RNAs Everywhere: Genome-Wide Annotation of Structured RNAs

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ABSTRACT Starting with the discovery of microRNAs and the advent of genome-wide transcriptomics, non-protein-coding transcripts have moved from a fringe topic to a central field research in molecular biology. In this contribution we review the state of the art of "computational RNomics", i.e., the bioinformatics approaches to genome-wide RNA annotation. Instead of rehashing results from recently published surveys in detail, we focus here on *the* open problem in the field, namely (functional) annotation of the plethora of putative RNAs. A series of exploratory studies are used to provide non-trivial examples for the discussion of some of the difficulties. *J. Exp. Zool.* (*Mol. Dev. Evol.*) 308B:1–25, 2007. © 2006 Wiley-Liss, Inc.

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INTRODUCTION

A series of recent studies of the mammalian transcriptome have dramatically changed our perception of genome organization. Experimental studies using a variety of different techniques, from tiling arrays (Bertone et al., 2004; Kampa et al., 2004; Cheng et al., 2005; Johnson et al.,

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2005), to cDNA sequencing (Okazaki et al., 2002; Imanishi et al., 2004; Carninci et al., 2005; Ravasi et al., 2006), and unbiased mapping of transcription factor binding sites (TFBSs) (Cawley et al., 2004) agree that a substantial fraction of the genome is transcribed and that non-protein-coding RNAs (ncRNAs), Table 1, are the dominating component of the transcriptome. It remains unclear, however, to what extent these ncRNAs are functional; alternatively they might be "transcriptional noise" (Hüttenhofer et al., 2005) or they could be the by-product of transcriptional activity that takes place in order to regulate gene expression at adjacent loci. As shown by Ravasi et al. (2006), however, many non-coding cDNA clones are "derived from genuine transcripts of unknown function whose expression appears to be regulated".

Non-coding RNAs form a very heterogeneous group of transcripts: Besides the well-characterized "ancient" classes (such as the spliceosomal RNAs and tRNAs), the function of several pol-III transcripts remains unknown. Vault RNAs (Mossink et al., 2003; van Zon et al., 2003) seem to play a critical role in multi-drug resistance (Gopinath et al., 2005) and Y RNAs (Maraia et al., '94; Farris et al., '99; Stein et al., 2005) control the activity of RNA chaperones as Ro60 and La (Belisova et al., 2005; Stein et al., 2005).

Several ncRNAs exhibit more or less strong similarity to retroelements. In mammals, SINEs are derived from tRNAs and 7SL RNAs and LINEs from tRNAs (Deininger and Batzer, 2002; Kramerov and Vassetzky, 2005). Both are able to serve as source for new ncRNAs, as shown for a set of microRNAs (Smalheiser and Torvik, 2005; Tanzer et al., 2005) as well as 4.5SH RNA (Gogolevskaya et al., 2005) and 4.5SI RNA (Gogolevskaya and Kramerov, 2002) in rodents. Interestingly, the ncRNAs are derived from the long terminal repeats of LINEs, not from their protein coding regions. The small RNA generating loci in Arabidopsis follow a similar principle: inverted duplication of target genes leads to new miRNAs (Allen et al., 2004a).

Genes annotated as, e.g., "Putative ORF" are good candidates for so-called mRNA-like-ncRNAs

(mlncRNAs). These transcripts are processed just as normal mRNAs, but carry only very small ORFs or no ORFs at all. Transcriptional control (Berteaux et al., 2004, 2005; Carninci et al., 2005).tissue-specific differential expression (French et al., 2001), alternative splicing and polyadenylation (Sawata et al., 2004) of mlncRNAs do not seem to differ from those of protein coding polymerase II products, but some of them remain in the nucleus (Sawata et al., 2004). If amino-acid sequences are predicted for such transcripts, they are usually not conserved within a genus (Inagaki et al., 2005). Several miRNAs (Rodriguez et al., 2004; Baskerville and Bartel, 2005) and snoRNAs (Pelczar and Filipowicz, '98; Makarova and Kramerov, 2005) reside in introns and even exons of mlncRNAs. A few examples of functional ncRNAs that changed their host genes have been reported, see e.g. (Rodriguez et al., 2004; Bompfunewerer et al., 2005). Only a hand full of mlncRNAs are annotated in databases as in Y2K (Erdmann et al., '99, 2000) or RNAdb (Pang et al., 2005), while most cDNAs that lack a coding sequence (CDS) remain functionally unassigned (Carninci et al., 2005).

Long ncRNAs, such as *rox* in Drosophila or *Xist* in mammals are key players in imprinting and gene dosage compensation (Akhtar, 2003). The observation that *cis*-regulatory trithorax response elements in the Drosophila *Ubx* gene are transcribed as kilobase-sized ncRNAs, and that transcription activation occurs by recruiting a non-DNA-binding epigenetic regulator to the template sequence suggest that direct DNA-RNA interactions may also have an important general function in gene regulation (Sanchez-Elsner et al., 2006).

