Protein structure alignment is a fundamental problem in computational structure biology. Many programs have been developed for automatic protein structure alignment, but most of them align two protein structures purely based upon geometric similarity without considering evolutionary and functional relationship. As such, these programs may generate structure alignments which are not very biologically meaningful from the evolutionary perspective. This paper presents a novel method DeepAlign for automatic pairwise protein structure alignment. DeepAlign aligns two protein structures using not only spatial proximity of equivalent residues (after rigid-body superposition), but also evolutionary relationship and hydrogen-bonding similarity. Experimental results show that DeepAlign can generate structure alignments much more consistent with manually-curated alignments than other automatic tools especially when proteins under consideration are remote homologs. These results imply that in addition to geometric similarity, evolutionary information and hydrogen-bonding similarity are essential to aligning two protein structures.
CLESUM is better than spatial proximity in that it favors the alignment of conserved loop regions especially those appearing at the two ends of a regular secondary structure segment and that it also disfavors the alignment of two evolutionarily-unrelated alpha helices. In addition to evolutionary distance, we also make use of hydrogen-bonding similarity to improve alignments for hydrogen bonds, which greatly helps generate biologically more meaningful alignments especially for beta-strands.

It is challenging to optimize a scoring function to find the optimal structure alignment. Most search algorithms can only solve scoring functions to suboptimal and thus, generate a suboptimal alignment, which may impact alignment accuracy. Along with the enlargement of the Protein Data Bank (PDB), it is now time-consuming to retrieve all the remotely-related protein structures in the PDB for a query protein structure. Therefore, a structure alignment tool shall also be computationally efficient. Different from other structure alignment methods, DeepAlign reduces its running time by carefully identifying a set of initial alignments. In particular, DeepAlign identifies evolutionarily-related structural fragment pairs using both amino acid and local substructure mutation scores and then build initial alignments based upon these fragment pairs. Starting from the initial alignments, DeepAlign iteratively optimizes the scoring function and finally produces high-quality alignments. By using the evolutionary information, DeepAlign can accurately identify a very small set of evolutionarily-related structural fragment pairs which are very likely contained in the optimal alignment. This greatly helps reduce the running time of DeepAlign since it avoids starting from many initial alignments that cannot lead to the optimal alignments.

We evaluate DeepAlign using several metrics including reference-independent and dependent alignment accuracy, evolutionary scores and superfamily/fold discrimination, based upon three manually-curated benchmarks and also SABMark. Experimental results indicate that DeepAlign outperforms several popular tools on all the datasets when judged by human-curated alignments. DeepAlign works particularly well when proteins under consideration are not very close. The DeepAlign alignments also have better mutation scores and thus, are more biologically meaningful from the evolutionary perspective. Even evaluated by the pure geometric measures such as TMscore and uGDT (un-normalized Global Distance Test), DeepAlign still compares favorably to other popular tools such as DALI, MATT and TMalign, all of which align protein structures based upon only geometric similarity.

**Results**

We evaluate our program DeepAlign using three manually-curated alignment databases and a few metrics: LALI (length of alignment), RefAcc (reference-dependent alignment accuracy), RMSD (root mean squared deviation), TMscore and mutation scores (i.e., BLOSUM and CLESUM). RefAcc is calculated as the percentage of correctly-aligned positions as judged by the gold standard (i.e., manual alignments), measuring consistency between automatic alignments and human-curated alignments. RMSD and TMscore, both of which are well-established, measure geometric similarity of protein structures, but TMscore is better than RMSD because TMscore is length-independent and not biased by few badly-aligned residue pairs. The evolutionary scores measure if one alignment is favorable or not from the evolutionary perspective. We calculate the evolutionary scores of a structure alignment using both the BLOSUM and CLESUM substitution matrices. Ideally, a good alignment shall have preferable performance regardless of the metrics.

