Clustering of common structured RNA domains by aligning basepair probabilities

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Abstract: RNA molecules often function by their RNA secondary structure. Many short RNAs have characteristic

global structures, such as the two sequential stem-loop structures of HACA snoRNAs. Long noncoding RNAs (lncRNAs) have important local structured RNA domains that are, for instance, recognized by RNA binding proteins. The discovery of functional RNA domains demands fast and accurate clustering approaches that are based on the structure landscape of RNA sequences. The gold standard of comparative RNA analysis, namely Sankoff-style simultaneous alignment and folding, is, however, not applicable. Here, we present a novel algorithm for RNA structure alignments that we named <code>DotAligner</code>. The presented method optimizes the alignment of two RNA sequences with respect to all possible structures simultaneously, which allows the alignment of similar structured RNA domains of low sequence similarity. The generated dissimilarity matrices are used to cluster RNA sequences in common structured RNA domains. The method was successfully tested on a selected set of HACA-box snoRNAs. It can contribute to the detection of functional RNA domains

on a selected set of HACA-box snoRNAs. It can contribute to the detection of functional RN, through its speed and very high specificity.

1 INTRODUCTION

The structure of RNA molecules is an essential functional criteria of many non-coding RNAs (ncRNAs), such as the stem-loop of microRNAs and the double stem-loop RNA motifs of the HOTAIR long ncRNA (Gupta et al., 2010). NcRNAs can be devided in RNA families of similar inherent functionality, structures, or composition. The largest collection of RNA families is the Rfam database with 2,208 families in its version 11.0 (Burge et al., 2013). However, highthroughut sequencing continuously uncovers novel non-coding RNA transcripts and genome-wide RNA structure predictions have revealed hundreds of thousands putative conserved RNA secondary structures. We hypothesize that the RNA secondary structure is the scaffold for intermolecular interactions of many ncRNA-driven regulatory pathways. Protein binding domains of RNA molecules may evolve totally independent from sequence and, instead, may be solely determined by structure. It has been shown that if the sequence similarity falls below 60% sequence comparison will not find anymore domain similarities that are based on structure (Gardner et al., 2005). In addition, competing structures and suboptimal structures

may support or even drive the functionality of an RNA domain. Hence, methods are needed that find structural similarity independent from sequence conservation and freed from one single optimal RNA secondary structure.

For clustering of RNA domains a dissimilarity measurement of all pairs of query structures is needed. The dissimilarity is described through a pairwise weighted string alignment with arbitrary pairwise dependencies (for base pairings). Needleman-Wunsch (2) algorithm solves the maximum weight string alignment problem by dynamic programming in $O(n^2)$ by preserving the sequence order and maximizing the similarity. The consideration of pairs of nucleotides in each sequence that form intra-molecular interactions extends the problem to pairwise dependencies among positions in each string. This problem variant is MAX-SNP-hard. However, the problem can be attacked by intelligent heuristics that avoid the examination of all possible aligning states.

Simultaneous alignment and folding (Sankoff, 1985) is the acknowledged gold standard to predict the consensus structure and alignment of a set of related RNA sequences. Because the Sankoff algorithm

is practically not applicable, the pre-calculation of the structure ensemble of each sequence, e.g. basepair probabilities in thermodynamically equilibrated RNA structure ensembles (McCaskill, 1990), is used by different methods to speed up the calculation of structure-based alignments. The programs pmcomp for pairwise and pmmulti for multiple alignments (Hofacker et al., 2004), as well as LocaRNA (Will et al., 2007) score the alignment based on the notion of a common secondary structure. Despite of the usage of the basepair probability matrices extract these methods the maximum-weight common secondary structure but do not explicitely consider suboptimal structures in the alignment. The pairwise alignment of basepair probability matrices (dot plots) has been first introduced by CARNA (Palù et al., 2010; Sorescu et al., 2012). CARNA finds iterativelly better alignments with an effective constraint programming technique using a branch and bound scheme (propagator).

