

METHOD

Identification and classification of common RNA structure motifs

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Abstract

The abundance and diversity of processed transcripts in eukaryotic genomes poses a challenge for the systemic classification of their biological functions. Sparse sequence conservation in non-coding sequences and the unreliable nature of RNA structure prediction algorithms further exacerbate this conundrum. Here, we present a unified computational approach for the unsupervised discovery and classification of homologous RNA structure motifs from a set of sequences of interest. At its core lies DotAligner, a heuristic pairwise RNA structure alignment algorithm that considers both sequence similarity and the ensemble of sub-optimal RNA base pairings. Our approach outperforms other state of the art tools at classifying known RNA structure families, both in time and accuracy. When combined to density-based clustering using an empirically measured threshold, this method identifies both known and novel RNA structure motifs from ENCODE immuno-precipitation data for 44 proteins, further expanding the lexicon of functional transcriptomic motifs.

Keywords: RNA structure clustering; Functions of RNA structures; RNA–protein binding; Regulation by non-coding RNAs

Background

The human genome is pervasively transcribed into RNA with less than 2% encoding protein sequences. As genomic technologies progress, an ever increasing amount of non-protein coding RNAs (ncRNA) are being discovered. Once class of ncRNAs – long noncoding RNAs (lncRNAs) – are of particular interest for functional genome annotation given the large expanses they encompass. Understanding the molecular mechanisms underlying the biological functions of lncRNAs – and how they are disrupted in disease – is required to improve the functional annotation of the human genome.

So far, only a relatively small quantity of lncRNAs have been functionally characterised, with regulation of gene expression and epigenetic states recurring as common biological functions [1, 2]. The higher-order structure of RNA molecules is an essential feature of non-coding RNAs that can be used for their classification and the inference of their biological function. We, and others, hypothesise that lncRNAs act as scaffolds for the recruitment of proteins and assembly of ribonucleoproteins (RNPs), mediated by the presence of modular RNA structures, akin to the domain organisation of proteins [3, 4, 5, 6, 7, 8, 9]. Protein-interacting regions of lncRNAs are likely to contain a combination of sequence and structure motifs that confer binding specificity.

Most small ncRNAs have well characterised secondary and tertiary structures, as evidenced in RFAM, the largest collection of curated RNA families (2,588 families as of version 12.2

[10]). However, determining the structural features of long ncRNAs (lncRNAs) is a complex problem given their size and, in general, faster evolutionary turnover. These challenges have raised doubts concerning the prevalence of functional structural motifs in lncRNAs [11, 12], despite evolutionary and biochemical support of conserved base-pairing interactions [7, 13, 14].

Perhaps this paragrapf should go to the discussion? As the increasing accessibility of next generation sequencing coalesces with precision medicine, in-depth transcriptome profiling will help elucidate the function and clinical impact of disease-associated, non-coding single nucleotide variants. Indeed, 80% of disease-associated single nucleotide polymorphisms occur in non-coding regions [15, 16]. Thus, elucidating the structural features of RNAs associated to RNA-binding proteins and ribonucleoprotein (RNP) complexes, combined to the systematic classification of their genome-wide occurrence, can identify novel riboSNitches (functional RNA structures that are disrupted in disease) and help pinpoint the molecular function of non-coding mutations.

Identifying RNAs with similar functions involves comparing both their primary sequence and higher-order structures simultaneously. If the sequence similarity falls below 60% sequence comparison will not find anymore domain similarities that are based on structure (Gardner et al., 2005). Hence, methods are needed that find structural similarity independent from sequence conservation and freed from one single optimal RNA secondary structure. The Sankoff algorithm resolves the optimal sequence-structure alignment of 2 RNAs [17], but its computational complexity limits its practicality. Alternative strategies often employ pre-calculated secondary structure ensembles for each sequence, e.g. basepair probabilities in thermodynamically equilibrated RNA structure ensembles [18]. The latter can substantially speed up the calculation of structure-based alignments [19], of which there are many variants. The programs PMcomp (Hofacker et al., 2004), LocaRNA (Will et al., 2007), and ProbAlign (PMID:16954142) use the pre-computed basepair probability matrices of both sequences and score the alignment based on the notion of a common secondary structure. The sequence-structure alignment problem is reduced to a two-dimensional problem by RNApaln (PMID:22115189) and StrAL (Dalli et al., 2006) which reduces base-pair probabilities to base specific probabilities (such as unpaired probability). All these methods do not explicitly 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 iteratively better alignments with an effective constraint programming technique using a branch and bound scheme. An alternative heuristic is pruning of the dynamical programming matrix without pre-folding constraints, which is implemented by FoldAlign (PMID:17937495, PMID:26704597).

Will et al. first showed that a (dis)similarity matrix can be constructed from all-vs-all pairwise RNA structure alignments with the pairwise alignment tool LocaRNA, identifying known and novel groups of homologous RNAs using hierarchical clustering [20]. However, this strategy involves applying a subjective threshold to the resulting dendrogram to extract structurally related sequences. Alternative methods to identify clusters of homologous RNAs include NoFold, which clusters query sequences based on their relative

similarity to reference structures [21], and GraphClust, an alignment-free approach that decomposes RNA structures into graph-encoded features [22]. RNAscClust, an extension of GraphClust, utilizes the evolutionary signatures of RNA structures as additional classification feature [23].

Here, we describe a computational pipeline for the identification and classification of homologous RNA structures from a large set of query sequences. At its core lies DotAligner, a heuristic pairwise sequence alignment algorithm that considers suboptimal base-pairing probabilities. We compare DotAligner with other pairwise RNA structure alignment algorithms to highlight its speed and accuracy at classifying known RNA families. We combine DotAligner with density based clustering for the impartial identification of RNA structural motifs, which can identify known RFAM families and novel RNA structural motifs from ENCODE eCLIP data. The resulting clusters of homologous RNA structures can then be used to search for homologous structures across reference genomes and transcriptomes.

Results and Discussion

Ensemble-guided pairwise RNA structure alignment

DotAligner leverages the diversity of suboptimal solutions from a partition function of RNA secondary structure predictions to identify an optimal sequence-structure alignment of two RNAs. The algorithm overcomes the limitations of comparing unique RNA secondary structures (such as minimum free energy predictions) to yield an optimal sequence alignment that considers mutual base pair probabilities. **Figure 1** illustrates a structural alignment performed with DotAligner in contrast to an alignment that considers only sequence composition.

A major criteria for the implementation was a fast running time to make DotAligner applicable for RNA structure clustering of large data sets. Consistently, the algorithm performs pairwise sequence-structure alignments from pre-calculated RNA dot plots and using alignment-envelope heuristic, which impose constraints on sub-optimal string alignments, and fold-envelope heuristics, which impose constraints to pre-calculated base pairing probabilities. The alignment procedure thus consists of two steps, each considering base pairing probabilities: (i) Computation of a partition function over all canonical pairwise string alignments, and (ii) structure-weighted stochastic backtracking of all string alignments. The detailed implementation and mathematical description of DotAligner can be found in the **Supplementary Information**.

