]) + D(i-3, j-1) + 2 \cdot fs_{open cost} \\
   12. \quad gap_{cost} + D(i-3, j) \\
   13. \quad D(i-1, j) + fs_{open cost} \\
   14. \quad s_{mv}(A[0, B[j-1]]) + s_{mv}(A[i-2, B[j]]) + D(i-2, j-3) + fs_{open cost} \\
   15. \quad s_{mv}(A[i, B[j-1]]) + D(i-1, j-3) + 2 \cdot fs_{open cost} \\
   16. \quad s_{mv}(A[i-2, B[j]]) + D(i-1, j-3) + fs_{open cost} \\
   17. \quad gap_{cost} + D(i, j-3) \\
   18. \quad D(i, j-1) + fs_{open cost} \\
   \end{cases}
   \]

2. If \(i \mod 3 = 0\) and \(j \mod 3 \neq 0\)

   \[
   D(i, j) = \max \begin{cases} 
   1. \quad s_{mv}(A[i-2, B[j]], D(i-3, j-3) + fs_{extend cost} \\
   \quad \quad + s_{mv}(A[i-1, B[j-1]]) + D(i-3, j-2) + fs_{extend cost} \\
   \quad \quad \quad + f_{j-1(\mod 3) \neq 0} \\
   2. \quad s_{mv}(A[i, B[j]]) + s_{mv}(A[i-2, B[j-1]]) + D(i-3, j-3) + fs_{extend cost} \\
   \quad \quad \quad + f_{j \mod 3 = 0} \\
   3. \quad s_{mv}(A[i, B[j]]) + s_{mv}(A[i-2, B[j-1]]) + D(i-3, j-2) + fs_{extend cost} \\
   \quad \quad \quad + f_{j \mod 3 = 0} \\
   4. \quad s_{mv}(A[i, B[j]]) + D(i-3, j-1) + fs_{extend cost} \\
   5. \quad s_{mv}(A[i, B[j]]) + D(i-1, j-3) + fs_{extend cost} \\
   6. \quad s_{mv}(A[i, B[j]]) + s_{mv}(A[i-2, B[j-1]]) + D(i-3, j-3) + fs_{extend cost} \\
   \quad \quad \quad + f_{j \mod 3 = 0} \\
   7. \quad s_{mv}(A[i, B[j]]) + D(i-3, j-1) + fs_{extend cost} \\
   8. \quad s_{mv}(A[i, B[j]]) + D(i-3, j-1) + fs_{extend cost} \\
   9. \quad gap_{cost} + D(i-3, j) \\
   10. \quad D(i-1, j) + fs_{open cost} \\
   11. \quad D(i-1, j) \\
   \end{cases}
   \]

3. If \(i \mod 3 \neq 0\) and \(j \mod 3 = 0\), the equation is symmetric to the previous case.

4. If \(i \mod 3 \neq 0\) and \(j \mod 3 \neq 0\)

   \[
   D(i, j) = \max \begin{cases} 
   1. \quad s_{mv}(A[i, B[j]]) + D(i-1, j-1) \\
   2. \quad D(i-1, j) \\
   3. \quad D(i, j-1) \\
   \end{cases}
   \]
The proof of Lemma 1 is given in the Additional file 1. Figure 3 illustrates the configurations of alignment considered in Lemma 1 for computing $D(i, j)$ for Cases 1 and 2.

**Figure 3** Illustration of the configurations of alignment considered in Lemma 1 for computing $D(i, j)$ in Cases 1 and 2. The right-most nucleotides of the sequences $A[1 .. i]$ and $B[1 .. j]$ are represented using the character $x$. The nucleotides are colored according to the type of the codon to which they belong: IM codons in blue color, FSext codons in red color, InDel codons in green color and FSinit codons in black color. The nucleotides that appear in gray color are those belonging to codons whose type has not yet been decided. In such case, the table $D_F$ is used in order to decide of the type of these codons subsequently and adjust the score accordingly.

**Lemma 2** (Filling up table $D_F$)

1. If $i \,(\text{mod } 3) = 0$ and $j \,(\text{mod } 3) = 0$

   $D_F(i, j) = D(i, j)$
If \(i \mod 3 = 2\) and \(j \mod 3 = 0\),

\[
D_f(i,j) = \max \begin{cases} 
\text{score}(A[i+1..i+3], B[j+1..j+3]) + D_f(i-1,j-1) + \text{fs_open cost}, & \text{if } i \equiv 1 \mod 3 \\
\text{score}(A[i+1..i+3], B[j+2..j+2]) + D_f(i-1,j) + \text{fs_open cost}, & \text{if } i \equiv 2 \mod 3
\end{cases}
\]

If \(i \mod 3 = 0\) and \(j \mod 3 = 2\), the equation is symmetric to the previous case.

If \(i \mod 3 = 1\) and \(j \mod 3 = 0\),

\[
D_f(i,j) = \max \begin{cases} 
\text{score}(A[i+2..i+2], B[j+1..j+1]) + D_f(i-1,j-1) + \text{fs_extend cost}, & \text{if } i \equiv 1 \mod 3 \\
\text{score}(A[i+2..i+2], B[j+2..j+2]) + D_f(i-1,j) + \text{fs_open cost}, & \text{if } i \equiv 2 \mod 3
\end{cases}
\]

If \(i \mod 3 = 0\) and \(j \mod 3 = 1\), the equation is symmetric to the previous case.

**Proof of Lemma 2.** The proof follows from Lemma 1.

1. If \(i \mod 3 = 0\) and \(j \mod 3 = 0\), this case is trivial.
2. If \(i \mod 3 = 2\) and \(j \mod 3 = 0\), then \(i + 1 \mod 3 = 0\) and \(j + 1 \mod 3 = 1 \neq 0\). The five cases follow from the application of Lemma 1, Case 2 for computing \(D(i+1,j+1)\), and by keeping only the cases where \(A[i+1]\) and \(B[j+1]\) are aligned together (cases 1, 2, 3, 4, 5 among the 11 cases). However, in each of the cases, we must subtract half of the score of aligning \(B[i+1]\) with \(A[j+1]\) \(
\frac{\text{score}(A[i+1..i+3], B[j+1..j+1])}{2}
\)

3. If \(i \mod 3 = 0\) and \(j \mod 3 = 2\), the proof is symmetric to the previous case.

4. If \(i \mod 3 = 1\) and \(j \mod 3 = 0\), then \(i + 2 \mod 3 = 0\) and \(j + 2 \mod 3 = 2 \neq 0\). Here again, the three cases follow from the application of Lemma 1, Case 2 for computing \(D(i+2,j+2)\) and by keeping only the cases where \(A[i+1], B[i+1]\) and \(A[i+2], B[i+2]\) can be aligned together (cases 1, 2, 5 among the 11 cases). However, in each of the cases, we must subtract half of the scores of aligning \(B[i+2]\) with \(A[j+2]\) and aligning \(B[i+1]\) with \(A[j+1]\) \(
\frac{\text{score}(A[i+2..i+2], B[j+2..j+2]) + \text{score}(A[i+1..i+1], B[j+1..j+1])}{2}
\)

5. If \(i \mod 3 = 0\) and \(j \mod 3 = 1\), the proof is symmetric to the previous case.

\[ \square \]

The alignment algorithm using Lemma 1 and 2 is described in the next theorem.

**Theorem 1 (Computing a maximum score alignment)**

Given two CDSs \(A\) and \(B\) of lengths \(n\) and \(m\), a maximum score alignment between \(A\) and \(B\) can be computed in time and space \(O(nm)\), using the following algorithm.