Small ncRNAs of only about 20 nt in length seem to serve as the exchangeable RNA module in protein complexes allowing them to bind DNA and RNA in a sequence-specific way. MicroRNAs, one of the most prominent classes of ncRNA, are found in plants (Jones-Rhoades et al., 2006) and animals (Berezikov and Plasterk, 2005; Massirer and Pasquinelli, 2006; Plasterk, 2006) and play a fundamental role in virus infections (Sullivan

Ancient RNAs	rRNAs, tRNAs, SRP RNA, RNase P
Repeat associated	miRNAs, rasiRNAs, 4.5SH RNA, 4.5SI RNA, LINEs, SINEs
mRNA-like	H19, AncR-1, Ntab, U87HG, BIC, Evf-1
mRNA-like associated	miRNAs, snoRNAs
Pol-III transcripts	snRNA, vRNA, Y RNA, tRNAs, MRP, U6, H1, 7SK, 7SL
Small RNAs	miRNAs, siRNAs, rasiRNAs, piRNAs

and Ganem, 2005; Nair and Zavolan, 2006). They differ slightly from siRNAs (Du and Zamore, 2005; Valencia-Sanchez et al., 2006). High expression of repeat-associated small RNAs (rasiRNAs) was detected during embryogenesis of *D. melanogaster* (Aravin et al., 2003) and later also in *Danio rerio* (Chen et al., 2005b).

A new class of presumably testes-specific small ncRNAs about 26-31 nt in length was detected in mouse, rat and human. Because of their association with members of the PIWI protein family. they have been termed "PIWI-interacting RNAs" (piRNAs) (Aravin et al., 2006; Girard et al., 2006; Grivna et al., 2006; Lau et al., 2006). Like siRNAs and miRNAs they show 5' phosphate and 3' hydroxyl groups indicating a specific but yet unknown maturation process from precursors rather than random degradation. Like miRNAs, piRNA clusters might be transcribed as long primary transcripts, which then are subjected to RNase III like enzymes. In contrast to microRNAs, however, piRNAs are not contained in miRNA like stem-loop structures or other conserved RNA secondary structures. The majority of piRNAs maps to dense, evolutionarily conserved intergenic regions. Notably, their genomic organization rather than the individual sequences seems to be conserved. Two functions were proposed so far: (1)a role in translational repression (Grivna et al., 2006: Lau et al., 2006) and (2) a possible function in pairing of homologous chromosomes and genome structure (Girard et al., 2006).

Recently, genome-wide surveys for non-coding RNAs have provided evidence for tens of thousands of previously undescribed evolutionary conserved RNAs with distinctive secondary structures (Washietl et al., 2005a; Pedersen et al., 2006). The conservation of structure indicates that the molecule functions (also) as an RNA. Taken together, both the experimental and computational data provide strong evidence that ncRNAs are an important, functional component of the mammalian transcriptome. The elucidation of these functions, however, remains elusive in almost all cases.

In this contribution, we discuss the currently available techniques for finding structured RNAs and we focus in particular on current approaches towards their annotation. In Figure 1, we propose a work flow for annotation of structured noncoding RNAs. The organization of this contribution largely follows the same outline. For quite a few of the individual tasks, there are no established software tools. In order to demonstrate the feasibility of this protocol, we therefore resort our own pilot studies. This manuscript is thus deliberately—organized as a somewhat unusual mixture of review (where possible) and original research. Additional material on these parts of the manuscript is provided in an electronic supplement.¹

COMPUTATIONAL ncRNA DETECTION

Large, highly conserved ncRNAs, in particular ribosomal RNAs, can easily be found using blast (Altschul et al., 1990). Similarly, blast can be used to find orthologous ncRNAs in closely related species, e.g. (Tanzer and Stadler, 2004; Weber, 2005). In most cases, however, this approach is limited by the relatively fast evolution of most ncRNAs. Since RNA sequence often evolves much faster than structure, the sensitivity of search tools can be greatly improved by using both sequence and secondary structure information.

Specialized search methods for particular RNA classes

Specialized programs have been developed to detect members of particular ncRNA families. Examples of this approach include miRseeker for microRNAs (Lai et al., 2003), BRUCE for tmRNAs (Laslett et al., 2002), tRNAscan for tRNAs (Lowe and Eddy, 1997), snoScan (Lowe and Eddy, 1999) and SNO.pl (Fedorov et al., 2005) for box C/D snoRNAs, fisher (Edvardsson et al., 2003) and snoGPS (Schattner et al., 2004) for box H/ACA snoRNAs, as well as a heuristic for SRP RNAs (Regalia et al., 2002; Rosenblad et al., 2004).

MicroRNAs in plants can be found by extracting those hairpin structures that contain sequence motifs complementary to an mRNA, which is then a putative target (Bonnet et al., 2004; Jones-Roades and Bartel, 2004; Adai et al., 2005). In animals, on the other hand, the situation is more complicated since miRNAs do not bind with perfect complementarity to their target. A large array of different approaches, summarized in Table 2, has recently been developed to detect microRNAs, among them our own tool RNAmicro that has been designed specifically to analyze large-scale comparative genomics data (Hertel and Stadler, 2006).

 $^{^1\}mathrm{URL:}$ www.bioinf.uni-leipzig.de/Publications/SUPPLEMENTS/ 06-005/



Fig. 1. Flow chart for the annotation work flow of structured non-coding RNAs. RNA families in the sense of the Rfam database are defined predominantly by sequence homology, while RNA classes are defined via functional and/or structural similarities that may or may not be the consequence of common ancestry. This diagram also serves as a rough outline of this contribution.