After initial parameter optimisation (see **Methods**), we applied DotAligner to BRAliBase 2.1 pairwise RNA structure alignments, a reference dataset specifically designed for algorithm benchmarking [24, 25]. In this application, DotAligner seldom produces alignments of better quality than those generated by three other state of the art algorithms, namely CARNA [26], FOLDALIGN [27, 28] and LocaRNA [20] (**Figure 2**).

Interestingly, many of the pairwise structure alignments produced Structural Conservation Index (SCI) scores above those from the BRAliBase 2.1 reference alignments (**Figure 2B**). The SCI represents the alignment consensus energy normalized by the average energy of the single sequences folded independently [29]; It has been shown to be one of

the most reliable metrics for conserved RNA structure detection [30]. With the exception of DotAligner, all other surveyed algorithm produce a substantial amount of alignments with SCI values above that of the reference alignment **Figure 2B**, suggesting that many optimization algorithms tend to overestimate the amount of paired bases in consensus RNA structure predictions. However, another possibility is that BRAliBase 2.1 alignments do not correctly depict several RNA families, which may have been automatically generated in RFAM based on similarity to a covariance model.

Fast and accurate classification of RNA structures

The intended application of DotAligner is the identification and classification of RNA structural motifs from a large and diverse set of sequences of interest. Therefore, we evaluated the ability of DotAligner to distinguish between distinct structured RNA species from a heterogeneous sample of known RNA structure families. We performed all versus all pairwise structure alignments of stochastically sampled RFAM sequences, which were selected with constraints on their sequence composition (PID) to control for and ascertain any sequence-dependent biases (see **Methods**). The alignment scores of DotAligner and other algorithms were then converted into a similarity matrix and compared to a binary classification matrix of sampled RFAM entries.

Despite the seemingly poor quality of pairwise alignments generated by DotAligner, it reproduces the known classification of RFAM structures more accurately than almost all other pairwise RNA structure alignment tools (**Figure 3A-C**). Only CARNA, another ensemble-based structural alignment algorithm, presents classification accuracies comparable to DotAligner. However, CARNA requires substantially more time to perform the comparisons (**Figure 3D**) as it will indefinitely continue to compute the alignment until it converges on an optimal result, or a hard time constraint is enforced. In this regard, DotAligner performs better than all other RNA structure alignment tools, highlighting the efficacy of the implemented heuristics it employs. In contrast, the heuristics implemented in RNAlign increased the speed of pmcomp by 2 orders of magnitude, but at a slight reduction in accuracy (Figure 3C,D). Only a C++ implementation of the Needleman-Wunsch Algorithm (NWA) [31]—a classical sequence alignment algorithm that ignores secondary structure information—performs faster than DotAligner on average, most likely due to the presence of stretches of homologous sequences within biologically related RNAs.

Density-based clustering of homologous RNA structures

Given DotAligner's accurate classification of known structured RNA, we subjected its output to cluster analysis to identify and extract input sequences which display common sequence-structure motifs. The previous work by Will et al. applied hierarchical clustering to the dissimilarity matrices produced by LocaRNA to organise sequences based on their structural homology [20], yet this does not apply a cut-off that can be used to extract meaningful clusters of structurally homologous sequences in an unsupervised manner. We attempted to achieve this by applying a statistical threshold derived from bootstrapping the underlying data using pvclust [32], but this generated clusters of variable size that often spanned across many disjoint families (data not shown).

We therefore opted for a density-based clustering strategy that, in theory, can decipher clusters of varying density (i.e. subsets of the data with greater sequence-structure homology). The OPTICS (Ordering Points To Identify the Clustering Structure) algorithm [33] was chosen for this purpose, as it has very few parameters to optimise. OPTICS is a derivative of the Density-BaSed Clustering for Application with Noise (DBSCAN) [34] algorithm that, as its name states, is suitable for noisy data, such as RNA immunoprecipitation followed by high-throughput sequencing (RIPseq).

We then benchmarked the two main OPTICS clustering parameters— ξ steepness threshold and the minimum number of points in a cluster (**Supplementary Figure 1**)—on a pooled set of 580 stochastically sampled RFAM sequences encompassing various ranges of sequence similarity, as well as a corresponding set of 580 dinucleotide shuffled controls (see **Methods**). After performing all vs all pairwise alignments with DotAligner, the alignment scores were normalised from 0 to 1 for the minimal and maximal values, respectively. The resulting similarity matrix was then used to evaluate the effect of OPTICS parameters on clustering performance, revealing that a minimum of 4 points (or sequences) and a steepness threshold of 0.006 gave the best results **Supplementary Figure 1A**. Extracting only the minimum cluster when

When compared to other state of the art methods, namely NoFold and GraphClust, the combination of DotAligner and OPTICS performs comparably well (**Figure 4, Table 1**). The default version of NoFold nonetheless outshines DotAligner at clustering known RFAM families. However, it intrinsically employs RFAM covariance models when comparing input sequences, therefore is likely to be subject to over-fitting in this example. We thus removed the CMs associated to the RFAM sequences in our benchmarking dataset from the NoFold algorithm, which yielded lower sensitivity and less accurate qualitative cluster metrics than the DotAligner and OPTICS combination. Interestingly, the specificity of NoFold increased slightly despite removing 72 CMs from its classification set.

Identifying protein-binding RNA motifs from eCLIP data

The optimised parameters for OPTICS clustering of DotAligner output were incorporated into a high-performance computing pipeline that extracts clusters of homologous RNA structural motifs from a set of input sequences (see **Methods**). This pipeline was applied to enhanced cross-linked RNA immunoprecipitation (eCLIP) sequencing data from 44 RNA binding proteins from the ENCODE consortium [35].

If a structural motif is targeted by a RBP, it is highly probable that the associated CLIPseq peaks do not encompass the entire sequence that forms the structural motif, since reverse transcription will terminate at the covalent bond formed between the protein and the RNA. Consequently, large RNA structures bound by RBPs are difficult to model without additional structure probing data. This experimental approach also introduces a 3' bias in the sequence-specificity of RBP binding sites given the nature of 3' to 5' reverse transcription required for cDNA synthesis.

To address these caveats, we surveyed the potential to form locally-stable RNA secondary structures in the regions directly flanking 36,161 filtered eCLIP peaks (see **Methods**). XXX% overlap a predicted RNA secondary structure, compared to XXX random peaks.

Conclusion

Given its relative speed and accuracy, DotAligner can be used to generate larger (dis)similarity matrices for cluster analysis than other pairwise structure alignment algorithms, or at least produce them with reasonable computational power.