**Algorithm Align(A,B)**

```plaintext
for i = 0 to n do 
  D(i,0) = floor(\(1/3\)) * gap_cost 

D_f(i,0) = D(i,0) + \begin{cases} 
\text{score}(A[i..i+1], B[j..j+1]) + \text{fs_open cost}, & \text{if } i \equiv 1 \mod 3 \\
\text{score}(A[i..i+1], B[j+2..j+2]) + \text{fs_open cost}, & \text{if } i \equiv 2 \mod 3
\end{cases}

for j = 0 to m do 
  D(0,j) = floor(\(1/3\)) * gap_cost 

for i = 0 to n do 
  for j = 0 to m do 
    D_f(i,j) = \begin{cases} 
\text{score}(A[i+2..i+2], B[j+1..j+1]) + D_f(i-1,j-1) + \text{fs_extend cost}, & \text{if } i \equiv 1 \mod 3 \\
\text{score}(A[i+2..i+2], B[j+2..j+2]) + D_f(i-1,j) + \text{fs_open cost}, & \text{if } i \equiv 2 \mod 3
\end{cases}
```

```
Proof of Theorem 1
The proof relies on two points: (1) The algorithm computes the maximum score of an alignment between \(A\) and \(B\) and (2) the algorithm runs with an \(O(nm)\) time and space complexity.

(1) The validity of the algorithm, i.e. the fact that it fills the cells of the tables \(D\) and \(D_F\) according to Definition 4, follows from five points.

- The initialization of the tables is a direct consequence of Definition 4.
- Lemmas 1 and 2.
- The couples \((i, j)\) of prefixes of \(A\) and \(B\) that need to be considered in the algorithm are all the possible couples for \(D(i, j)\) and only the couples such that \(i (\text{mod } 3) = 0\) or \(j (\text{mod } 3) = 0\) for \(D_F(i, j)\) (see all the cases in which the table \(D_F\) is used in Lemmas 1 (7 cases) and 2 (3 cases)).
- The couples \((i, j)\) of prefixes of \(A\) and \(B\) are considered in increasing order of length and \(D[i, j]\) is computed before \(D_F[i, j]\) in the cases where \(i (\text{mod } 3) = 0\) or \(j (\text{mod } 3) = 0\).
- A backtracking of the algorithm allows to find a maximum score alignment between \(A\) and \(B\).

(2) The time and space complexity of the algorithm is a direct consequence of the number of cells of the tables \(D\) and \(D_F\), \(2 \times (n + 1) \times (m + 1)\). Each cell is filled in constant time. The exact formula for the computational complexity of the algorithm is computed below.

\[
\begin{align*}
18 & \times \frac{nm}{n} \text{ for } \frac{nm}{n} \text{ calls of the Case 1 of Lemma 1} \\
+ 11 & \times \frac{2nm}{m} \text{ for } 2 \times \frac{nm}{m} \text{ calls of the Cases 2 or 3 of Lemma 1} \\
+ 3 & \times \frac{4nm}{m} \text{ for } 4 \times \frac{nm}{m} \text{ calls of the Case 4 of Lemma 1} \\
+ 1 & \times \frac{nm}{m} \text{ for } \frac{nm}{m} \text{ calls of the Case 1 of Lemma 2} \\
+ 5 & \times \frac{2nm}{m} \text{ for } 2 \times \frac{nm}{m} \text{ calls of the Cases 2 and 3 of Lemma 2} \\
+ 3 & \times \frac{2nm}{m} \text{ for } 2 \times \frac{nm}{m} \text{ calls of the Cases 4 and 5 of Lemma 2} \\
\text{Total} & = 12.55nm
\end{align*}
\]

Results and discussion
We implemented the present CDS alignment algorithm with an affine gap penalty scheme [25] such that the penalty for a concatenation of \(k\) inserted (resp. deleted) codons is \(\text{gap\_open\_cost} + k \times \text{gap\_cost}\), such that \(\text{gap\_open\_cost}\) is a negative penalty cost for gap initiations. This was done by adding two dynamic programming tables \(G_A\) and \(G_B\) such that the cell \(G_A(i, j)\) (resp. \(G_B(i, j)\)) contains the maximum score of an alignment between the prefixes \(A[1 .. i]\) and \(B[1 .. j]\) where the codon \(A[i - 2 .. i]\) (resp. \(B[j - 2 .. j]\)) is an InDel codon.

Data
We evaluated the algorithm through applications on a mammalian dataset containing CDS sequences from ten gene families obtained from the database Ensembl-
Compara version 83 [26]. The first gene family named “FAM86” is such that three CDS from three of its paralogous human genes were shown in [6] to share a common FS region translated in three different frames in the three CDS (see Figure 4 for an illustration of the multiple alignment of these three CDS). The nine other families are the nine smallest (in term of the overall length of CDS) of fifteen gene families listed in [12] where they were shown to display one FS translation region between some pairs of CDS. For each gene family, the CDS of all human, mouse and cow genes belonging to the family and satisfying Definition 1 were downloaded. The overall number of distinct pairs of CDS within the ten gene families is 4011. Table 1 gives the details about the content and size of the ten gene families (The CDS of the ten gene families are provided in the Additional file 2).

Table 1 Detailed description of the ten gene families of the mammalian dataset.

| Gene family | Human gene | # of genes | # of CDS | Length  | $N(N−1)$ |
|-------------|------------|------------|----------|---------|----------|
| I (FAM86)   | ENSG00000118894 | 6          | 14       | 10335   | 91       |
| II (HBG017385) | ENSG00000143867 | 6          | 10       | 8988    | 45       |
| III (HBG020791) | ENSG00000179526  | 6          | 10       | 11070   | 45       |
| IV (HBG04532)  | ENSG00000173020  | 17         | 33       | 52356   | 528      |
| V (HBG016641)  | ENSG00000147041  | 13         | 33       | 64950   | 528      |
| VI (HBG014779) | ENSG00000233803  | 28         | 44       | 45813   | 946      |
| VII (HBG012748) | ENSG00000134545  | 24         | 44       | 28050   | 946      |
| VIII (HBG015928) | ENSG00000178287 | 5          | 19       | 5946    | 171      |
| IX (HBG004374)  | ENSG00000140519  | 13         | 30       | 36405   | 435      |
| X (HBG000122)   | ENSG00000150717  | 11         | 24       | 27081   | 276      |
| Total number of pairs of CDS | 4011 |

For each gene family, the family identifier used in [6] or [12], the Ensembl identifier of a human gene member of the family, the number of human, mouse and cow genes in the family, the total number of CDS of these genes, the total sum of lengths of these CDS and the number of distinct pairs of CDS are given.

Evaluation strategies

We compared the accuracy of five pairwise global alignment methods, including the present method, for computing CDS alignments in the presence or absence of FS translation between the compared CDS. The five methods vary according to the alignment algorithm used, either the present CDS alignment algorithm called FsePSA allowing to penalize both FS translation initiation and extension, or the CDS alignment algorithm called MACSE [18] penalizing FS translation initiation, or the Needleman-Wunsch (NW) sequence alignment algorithm [24] penalizing neither. Table 2 summarizes the alignment algorithm and the values of parameters used for each of the five methods.

The present CDS alignment algorithm is used in two of the five methods, namely fse and fse0. These two methods differ according to the value given to the parameter $fs\_ext\_cost$, either $fs\_ext\_cost < 0 (−1, −0.5 or −0.2)$ for the method fse penalizing FS translation extension, or $fs\_ext\_cost = 0$ for the method fse0 not penalizing FS translation extension. The pairwise version of MACSE [18] is used in the method called macae.p. The NW alignment algorithm is used in the last two methods, the method called needlenuc computing scores and alignments at the nucleotide level and the method called needleprot at the AA level. For all methods using both the amino acid and nucleotide scoring functions $s_{aa}$ and $s_{an}$, $s_{an}$ was fixed to $+1/−1$ for match/mismatch, so that the overall score of 3 consecutive nucleotide identities in an alignment scores less than the smallest identity.
score in $s_{\text{hit}}$. All other parameters shared by several methods were given the same value for all methods. In particular, for the three methods $f_{\text{s}}$, $f_{\text{se}}$ and $\text{macae}_{\text{p}}$ penalizing FS translation initiation, the parameter $f_{\text{s open cost}}$ was given the values $-10$, $-20$ or $-30$. All other parameters were fixed to the default values for the NW algorithm implementation of NCBI Blast at the nucleotide and AA levels [28].