**The programs to be compared.** We compare DeepAlign with several popular structure alignment tools such as DALI, TMalign, MATT, Format, which represents four very different methods. DALI is a distance matrix based approach, aligning two structures by matching their distance matrices. TMalign aligns two protein structures by maximizing the TMscore, using a similar search algorithm as STRUCTAL. MATT aligns protein structures by concatenating the alignments of some short structural fragments. MATT also calculates a p-value to indicate the degree to which two proteins are structurally similar. Format is an extension of MATT by taking into consideration secondary similarity in aligning two structures.

**The benchmarks.** We use three manually-curated benchmarks: (i) A subset of CDD (Conserved Domain Database) used in; (ii) MALIDUP; and (iii) MALISAM. The CDD set contains 3591 manually-curated pairwise structure alignments. The human-curated alignments for CDD contain only the alignments of core residues. The CDD set has already been used to evaluate a bunch of pairwise structure alignment algorithms, including CE, FAST, LOCK2, MATRAS, VAST and SHEBA. MALIDUP has 241 manually-curated pairwise structure alignments for homologous domains originated from internal duplication within the same polypeptide chain. About half of the pairs in MALIDUP are remote homologs. MALISAM contains 130 protein pairs and the two proteins in any pair are structural analogs with different SCOP folds. There is strong evidence indicating that proteins in a MALIDUP pair are not homologs. Therefore, MALIDUP are the most challenging benchmark among these three. The alignments in these three databases are manually-curated, taking into consideration not only geometric similarity, but also evolutionary and functional relationship. Therefore, the manually-curated alignments make more biological sense and it is reasonable to use them as reference to judge automatically-generated alignments.

**Performance on CDD.** DeepAlign obtains the highest reference-dependent alignment accuracy of 93.8% among the five automatic structure alignment methods (in Table 1). DeepAlign also outperforms the methods evaluated in terms of ref-dependent alignment accuracy. That is, DeepAlign is more consistent with human experts than the other programs. In terms of TMscore and RMSD, the TMalign alignments are slightly better than the DeepAlign alignments, but the former are less consistent with manual alignments than the latter. This implies that the geometric similarity score used by TMalign (i.e., TMscore) does not accurately reflect the alignment criteria used by human experts. The DeepAlign alignments also have much better evolutionary scores than the other three programs no matter how the mutation scores are calculated. As a control, we also calculate the evolutionary scores of the manual alignments. The manual alignments have much lower mutation score per alignment because only core residues are aligned. However, the manual alignments have the best average mutation scores per aligned position. Note that the manual alignments are not explicitly driven by a specific mutation score. This confirms that human experts indeed take into consideration evolutionary relationship in aligning two protein structures and that TMalign may align many more evolutionarily-unrelated residues together than DeepAlign. Format has a similar mutation (i.e., BLOSUM/CLESUM) score per alignment as DeepAlign, but Format has a better average mutation score per aligned position than DeepAlign because Format has a smaller LALI. In terms of reference-independent or dependent alignment accuracy, DeepAlign is much better than Format partially because the latter has a much smaller LALI.