In addition to its speed, DotAligner's strength lies in it's capacity to accurately score structurally homologous RNA sequences. The algorithm appears to generate pairwise alignments that differ somewhat to the reference structural alignments. Despite this, DotAligner can harness the information content of base pair probability ensembles to output a reliable structural similarity score of two RNA sequences. Our results show that this can nonetheless be sufficient to extract structurally and functionally related sequences from a large amount of noisy input; an ideal application for screening high-throughput sequencing data, such as RNA immunoprecipitation data, for common structural motifs.

weaknesses: boundaries, parameter-choice of algorithms. applications: riboswitches, ...

Methods

Benchmarking and parameter optimisation

The DotAligner algorithm implements several theoretical parameters that first need to be tuned before being applied to biological sequence analysis. All combinations of core parameters were tested on the 8,976 pairwise RNA structure alignments curated in the BRAliBase 2.1 reference dataset [25]. For each set of parameter combinations, the amount of alignments producing identical structural topologies to the reference alignment was determined using *RNAdistance*. The Structural Conservation Index (SCI), a robust measure of RNA structural alignment integrity [30], and the Matthews Correlation Coefficient (MCC) were also calculated (see below) for all resulting alignments.

Baseline parameters were then selected via a product rank these 2 metrics (**supplementary data?**).

$$\text{Matthews Correlation Coefficient (MCC)} = \frac{(TP*TN)-(FP*FN)}{\sqrt{(TP+FP)(TP+FN)(TN+FP)(TN+FN)}}$$

$$\text{Structural Conservation Index (SCI)} = MFE_{consensus}/\bar{MFE}_{single}$$

$$\delta SCI = SCI_{observed}/SCI_{reference}$$

Classification of RNA secondary structure families

Further refinement of the optimal parameters was performed using a binary classifier two sets of 200 stochastically sampled RFAM entries with published structures: (i) a low pairwise sequence identity set, and (ii) a high pairwise sequence identity set, where any two sequences from the same family share between 0-55% and 56-95% pairwise sequence identity, respectively. A binary classification matrix was then constructed, where sequences x and y present a score of 1 if they belong to the same RFAM family, versus a score of 0 if they do not. The similarity matrix resulting from all-vs-all pairwise comparisons with DotAligner was tested for accuracy using the Area Under the Curve AUC of the ROC, as calculated R package pROC [36]. A more restricted range of parameter values were then tested on both datasets, namely t k e o, which had the highest impact on alignment accuracy in preliminary testing (SUPP FIG X??). Finally, a ranked sum for both datasets of the AUC (first) and lowest average runtime (second) was performed to determine the default runtime

parameters for DotAligner [Supplementary table X](#).

This was achieved by sampling the entire collection of RFAM entries with published structures in a stochastic manner, while ensuring that all sampled sequences respected constraints on their sequence composition. Specifically, we extracted a high Pairwise sequence IDentity (PID) and a low PID set, where any two sequences from the same set present greater than or less than 55% PSI. The

Clustering RNA structures with randomised controls

OPTICS benchmarking was performed by stochastically sampling the collection of RFAM 12.0 seed alignments using the accompanying JAVA program `GenerateRFAMsubsets.java` (see **Supplementary Information**) with parameters [GET PARAMETERS FROM SERVER](#), [how many sequences minimum/maximum](#), with 3 ranges of pairwise sequence identity: 1-55%, 56-75%, and 75-95%. The resulting 580 unique sequences were then randomised while controlling their dinucleotide content with the `easel` program included in the `Infernal` (v1.1.2) software package [37] with option "-k 2". The 1160 sequences were submitted to all-vs-all pairwise comparisons with `DotAligner` and the scores were inverted and a normalised (min=1, max=0) into a dissimilarity matrix, which was then imported into the R statistical programming language, converted into a 'dist' object without transformation, and subjected to OPTICS clustering as implemented in the 'dbscan' CRAN repository with a range of parameters (see **Figure 4A,B**).

Clustering performance metrics were calculated like so:

- True Positives (TP) = Number of representatives from the dominant RFAM family in a cluster;
- False Positives (FP) = Number of non-dominant RFAM family representatives in cluster, or clusters where there is no dominant RFAM family (i.e. equally represented families), or clusters where dominant sequence is a negative control;
- False Negatives (FN) = RFAM sequences that fail to cluster;
- True Negatives (TN) = Negative control sequences that fail to cluster.
- Sensitivity (recall) = $TP / (TP + FN)$;
- Specificity = $TN / (TN + FP)$;
- False positive rate = $1 - \text{Specificity}$;
- Precision = $TP / (TP + FP)$;
- Accuracy = $(TP + TN) / (TP + TN + FP + FN)$;

Clustering of protein-bound evolutionarily-conserved RNAseq reads

The genomic coordinates of ENCODE eCLIP peaks were downloaded in .bed format from the April 2016 release via the ENCODE portal (<https://www.encodeproject.org/search>). The resulting 5,040,096 peaks were filtered to keep only those with $\geq 8x$ fold enrichment over the total input background and an associated P-value $\leq 10^{-4}$. Furthermore, peaks were merged if they overlapped by more than 50nt to avoid over-representing the same sequence (see **Supplementary Information** section 2). The remaining peaks were subsequently filtered by retaining only those that present same-strand overlap with any evolutionarily conserved structure (ECS) predictions from [7]. Finally, the associated genomic sequences were extracted into a .fasta file, which was supplemented with 100

reference RNA structures from 11 RFAM families (see Supplementary Table 1). Merging, overlap, and sequence extraction operations were performed with bedtools version v2.26.0.

The normalised similarity matrix resulting from all vs all pairwise comparisons with DotAligner was then subjected to clustering with the dbscan 1.1-1 R package from Michael Hahsler <https://github.com/mhahsler/dbscan> using the command ‘opticsXi (optics(D, eps=1, minPts=4, search="dist"), xi = 0.006, minimum=T)’. The sequences for each cluster were then extracted and submitted to multiple structure alignment with mLocaRNA version 1.9.1 using parameters ‘--probabilistic --iterations=10 --consistency-transformation --noLP’.

The complete analytical pipeline is available at <https://github.com/noncodo/BigRedButton>, which will identify and extract homologous RNA structural motifs from a set of input sequences in fasta format.

Competing interests

The authors declare that they have no competing interests.

Author's contributions

Text for this section ...