Beside of LocaRNA and a method based on directed acyclic graph kernels (Sato et al., 2008), the alignment-free approach ClustGraph (Heyne et al., 2012) has been used to cluster RNA structure in common domains. Here, we propose an alternative heuristic for the pairwise weighted string alignment with arbitrary pairwise dependencies that can deliver dissimilarity scores of dot plots in time close to an Needleman-Wunsch alignment which makes the approach applicable for clustering of large numbers of putative RNA domains.

2 IMPLEMENTATION

The proposed algorithm makes the computationally complex problem of aligning two dotplots available by the iteration through a two step approach: (1) find dissimilarity (distance) of basepair probabilities of each nucleotide in sequence S_a to each nucleotide in sequence S_b ; and (2) find best path through the distance matrix generated in 1. The iteration is stopped when no more gaps are introduced in step 2. This algorithm runs in $O(n^2)$, hence, in the same time complexity as the sequence-based alignments. In the following we discuss in detail how the algorithm works.

As described in (Palù et al., 2010) the weight Z of alignment A of two arc-annotated sequences (S_a, P_a) and (S_b, P_b) is defined by

$$Z(A) = \sum_{(i,i') \in A} \sigma(i,i') + \sum_{\substack{(i,j) \in P_a, \\ (i',j') \in P_b, \\ (i,i') \in A, (j,j') \in A}} \tau(i,j,i',j') + \gamma \times L,$$
(1)

where S is a sequence and P is a base pairing probability matrix, $\sigma(i,i')$ is the similarity of sequence positions $S_a[i]$ and $S_b[i']$, $\tau(i,j,i',j')$ is the similarity of arcs $(i,j) \in P_a$ and $(i',j') \in P_b$, and γ is the gap cost associated with each sequence position that is not matched $(L = |S_a| + |S_b| - 2|A|)$. The alignment problem finds the maximal Z(A). As its solution is MAX-SNP-hard we implemented a heuristic of the alignment problem in DotAligner which is summarized in the following pseudocode:

Pseudocode

Get Alignment A of the two dotplots P_a and P_b

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Require: (S_a, P_a) of length N, (S_b, P_b) of length M repeat \{STEP\ 1:\ global\ alignment\ of\ pairing\ probabilities\ of\ each\ base\ in\ <math>S_a\ and\ S_b\} for i=1 to N do for i'=1 to M do Z(A|i,i')=\kappa\times\sigma(i,i')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i,j)\in P_a,\ (j,j')\in A}}\tau(i,j,i',j')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in P_b,\ (j,j')\in A}}\tau(i,j,i',j')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in P_b,\ (j,j')\in A}}\tau(i,j,i',j')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in P_b,\ (j,j')\in A}}\tau(i,j,i',j')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in P_b,\ (j,j')\in A}}\tau(i,j,i',j')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in A}}\tau(i,j,i',i')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in A}}\tau(i,j,i',i')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in A}}\tau(i,j,i',i')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in A}}\tau(i,j,i')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in A}}\tau(i,j,i')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in A}}\tau(i,j,i')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in A}}\tau(i,j,i')+\frac{1-\kappa}{\min(N,M)}\sum_{\substack{(i',j')\in A}}\tau(i,j,i')+\frac{1
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 $H_{ij} = max \begin{cases} 0 \\ H_{i-1j} + \gamma \\ H_{i-1j-1} + Z(A|i-1, j-1) \\ H_{ij-1} + \gamma \end{cases}$ (3)

end for end for $A(S_a, S_b) = BACKTRACKING(H)$ Remove bases in (S_a, P_a) and (S_b, P_b) that align to gaps in $A(S_a, S_b)$ until gaps $\notin A(S_a, S_b)$