**Performance on MALIDUP.** DeepAlign obtains a reference-dependent alignment accuracy of 92%, greatly exceeding the other three tools (in Table 1). DeepAlign is 6% better than the second best algorithm DALI. Although the TMalign alignments have a longer alignment length and the MATT alignments have a smaller RMSD, both TMalign and MATT have much lower reference-dependent alignment accuracy. This again implies that the TMalign and MATT scoring functions greatly deviates from what are implicitly
Assessment of Structure Prediction to evaluate the quality of a SCIENTIFIC REPORTS Performance on MALISAM Formatt performs similarly on this dataset as on CDD. many more evolutionarily unfavorable residues than DeepAlign. on average are longer, this again confirms that TMalign may align mutation score per aligned position. Since the TMalign alignments only second to the Formatt alignments in terms of the average the DeepAlign alignments have much better evolutionary scores, slightly second to TMalign, but better than the others. However, used by human experts. In terms of TMscore, DeepAlign is only slightly second to TMalign, but better than the others. However, the DeepAlign alignments have much better evolutionary scores, only second to the Formatt alignments in terms of the average mutation score per aligned position. Since the TMalign alignments on average are longer, this again confirms that TMalign may align many more evolutionarily unfavorable residues than DeepAlign. Formatt performs similarly on this dataset as on CDD. Performance on MALISAM. DeepAlign obtains the highest ref-dependent alignment accuracy of 77.5% among all the five computer programs (in Table 1). DeepAlign is 10% better than the second best algorithm DALI. MALISAM is much more challenging than CDD, MALIDUP and MALISAM. In MALISAM, 80 pairs (i.e., 61.5% of the total) contain proteins with different SCOP folds31. The DALI and TMalign alignments have similar average alignment lengths as the DeepAlign alignments, but slightly higher RMSD. Furthermore, the DALI and TMalign alignments have similar average mutation scores per aligned position while the DeepAlign alignments have much better evolutionary scores, slightly second to TMalign, which aligns two protein structures by optimizing only TMscore. The manual alignments tend to have better mutation scores but much lower geometric similarity score (e.g., TMscore) partially because human experts tend to align only those evolutionarily conserved residues instead of spatially close residues (after rigid-body transformation). By contrast, DeepAlign achieves a good balance between geometric similarity and evolutionary conservation.

| Method     | LALI | RMSD | TMscore | RefAcc | Blosum1 | Clesum1 | Blosum2 | Clesum2 |
|------------|------|------|---------|--------|---------|---------|---------|---------|
| CCD [3591] |      |      |         |        |         |         |         |         |
| DeepAlign  | 134.8| 2.86 | 0.667   | 93.8   | 0.261   | 1.782   | 43.45   | 243.71  |
| DALI       | 130.8| 2.75 | 0.663   | 92.8   | 0.165   | 1.684   | 28.78   | 225.15  |
| MATT       | 128.6| 2.53 | 0.655   | 91.4   | 0.152   | 1.728   | 30.19   | 229.59  |
| Formatt    | 112.3| 2.32 | 0.566   | 86.4   | 0.343   | 1.983   | 44.11   | 235.64  |
| TMalign    | 138.4| 2.84 | 0.686   | 85.6   | 0.047   | 1.531   | 15.25   | 211.88  |
| Manual     | 62.6 | 1.66 | 0.345   | 100.0  | 0.677   | 2.499   | 43.89   | 157.67  |
| MALIDUP [241] |     |      |         |        |         |         |         |         |
| DeepAlign  | 85.5 | 2.61 | 0.622   | 92.0   | 0.314   | 1.872   | 29.31   | 158.28  |
| DALI       | 83.5 | 2.65 | 0.600   | 86.4   | 0.172   | 1.700   | 18.63   | 147.53  |
| MATT       | 82.3 | 2.47 | 0.608   | 79.8   | 0.178   | 1.824   | 18.84   | 150.00  |
| Formatt    | 70.6 | 2.19 | 0.542   | 86.2   | 0.344   | 2.196   | 28.62   | 154.66  |
| TMalign    | 87.0 | 2.62 | 0.631   | 81.0   | 0.110   | 1.600   | 12.50   | 137.64  |
| Manual     | 77.9 | 2.49 | 0.587   | 100.0  | 0.294   | 1.853   | 27.67   | 154.81  |
| MALISAM [130] |    |     |         |        |         |         |         |         |
| DeepAlign  | 61.3 | 2.96 | 0.521   | 77.5   | −0.601  | 1.108   | −36.48  | 67.66   |
| DALI       | 61.0 | 3.11 | 0.515   | 67.7   | −0.595  | 0.925   | −35.52  | 56.28   |
| MATT       | 56.2 | 2.74 | 0.486   | 51.7   | −0.625  | 1.013   | −34.05  | 56.98   |
| Formatt    | 44.9 | 2.42 | 0.411   | 56.3   | −0.486  | 1.489   | −21.1   | 65.69   |
| TMalign    | 61.1 | 3.06 | 0.517   | 53.7   | −0.684  | 0.739   | 40.04   | 45.65   |
| Manual     | 56.7 | 2.92 | 0.488   | 100.0  | −0.556  | 1.240   | 31.58   | 70.75   |