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Figures

Tables

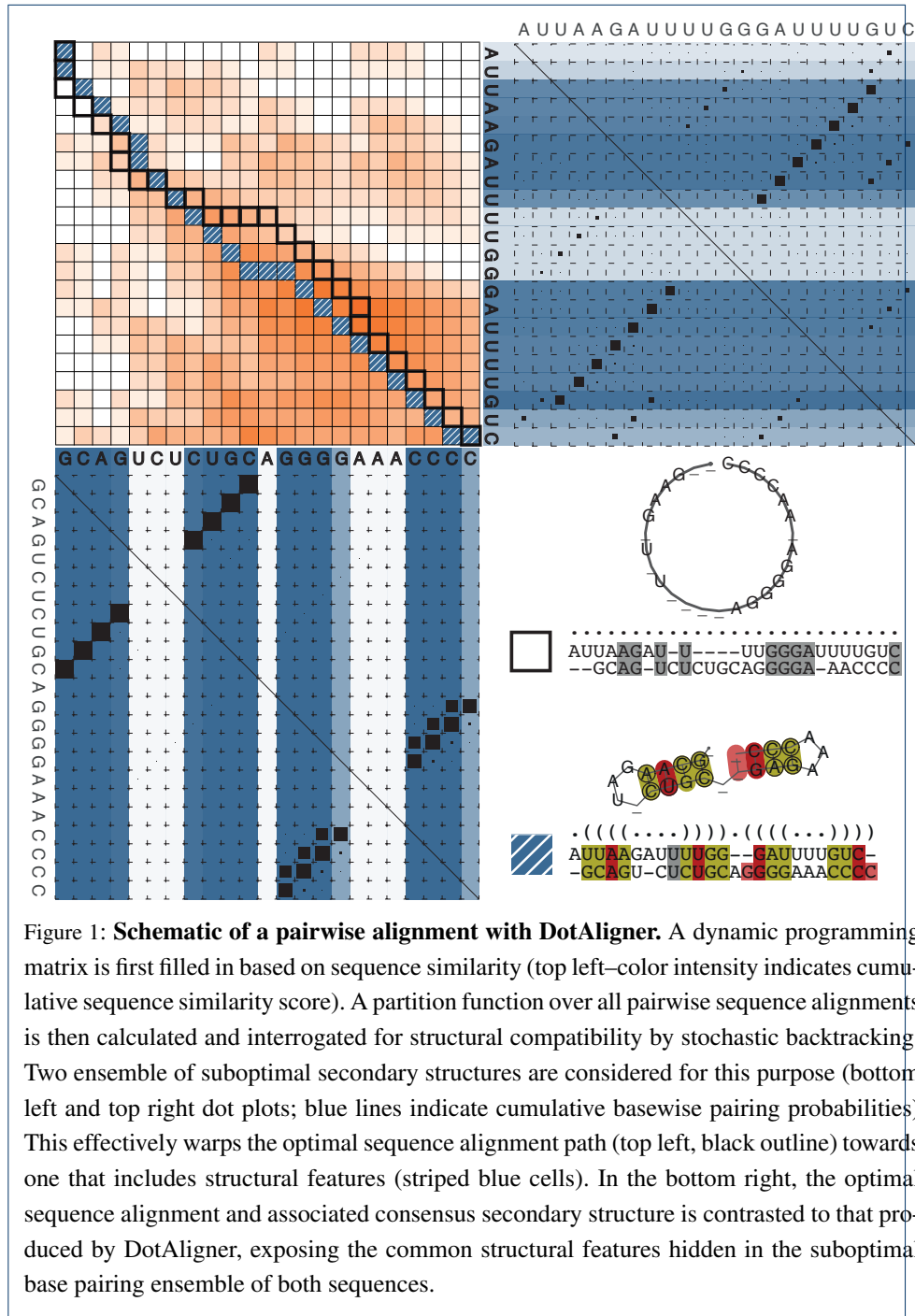
Table 1: Comparative clustering performance.

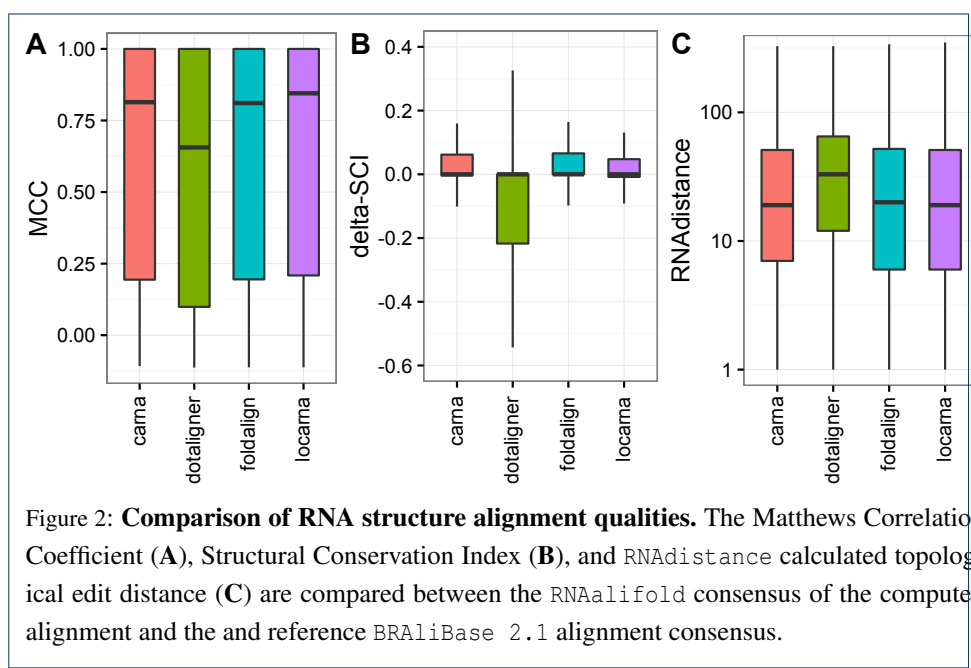
Algorithm	# Clusters	Sensitivity	Specificity	Accuracy
DotAligner+OPTICS	53	0.716	0.886	0.802
GraphClust	201	0.990	0.110	0.635
NoFold (all CMs)	62	0.866	0.965	0.916
NoFold (filtered)	45	0.674	0.976	0.826

Additional Files

Additional file 1 — Supplemental Material

More detailed description of the DotAligner implementation, RNA structure clustering and eCLIP data processing, and Suppemetla Tables and Figures summarizing study results.