General methods for structure-based searches

A wide variety of different approaches to perform homology searches based on both sequence and structure have been proposed in the last few years in order to utilize the strong conservation of secondary structure in many ncRNA families, see Bompfünewerer et al. (2005) for a recent more extensive summary of this topic. Stochastic context-free grammars (SCFGs) can be used to construct covariance models from a multiple alignment with structural annotation as in infernal (Eddy, 2002). The consensus model can then be used to search for homologs. Similar in spirit, ERPIN also uses multiple structureannotated alignments as input to construct a descriptor for homology search. Rsearch (Klein and Eddy, 2003) is a local alignment algorithm which considers structural and sequence

Approach	miRNA only	Sequence	Structure	Homology	Mach.learning	Web tool	Download	Ref.	Remark
				;					
MiRseeker	Υ	Υ	Y	Υ	u	u	u	Lai et al. (2003)	Drosophilids only
MiRscan	Υ	Υ	Υ	Υ	n	Υ	n	Lim et al. (2003a,b)	Scores hairpins
ERPIN	n	Υ	Υ	Υ	I	Υ	Υ	Gautheret and Lambert (2001),	Scores structure profiles
								Legendre et al. (2005)	
infernal	u	Υ	Υ	Υ	SCFG	u	Υ	http://www.genetics.wustl.edu/ eddy/infernal/	1
HARVESTER	Υ	Υ	Υ	Υ	u	Υ	Υ	Dezulian et al. (2006)	Plants only
ProMiR	Υ	Υ	Υ	Υ	HMM	u	u	Nam et al. (2005)	
PalGrade	Υ	u	Υ	u	u	u	u	Bentwich et al. (2005)	Scores hairpins
mir-abela	Υ	Υ	Υ	*u	NVN	Υ	u	Sewer et al. (2005)	Scores hairpins, clusters
Vmir	Υ	u	Υ	u	I	u	u	Grundhoff et al. (2006)	Scores hairpins, clusters
	Υ	Υ	Υ	Υ	n	u	u	Berezikov et al. (2005)	Phylogenetic shadowing
BayesMIRfinder	Y	Υ	Υ	Υ	NBS	Υ	u	Yousef et al. (2006)	NBS first, then comparative
RNAz+RNAmicro	*	Υ	Υ	Υ	NVN	u	Υ	Washietl et al. (2005b),	Investigates alignments
								Hertel and Stadler (2006)	
HMM: Hidden M ¹ Infernal is used	arkov Model, NB egularly for hom	S: Naïve Bay nology search	es Score, SCF with the Rfa	FG: Stochastic m database.	: Context Free Gr	ammar, SVI	1: Support V€	sctor Machine.	

constraints. It uses both single nucleotide and base pair substitution matrices to define alignment scores. Rsearch operates on a single input sequence rather than on an alignment. FastR (Bafna and Zhang, 2004) combines a pairwise alignment algorithm with a filtering step to improve performance. Backofen and Will (2004) introduced an efficient local sequence-structure alignment method based on predicted structures. Beside sequence-local motifs (i.e., motifs that consist of a subsequence in each molecule), it is able to find also structure-local motifs, i.e. motifs that are connected substructures such as a helix without the connecting hairpin loop.

Simple description languages have been proposed to allow users to define combined sequence/ structure for genome-wide searches. Such approaches are implemented e.g. in RNAmot (Gautheret et al., '90) and Sean Eddy's rnabob. Hybrid languages, like HyPaL (Gräf et al., 2001) or the language used in RNAMotif (Macke et al., 2001), connect pattern languages with userdefined approximate rules, which rank the results according to their distance to the motif.

A number of large-scale surveys have been performed using one or more of the generalpurpose tools mentioned above, including a micro-RNA survey using ERPIN (Legendre et al., 2005), a search for U5 snRNA and RNase P using RNAmotif (Collins et al., 2004), and a survey of RNase P RNAs in bacterial genomes (Li and Altman, 2004).

Alignment-based de novo prediction of structured RNA motifs

Attempts to predict novel functional RNAs are in general based on predicted secondary structures. However, since most RNA sequences will form extensive structures, the problem of distinguishing incidental from functional structures is non-trivial. It was first suggested by Maizel and co-workers that functional RNA elements should have a secondary structure that is energetically more stable than expected by chance (Le et al., 1988). However, Rivas and Eddy had to conclude in an in-depth study on the subject that thermodynamic stability alone is generally not statistically significant enough for reliable ncRNA detection (Rivas and Eddy, 2000).

Therefore, all current approaches to de novo prediction of structured RNAs work comparatively, requiring two or more related sequences as input, typically in the form of a multiple sequence alignment. The first reasonably successful attempt to predict structured RNAs from sequence alignments was grna (Rivas and Eddy, 2001). This program compares the score of three distinct models of sequence evolution to decide which one describes best the given alignment: a pair SCFG is used to model the evolution of secondary structure, a pair hidden Markov model (HMM) describes the evolution of protein coding sequence, and a different pair HMM implements the null model of a non-coding sequence. Qrna was successfully used to predict ncRNAs candidates in E. coli and S. cerevisiae (Rivas et al., 2001; McCutcheon and Eddy, 2003), some of which could be verified experimentally. EvoFold (Pedersen et al., 2006) is essentially an extension of the grna approach to multiple sequence alignments. The program combines SCFGs for RNA structure modeling with phylogenetic models that describe the substitution process along the branches of a tree.