In summary, our method DeepAlign performs significantly better than other popular tools when judged by manually-curated alignments and from the evolutionary perspective. Even if only geometric similarity measures such as TMscore and uGDT are considered, DeepAlign still compares favorably to other tools, only slightly second to TMalign, which aligns two protein structures by optimizing only TMscore. The manual alignments tend to have better mutation scores but much lower geometric similarity score (e.g., TMscore) partially because human experts tend to align only those evolutionarily conserved residues instead of spatially close residues (after rigid-body transformation). By contrast, DeepAlign achieves a good balance between geometric similarity and evolutionary conservation.

Discrimination of distant homologs and structural analogs. We use SABmark to test the performance of DeepAlign in identifying distant homologs and structural analogs27. SABmark-sup is the superfamily set in SABmark (version 1.65), containing 425 protein groups with low to intermediate sequence identity. SABmark-tw is the twilight set in SABmark, containing 209 groups with low sequence identity. Each SABmark-sup (-twi) group contains at most 25 structures sharing a SCOP superfamily (fold). It is believed that if two proteins are in the same SCOP superfamily, it is likely these two proteins are remote homologs. If two proteins share only the same SCOP fold, it is very likely that they are structural analogs instead of remote homologs. Given a protein structure, we align it to all the proteins in the benchmarks and then rank all the alignments by certain criteria. We examine if the top-ranked protein structures are in the same group as the query protein or not. DeepAlign uses its scoring function to rank the proteins. Similarly, TMalign, MATT and DALI use TMscore, P-value and Z-score to rank the alignments, respectively. The ranking results are evaluated by ROC (receiver operator curve) and AUC (area under curve). Formatt has a very similar result as MATT.

As shown in Figure 1(A), tested on SABmark-sup, DeepAlign has the best ROC curve, especially at the high specificity area. For
example, at the specificity level 0.99, DeepAlign has sensitivity around 0.4 while the other three have sensitivity only around 0.2. As shown in Table 2, DeepAlign, DALI, MATT and TMalign have AUCs 0.970, 0.956, 0.933 and 0.960, respectively. We also observe the same trend on the SABmark-twii set. SABmark-twii is more challenging because each group in this set consists of proteins similar at only SCOP fold level. However, DeepAlign outperforms others by an even larger margin. As shown in Figure 1(B), DeepAlign has sensitivity 0.6 at specificity 0.96 while the second best algorithm DALI has sensitivity only 0.4 at the same specificity level. MATT and TMalign have only sensitivity around 0.2 at this specificity level. As shown in Table 2, DeepAlign, DALI, MATT and TMalign have AUCs 0.933, 0.908, 0.873 and 0.903 on SABmark-twii, respectively. These results imply that DeepAlign scoring function is better than DALI’s Z-score, MATT’s P-value and TMscore in detecting the superfamily relationship of proteins.

Relative importance of the four score items. Our scoring function consists of four items: BLOSUM, CLESUM, the hydrogen-bonding score and TMscore. In order to estimate the impact of each item on structure alignment, we exclude each of them from the DeepAlign scoring function and regenerate the alignments for the protein groups in the three manually-curated benchmarks using the new scoring functions. Table 3 illustrates the ref-dependent alignment accuracy (RefAcc) that can be obtained by the four new scoring functions.