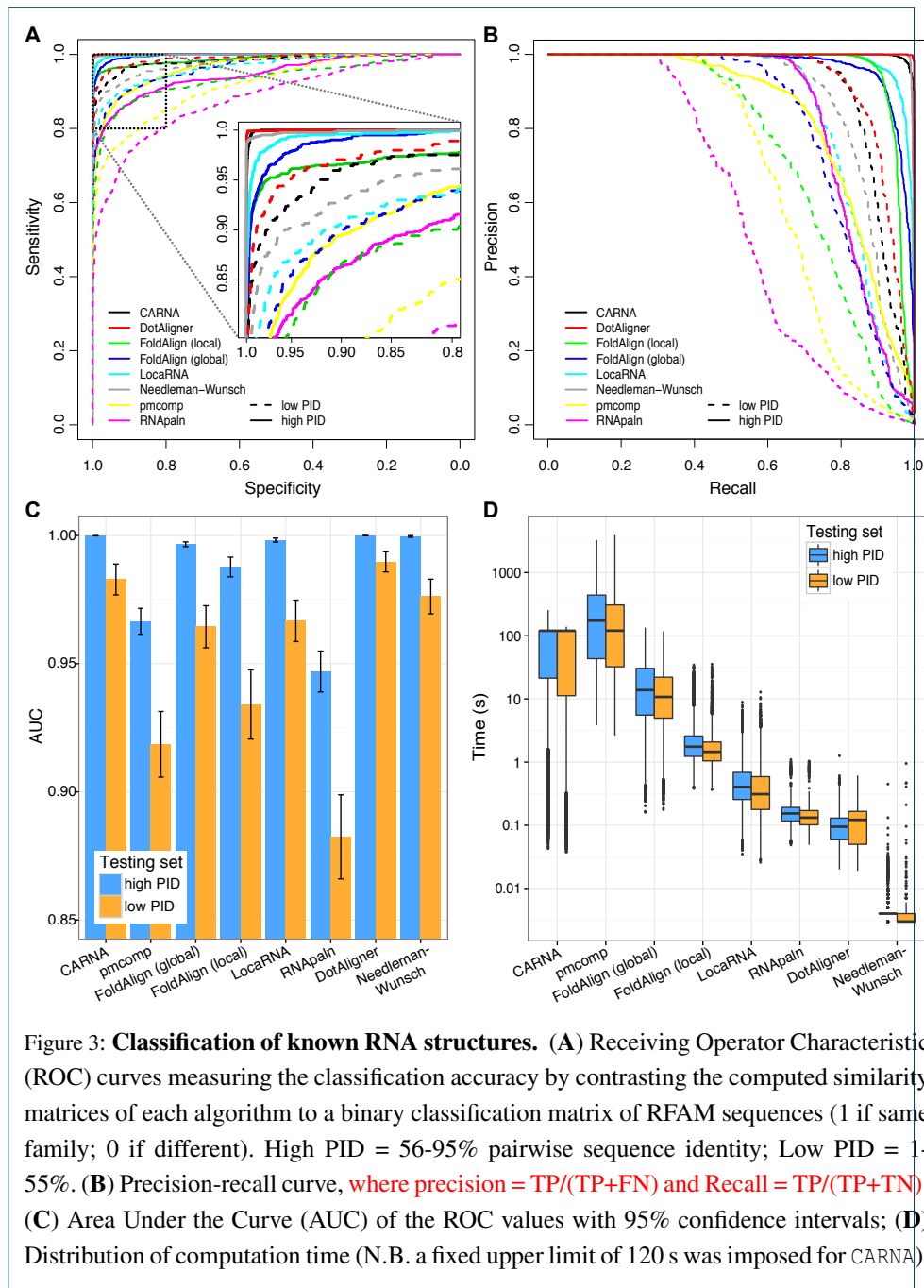
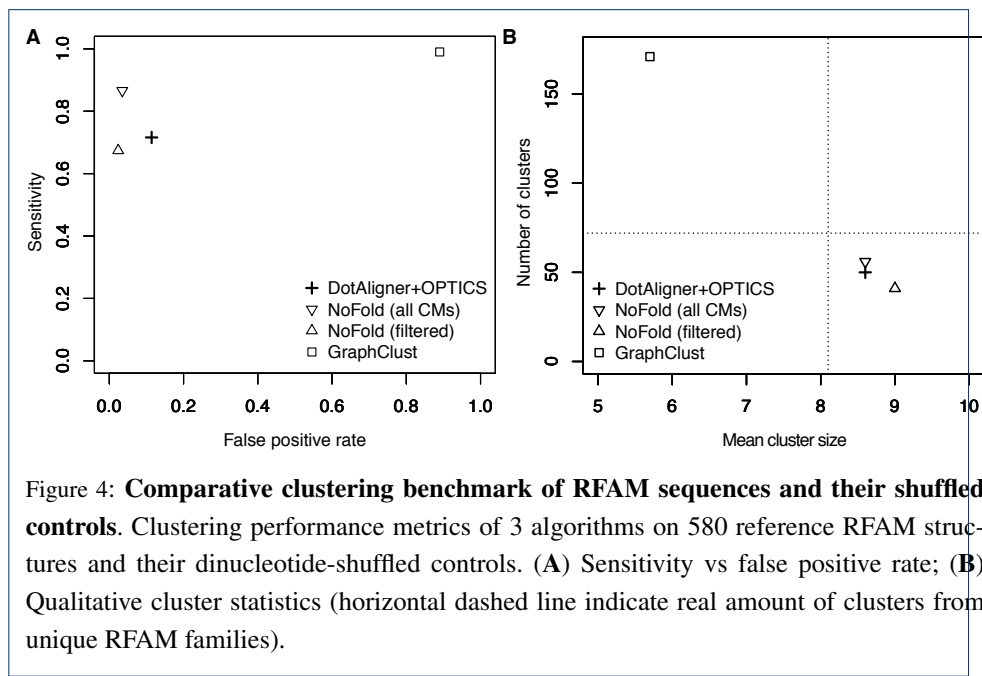
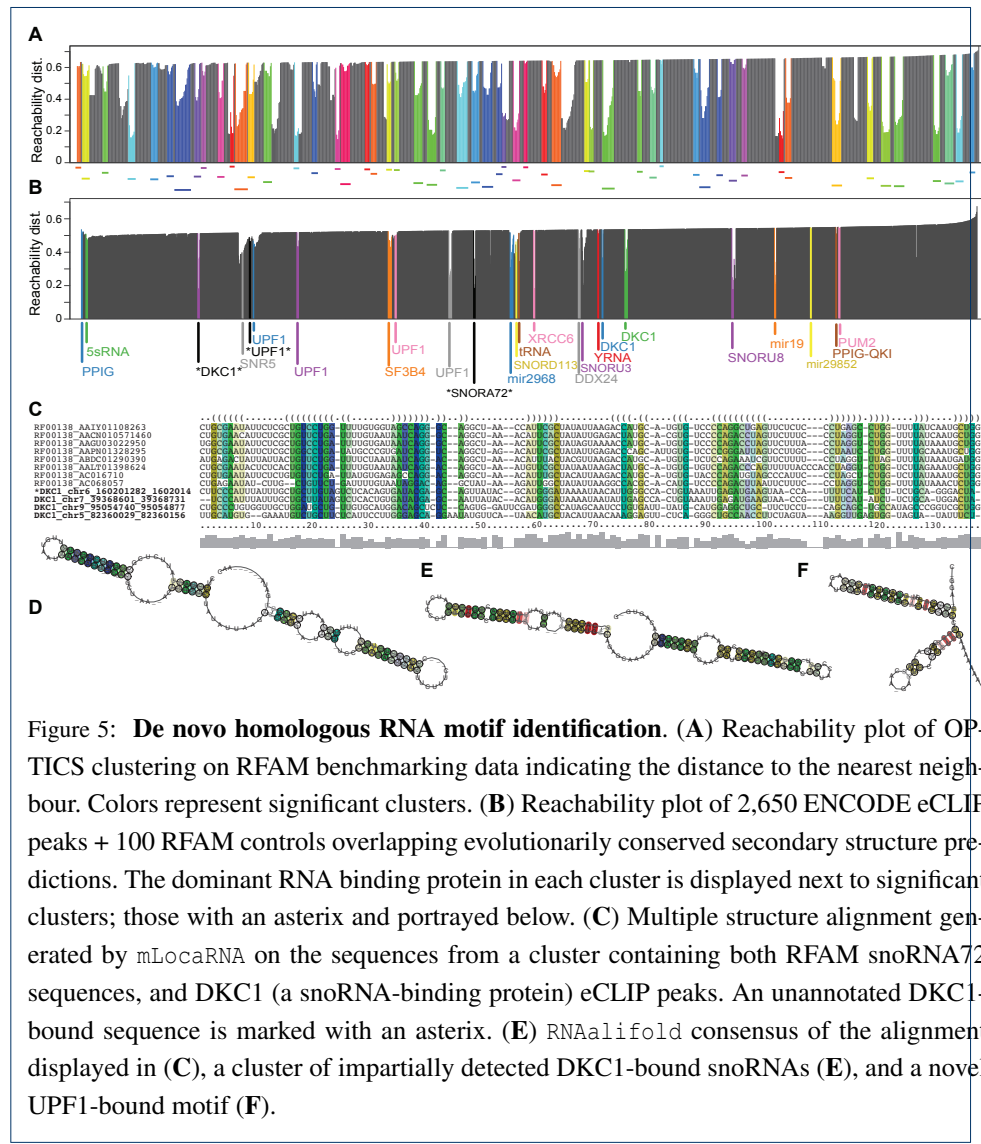