The RNAz algorithm, in contrast, is based on thermodynamic RNA folding (Washietl et al., 2005b). It uses two independent criteria for classification: a z-score measuring thermodynamic stability of individual sequences and a structure conservation index (SCI) obtained by comparing folding energies of the individual sequences with the predicted consensus folding. The two criteria are combined by a support vector machine that detects conserved and stable RNA secondary structures with high sensitivity and specificity. Other recent programs for detecting conserved RNA secondary structures include ddbRNA

(di Bernardo et al., 2003) and MSARi (Coventry et al., 2004).

Both RNAz and EvoFold have been applied to surveying the human genome providing evidence for tens of thousands of genomic loci with signatures of evolutionarily conserved secondary structure (Washietl et al., 2005b; Pedersen et al., 2006). Further RNAz surveys have been conducted for urochordates (Missal et al., 2005), nematodes (Missal et al., 2006) and yeasts (Steigele et al., 2006). These investigations have produced extensive lists of candidates for functional RNAs without using (or providing) information on membership in a particular class of RNAs, see Figure 2.

Alignment-free approaches

Approaches based on pairwise or multiple sequence alignments are of course limited by the quality of the input alignment. In regions with sequence similarity below some 50–60%, sequence alignments will in general not be structurally correct, making accurate prediction of consensus structures impossible (Washietl and Hofacker, 2004). This problem can in principle be overcome by computing structural alignments, albeit at significantly higher computational cost. Most recently, Uzilov et al. (2006) presented a classification method based on an updated version of dynalign (Mathews and Turner, 2002), a restricted variant of the Sankoff algorithm for the simultaneous computation of alignment and consensus structure (Sankoff, 1985). Using a similar



Fig. 2. Summary of a comparative screen of vertebrate genomes, which evaluated conserved genomic DNA sequences for signatures of structural conservation of base pairing patterns and exceptional thermodynamic stability using the RNAz program. (Adapted from Washietl et al., 2005a). About half of the structured RNA motifs are found far away from known coding regions, the other half are located within known protein-coding genes. Two thirds of the latter motifs are intronic, one sixth each is located in the 5'-UTRs and 3'-UTR of the mRNA, respectively.

approach based on their foldalign variant of the Sankoff algorithm, Torarinsson et al. (2006) screened a significant fraction of the non-alignable DNA that could be identified as homologous between man and mouse by virtue of alignable flanking sequences. These authors reported several thousands of regions without significant sequence conservation that show evidence for a conserved RNA secondary structure.

A major limitation of these "Sankoff-based" algorithms is their enormous computational cost. The computations required to evaluate the \approx 100,000 genomic regions as described by Torarinsson et al. (2006) took 5 months on 70 CPUs with 2 gigabytes of RAM. Uzilov et al. (2006) estimate that it would take several months on a similar sized computing cluster to screen all human/mouse regions with pairwise identity below 50% using dynalign. Although such approaches are feasible given sufficient computational resources, the CPU requirements render them impracticable for extended analysis tasks. Moreover, since both dynalign and foldalign perform pairwise alignments only, one would have to screen many different pairwise combinations (e.g. human/mouse and human/rat). A related approach (Bafna et al., 2006) matches entire stacks instead of individual base pairs.

Several faster methods for performing structural alignments exist. These approaches often start from predicted structures in the form of minimum energy structures, e.g. RNAforester (Höchsmann et al., 2003) or MARNA (Siebert and Backofen, 2005), or from lists of likely helices such as SCARNA (Tabei et al., 2006). Methods that rely of predicted structures as input are of course plagued by the inevitable inaccuracies of RNA folding algorithms. In contrast, the RNAcast program (Reeder and Giegerich, 2005) is based on coarse grained structures, so-called abstract shapes, and avoids the problem of aligning the sequences altogether. For each sequence it computes near-optimal shapes and then selects the best shape common to all sequences as consensus shape.

None of these approaches are feasible at genomic scales, however, without first identifying homologous regions of moderate size. This could be achieved for instance by extracting intergenic regions between pairs of homologous flanking genes. Alignment-based methods such as qrna, RNAz and EvoFold are of the same algorithmic time complexity (essentially cubic with alignment length). Although effective run time of the different programs may vary considerably, all three programs seem to be fast enough to allow routine analysis of even large mammalian genomes.

LIMITATIONS OF SEQUENCE ALIGNMENTS: AN RNAZ SCREEN OF STRAMENOPILES

To-date, genomic screens for non-coding RNAs have been applied mostly to fairly closely related organisms, e.g. vertebrates (with a focus on mammals), rhabditid nematodes or ascidians. In principle, however, the applicability of RNAz is limited only by the quality of the input alignments, so that highly conserved structures from distant organisms might still be detectable.