As shown in Table 3, when BLOSUM is excluded, the RefAcc on relatively easy benchmarks (i.e., CDD and MALIDUP) is decreased while that on the more challenging benchmark MALISAM is improved. This implies that evolutionary information at the amino acid level is useful for the alignment of homologous proteins, but may slightly hurt that of structurally analogous proteins. When CLESUM is excluded, the performance on CDD is slightly improved while on the two more challenging benchmarks MALIDUP and MALISAM dramatically decreased. That is, CLESUM is particularly useful for the alignment of structurally analogous proteins and remote homologs. Table 3 also shows that the TMscore and the hydrogen bonding score are useful across all similarity levels.

The hydrogen-bonding score has large impact on beta-containing proteins. For example, tested on the 94 beta-containing protein pairs in MAILSAM each of which contains a large portion of beta-strands, the overall accuracy is only 73.7% when the hydrogen-bonding score is not used. This is significantly worse than what can be obtained (79.3%) when the hydrogen-bonding score is used. Further, among the 334 CDD beta-containing protein pairs, using the hydrogen-bonding score can improve the alignment for 212 protein pairs by at least 5%.

In summary, the 3D geometric similarity (e.g., TMscore) is the major factor determining if two proteins are similar or not. However, in order to generate biologically meaningful structure alignments, other factors are also indispensable. The local structure similarity is especially important for distantly-related proteins while the hydrogen-bonding score is important for beta-containing proteins.

Specific examples. Case study 1: d1h99a1 and d1h99a2. Both domains are in the same SCOP family (a.142.1.1) and they are mutants. Table S2 shows the evaluation of the structural alignments of these two domains generated by DeepAlign, DALI, MATT, Formatt and TMalign. DeepAlign, MATT and Formatt generate almost the same alignment as the human-curated and the alignments also have very good mutation scores. Although the DALI and TMalign alignments have slightly higher TMscore, but they have much worse mutation scores, which indicate that these alignments contain some

**Table 2** | The AUCs obtained by DeepAlign, DALI, MATT/Formatt and TMalign on SABmark-sup and SABmark-twii

| Method     | SABmark-sup | SABmark-twii |
|------------|-------------|--------------|
| DeepAlign  | 0.970       | 0.933        |
| DALI       | 0.956       | 0.908        |
| Formatt    | 0.934       | 0.874        |
| MATT       | 0.933       | 0.873        |
| TMalign    | 0.960       | 0.903        |

**Table 3** | Impact of the four score items on alignment. The numbers in the table are the reference-dependent alignment accuracy (RefAcc). “All” indicates all score items are used. “—” indicates one score item is excluded.

| Dataset | All | BLOSUM | CLESUM | Hydrogen | TMscore |
|---------|-----|--------|--------|----------|---------|
| CDD     | 93.8 | 92.1   | 94.2   | 92.2     | 75.7    |
| MALIDUP | 92.0 | 90.7   | 87.3   | 89.5     | 78.3    |
| MALISAM | 77.5 | 78.5   | 65.4   | 74.4     | 48.1    |
evolutionarily unfavorable residue pairs. DALI misaligns one helix, as shown in Figure S1. TMalign misaligns one helix either. Figure S1 displays the DeepAlign and DALI alignments, but not the TMalign alignment due to space limit.

Case study 2: d1neck_ and d1nekd_. These two domains are taken from the CDD database (ID: cd03493). Table S3 shows the evaluation of the structural alignments generated by different alignment tools for these two domains. The DeepAlign and DALI alignments are highly consistent with the human-curate alignment. MATT (as well as Format) mistakenly aligns the 1st and 2nd helices and the linker loop region, as shown in Figure S2. The DeepAlign and DALI alignments have very good TMscore, only slightly second the TMalign alignment which is generated by optimizing only TMscore. The DeepAlign alignment also has much better mutation scores than the MATT alignment, which even has a negative BLOSUM score. Overall, the MATT alignment has bad values in three performance metrics including TMscore, RefAcc and mutation scores.