Figure 3: Classification of known RNA structures. (A) Receiving Operator Characteristic (ROC) curves measuring the classification accuracy by contrasting the computed similarity matrices of each algorithm to a binary classification matrix of RFAM sequences (1 if same family; 0 if different). High PID = 56-95% pairwise sequence identity; Low PID = 1-55%. (B) Precision-recall curve, where precision = $TP/(TP+FN)$ and Recall = $TP/(TP+TN)$; (C) Area Under the Curve (AUC) of the ROC values with 95% confidence intervals; (D) Distribution of computation time (N.B. a fixed upper limit of 120 s was imposed for CARNA).





Additional file 1 — Supplemental Material

1. DotAligner implementation

As described in [38] the weight W of alignment A of two arc-annotated sequences (S_a, P_a) and (S_b, P_b) is defined by

$$\begin{aligned} W(A) &= \sigma(A) + \tau(A) + \gamma(A) \\ &= \sum_{(i,i') \in A} \sigma(i, i') + \sum_{\substack{(i,j) \in P_a, \\ (i',j') \in P_b, \\ (i,i') \in A, \\ (j,j') \in A}} \tau(i, j, i', j') + \gamma \times N \end{aligned} \quad (1)$$

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 ($N = |S_a| + |S_b| - 2|A|$). The alignment problem finds the maximal $W(A)$. As its solution is MAX-SNP-hard, in praxis heuristics are used to find near-optimal solutions.

DotAligner solves the related problem of aligning two basepair probability matrices (dot plots). A major criteria for the implementation was a fast running time enable all-vs-all pairwise structural alignments and the associated distance (dissimilarity) matrix, which can be used for cluster analysis of large data sets [20]. Consequently, it employs a heuristic alignment-envelope, which imposes constraints to sub-optimal string alignments, and a fold-envelope, which imposes constraints to pre-calculated base pairing probabilities, to build pairwise sequence-structure alignments. DotAligner makes use of the observation that large samples from the ensemble of stochastic sequence alignments contain the correct structure-based alignment with significant probability, even though the optimal sequence alignment deviates significantly from the structural alignment [39].

Below, we describe the alignment procedure and weight functions. The alignment procedure consists of two steps:

- 1 Pairwise probabilistic string alignments;
- 2 Stochastic backtracking of string alignments and combining weights of base pairing probability matrices.

1.1 Pairwise probabilistic string alignments

In step 1 the computation of the partition function over all canonical pairwise string alignments is adapted from probA [39]. The probability of an alignment A in the ensemble of all alignments $Z(T)$ is

$$Pr(A; T) = \frac{1}{Z(T)} \exp(\beta W(A)), \quad (2)$$

where $\beta = 1/T$. The parameter T is analogous to the temperature in the thermodynamic interpretation of the alignment problem and determines the relative importance of the optimal string alignment. If $T = 1$ then we recover the 'true' probability, if $T \rightarrow 0$ then $Pr(A; 0) = 0$ for all alignments with a score $W(A)$ less than the score of the optimal string alignment,

and if $T \rightarrow \infty$ then all alignments have the same $Pr(A, \infty) = 1/Z(\infty)$. Hence, T controls the search space of suboptimal alignments for step 2. The algorithm runs in $O(N^3)$ for calculating the partition function. The weight function $W(A)$ of the probA implementation is changed to explore the ensemble of dot plot alignments. We reduce the sequence-structure alignment problem to a two-dimensional problem similar to the metric introduced in StrAL [40]. Hence, step 1 considers only the similarity σ and the gap cost γ described in equation 1:

$$W_{\text{Step1}}(A) = \sigma(A) + \gamma(A) \quad (3)$$

The similarity $\sigma(i, i')$ for matched sequence positions $S_a[i]$ and $S_b[i']$ takes into account sequence similarity M_{Seq} and the similarity in their unpaired probabilities $\Delta\omega(i, i')$ weighted by the parameter θ :

$$\sigma(i, i') = \theta \times M_{Seq}^{(i, i')} + (1 - \theta) \times \Delta\omega(i, i') \quad (4)$$

$M_{Seq}^{(i, i')}$ is 1 if sequence positions $S_a[i]$ and $S_b[i']$ match and else 0. The similarity of unpaired probabilities is defined as

$$\Delta\omega(i, i') = \begin{cases} 0 & \text{if } \omega(i) == 0 \\ & \text{and } \omega(i') == 0 \\ 1 - |\omega(i) - \omega(i')| & \text{else} \end{cases} \quad (5)$$

so that $\Delta\omega = (0, 1)$. Alternatively a statistical substitution model R_{Seq} replaces the sequence similarity and is multiplied with the ζ weighted sum of $\Delta\omega$ and the similarity in ratios of upstream pairing probability $\Delta\omega^{up}$:

$$\sigma(i, i') = R_{Seq}^{(i, i')} \times \zeta \times \Delta\omega(i, i') + R_{Seq}^{(i, i')} \times (1 - \zeta) \times \Delta\omega^{up}(i, i') \quad (6)$$