As an example of an RNAz screen of phylogenetically very distant organisms, we summarize a survey of the three currently available stramenopile sequences. Heterokonts, or stramenopiles, form a major clade within the eukaryote kingdom chromista, see e.g. Yoon et al. (2002). Most are algae, ranging from the giant multicellular kelp to the unicellular diatoms. However, some are colorless and superficially resemble fungi. Three complete genomes have been sequenced: data are available for two closely related oomycetes Phytophthora sojae,² Phytophthora ramorum³ (Gajendran et al., 2006) and the diatom Thalassiosira pseudonana⁴ (Armbrust et al., 2004). Our protocol closely follows the approach taken in Missal et al. (2005, 2006).

In the first step an annotation track for *P. sojae* was constructed by mapping the available mRNA and protein sequences back to the genome using blat. Using blast with a $E < 10^{-10}$, all nonprotein-coding loci were compared to the entire P. ramorum genome. We combine blast alignments that are separated not more than 30 nt provided they pass several consistency checks detailed in Missal et al. (2005, 2006). This leaves 149,375 conserved loci with an average length of 195 nt. Since the two phytophthora sequences are too similar, we estimate an unacceptably high estimated false discovery rate for the pairwise RNAz screen. We therefore compare these loci to the much more distant diatom T. pseudonana and obtain 903 homologous non-coding loci with an average length of about 80 nt. We realigned the blast hits using clustalw (Thompson et al., 1994) and screened them using RNAz with window-length

²http://genome.jgi-psf.org/sojae1/sojae1.home.html

³http://genome.jgi-psf.org/ramorum1/ramorum1.home.html

⁴http://genome.jgi-psf.org/thaps1/thaps1.home.html

120 nt in steps of 50 nt (for alignments longer than 120 nt). For some loci more than one alignment is found. These are combined if possible; otherwise only the alignment with the largest RNAz P-score is retained so that each genomic locus is covered by at most one RNA prediction. Details of this procedure are given in Missal et al. (2006). In order to estimate the false positive rate and the false discovery rate, the RNAz screen was repeated with shuffled alignments as described by Washietl et al. (2005a) and Missal et al. (2006). The results are summarized in Table 3.

The 115 RNAz slices that are classified with $p_{\text{RNAz}} > 0.5$ map to only 44 distinct loci in the *P. sojae* genome. Twenty of these can be identified as tRNA genes. A comparison with the updated annotation at the JGI *P. sojae* site shows that the remaining loci map to protein-coding regions. Given the data in Table 3, we expect a substantial false discovery rate. Furthermore, there is growing evidence for evolutionarily conserved secondary structure also within the coding parts of mRNAs (Meyer and Miklós, 2005; Steigele et al., 2006), so that some of these signals could well be real.

Due to the high degree of sequence divergence, most of the known ncRNAs do not lead to significant alignments of sufficient length between *P. sojae* and *T. pseudonana*. This set includes about 60 loci in the *P. sojae* genome that can be identified by comparison with the noncode database. Among them are 13 U2, 30 U4, 1 U5, 1 U6 snRNA and 1 SRP RNA. In addition, tRNAscanSE predicts 235 tRNA loci.

The low sensitivity of the screen on this data set highlights the limitations of approaches that are based on sequence alignments. With genome sizes of 33–87 Mb, using a structure-based approach (e.g. dynalign or foldalign) requires excessive computational resources. As more sequenced genomes become available, however, the scope of sequence-alignment-based methods expands for two reasons: (1) The specificity of methods such as

 TABLE 3. Summary statistics of the RNAz screen of stramenopiles

Threshold	$P \! > \! 0.50$	P > 0.90	$P \! > \! 0.98$	P > 0.99
Specificity per test	0.9861	0.9949	0.9985	0.9996
Candidate alignments	115	60	42	35
Randomized	37	14	4	1
False discovery rate (%)	32	23	10	3
Distinct loci (P. sojae)	44	17	12	11

RNAz increases dramatically with the number of aligned sequences. (2) Additional genomes in a suitable evolutionary distance from the currently available ones can give very good results already from pairwise comparisons as demonstrated in the case of ascidians (Missal et al., 2005) and nematodes (Missal et al., 2006).

THE IMPORTANCE OF BEING LOCAL

A comprehensive understanding of structured RNAs requires the analysis not only of ncRNAs with an often globally conserved structure, but also of local RNA motifs in larger molecules. Examples of the latter class are IRES (internal ribosome entry sites), selenocystein insertion elements or the *Rho*-independent termination signal in *E.coli*.

From a computational genomics point of view, there is actually little difference between these two classes of RNA structures. In a large-scale screen of genomic sequence, the transcript structure is typically unknown. As a consequence, both ncRNAs and structural mRNA motifs appear as local features in the genomic input sequence. The ability to compute *locally stable* secondary structures is thus a necessary prerequisite for any genome-wide analysis of structured RNA for both computational and biological reasons: (i) Longrange base pairs in large transcripts are disfavored kinetically (Flamm et al., 2000) (ii) even in long RNAs most base pairs are local (in 16S and 23S rRNA 75% of pairs span less than 100 nt) and longrange base pairs predicted by energy minimization are very inaccurate (Doshi et al., 2004). (iii) Global approaches to RNA folding are limited to sequence length $\leq 20,000$ on most hardware because of memory consumption. (iv) In general, the exact boundaries of the transcript are unknown, so that global folds cannot add to the accuracy of the structure prediction relative to folding individual sequence windows.