Case study 3: d1ef5a_ and d1nda_. in the CDD database (ID: cd00196). DALI, MATT, Formatt and TMalign misalign the loop region between the 1st alpha helix and the 3rd beta strand. MATT and TMalign also mistakenly align the 3rd beta strand (see Figure S3). Table S4 shows the evaluation of the structural alignments for these two domains. The DeepAlign alignment has the best ref-dependent alignment accuracy (RefAcc), better than DALI and much better than MATT and TMalign. In terms of TMscore, DeepAlign is slightly worse than DALI and TMalign, but much better than MATT. In terms of the CLESUM scores, DeepAlign significantly outperforms the other three programs. According to the BLOSUM scores, DeepAlign is not very different from the others. This is because that the two domains are not close homologs and BLOSUM is not suitable to measure their evolutionary relationship. Since BLOSUM does not work for this case, we can calculate the sequence profile score at each position for the sub-alignments between the 38-th and the 53-th residues of d1ef5a_, which corresponds to the region between the 1st alpha helix and the 3rd beta strand (inclusive), as shown in Table S5. The protein sequence profile and the corresponding profile score is generated using the HHpred package. The profile score usually ranges between -1.0 and 1.0. The higher the profile score, the better. Note that profile score is not used in the DeepAlign scoring function, but the DeepAlign alignment has positive profile scores at almost all the aligned positions. By contrast, the other three alignments have negative profile scores at some positions. This further confirms that DeepAlign aligns evolutionarily-related residues together even if the proteins are not close homologs.

Running time analysis. We measure the running time on an Ubuntu Linux PC with 2 GB RAM and Intel®Core™2 Quad CPU T5600 @ 1.83 GHz. The performance and running time of DeepAlign depend on three parameters TopK, TopJ and M. DeepAlign is run using default parameters (i.e., TopK = 100, TopJ = 20, M = 10), which are also used to generate the alignments discussed in section RESULTS. Tested on the 23074 protein pairs in SABmark-twi, DeepAlign, TMalign, MATT, Formatt and DALI have running times of 1878, 1073, 29192, 35138 and 54297 seconds, respectively. That is, DeepAlign is much faster than MATT, Formatt and DALI, but (not much) slower than TMalign. DeepAlign is slower than TMalign partially because that DeepAlign uses a scoring function of four items while TMalign uses only one of them (i.e., TMscore). That is, it takes a much longer time for DeepAlign to calculate its scoring function than TMalign.

Discussion
This paper has presented a novel method DeepAlign for automatic protein structure alignment, which can generate alignments highly consistent with manually-curated alignments. Manually-curated alignments usually make much more biological sense since they are built by human experts taking into consideration evolutionary and functional relationship, in addition to geometric similarity. The novelty of DeepAlign lies in its scoring function, which considering not only 3D geometric similarity, but also evolutionary information at the sequence and local substructure levels as well as hydrogen-bonding similarity. Note that the DeepAlign scoring function is the natural combination of four different items and there are no parameters to be fine-tuned. Therefore, we do not bias DeepAlign towards a specific performance metric.

We have tested DeepAlign with four widely-used structure alignment tools TMalign, MATT, Formatt and DALI on three manually-curated benchmarks. These benchmarks contain many distantly-related protein structures (which are remote homologs or structural analogs) and are very challenging for any automatic alignment tools. Many proteins in the most challenging benchmark MALISAM are not even in the same SCOP fold although they share structurally analogous motifs. Nevertheless, DeepAlign can still align two protein structures consistently with human experts. DeepAlign also tends to align many more evolutionarily-related residues together than the other tools and open much fewer gaps in an alignment. In addition, DeepAlign compares favorably to other tools when evaluated by purely geometric similarity measures such as TMscore and uGDT. DeepAlign is also better than the other tools in discriminating two proteins similar at the superfamily or fold level.