In step 1 we calculate weights Z(A|i,i') as defined in equation 2 for all combinations of fixed positions i in sequence S_a and i' in sequence S_b . The only difference between equations 2 and 1 is the fixation of iand i', and the introduction of parameter κ setting the impact of sequence conservation. Thus, we globally align two vectors of probabilities instead of two matrices which can be done by the Needleman-Wunsch algorithm. The actual alignments aren't needed in step 2, instead the two sequences are locally aligned by using the weights Z(A|i,i') from step 1 as similarity scores, see equation 3. This can be done by the Smith-Waterman algorithm. Both steps are repeated until no more gaps are introduced in step 2, thus, the same nucleotides in S_a and S_b are aligned in the last two iterations. Then the final similarity of the two dotplots is calculated by

$$Z(A) = \frac{1}{|A|} \left(\sum_{\substack{i=-\text{gap}, i'=-\text{gap}}} Z(A|i, i') + \gamma \times L \right)$$
(4)

where Z(A|i,i') is equal to equation 2 without the term for gaps, |A| is the length of the local alignment and $L = |S_a| + |S_b| - 2|A|$.

The robustness of the alignment is improved by applying log-odds scores of having a specific base pairing against the null model of a random pairing (Will et al., 2007). Here, we replace P(i, j) with

$$\Psi_{i,j} = log \frac{P(i,j)}{p_0(i,j)} / log \frac{1}{p_0(i,j)}$$
 (5)

where $p_0(i,j)$ is the expected probability for a pairing to occur at random (use ribosomal paired probability matrix (Knudsen and Hein, 1999)). The term $log \frac{1}{p_0(i,j)}$ is a normalization factor that transforms the scores to a maximum of 1. P == 1 results in $\Psi = 1$, $P > p_0$ results in $\Psi > 0$, $P == p_0$ results in $\Psi = 0$, and $P < p_0$ results in $\Psi^* \leq \Psi < 0$ where Ψ^* is a lower boundary. This transformation gives weaker similarities if low basepair probabilities are compared, but stronger similarities for high basepair probabilities. The similarity τ is then calculate by

$$\tau(i, j, i', j') = \begin{cases} 1 - \delta(i, j, i', j') & \text{if } \delta(i, j, i', j') < 1 \\ 0 & \text{else} \end{cases}$$

where $\delta(i,j,i',j') = |\Psi(i,j) - \Psi(i',j')|$ so that $\tau = (0,1)$.

Unpaired probabilities are handled in a similar way by

$$\omega(i) = log \frac{1 - \sum_{k} P(i, k)}{p_0(i)} / log \frac{1}{p_0(i)}$$
 (7)

where $p_0(i)$ is the expected probability for an unpaired base to occur at random (use ribosomal unpaired probabilities (Knudsen and Hein, 1999)). Our model is based on structure similarity, however, the sequence similarity σ may be especially important in unpaired regions, *e.g.* as accessible sequence-specific binding motif. Therefore, we weight matching nucleotides by the similarity of their unpaired probabilities:

$$\sigma(i,i') = \begin{cases} 1 - \delta(i,i') & \text{if } i == i' \text{ and } \delta(i,i') < 1 \\ 0 & \text{else} \end{cases}$$
(8)

where $\delta(i,i') = |\omega(i) - \omega(i')|$ so that $\sigma = (0,1)$. By doing so, sequence similarity gets highest weight if the base in both sequences is likely to be unpaired.

Finally, the proposed algorithm can be optimized by different parameters which will be evaluated in the result section:

- 1. weight of sequence similarity (optimize κ)
- 2. replace $\gamma \times L$ with affine gap costs $l \times \alpha + k \times \beta$ where l is number of initiation gaps and k is the number of all gaps (optimize α and β)

2.1 Speed up

To achieve a very fast method that can be applied on a large amount of pairwise comparisons, *e.g.* a set of 2,000 RNA sequences requires 2×10^6 comparisons, we implemented two complementary strategies:

- 1. set maximal allowed shift of two input sequences in the final alignment
- 2. random seed alignments

The first strategy reduces the amount of comparisons in step 1 by ignoring pairs of distant nucleotides which will normally never align. This restriction will always find the best global alignment but may miss local alignments of long input sequences if the maximal shift is set to small. The second strategy randomly selects short seed sequences (5 nucleotides) in sequence 1 and aligns them to sequence 2. As soon as the first seed alignment has a gap-free alignment above a given threshold then the entire sequences are aligned around the already calculated similarities. If all seed alignments fail the program is stopped and a similarity of 0 is returned.