Local folds can trivially be obtained by folding subsequences of length L in a window sliding along the genomic sequence nucleotide by nucleotide. In practice, however, the sequence windows have to be shifted by a substantial fraction of L in order to keep the CPU requirements manageable. It is well known, however, that the predicted structures depend strongly on the flanking context, i.e., on the exact window position. In fact, a recent algorithm for microRNA detection is based upon the idea to consider the stability of secondary structure against changes in the immediate environment (Sewer et al., 2005). A large step size for the window implies poor sampling of the plausible local structures, hence small step sizes are important for accuracy.

Combining a global folding algorithm with a sliding window is also problematic in the context of ncRNA detection using tools such as RNAz: Large window sizes are preferable in order to detect larger ncRNAs, but may actually be detrimental for detecting small RNA structures, since the flanking regions interfere with the signal from the small structured RNA.

Two modifications of the global RNA folding algorithm have been developed to address this problem. RNALfold computes local minimum free energy structures with base pairs spanning no more than L bases in $\mathcal{O}(N \times L^2)$ time (Hofacker et al., 2004b). This is equivalent to folding all windows of size L, while saving a factor L in CPU time compared to the naive approach. A partition function variant with the same time complexity, RNAplfold, computes the probability of a base pair (i, j) occurring in the structural ensemble, averaged over all sequence windows with a given size W (Bernhart et al., 2006). To get robust statistics, the size of the averaging window W should be chosen somewhat larger than the maximum span L, resulting in an algorithm with complexity $\mathcal{O}(N \times W^2)$, see Figure 3.

Both RNALfold and RNAplfold are true "scanning algorithms"; requiring only $\mathcal{O}(N + L^2)$ memory, and are therefore suitable for genome-wide surveys. Together with a *z*-score for the energy of a sequence window, which could be cheaply computed in the course of the algorithm (Washietl et al., 2005b), RNAplfold may be used as a first crude indicator whether stable RNA secondary structures can be expected in a given part of the genome.

In principle, it is straight-forward to generalize the prediction of consensus structures from aligned sequences from global structures to local structures. RNAalifold (Hofacker et al., 2002) computes the most stable structure that is common to a collection of aligned sequences. Algorithmically, only the energy model changes: one simply has to add up the contributions of aligned sequence intervals instead of evaluating a single sequence. This modification can easily be implemented also in the scanning programs described in the previous paragraph. The resulting RNALalifold program will be available with the next release of the Vienna RNA package. An example for the partition function case is given



Fig. 3. (a) Local pair probabilities of a human miRNA cluster. (b) Homologs in dog (upper), opossum (middle) and mouse (lower) to the miRNA cluster in (a). MiRNAs annotated in miRBase are outlined in full lines, putative miRNAs un-annotated in dashed lines. Note that every putative miRNA almost perfectly aligns to the human counterpart (with one mutation at most). Dog, opossum: W = L = 100, Mouse: W = 150, max. base pair span L = 100. The base pairs on the upper edge of the mouse plot (long-range base pairs) are less probable than in the other three because of the more robust statistics. (c) RNALalifold partition function of an alignment of the homologous miRNA clusters in (a) and (b). The noise is considerably reduced as opposed to the single folds shown above.

in Figure 3, where the conservation of the miRNA precursor structures, as opposed to any other structural features present is shown.

Foldalign (Hull Havgaard et al., 2005; Torarinsson et al., 2006) implements a scanning local alignment algorithm. More precisely, this version of the Sankoff algorithm mutually scans two sequences of arbitrary length for common local structures with a maximum motif length. While the restriction to local motifs speeds up the algorithm, it is still computationally demanding.

In principle, also the SCFG-based algorithms QRNA and EvoFold predict local secondary structures. However, both are not implemented as true "scanning algorithms", and thus still require a sliding window approach.

WHO IS WHO? — APPROACHES TOWARDS RNA ANNOTATION

The problem

With the exception of a small number of evolutionarily very well conserved RNAs (in particular rRNAs, tRNAs (Lowe and Eddy, 1997), the U5 snRNA (Collins et al., 2004), RNAse P and MRP (Piccinelli et al., 2005)), most ncRNAs are not only hard to discover de novo in large genomes, but they are also surprisingly hard to recognize if presented without annotation. While it is often impossible to use relatively faint sequence homologies to find homologs of known RNAs in a whole genome, it is usually rather easy to recognize the same sequences in the output of genomic ncRNAs screens, due to the enrichment of true RNAs by several orders of magnitude. As a consequence, we found in all our RNAz screens, that a comparison with Rfam alignments using infernal identifies very few RNAz hits that are not already recognizable by blast. While homologs of known sequences can usually be reliably recognized, determining class membership of novel examples is a much harder problem.

Given an alignment not more than a few hundred nucleotides in length that is known to contain a conserved secondary structure, it should be very easy to decide whether these sequences belong to a known class of ncRNAs or not. Conceptually, this is a very simple classification task that should be solvable efficiently by most machine learning techniques.