We used the five methods to compute pairwise alignments between the pairs of CDS within each of the ten gene families of our dataset, yielding 4011 alignments in total for each of five methods. In the absence of available benchmarks for the direct evaluation of the accuracy of CDS alignments, we base our evaluation on four indirect strategies.
Table 2 Description of the five methods considered in the experiment.

| Method   | Alignment approach & specific parameters | FS initiation cost | Other parameters |
|----------|-----------------------------------------|--------------------|-----------------|
| fse      | Present approach                        | fs_open_cost = -11  | AA gap_open_cost = -11 |
|          |                                         | fs_extend_cost = 0  | AA gap_cost = -1  |
| fse0     | Present approach                        |                     | ssa = BLOSUM62 matrix |
|          |                                         |                     | sm = +1/-1        |
| macse:p  | Manveez et al. [18]                     |                     |                  |
|          |                                         | stop_cost = -100    |                  |
| needleprot| NW [24] at AA level                     | not applicable      |                  |
| needlenuc| NW [24] at NT level                     | not applicable      | NT gap_open_cost = -5 |
|          |                                         |                     | NT gap_cost = -2   |
|          |                                         |                     | sm = +2/-3        |
|          |                                         |                     | match/mismatch    |

For each method, the alignment approach and the values of specific and common parameters are given.

In the first strategy, we consider the CDS multiple alignment of each gene family obtained using MACSE [18] as a benchmark. This strategy exploits the fact that multiple alignments are usually more accurate than pairwise alignments. It then assumes that the MACSE multiple alignments are closer to the reality than the pairwise alignments obtained using the five methods. Note that all the pairwise alignment methods included in the comparison can be extended to multiple sequence alignment methods using classical strategies. Thus, the more accurate pairwise alignment methods should lead to more accurate multiple alignment methods. Here, we focus on the comparison of the pairwise versions of the methods. In the second strategy, we consider six composition criteria for a CDS pairwise alignment called Identity NT, Identity AA, Gap_init, Gap_length, FS_init, FS_length. The definitions of these criteria are given below, and used to compare the five methods. In the third strategy, we manually build and use as a benchmark, the real multiple alignment of three CDS from three paralogous human genes of the gene family I (FAM86). In the fourth strategy, we generate and use a set of three CDS splicing orthology groups, each group containing seven existing or putative CDS from seven genes of gene family I (FAM86).

Based on the results of the large-scale experiments discussed in the following, the best compromise for default values of FsePSA parameters are -30 for fs_open_cost and -1 for fs_extend_cost.

Discussion

First strategy: Using MACSE multiple alignments as benchmark

MACSE [18] was used with its default parameters (fs_open_cost = -30, stop_cost = -100, ssa = BLOSUM62 matrix, gap_open_cost = -7, gap_cost = -1) to compute the CDS multiple alignment of each of the ten gene families. For each MACSE multiple alignment of N CDS, we consider the \( \frac{N(N-1)}{2} \) induced pairwise alignments as a benchmark. In total, we then obtained a benchmark composed of 4011 pairwise alignments. In order to compare an alignment \((A', B')\) obtained with one of the five methods to the corresponding alignment \((A'', B'')\) in the benchmark, we computed the number of nucleotides aligned in \((A', B')\) with the same partner as in the benchmark alignment \((A'', B'')\).

Table 3 shows the overall percentage of nucleotides aligned with the same partners as in the benchmark for each of the compared methods, for varying
fs_open_cost (−10, −20 and −30) and fs_extend_cost (−1, −0.5 and −0.2). It shows that the different versions of the fse method and the fse0 method have the best scores greater than 79.4%, followed by the needleprot method with a score of 78.82%. On the opposite, the needleuc and macse_p method with fs_open_cost=−30 return the worst scores, respectively 50.95% and 47.35%. These results also show that the fse method is more robust to the fs_open_cost parameter changes as compared to the macse_p method, whose scores show a large variation between 47.35% and 78.29%. Note that the needleuc and needleprot do not account for the fs_open_cost parameter.
Table 3 Comparison with MACSE multiple alignments benchmark.

| fs_open_cost | fse0 | fse (-1) | fse (-0.5) | fse (-0.2) | macse | needlenuc | needleprot |
|--------------|------|----------|------------|------------|-------|-----------|------------|
| -10          | 79.58 (1404) | 79.40 (1364) | 79.55 (1415) | 79.58 (1433) | 77.17 (1076) | 50.95 (255) | 78.82 (972) |
| -20          | 79.68 (1550) | 79.66 (1526) | 79.65 (1558) | 79.67 (1552) | 78.29 (1389) |          |            |
| -30          | 79.75 (1558) | 79.47 (1529) | 79.60 (1546) | 79.63 (1547) | 47.35 (742)  |          |            |

Percentage of nucleotides aligned with the same partner as in the benchmark alignments induced by the MACSE multiple alignments, for each method for varying fs_open_cost (-10, -20 and -30) and fs_extend_cost (-1, -0.5 and -0.2). In each case, the number of CDS pairs with an alignment that presents the highest similarity with the corresponding benchmark alignment as compared to the other methods is given in parenthesis. The best results are indicated in bold.

Table 4 Values of the six criteria for the noFS dataset (variations as compared to needleprot).

| fs_open_cost (# CDS pairs) | Method   | Identity_NT | Identity_AA | Gap_init | Gap_length | FS_init | FS_length |
|---------------------------|----------|-------------|-------------|----------|------------|---------|----------|
| -10 (1672)                | fse0     | 3281 (1158) | 5376 (1222) | -495 (1606) | -2718 (1521) | 0 (1672) | 0 (1672) |
|                           | fse      | 8120 (955)  | 27942 (676) | 3701 (711)  | 9618 (1102)  | 0 (1672) | 0 (1672) |
|                           | macse_p  | 170239 (156) | -82002 (442) | 104811 (218) | 21422 (427)  | 44488 (256) | 263365 (256) |
|                           | needlenucl| 1090957 | 2047608 | 10230 | 530688 | 0 | 0 |
|                           | needleprot| 2000228 | 3494760 | 31793 | 1313658 | 0 | 0 |
| -20 (3441)                | fse0     | 1409 (2612) | -8622 (2672) | -3564 (3169) | -9984 (3057) | 0 (3441) | 0 (3441) |
|                           | fse      | 24909 (1437) | 95844 (1011) | 13787 (1076) | 30884 (1791) | 0 (3441) | 0 (3441) |
|                           | macse_p  | 177203 (176) | -177203 (680) | 317250 (219) | 12210 (552) | 138204 (257) | 644401 (257) |
|                           | needlenucl| 2000228 | 3494760 | 31793 | 1313658 | 0 | 0 |
|                           | needleprot| 2143630 | 3715632 | 352396 | 1439764 | 0 | 0 |
| -30 (3740)                | fse0     | 1368 (2834) | -10788 (2912) | -4047 (3448) | -11316 (3321) | 0 (3740) | 0 (3740) |
|                           | fse      | 27840 (1547) | 106516 (1076) | 15561 (1117) | 34726 (1846) | 0 (3740) | 0 (3740) |