Currently DeepAlign uses only the local conformation and the amino acid substitution matrices to measure equivalence of two residues. We can also use other information such as sequence profile, represented by a position-specific scoring matrix or Hidden Markov Model, to measure evolutionary distance, which shall be more sensitive than BLOSUM62. DeepAlign uses dynamic programming to generate a complete alignment and thus, cannot generate sequence order-independent alignments. We will extend our algorithm to deal with non-sequential alignments. We will also extend DeepAlign to some other applications such as binding-site recognition and protein-interface alignment.

Methods
The scoring function. The scoring function is used to determine how likely two residues, of the two proteins under consideration, shall be aligned. Our scoring function is composed of amino acid mutation score, local substructure substitution potential, hydrogen-bonding similarity and geometric similarity. In particular, the equivalence of two residues i and j is estimated by the following scoring function.

\[ \text{Score}(i,j) = (\max(0, \text{BLOSUM}(i,j)) + \text{CLESUM}(i,j)) \times v(i,j) \times d(i,j) \]  

(1)

Meanwhile, BLOSUM and CLESUM measure the evolutionary distance of two proteins at the sequence and local substructure levels, respectively. BLOSUM is the widely-used amino acid substitution matrix BLOSUM62. CLESUM is the local structure substitution matrix. \( v(i,j) \) measures the hydrogen-bonding similarity.
and \(d(i,j)\) measures the spatial proximity of two aligned residues (after rigid-body superposition).

In addition to the BLOSUM62 substitution matrix, other matrices (e.g., PAM250) can also be used to measure the evolutionary distance of two proteins at the sequence level. The \(d(i,j)\) function in Eq. (1) is used to handle the situation where two proteins to be aligned are distantly-related. In this case, we will only rely on CLESUM to measure the evolutionary distance. In the future we may use sequence profile similarity to measure evolutionary distance, which usually is more sensitive than BLOSUM matrices. One issue of using sequence profiles lies in that it is time-consuming to generate sequence profiles.

CLESUM is the substitution matrix for the 17 local conformation letter defined in \(\text{C} \alpha \text{b}\). Each letter represents one typical \(\text{C} \alpha \text{b}\) conformation of a 4-mer protein segment. The 17 representative local conformations are generated by clustering all the 4-mer structural fragments in a subset of non-redundant protein structures. CLESUM is derived from a representative set of pairwise structure alignments in the FSSP database. CLESUM disfavors the match of two unrelated helices but favors the alignment of two evolutionarily related loop regions. Loop regions are usually harder to align than alpha helices and beta strands if only spatial proximity is used in the scoring function.

In Eq. (1), \(d(i,j)\) is the spatial proximity of two aligned (or equivalent) residues after the rigid-body superimposition. Here we use TMscore to calculate \(d(i,j)\), which is defined as follows.

\[
d(i,j) = \frac{1}{1 + (|A_i - B_j| / d_0)}
\]

Where \(A_i\) and \(B_j\) are the (transformed) 3D coordinates of the two \(\text{C} \alpha\) atoms and \(d_0 = 1.24 \times \sqrt{L} \times 1.5 - 1.8\) is a length-dependent normalization factor, which is used to offset the impact of protein length. TMscore is a widely-used measure in the field of protein structure prediction that has demonstrated excellent performance in identifying structurally similar proteins. Generally speaking, when two protein structures have a TMscore larger than 0.6, it is highly likely they have similar folds. As shown in Eq. (2), an aligned fragment pair (AFP) shall not be too short (say less than 4 residues). However, since our scoring function does not explicitly penalize gap openings, the resultant alignment may have more gap openings than desirable. To deal with this, we use some heuristics to merge one very short AFP (less than 4 residues) to its neighboring AFPs to reduce the number of gap openings.

**Availability.** DeepAlign is available at http://ttic.uchicago.edu/~jinbo/software.htm.

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**Author contributions**

S.W. conceived, designed and implemented the algorithm. J.M., J.P. and J.X. helped design the algorithm and interpret the results. S.W., J.M. and J.X. wrote the manuscript. All authors read and approved the final manuscript.

**Additional information**

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