In the case of non-coding RNAs, however, machine learning approaches severely suffer from the very limited amount of available positive training data and the fact that negative training data are almost never known at all. Even for the most benign case, microRNA precursors, there is only a few hundred independent known examples, namely the miRNA families listed in the mir-base (Griffiths-Jones, 2004; Griffiths-Jones et al., 2005; Griffiths-Jones, 2006; Hertel et al., 2006). Overtraining is thus a serious problem. As a consequence, it is necessary to restrict oneself to a small set of descriptors. These constraints, however, make the choice of the descriptors a crucial task.

Which direction?

A relatively simple example for such a classification task is the problem of strand prediction: Large parts of the RNA energy model, in particular the stacking energies for Watson-Crick pairs, are symmetric when forming the reverse complement of a sequence and its structure. Asymmetry is introduced in particular by GU pairs that map to a non-canonical AC in the reverse complement. Nevertheless, plus and minus strand of a sequence often exhibit similar folding energies. In computational screens for ncRNAs one will usually look at both the forward and reverse version of any given alignment, and often a significant signal for a structural RNA is detected on both strands. RNAz so far simply estimated the reading direction as the one that achieved a higher classification probability for "structured RNA". This method is quite inaccurate, however, in particular when the differences in classification probability are small.

An efficient strand detector to be used in conjunction with RNAz can be constructed from only six descriptors which moreover are already computed by RNAz: The difference of the SCI value for plus and minus strand, the difference of the RNAalifold consensus energy, the difference in mean folding energies, and the difference in mean z-score. In addition, the average sequence conservation and the length of the alignment is used. A support vector machine can then predict the correct strand with over 96% accuracy. This method is implemented also as a stand-alone tool RNAstrand (Missal and Stadler, submitted) that can be used to re-evaluate earlier ncRNA screens. Figure 4 shows one such example.

Family membership: H/ACA-Box snoRNAs

In order to assign predicted ncRNAs to a particular ncRNA family, it seems natural to include structural descriptors in the classification procedure. RNA structure prediction, however, is less than perfect even when co-variation information from an alignment can be used (Hofacker et al., 2002). This is true in particular when the exact ends of structured sequences within the multiple sequence alignment are not known. Furthermore, most ncRNAs can tolerate deviations from the "typical" structure without loss of function. The microRNA precursor structure may for example contain small branching helices, instead of forming a single stem-loop. These limitations restrict the usefulness of structure description languages, in particular, when one is interested in ncRNAs that are not members of one of the few well-known families.

Thus, structural descriptors have to be sufficiently fuzzy to allow for imperfect structure prediction and structural variation. The RNAmicro program for example uses 12 descriptors, only two of which are derived from the structure, namely the length of the stem and hairpin loop region of the miRNA stem-loop structure. Four descriptors measure sequence conservation in the loop and stem regions (the loop tends to be very variable, while the mature miRNA is highly conserved), another five descriptors measure the thermodynamic stability, and one measures sequence composition. This approach was quite successful, see e.g. Figure 5 for an example.

It seems thus natural to extend this approach to other classes of ncRNAs. During the work on RNAmicro we observed that H/ACA-box snoRNAs, which also form hairpin-like structures, formed a particularly resilient group of false positives. This suggests to use the same set of descriptors and simply train the system with multiple sequence alignments of H/ACA-box snoRNAs as positive training set, while a sample of randomized hairpins, microRNAs, as well as known stem-loop structures from other ncRNA classes are used as a negative training set. This yields a sensitivity of only about 63% at a specificity of about 75%. This suggests to include a small number of additional descriptors that are geared towards specific structural properties of box H/ACA snoRNAs discussed e.g. in (Henras et al., 2004).



Fig. 4. The RNAz prediction *Ce-512233* (Missal et al., 2006) coincides with *pre-mir-79*. The RNAz prediction favors the minus strand (top). The correct reading direction is on the plus-strand, however. RNAstrand computes a score of D = -0.82 for the RNAz hit, indicating that the direction predicted by RNAz is incorrect. The RNAstrand classification coincides here with the correct reading direction of the microRNA precursor.



Fig. 5. RNAmicro annotation (Hertel and Stadler, 2006) of a RNAz survey of nematode genomes (Missal et al., 2006). About half of the known *C. elegans* microRNAs are not conserved in *C. briggsae* and are hence not detected by comparative genomics. (Adapted from Hertel and Stadler, 2006).

The snowReport classificator uses nine descriptors, among them the same quantities for assessing folding thermodynamics as in RNAmicro: energy z-score, SCI, average folding energy of the individual aligned sequences, ratio of folding energy and GC content. The stem-loop structures of snoRNAs are significantly shorter than those of miRNAs. Thus we include the number of stacked pairs and the length of the hairpin loop. Furthermore, H/ACA box snoRNAs have a single large interior loop which is (nearly) symmetric. We hence add the average symmetry (absolute value of the length difference between the 3' and 5'unpaired stretches of all interior loops) as well as the length of the longest interior loop as additional descriptors. We use libsvm 2.8.2 with the same settings as RNAmicro, RNAz and RNAstrand.