|                           | macse_p  | 613035 (177) | -102231 (709) | 351748 (219) | 47356 (573) | 154255 (257) | 948418 (257) |
|                           | needlenucl| 2143630 | 3715632 | 352396 | 1439764 | 0 | 0 |
|                           | needleprot| 2143630 | 3715632 | 352396 | 1439764 | 0 | 0 |

For varying values of the parameter fs_open_cost, the number of CDS pairs in the dataset is given. The values of the criteria for the reference method “needleprot” are indicated in bold characters. For each of the other methods (fse, fse0, macse_p, needlenucl), the variations of the criteria values as compared to the reference values are given. For each criteria and each method, the number of CDS pairs that have the closest value to the reference needleprot value is given in parentheses.
**Second strategy: Using six composition criteria for CDS pairwise alignment**

Six criteria were defined and used to compare the five pairwise alignment methods. Given a pairwise CDS alignment, the first criterion $\text{Identity}_{\text{NT}}$ counts the number of gap-free columns in the alignment containing a nucleotide match. The second criterion $\text{Identity}_{\text{AA}}$ counts the number of $\text{IM}$ and $\text{FSext}$ codons $c$ in the alignment that are aligned with a triplet of nucleotides yielding the same amino acid as $c$. The third criterion $\text{Gap}_{\text{init}}$ is the number of gap-containing columns in the alignment, either insertion or deletion columns that are preceded by a different type of column. The fourth criterion $\text{Gap}_{\text{length}}$ is the overall number of gap-containing columns in the alignment. The fifth criterion $\text{FS}_{\text{init}}$ is the number of FS translation segments found in the alignment. The last criterion $\text{FS}_{\text{length}}$ is the overall number of columns in the alignment intersecting a $\text{FSext}$ codon.

Note that the definitions of the six criteria exploit the definitions of codon sets used in Definition 3 but they are independent of any alignment scoring scheme. For example, for the alignment depicted in Figure 2, $\text{Identity}_{\text{NT}} = 28$, counting all gap-free columns except the five columns at the positions $\{9, 12, 15, 42, 45\}$ containing a nucleotide mismatch. $\text{Identity}_{\text{AA}} = 14$, counting all $\text{IM}$ and $\text{FSext}$ codons except the two $\text{IM}$ codons $\text{AAG}$ and $\text{AAT}$ ending at position 15 yielding two different amino acids $K$ and $N$, and the $\text{FSext}$ codon $\text{AAT}$ ending at position 42 yielding the amino acid $N$ different from the amino acid $K$ yielded by the triplet $\text{AAG}$. $\text{Gap}_{\text{init}} = 7$, counting the positions $\{4, 16, 22, 27, 31, 36, 44\}$. $\text{FS}_{\text{init}} = 3$, counting the positions $\{18, 28, 39\}$. The two last criteria have the values $\text{Gap}_{\text{length}} = 15$ and $\text{FS}_{\text{length}} = 11$.

For each of the nine cases obtained by combining the values of the parameters $\text{fs}_{\text{open\_cost}} (-10, -20 \text{ or } -30)$ and $\text{fs}_{\text{extend\_cost}} (-1, -0.5 \text{ or } -0.2)$, we considered the 4011 pairs of CDS from the ten gene families dataset, and partitioned them into three sets. For each case, the first set called the noFS dataset is composed of the pairs of CDS for which the pairwise alignments obtained using the $\text{fse0}$, $\text{fse}$ and $\text{macse\_p}$ methods all have the criteria $\text{FS}_{\text{init}} = 0$. The second set called the FS dataset is composed of the pairs of CDS for which the alignments obtained using the $\text{fse0}$, $\text{fse}$ and $\text{macse\_p}$ methods all have the criteria $\text{FS}_{\text{init}} > 0$. The third set called the ambiguFS dataset is composed of the remaining pairs of CDS.

Note that, in all nine cases, the set of CDS pairs for which $\text{FS}_{\text{init}} = 0$ with the $\text{macse\_p}$ method was strictly included in the set of CDS pairs for which $\text{FS}_{\text{init}} = 0$ with the $\text{fse}$ method. For each of the nine cases, we computed the overall value of the six criteria for each method ($\text{fse0}$, $\text{fse}$, $\text{macse\_p}$, $\text{needlenuc}$ and $\text{needleprot}$) and each dataset (noFS, FS and ambiguFS). Tables 4, 5 and 6 present the results.

**Results for the noFS datasets.** For the noFS datasets, we assume that the real alignments should not contain FS translations. So, the $\text{needleprot}$ method most likely computes the more accurate alignments since it does not allow any FS translation in the alignments. Indeed, it computes a maximum score NW alignment at the AA level and back-translates this alignment at the nucleotide level. We then take the $\text{needleprot}$ result as a reference for the noFS dataset, in all cases. By construction of the noFS dataset, for a fixed value of the parameter $\text{fs\_open\_cost}$, the $\text{fse0}$ and $\text{fse}$ methods necessarily return two alignments with the same similarity...
score for each pair of CDS of the dataset. Indeed, we observed that, for each value of \textit{fs\_open\_cost} (\(-10, -20\) or \(-30\)), the alignments obtained using the methods \textit{fse0} or \textit{fse} with varying values of the parameter \textit{fs\_extend\_cost} are unchanged.

Table 4 summarizes the results for \textit{fs\_open\_cost} = \(-10, -20\) and \(-30\), presenting the results of the varying versions of \textit{fse} and \textit{fse0} in a single line in the 3 cases. It shows that the results of the \textit{fse} and \textit{fse0} methods are the closest to the reference for all the six criteria in all cases. However, they slightly overestimate or underestimate the criteria. The tendency of overestimating the Identity\_AA and all other criteria is particularly accentuated for the \textit{macse} method as compared to the \textit{fse} and \textit{fse0} methods, in all cases. On the opposite, the \textit{needlenuc} method always largely underestimates the Identity\_AA, while overestimating all other criterion.

**Results for the FS datasets.** For the FS datasets, we assume that the real alignments must contain FS translations. So, the \textit{needleprot} method can no longer produce the most accurate results. On the contrary, it is most likely that it underestimates the Identity\_AA criterion. Indeed, it correctly aligns AA in CDS regions that are free of FS translation, but in FS translation regions, it either leads to several AA mismatches in the case of high mismatches scores, or to an overestimation of the Gap\_init criterion. As expected, we observed that the value of Identity\_AA for the \textit{needleprot} method was always the lowest (data shown in the Additional file 3). We focus on the four other methods.

Table 5 summarizes the results for the nine cases considered. For the Identity\_NT and Identity\_AA criteria, the differences between the values for the four methods are negligible. The main differences between the results reside in the values of the Gap\_init and FS\_init criteria. In particular, the FS\_init criterion is useful to compare the accuracy of the methods for correctly identifying real FS translation regions. In [6] (for family I) and [12] (for families II to X), at most one FS translation region was detected and manually validated for each pair of CDS of the ten gene families. So, the expected number of FS translation regions per alignment in the FS data is 1. In Table 5, we observe that, in all cases, the \textit{fse} and \textit{fse0} methods are the only methods for which the average numbers of FS\_init are close to 1 with +/- standard error values smaller than 1. The \textit{macse} method and especially the \textit{needlenuc} method overestimate the number of FS translation regions per alignment with large standard error values in all cases.