R_{Seq} is a 4×4 matrix of probabilities for observing a given substitution relative to background nucleotide frequencies. We use the log-odd scores L from the RIBOSUM85-60 matrix introduced in [41] which are transformed to probabilities R_{Seq} by $2^{L(i, i')}/(1 + 2^{L(i, i')})$. The ratio of upstream pairing probability ω^{up} is defined as

$$\omega^{up}(i) = \sum_{k=1}^{i-1} \psi(k, i) / \sum_{k=1}^{|S|} \psi(k, i) \quad (7)$$

where $i \in S$, $|S|$ is the length of sequence S , and $\psi(k, i)$ is the pairing probability of sequence positions $S[k]$ and $S[i]$. The downstream pairing probability is implicitly considered in the

weight function through the usage of unpaired probability and upstream pairing probability. The gap term in equation 1 is replaced with affine gap costs:

$$\gamma(A) = l \times g_o + (N - l) \times g_{ext} \quad (8)$$

where l is the number of initiation gaps, N is the number of all gaps, g_o is the penalty for opening a gap and g_{ext} is the penalty for gap extensions. Start and end gaps are considered as free.

1.2 Stochastic backtracking and combined weight of dot plot alignments

Here, a properly weighted sample of stochastic pairwise string alignments in the alignment ensemble is examined across both sequences for sequence-structure similarity. The stochastic backtracking is adapted from probA [?] for selecting s suboptimal string alignments A_s . The combined weight W_{Step2} is a variant of equation 1 to explore the similarity of the corresponding dot plot alignments:

$$W_{\text{Step2}}(A_s) = \kappa \times \frac{W_{\text{Step1}}(A_s)}{|A_s|} + (1 - \kappa) \times \frac{\tau(A_s)}{|\text{Match}_{A_s}|^2} \quad (9)$$

where the parameter κ weights for each alignment A_s between the sequence-based similarity $W_{\text{Step1}}(A_s)$ normalized by alignment length $|A_s|$ and dot plot similarity $\tau(A_s)$ normalized by the number of aligned bases $|\text{Match}_{A_s}|$ in alignment A_s . Similar to equation 4 the dot plot similarity τ sums the parameter θ weighted similarity of aligned basepairs M_{paired} and the similarity in their pairing probabilities $\Delta\psi$:

$$\tau(i, j, i', j') = \theta \times M_{\text{paired}}^{(i, j, i', j')} + (1 - \theta) \times \Delta\psi(i, j, i', j') \quad (10)$$

where $M_{\text{paired}}^{(i, j, i', j')}$ is 1 if $S_a[i]$ and $S_a[j]$ as well as $S_b[i']$ and $S_b[j']$ form canonical basepairs (G-C, C-G, A-U, U-A, G-U or U-G) and else 0. The similarity in pairing probabilities $\Delta\psi$ is then calculate by

$$\Delta\psi(i, j, i', j') = \begin{cases} 0 & \text{if } \psi(i, j) == 0 \text{ and } \psi(i', j') == 0 \\ 1 - |\psi(i, j) - \psi(i', j')| & \text{else} \end{cases} \quad (11)$$

Similar to M_{Seq} in equation 4, the basepair similarity matrix M_{paired} can be replaced by a statistical substitution model R_{paired} which describes the probability for observing a given basepair substitution relative to background nucleotide frequencies:

$$\tau(i, j, i', j') = R_{\text{paired}}^{(i, j, i', j')} \times \Delta\psi(i, j, i', j') \quad (12)$$

Again, the log-odd scores L from the RIBOSUM85-60 matrix [41] are transformed to probabilities R_{paired} .

For both sequences S_a and S_b , the pairing probability matrices P_a and P_b are computed in advance using McCaskill's algorithm, implemented in RNAfold or RNAlignfold. The robustness of the alignment is improved by applying log-odds scores ψ of having a specific base pairing against the null model of a random pairing [20]:

$$\psi(i, j) = \max \left(0, \log \frac{P(i, j)}{p_0} / \log \frac{1}{p_0} \right) \quad (13)$$

where p_0 is the expected probability for a pairing to occur at random. The term $\log \frac{1}{p_0}$ is a normalization factor that transforms the scores to a maximum of 1. $P == 1$ results in $\psi = 1$, $P > p_0$ results in $\psi > 0$, and $P \leq p_0$ results in $\psi = 0$. This transformation gives weaker similarities if low basepair probabilities are compared, but stronger similarities for high basepair probabilities. Unpaired probabilities are handled in a similar way by

$$\omega(i) = \max \left(0, \log \frac{1 - \sum_k P(i, k)}{p_0} / \log \frac{1}{p_0} \right) \quad (14)$$

where p_0 is the expected probability for an unpaired base to occur at random.

2. Clustering RNA structures with randomised controls

Below is the code used to calculate the accuracy and other performance metrics of the clustering benchmark of stochastically sampled RFAM entries. All files can be found on the associated GitHub repository <https://github.com/noncodo/BigRedButton>.

```
cat("File name", "TP", "TN", "FP", "FN", "SENS", "SPEC", "ACC", "\n", sep="\t",
    file="accuracies.tsv")
file.names <- dir(pattern="*_clust.tsv$")
for(x in 1:length(file.names)){
  gc <- read.delim(file.names[x], header=F)
  # for 1 - max V2
  TP=0
  FP=0
  NumClust <- max(gc$V2)
  for ( cl in 0:NumClust) {
    if ( cl %in% gc$V2 ) {
      v <- as.vector( gc$V1[ gc$V2 == cl ] )
      t <- sort( table( v ), decreasing=T )
      best <- as.integer( t[1] )
      cID <- names( t[ 1 ] )
      if ( cl == 0 ) {
        if ( cID == "shuffled" ) {
          FN <- length(v)-best
          TN <- best
        }
        else
          cat("Houston, we have a TN problem")
      }
      else {
        if ( cID == "shuffled" ) {
```

```

        FP = FP + length(v)
    }
    if ( is.na( as.integer( t[2] ) ) || as.integer( t[2] ) < best ) {
        TP = TP + best
        FP = FP + length(v)-best
    }
    else if ( as.integer( t[2] ) == best ) {
        # treat both as false positives
        FP = FP + length(v)
    }
}
}
TP
TN
FP
FN
SENS=TP / (TP + FN )
SENS
SPEC=TN / ( TN + FP )
SPEC
ACC=(TP + TN) / ( TP + TN + FP + FN )
ACC
cat(file.names[x],TP,TN,FP,FN,SENS,SPEC,ACC, "\n",sep="\t",
     file="accuracies.tsv", append=T)
}