2.2 Local alignments

The so far presented approach finds only semi-local alignments, meaning the heuristic in the first step of the algorithm gives global alignments, whereas the second step provides a final local alignment. This strategy is applicable for input sequences of similar lengths, however, a real local alignment is necessary if input sequences are very long or have different lengths. Finding the optimal local alignment common to all pairs of pairing probabilities in step 1 is again NP-hard, why we apply another heuristic. In step 1 we find the optimal local alignment for each pair of pairing probabilities that includes the two reference bases. Then for each optimal local alignment

we run for its two subsequences step 1 with global alignments, and after that step 2 on the distances of the two subsequences. This results in a number of local alignments that can be sorted by their similarity.

3 RESULTS

The accuracy of the proposed algorithm is assessed using the specificity (SP) and the sensitivity (SN), which are defined as follows:

$$SP = \frac{TN}{TN + FP}, SN = \frac{TP}{TP + FN}$$
 (9)

where TP is the number of correctly predicted positives, FP is the number of incorrectly predicted positives, TN is the number of correctly predicted negatives, and FN is the number of incorrectly predicted negatives. Furthermore, the area under the receiver operating characteristic (ROC) curve was used to optimize κ , α , and β . The ROC curve plots the true positive rates (SN) as a function of the false positive rates (1 - SP) for varying parameters.

As benchmark data set we selected 300 sequences of 10 H/ACA-box snoRNA families from Rfam version 11.0 seed alignments with average pairwise sequence identity (APSI) < 90% and sequence lengths of > 130bp and < 140bp: SNORA1, SNORA13, SNORA14, SNORA15, SNORA16, SNORA17, SNORA18, SNORA19, SNORA2, SNORA22. We chose only sequences of similar length because step 1 of DotAligner performs global alignments.

Martins benchmark for different APSIs. Compare with reference alignments by the SPS measure introduced in Bralibase 2.1 (see CARNA paper). Compare Rfam families with significant clusters generated by pvclust.

3.1 Parameter optimization

We tried gap costs γ of 3, 4, 5 and 6, which are used unweighted in step 1 of the algorithm (equation 2) and weighted in step 2 of DotAligner (equation 3) by the factor

$$\frac{1}{N \times M} \sum_{i=1}^{N} \sum_{i'=1}^{M} Z(A|i,i')$$
 (10)

where N and M are lengths of S_a and S_b , respectively. The ROC curve in Figure 1 shows the lowest γ as most sensitive (SN = 0.61) and the highest γ as most specific (SP = 1.0) for correctly clustering the selected Rfam families. In the following we choose $\gamma = 4$, whereas the optimal gap cost lies somewhere between 3 and 4.

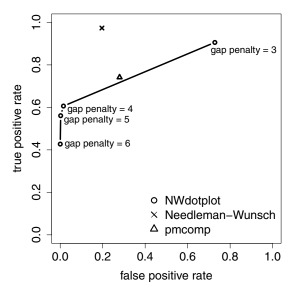


Figure 1: Performance comparison of hierarchical cluster analysis: Degree of agreement between the 10 tested Rfam families and the automated clustering based on distance scores from DotAligner with different gap penalties, Needleman-Wunsch algorithm and pmcomp.

3.2 Comparison with other methods

We compare DotAligner with sequence alignments (in-house implementation of Needleman-Wunsch algorithm with the blastn parameters match = 2, mismatch = -3 and gap penalty = 5 which are optimized for sequence identity of 90%) and the structure alignment tools pmcomp (using default parameters or larger values for parameter -D if the length difference of two sequences is > 5 bp), CARNA, and LocaRNA. Figure 1 shows that the sequence aligner (SP = 0.80, SN=0.97) performs very well on our benchmark set with a very high sensitivity which is most likely due to the fact that the input sequences have some degree of sequence information. pmcomp (SP = 0.72, SN = 0.74) performed with a medium sensitivity and specificity. With DotAligner we are able to find very well defined clusters (SP = 0.99), however, at the cost of sensitivity (SN = 0.61), see Figure 2.