**Results for the ambiguFS datasets.** For the ambiguFS datasets, all methods do not agree for the presence or absence of FS translation regions between the pairs of CDS. Note that the \textit{needlenuc} method reports FS translations for all pairs of CDS, with the highest average number of FS translation regions per alignment in all cases (data shown in the Additional file 3). As \textit{needlenuc} is already shown to perform poorly in both the absence and the presence of FS translation regions, we focus on the four other methods. Table 6 summarizes the results. We observe that, for all criteria, \textit{macse}\_p has higher values than \textit{fse0}, \textit{fse} and \textit{needleprot} that have similar values. The most significant difference between the results resides in the values for the FS\_init and FS\_length criteria. The \textit{fse} method always reports a null or a very small number of FS regions with an average FS\_init equals to 1 as expected. In all cases, the \textit{fse0} and \textit{macse} methods overestimate the number of FS translation regions per alignment.
### Table 5: Values of the six criteria for the FS dataset.

| fs.open_cost | fs.extend_cost (# CDS pairs) | Method | Identity_(init) | Identity_(NT) | Gap_length | FS_init (avg) | FS_length | Gap_init |
|--------------|-------------------------------|--------|-----------------|---------------|------------|--------------|-----------|----------|
|              |                               | fse0   | 156002          | 32512         | 895        | 60662       | 326 (1.00 ± 0.25) | 60219    |
|              |                               | fse    | 185720          | 32026         | 901        | 60624       | 216 (1.01 ± 0.14) | 18705    |
| -1 (212)     |                               | macse_p| 156167          | 324999        | 1445       | 61562       | 432 (2.03 ± 0.36) | 22742    |
|              |                               | needlenc| 172599         | 321348        | 5053       | 60038       | 2103 (9.91 ± 2.67) | 29616    |
| -10          |                               | fse0   | 252590          | 464172        | 2400       | 114859      | 482 (1.24 ± 0.47) | 31777    |
|              |                               | fse    | 251047          | 405387        | 2387       | 115269      | 401 (1.03 ± 0.19) | 29842    |
|              |                               | macse_p| 255715          | 405394        | 4161       | 117165      | 1306 (3.38 ± 4.53) | 41742    |
|              |                               | needlenc| 279862         | 452673        | 19408      | 113195      | 6832 (20.80 ± 31.02) | 68226    |
| -0.5 (386)   |                               | fse0   | 371062          | 641748        | 5334       | 204370      | 805 (1.30 ± 0.52) | 43381    |
|              |                               | fse    | 370560          | 640377        | 5270       | 204806      | 688 (1.11 ± 0.33) | 37376    |
|              |                               | macse_p| 374729          | 646893        | 9308       | 208344      | 2893 (4.67 ± 5.34) | 72030    |
|              |                               | needlenc| 442564         | 618270        | 48799      | 209420      | 19751 (31.90 ± 34.48) | 141217   |
| -0.2 (619)   |                               | fse0   | 133814          | 244350        | 461        | 40315       | 166 (1.04 ± 0.29) | 17770    |
|              |                               | fse    | 133610          | 243149        | 468        | 40195       | 164 (1.01 ± 0.14) | 16924    |
|              |                               | macse_p| 133541          | 243591        | 709        | 40585       | 223 (1.38 ± 0.13) | 18119    |
|              |                               | needlenc| 132545         | 242742        | 1493       | 39031       | 650 (4.03 ± 3.85) | 19405    |
| -20          |                               | fse0   | 147476          | 290147        | 549        | 49485       | 197 (1.04 ± 0.20) | 19599    |
|              |                               | fse    | 147401          | 290148        | 557        | 49383       | 194 (1.02 ± 0.16) | 19279    |
|              |                               | macse_p| 147143          | 290271        | 838        | 49841       | 260 (1.37 ± 0.96) | 19976    |
|              |                               | needlenc| 149551         | 288086        | 1872       | 47515       | 808 (4.27 ± 6.17) | 21440    |
| -0.2 (216)   |                               | fse0   | 161906          | 318117        | 723        | 55383       | 225 (1.04 ± 0.20) | 21300    |
|              |                               | fse    | 161885          | 318099        | 732        | 55393       | 223 (1.03 ± 0.18) | 21115    |
|              |                               | macse_p| 161822          | 317205        | 1061       | 55715       | 306 (1.41 ± 0.99) | 21997    |
|              |                               | needlenc| 165260         | 315531        | 2851       | 53613       | 1186 (5.49 ± 6.82) | 24403    |
| -1 (161)     |                               | fse0   | 128014          | 279641        | 440        | 39803       | 148 (1.03 ± 0.28) | 13845    |
|              |                               | fse    | 128030          | 279870        | 443        | 39864       | 148 (1.03 ± 0.28) | 13845    |
|              |                               | macse_p| 128030          | 279870        | 443        | 39864       | 148 (1.03 ± 0.28) | 13845    |
|              |                               | needlenc| 128030         | 279870        | 443        | 39864       | 148 (1.03 ± 0.28) | 13845    |
| -0.5 (189)   |                               | fse0   | 120558          | 237678        | 445        | 37975       | 159 (1.03 ± 0.18) | 17554    |
|              |                               | fse    | 120504          | 237678        | 452        | 37851       | 157 (1.01 ± 0.14) | 17319    |
|              |                               | macse_p| 120336          | 237684        | 691        | 38047       | 222 (1.37 ± 0.96) | 17926    |
|              |                               | needlenc| 120284         | 236604        | 1321       | 37531       | 575 (3.73 ± 5.14) | 18877    |
| -0.2 (216)   |                               | fse0   | 137451          | 271041        | 525        | 46049       | 184 (1.03 ± 0.18) | 18995    |
|              |                               | fse    | 137440          | 271008        | 531        | 45917       | 183 (1.02 ± 0.17) | 18972    |
|              |                               | macse_p| 137175          | 270258        | 803        | 46187       | 244 (1.37 ± 0.97) | 18935    |
|              |                               | needlenc| 139489         | 290399        | 1803       | 44303       | 779 (4.38 ± 6.52) | 20859    |

For varying values of the parameters fs.open_cost and fs.extend_cost, the number of CDS pairs in the dataset is given. The values of the criteria for the fse, fse0, macse_p, needlenc methods are indicated. For each method, the average number of FS.init per alignment, with corresponding standard error values are also indicated.
Table 6: Values of the six criteria for the ambiguFS dataset.

| fs_open_cost | fs_extend_cost (# CDS pairs) | Method | Identity_NT | Identity_AA | Gap_init | Gap_length | FS_init (avg) | FS_length |
|--------------|-----------------------------|--------|-------------|------------|----------|------------|--------------|-----------|
| -10          | -1 (2127)                   | fse0 (862)          | 1095102     | 1737105    | 24469    | 906218     | 1111 (1.28 ± 0.54) | 42730     |
|              |                             | fse     | 1085946     | 1719774    | 23483    | 906540     | 0            | 0         |