```

3. eCLIP data processing

Data in .bigBed format was acquired from the ENCODE data hub from the following link:
https://www.encodeproject.org/search/?type=Experiment&assay_term_name=eCLIP&files.file_type=bigBed+narrowPeak&month_released=April%2C+2016

```

# Convert accessions to protein IDs
cut -f 1,16,29 metadata.tsv | sed 's/-human _/g' | while read line
do
    F1=$(echo $line | awk '{print $1".bed"}')
    F2=$( echo $line | awk '{ print $2".bed"}')
    cp $F1 $F2
done

# Rename files accordingly
for file in *bed
do
    mv $file $(head -n 1 $file | cut -f 4).bed
done

# Filter for greater than or equal to 8x fold enrichment
# And -log10( P-value ) greater than or equal to 4
for file in *rep0?.bed
do
    awk '{if ($7 >= 4 && $8 >= 4) print }' $file > ../filtering/${file}_filt3
done

#Intersect both replicates (>1 overlap)
for file in *rep01.bed_filt3 ; do
    >&2 echo "Processing "$file
    bedtools intersect -s -u -f 0.5 -a <( cut -f 1-6 $file ) -b <( cut -f 1-6
    ${file//rep01/rep02} ) > ${file}_1

```

Supplementary Table 1. List of RFAM families from benchmark that did not cluster

Sequence count	RFAM ID	RFAM family
2	RF00005	tRNA
5	RF00015	U4 spliceosomal RNA
8	RF00020	U5 spliceosomal RNA
5	RF00021	Spot 42 RNA
1	RF00026	U6 spliceosomal RNA
10	RF00059	TPP riboswitch (THI element)
5	RF00167	Purine riboswitch
11	RF00169	Bacterial small signal recognition particle RNA
13	RF00199	SL2 RNA
4	RF00374	Gammaretrovirus core encapsidation signal
11	RF00378	Qrr RNA
6	RF00386	Enterovirus 5' cloverleaf cis-acting replication element
6	RF00389	Bamboo mosaic virus satellite RNA cis-regulatory element
4	RF00444	PrrF RNA
17	RF00494	Small nucleolar RNA U2-19
2	RF00515	PyrR binding site
4	RF00550	Hepatitis E virus cis-reactive element
7	RF01685	6S-Flavo RNA
7	RF01697	Chlorobi-RRM RNA
6	RF01705	Flavo-1 RNA
4	RF01725	SAM-I/IV variant riboswitch
2	RF01728	STAXI RNA
7	RF01734	crcB RNA
1	RF01750	pfl RNA
6	RF01754	radC RNA
4	RF01764	yjdF RNA
5	RF02033	HNH endonuclease-associated RNA and ORF (HEARO) RNA

```

bedtools intersect -s -u -f 0.5 -b <( cut -f 1-6 $file ) -a <( cut -f 1-6
${file//rep01/rep02} ) > ${file}_2

# merge peaks if they are close together
bedtools merge -d 50 -s -delim " | " -c 4,5,6 -o first,count,first -i <( cat
${file}_1 ${file}_2 | sort -k 1,1 -k 2,2n ) >
${file%*.bed_filt3}_filt_0.5_merged_50_s.bed

# intersect with ECS (in file ECS_congruous_sorted.bed6)
bedtools intersect -wo -s -b ${file%*.bed_filt3}_filt_0.5_merged_50_s.bed
-a ECS_congruous_sorted.bed6 >
${file%*.bed_filt3}_filt_0.5_merged_50_s_anyECS.bed
done

#merge all files into one
cat *_50_s_anyECS_merged.bed > All_ECS_merged_50nt_peaks.bed
# wc -l All_ECS_merged_50nt_peaks.bed
## 2650

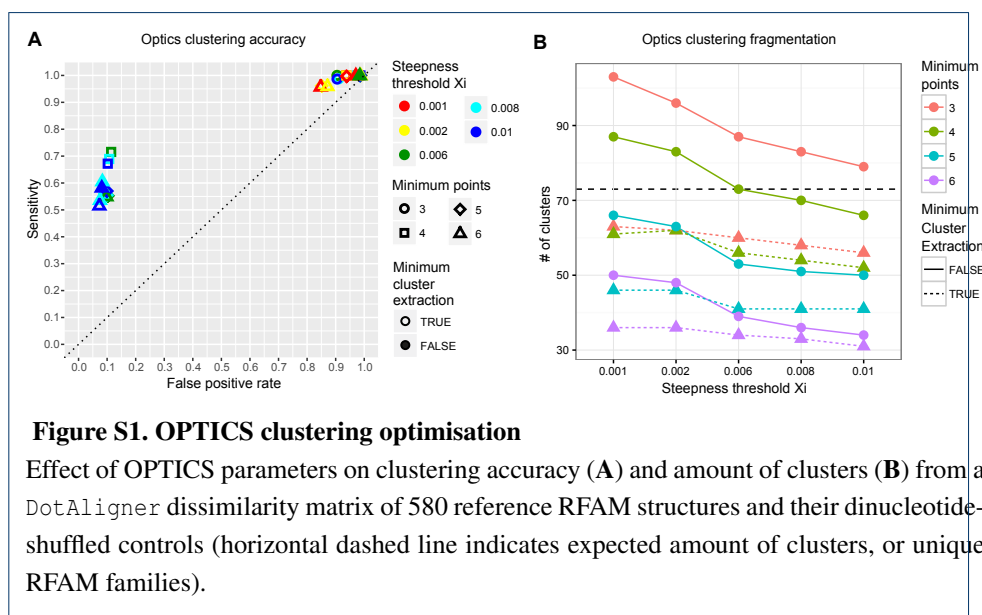
#edit sequence names and get sequence from reference genome (hg19)
awk 'OFS="\t">{print $1,$2,$3,$4"_"$1"_"$2"_"$3"_"$6,$5,$6}'
./All_ECS_merged_50nt_peaks.bed > ./All_ECS_merged_50nt_peaks_renamed.bed
bedtools getfasta -s -name -fi ~/data/fasta/hg19.fa -bed
./All_ECS_merged_50nt_peaks_renamed.bed -fo
./All_ECS_merged_50nt_peaks_renamed.fasta

#combine with known control RNA structure
cat All_ECS_merged_50nt_peaks_renamed.fasta spike-ins.fasta >
All_ECS_merged_50nt_peaks_renamed_spikeIns.fasta

```

Supplementary Table 2. List of control RNA structures

Sequences	RNA family	RFAM ID
5	5SRNA	RF00002
8	SNORAA72	RF00138
10	SNORD113	RF00181
10	SNORU3	RF00012
10	SNORU8	RF00096
8	SNR5	RF01252
9	YRNA	RF00019
10	mir19	RF00245
7	mir2968	RF02093
6	mir29852	RF02095
17	tRNA	RF00005

**Figure S1. OPTICS clustering optimisation**

Effect of OPTICS parameters on clustering accuracy (A) and amount of clusters (B) from a DotAligner dissimilarity matrix of 580 reference RFAM structures and their dinucleotide-shuffled controls (horizontal dashed line indicates expected amount of clusters, or unique RFAM families).