3.3 Clustering methodology

The reliability of our pairwise structure alignment algorithm at clustering homologous RNA structures was tested on a curated database of RNA structure families (cf RFAM). This enables both qualitative and quantitative performance evaluation using a gold-standard reference. We compared <code>DotAligner</code> to other RNA structure alignment and clustering tools using the following framework:

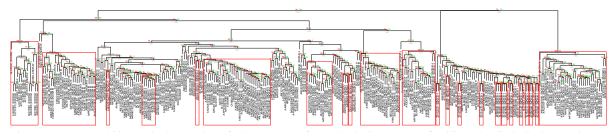


Figure 2: Automated hierarchical clustering of 300 sequences from 10 H/ACA snoRNA families. The dissimilarity matrix was calculated through DotAligner with gap penalty 4. The clustering was conducted by the R-package pvclust with multiscale bootstrap resampling with number of bootstrap 1000. We define clusters (red rectangles) as Approximately Unbiased (AU) p-values > 0.95 rejecting the hypothesis that "the cluster does not exist" with significance level 0.05.

- 1. Generate dissimilarity matrix dM_A from $\frac{n(n-1)}{2}$ pairwise structure comparisons with each algorithm
- 2. Hierarchical clustering of RNA secondary structures and significance testing with pvclust (Suzuki R and Shimodaira H. Bioinformatics 2006).
- 3. Generate dissimilarity matrix dM_R from scoring metric of (1.) from curated RFAM alignments (constrained alignment).
- 4. Calculate the correlation coefficient between dM_A and dM_R using the Mantel correlation statistic (the cross-product between the standardised distances).

Benchmarking was performed on both complete RNA structures (global alignment) and randomly selected subsequences (local alignment) for various RFAM families, as described below.

3.4 Benchmark data generation

xx RNA families were manually selected from the seed alignments of RFAM 11 (REF). How should we limit the mean pairwise identity? All structures must be within a given range and perform several independent comparisons, i.e. one per SeqID range? Then compare the individual SeqID ranges to a sample of variable SeqIDs (without selection)?

We employed the BuildRfamBenchmark JAVA program from (Smith M et al. NAR 2013) to generate the sample alignments for the RFAM entries listed in TableXX. The tRNA sample includes special tRNAs, like ser-tRNA with a 5th hairpin to see how the latter gets clustered by the algorithms.

RFAM ID	RNA class	average length
	5s rRNA	
	SRP	
	tRNA	
	HaCa snoRNA	
	pre-miRNA	

3.5 Complete RFAM sequences

Global alignment. More emphasis on quantitative clustering, accuracy, and correlation with control.

3.6 Fragmented RFAM sequences

Local alignment, simulating genomic screens. More emphasis on qualitative clustering

4 DISCUSSION

We plan to integrate the proposed method in a pipeline that screens regions of interest for structured RNA domains in a collection of RNA molecules. The pipeline may comprise window based thermodynamic folding, e.g. by RNAplfold (Bernhart et al., 2006), the identification of regions of high intra-molecular binding probabilities, e.g. RNAlocal (Dotu et al., 2010), followed by the presented alignment tool DotAligner. The pre-selection of local structural potential is necessary because DotAligner finds only semi-local alignments, meaning the heuristic in the first step of the algorithm gives global alignment, whereas the second step provides a final local alignment.

DotAligner can also be extended for multiple alignments, similar to the strategy implemented in pmmulti (Hofacker et al., 2004), and the generation of phylogenetic trees. This may replace or support the hierarchical clustering approach used here. In addition, both may serve as input for RNA secondary structure predictors, such as PETfold (Seemann et al.,

2008) unifying thermodynamic and evolutionary information.

The first step of the algorithm could also implement local alignments. In this case, however, the input basepair probabilities should be calculated in a sliding-window approach, such as provided by RNAplfold. Then step 2 has to find the best local alignment of local pairwise alignments from step 1 where the paired bases are overlapping. The search for overlapping local alignments from step 1 will increase the running time exponentially, but local alignments may improve the boundaries of common structured RNA domains.

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