|              |                             | macse_p (2076)     | 1124316     | 1790199    | 45335    | 936002     | 12436 (5.99 ± 4.76) | 216772    |
|              |                             | needleprot        | 1085007     | 1723950    | 25288    | 916518     | 0            | 0         |
| -0.5 (1953)  | -1 (2127)                   | fse0 (688)          | 1009851     | 1597605    | 22984    | 854021     | 855 (1.24 ± 0.53)  | 31172     |
|              |                             | fse     | 1003293     | 1587258    | 22102    | 853793     | 2 (1.0 ± 0.0)    | 80        |
|              |                             | macse_p (1902)     | 1036768     | 1649004    | 42019    | 880399     | 11562 (6.07 ± 4.91) | 197772    |
|              |                             | needleprot        | 1001357     | 1591134    | 23990    | 863199     | 0            | 0         |
| -0.2 (1720)  | -1 (2127)                   | fse0 (455)          | 890042      | 1420569    | 20050    | 764510     | 532 (1.36 ± 0.53)  | 19568     |
|              |                             | fse     | 887372      | 1415403    | 19465    | 764162     | 3 (1.0 ± 0.0)    | 92        |
|              |                             | macse_p (1669)     | 915754      | 1468305    | 37472    | 789220     | 9975 (5.97 ± 4.75) | 167484    |
|              |                             | needleprot        | 886178      | 1418748    | 20955    | 772272     | 0            | 0         |
| -20          | -1 (409)                    | fse0 (100)          | 219277      | 358554     | 3633     | 153487     | 120 (1.2 ± 0.40)  | 6937      |
|              |                             | fse     | 216936      | 353586     | 3619     | 152391     | 0            | 0         |
|              |                             | macse_p (403)      | 223976      | 374391     | 6009     | 156165     | 1348 (3.34 ± 3.00) | 36179     |
|              |                             | needleprot        | 216842      | 355656     | 4172     | 153957     | 0            | 0         |
| -0.5 (381)   | -1 (409)                    | fse0 (72)           | 195615      | 311757     | 3545     | 144317     | 91 (1.26 ± 0.44)  | 5108      |
|              |                             | fse     | 194048      | 308448     | 3505     | 144045     | 0            | 0         |
|              |                             | macse_p (375)      | 202374      | 327711     | 6380     | 146909     | 1311 (3.49 ± 3.05) | 34322     |
|              |                             | needleprot        | 193980      | 310632     | 4051     | 145563     | 0            | 0         |
| -0.2 (354)   | -1 (409)                    | fse0 (45)          | 181185      | 284787     | 3371     | 138419     | 63 (1.4 ± 0.49)  | 3407      |
|              |                             | fse     | 180151      | 285933     | 3344     | 138217     | (1.0 ± 0.0)    | 40        |
|              |                             | macse_p (348)      | 187895      | 300777     | 6157     | 143035     | 1265 (3.63 ± 3.11) | 32301     |
|              |                             | needleprot        | 180116      | 284946     | 3883     | 139731     | 0            | 0         |
| -30          | -1 (200)                    | fse0 (119)          | 151090      | 289617     | 805      | 42434      | 120 (1.01 ± 0.09) | 6818      |
|              |                             | fse     | 147509      | 282018     | 852      | 40221      | 0            | 0         |
|              |                             | macse_p (200)      | 152626      | 292254     | 1309     | 43404      | 378 (1.89 ± 2.16) | 14515     |
|              |                             | needleprot        | 147228      | 281472     | 933      | 40455      | 0            | 0         |
| -0.5 (117)   | -1 (200)                    | fse0 (36)          | 77603       | 143115     | 590      | 30765      | 37 (1.07 ± 0.16) | 2109      |
|              |                             | fse     | 76678       | 141108     | 626      | 29913      | 0            | 0         |
|              |                             | macse_p (117)      | 79224       | 146046     | 990      | 31321      | 284 (2.42 ± 2.65) | 9731      |
|              |                             | needleprot        | 76561       | 141036     | 703      | 30099      | 0            | 0         |
| -0.2 (93)    | -1 (200)                    | fse0 (12)          | 60770       | 109842     | 510      | 22091      | 12 (1.0 ± 0.0)  | 668       |
|              |                             | fse     | 60407       | 109170     | 518      | 22491      | 0            | 0         |
|              |                             | macse_p (93)       | 62387       | 112872     | 878      | 23181      | 252 (2.70 ± 2.89) | 8262      |
|              |                             | needleprot        | 60270       | 109122     | 581      | 22677      | 0            | 0         |

For varying values of the parameters fs_open_cost and fs_extend_cost, and for each method, the number of CDS pairs displaying a FS translation is given. The values of the criteria for each method are indicated. For each method, the average number of FS_init per alignment, with corresponding standard error values are also indicated.
Third strategy: Using a 3-CDS manually-built benchmark

We manually built the real pairwise alignments of three CDS from three paralogous human genes of gene family I, the CDS FAM86C1-002 coding for protein ENSP00000352182.4, FAM86B1-001 coding for protein ENSP00000431362.1 and FAM86B2-202 coding for protein ENSP00000311330.6. The real multiple alignment of the three CDS is roughly depicted and detailed in Figure 4. From Figure 4, we observe that FAM86C1-002 shares with FAM86B1-001 a nucleotide region of length 159 translated in the same frame and a nucleotide region of length 89 with FS translation, while it only shares with FAM86B2-202 a nucleotide region of length 238 (89 + 149) entirely under FS translation. It is then clear that CDS FAM86C1-002 and FAM86B1-001 are the most similar. Figure 4 also shows that each pair of CDS shares a single FS translation region.

Table 7 shows the normalized pairwise similarity scores and the number of FS translation regions computed by the five alignment methods (the pairwise alignments computed by the five methods with varying fs_open_cost and fs_extend_cost are given in the Additional file 4). It shows that needleprot and fse (in all cases where fs_extend_cost = -1) are the only two methods that allow to infer that FAM86C1-002 and FAM86B1-001 are the most similar. Table 7 also illustrates the fact that needlenuc and macse_p strongly overestimate the number of FS translation regions per alignment in all cases. The fse method with the parameters fs_open_cost = -10 and fs_extend_cost = -1 is the only method that allows to infer that FAM86C1-002 and FAM86B1-001 are the most similar and to detect a single FS translation region for each alignment.

**Table 7** Pairwise similarity scores and number of FS translation regions computed by the methods.

| fs_open_cost | Method  | C1-002 vs B1-001 | C1-002 vs B2-202 | B1-001 vs B2-202 |
|-------------|---------|-----------------|-----------------|-----------------|
| -10         | fse0    | 0.42 (1)        | 0.58 (2)        | 0.45 (1)        |
|             | fse (-1) | 0.34 (1)        | 0.27 (1)        | 0.19 (1)        |
|             | fse (-0.5) | 0.37 (1)      | 0.43 (1) | 0.31 (1)        |
|             | fse (-0.2) | 0.40 (1)      | 0.52 (1) | 0.39 (1)        |
|             | macse_p  | 0.40 (4)        | 0.54 (6)        | 0.44 (1)        |
| -20         | fse0    | 0.39 (1)        | 0.54 (1)        | 0.41 (1)        |
|             | fse (-1) | 0.36 (0)        | 0.24 (1)        | 0.14 (1)        |
|             | fse (-0.5) | 0.39 (1)      | 0.39 (1) | 0.28 (1)        |
|             | fse (-0.2) | 0.37 (1)      | 0.48 (1) | 0.36 (1)        |
|             | macse_p  | 0.54 (4)        | 0.47 (6)        | 0.59 (1)        |
| -30         | fse0    | 0.35 (1)        | 0.50 (1)        | 0.38 (1)        |
|             | fse (-1) | 0.36 (0)        | 0.20 (1)        | 0.11 (1)        |
|             | fse (-0.5) | 0.36 (0)      | 0.35 (1) | 0.25 (1)        |
|             | fse (-0.2) | 0.33 (1)      | 0.44 (1) | 0.33 (1)        |
|             | macse_p  | 0.27 (4)        | 0.39 (6)        | 0.29 (1)        |
|             | needlenuc | 0.16 (23)    | 0.35 (15)       | -0.36 (1)       |
|             | needleprot | 0.38 (0)     | -0.12 (0)       | -0.13 (0)       |

Normalized pairwise similarity scores and number of FS translation regions computed by the five methods for the 3-CDS manually-built benchmark composed of CDS FAM86C1-002, FAM86B1-001 and FAM86B2-202 (Similarity scores are normalized by dividing them by the lengths of alignments).

Fourth strategy: Inferring CDS splicing orthology groups and protein phylogenies

Based on the three CDS used in the previous strategy, CDS FAM86C1-002 from human gene ENSG00000158483, FAM86B1-001 from human gene ENSG00000186523 and FAM86B2-202 from human gene ENSG00000145002, we generated a dataset
of three CDS splicing orthology groups composed of 21 homologous CDS. Each
group contains one of the three initial CDS and its six splicing orthologs in the
following set of seven genes from gene family I: human genes ENSG00000158483 denoted H1, ENSG00000186523 denoted H2 and ENSG00000145002 denoted H3,
each containing one of the initial CDS, chimpanzee gene ENSPTRG00000007738 denoted Ch, mouse gene ENSMUSG00000022544 denoted M, rat gene ENSRNOG0-
0000002876 denoted R and cow gene ENSBTAG00000008222 denoted Co. The CDS
splicing orthologs were predicted based on the spliced alignment tool Splign [29]
as follows: for each initial CDS A1 of a gene A and each gene B different from A,
A1 was aligned to B and a putative or existing CDS of B ortholog to A1 with the
same splicing structure was inferred. The 21 resulting CDS are given in Additional
file 5.

We computed the normalized pairwise similarity scores between the CDS, us-
ing the five alignment methods (the pairwise alignments computed by the five
methods with varying f_open_cost and f_extend_cost are given in the Addi-
tional file 5). For each method, we constructed a phylogeny using an UPGMA and
a Neighbor-Joining (NJ) algorithm, based on the computed CDS similarity matrix.
The UPGMA algorithm was used to classify the CDS into three groups and infer
the similarity relationships between the groups independently of any rate of evo-
lution. The NJ algorithm was used to reconstruct the phylogeny inside each group.
Table 8 summarizes the results. The three splicing orthology groups are denoted
G1 (containing CDS C1-002), G2 (containing CDS B1-001) and G3 (containing CDS
B2-202).

All methods allow to correctly classify the CDS into the three initial splic-
ing orthology groups G1, G2, and G3. However, the needleprot and fse meth-
ods are the only methods that allow to infer the correct similarity relationships
((G1,G2),G3) between the groups, confirming the results of the third evaluation
strategy. For all methods, the CDS phylogeny reconstructed inside the group G2
is (Co,((M,R),((H1,Ch),(H2,H3)))) inducing an evolution of the seven genes with a
speciation event at the root of the gene tree. The phylogeny reconstructed for the
groups G1 and G3 is ((M,R),(Co,((H1,Ch),(H2,H3)))), inducing an evolution of the
genes with a duplication event at the root of the phylogeny.

Comparing of the running times
Table 9 shows the running times for each of the five methods on the three first
gene families of our dataset on a 24 × 2.1GHz processor with 10GB of RAM. The
needleprot method is the fastest, followed by macse_p and then needlenuc, while
fse and fse0 are the slowest methods.

Note that for fse, fse0, needlenuc and needleprot, the used implementations
are in Python, while we used a JAVA implementation for macse_p provided by its
authors. This explains the fact that macse_p is unexpectedly faster here than fse,
fse0, and even needlenuc. Indeed, the five methods share the same asymptotic
time complexity, but the exact complexity of each of them is dependent on the
number of calls of the main recurrence formulas in an execution, and the number of
cases considered in each recurrence formula. The exact computational complexity
of the five methods in terms of the lengths n and m of two compared CDS are
Table 8 Similarity relationships between the groups G1, G2 and G3 for the five methods.

| fs_open_cost | Method | ((G1,G3),G2) | ((G1,G2),G3) |
|--------------|--------|--------------|--------------|
| -10          | fse (-1) | X            |              |
|              | fse (-0.5) | X            |              |
|              | fse (-0.2) | X            |              |
|              | fse0    | X            |              |
|              | macse_p | X            |              |
| -20          | fse (-1) | X            |              |
|              | fse (-0.5) | X            |              |
|              | fse (-0.2) | X            |              |
|              | fse0    | X            |              |
|              | macse_p | X            |              |
| -30          | fse (-1) | X            |              |
|              | fse (-0.5) | X            |              |
|              | fse (-0.2) | X            |              |
|              | fse0    | X            |              |
|              | macse_p | X            |              |
|               | needleprot | X            |              |

Similarity relationships between the splicing orthology groups G1, G2 and G3 computed using the similarity matrices of the five methods for the 21-CDS dataset.

12.55 × nm for fse and fse0 (as shown in the proof of Theorem 1), 15 × nm for macse_p, 3 × nm for needlenuc and 0.33 × nm for needleprot.

Table 9 Running time in seconds for each method.

| Gene family | fse0 | fse | macse_p | needlenuc | needleprot |
|-------------|------|-----|---------|-----------|-----------|
| I           | 299  | 291 | 53      | 97        | 22        |
| II          | 270  | 260 | 45      | 93        | 20        |
| III         | 377  | 389 | 54      | 62        | 20        |

For each method and gene families I, II, and III, the running time was calculated on the same computer (24 processors of 2.1GHz each and 10GB of RAM) with the parameters fs_open_cost=−20 and fs_extend_cost=−0.2.

Conclusions

In this paper, we introduce a new scoring model for the alignment of CDS accounting for frameshift translation length. The motivation for this new scoring scheme is the increasing evidence for protein divergence through frameshift translation in eukaryotic coding gene families, calling for automatic methods able to compare, align and classify CDS while accounting for their codon structure. The aim of this paper is to validate the necessity of accounting for frameshift translation length when comparing CDS and show that computing a maximum score pairwise alignment under the new scoring scheme is possible in quadratic time complexity. The results of comparing five CDS alignment methods for the pairwise alignment of CDS from ten eukaryotic gene families show that our method is the best compromise for sets of CDS in which some pairs of CDS display FS translations while some do not. Future work will make use of benchmarks of CDS alignments generated manually and by simulation in order to confirm these experimental results. We also defer to a future work the extended study of our model’s robustness to parameter changes and the calibration of its parameters using real data benchmarks.

The perspectives of this work also include the design of a heuristic algorithm using local alignment that will achieve scalability for large datasets while keeping high accuracy, and the extension of the method toward multiple alignment. Finally, we plan to apply the algorithms for the discovery of non-annotated frameshifts, and the evaluation of the extent of frameshifts in eukaryotic gene families.
Availability of supporting data
An implementation of the pairwise alignment method in Python is available at
https://github.com/UdeS-CoBIUS/FsePSA. The dataset used in section Results is available in the Additional files.

List of abbreviations
CDS: Coding DNA Sequence; FS: Frameshift; NT: nucleotide; AA: amino acid; NW: Needleman-Wunsch.

Declarations

Author's contributions
SJ, EK, FB and AO wrote the program and its documentation. SJ and AO conceived the study and its design. SJ, EK and AR ran the experiments. SJ, EK and AO analyzed and interpreted the data. SJ and AO wrote the manuscript. SJ, EK, MS and AO critically revised the manuscript. All authors read and approved the final manuscript.

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References
1. Zambelli, F., Pavesi, G., Gissi, C., Horner, D.S., Pesole, G.: Assessment of orthologous splicing isoforms in human and mouse orthologous genes. BMC genomics 11(1), 1 (2010)
2. Barbosa-Morais, N.L., Irimia, M., Pan, Q., Xiong, H.Y., Gueroussov, S., Lee, L.J., Slobodeniuc, V., Kutter, C., Watt, S., Çolak, R., et al.: The evolutionary landscape of alternative splicing in vertebrate species. Science 338(6114), 1587–1593 (2012)
3. Christinat, Y., Moret, B.M.: A transcript perspective on evolution. IEEE/ACM Transactions on Computational Biology and Bioinformatics (TCBB) 10(6), 1403–1411 (2013)
4. Kuitche, E., Lafond, M., Ouangraoua, A.: Reconstructing protein and gene phylogenies by extending the framework of reconciliation. To appear in Proceedings of International Conference on Bioinformatics and Computational Biology (BICOB) (2017). arXiv preprint arXiv:1610.09732
5. Pruitt, K.D., Harrow, J., Harte, R.A., et al.: The consensus coding sequence (CCDS) project: Identifying a common protein-coding gene set for the human and mouse genomes. Genome Research 19(7), 1316–1323 (2009)
6. Okamura, K., Feuk, L., Marquis-Bonet, T., Navarro, A., Scherer, S.W.: Frequent appearance of novel protein-coding sequences by frameshift translation. Genomics 88(6), 690–697 (2006)
7. Barmak, M., Christopher, L.: A genomic view of alternative splicing. nature genetics 30, 13–19 (2003)
8. Stoffers, D., Zinkin, N., Stanojevic, V., Clarke, W., Habener, J.: Pancreatic agenesis attributable to a single nucleotide deletion in the human ipf1 gene coding sequence. nature genetics 15(1), 106–110 (1997)
9. IKUO, Y., YUICHI, M., HISAO, S., YOSHIFUMI, H., SHUJI, I., YOSHIKOE, M., NOBUO, M., YUTAKA, O.: Nucleotide deletion resulting in frameshift as a possible cause of complete thyroxine-binding globulin deficiency in six japanese families. nature genetics 71(2), 262–267 (1991)
10. Robin, K.: On programmed ribosomal frameshifting: the alternative proteomes. Front Genet 3(242), 1–10 (2012)
11. Wei, S., Valerie, B., Jonathan, S., Mary, K., Frank, M., John, M., Claudia, S., Natalia, V., Alexander, L., Robert, S., John, C.: Analysis of 454 sequencing error rate, error sources, and artifact recombination for detection of low-frequency drug resistance mutations in hiv-1 dna. Retrovirology 10(18), 1–16 (2013)
12. Raes, J., Van de Peer, Y.: Functional divergence of proteins through frameshift mutations. Trends in Genetics 21(8), 428–431 (2005)
13. Ogura, Y., Bonen, D., Inohara, N., Nicolae, D., Chen, F., Ramos, R., Britton, H., Moran, T., Karaliuskas, R., Duerr, R., Aichler, J., Brant, S., Bayless, T., Kirschner, B., Hanauer, S., Nunez, G., Cho, J.: A frameshift mutation in NOD2 associated with susceptibility to Crohn’s disease. Nature 411, 601 (2001)
14. Abascal, F., Zardoya, R., Telford, M.J.: TranslatorX: multiple alignment of nucleotide sequences guided by amino acid translations. Nucleic Acids Research, 291 (2010)
15. Morgenstern, B.: DIALIGN: multiple DNA and protein sequence alignment at BiBiServ. Nucleic Acids Research 32(suppl 2), 33–36 (2004)
16. Girdea, M., Noé, L., Kucherov, G.: Back-translation for discovering distant protein homologies in the presence of frameshift mutations. Algorithms for Molecular Biology 5(1), 1 (2010)
17. Moreira, A., Maass, A.: TIP: protein backtranslation aided by genetic algorithms. Bioinformatics 20(13), 2148–2149 (2004)
18. Ranwez, V., Harispe, S., Delsuc, F., Douzery, E.J.: MACSE: Multiple Alignment of Coding SEquences accounting for frameshifts and stop codons. PLoS One 6(9), 22594 (2011)
19. Danny, B., Catherine, L., Cynthia, B., Guillaume, T., Julie, M., Xavier, R.: An out-of-frame overlapping reading frame in the ataxin-1 coding sequence encodes a novel ataxin-1 interacting protein. THE JOURNAL OF BIOLOGICAL CHEMISTRY 288(30), 21824–21835 (2013)
20. Hein, J.: An algorithm combining DNA and protein alignment. Journal of Theoretical Biology 167(2), 169–174 (1994)
21. Pedersen, C.N., Lyngsø, R., Hein, J.: Comparison of coding DNA. In: Combinatorial Pattern Matching, pp. 153–173 (1998). Springer
22. Arvestad, L.: Aligning coding DNA in the presence of frame-shift errors. In: Combinatorial Pattern Matching, pp. 180–190 (1997). Springer
23. Sankoff, D., Kruskal, J.B.: Time warps, string edits, and macromolecules: the theory and practice of sequence comparison. Reading: Addison-Wesley Publication, 1983, edited by Sankoff, David; Kruskal, Joseph B. 1 (1983)
24. Needleman, S.B., Wunsch, C.D.: A general method applicable to the search for similarities in the amino acid sequence of two proteins. Journal of Molecular Biology 48(3), 443–453 (1970)
25. Altschul, S.F., Erickson, B.W.: Optimal sequence alignment using affine gap costs. Bulletin of mathematical biology 48(5-6), 603–616 (1986)
26. Cunningham, F., Amode, M.R., Barrell, D., et al.: Ensembl 2015. Nucleic Acids Research 43(D1), 662–669 (2015)
27. Gouy, M., Guindon, S., Gascuel, O.: Seaview version 4: a multipurpose graphical user interface for sequence alignment and phylogenetic tree building. Molecular biology and evolution 27(2), 221–224 (2010)
28. Johnson, M., Zaretzky, I., Raytserlis, Y., Merezhuk, Y., McGinnis, S., Madden, T.L.: NCBI BLAST: a better web interface. Nucleic Acids Research 36(suppl 2), 5–9 (2008)
29. Kapustin, Y., Souvorov, A., Tatusova, T., Lipman, D.: Splign: algorithms for computing spliced alignments with identification of paralogs. Biology direct 3(1), 20 (2008)

Additional Files
Additional file 1 – Proof of Lemma 1
PDF file containing the detailed proof of Lemma 1.

Additional file 2 – CDS of the ten gene families
Zip file containing the CDS files at the fasta format for each of the ten gene families considered in the Results section.

Additional file 3 – Additional lines for Tables 5 and 6
PDF file containing additional lines for Tables 5 (for needleprot) and 6 (for needlenc) of the Results section.

Additional file 4 – Pairwise alignments for the 3-CDS benchmark
Zip file containing the sequence file and the pairwise alignment files at the fasta format for the manually-built 3-CDS benchmark considered in the Results section, for each of the five methods and each parameter configuration.

Additional file 5 – Pairwise alignments for the 21-CDS dataset
Zip file containing the sequence file and the pairwise alignment files at the fasta format for the 21-CDS benchmark considered in the Results section, for each of the five methods and each parameter configuration.