In contrast to the microRNAs, a sufficiently large set of snoRNA alignments is not available, albeit there are several examples in the Rfam (Griffiths-Jones et al., 2005) and snoRNA-LBME-db (Lestrade and Weber, 2006) databases. We thus searched all available vertebrate genomes for homologs on the known human H/ACA-box snoR-NAs following the protocol for microRNA homology search used by Hertel et al. (2006). This yields 395 alignments containing 2–18 sequences per alignment.

Using half of this set as positive training set and half of the microRNA alignments reported by Hertel et al. (2006) as negative training set resulted in a sensitivity of 81% and a specificity of 87%. Afterwards, the test data were used for retraining the SVM and the training data for testing it. This resulted in a sensitivity of 91% and a specificity of 81%. For application to previously reported RNAz screens, we used the full sets of positive and negative examples for retraining. Unfortunately, snowReport misclassifies a large number of tRNAs and 5S rRNAs as snoRNAs. Closer inspection shows that in these cases only structures are recognized. Since the false positives appear to be restricted almost exclusively to these well-known ncRNAs, they do not present a serious problem since they are reliably identified by sequence homology or tRNAscanSE.

Metazoan box H/ACA usually are composed of two stem-loop structures (Henras et al., 2004); we hence classify only those RNAz hits as putative box H/ACA snoRNAs in which (1) two hairpins separated by not more than 20 nt are classified positively by snowReport, and which (2) contain the H-box motif. Results are summarized in Table 4. The most interesting observation of this

 TABLE 4. Application of snowReport on sea squirts, nematodes and vertebrates

Candidates	Urochordata	Nematoda	Vertebrata
SnowReport	1,553	1,833	5,519
True negatives	203	310	3
Distance constraint	14	255	17
H-box	14	204	12
Known HACA snoRNA	13	10	65
SnowReport	9	9	24

Numbers to distinct loci in the genomes of *C. intestinalis*, *C. elegans* and *H. sapiens*, resp. The false positives in urochordates and nematodes are tRNAs and rRNAs, which were excluded from the training data.

preliminary screen is the large number of plausible candidates in nematodes in contrast to both the urochordate and mammalian data. It is interesting to note in this context that the recent experimental screen by Deng et al. (2006) identified dozens of putative snoRNAs in *C. elegans*.

In contrast to snoGPS (Schattner et al., 2004), we do not rely on the existence of a known or suspected target site in an rRNA or snRNA. Our approach thus predicts a few plausible candidates of "orphan" snoRNAs, i.e., snoRNAs within unknown modification target site.

In contrast to RNAmicro, the SVM-based classification of box H/ACA snoRNAs was only moderately successful. The most significant problem appears to be the generally low quality of the predicted consensus structures, which seems to be at least in part a consequence of problems in the underlying sequence alignments. Reliable methods for structure-based or structure-assisted multiple sequence alignments are thus a necessary pre-requisite for the successful application of structural descriptors in automatic ncRNA annotation. Although several approaches exist, reviewed e.g. in a comparison of techniques for consensus structure prediction by Gardner and Giegerich (2004), their suitability for the purpose of ncRNA annotation has not vet been studied systematically.

snRNA-like candidates

A recent experimental survey of *C. elegans* genome (Deng et al., 2006) identified a class of snRNA-like ncRNAs that are characterized by a recognizable SMN-binding site. We have therefore reanalyzed the results of the RNAz screen of urochordates (Missal et al., 2005) to identify



Fig. 6. Secondary structure (top) and genomic location (bottom) of a putative snRNA-like RNA in *Ciona intestinalis*. The RNAz predictions 555,803 and 555,804 are located within two introns of an ENSEMBL gene, which match a single locus in the *Ciona savignyi* genome. It is the reverse complement of these two sequences, however, which contains the putative SMN binding site, which is highlighted in the secondary structure. Trimming the alignment to the three distinct sequences, two from *C. intestinalis* and a single one from *C. savignyi* so that only the well-conserved region is retained and rescoring with RNAz yields $p_+ = 0.709774$ and $p_- = 0.961678$. RNAstrand returns a decision of p = -0.999999, i.e., an unambiguous vote for the negative strand.

potential SMN binding sites in these structured RNA candidates.

We use RNAbob to search for the sequence motif AUUUYUS followed by a hairpin of rather variable stem and loop length. This pattern is a common generalization of the SMN binding sites in the known *Ciona intestinalis* snRNAs. In our analysis we require that the pattern occurs in aligned positions of the *C. intestinalis* and *Ciona savignyi* ncRNA candidates. This procedure recovers many of the known snRNAs that we found by the RNAz screen and in additions identifies 28 plausible candidates (as well as five copies of tRNA-Ile and one probable protein coding transcript). One example is described in some detail in Figure 6.

NEW KIDS ON THE BLOCK

Sequence-based clusters

The simplest approach to identifying multi-gene families is blastclust. A reinvestigation of the urochordate RNAz screen (Missal et al., 2005) shows that about a third of the candidates have at least one related sequence in the candidate set, Table 5. As one would expect, individual tRNA and snRNA families are identified by this approach. In addition, however, we find three very large families of candidates. They do not show significant homology outside the Ascidians and they are not associated with a known or predicted family of protein coding genes. The cluster members are not uniformly distributed across the genome but appear concentrated at a few genomic loci. This pattern is reminiscent of many groups of vertebrate ncRNAs, including in particular tRNAs and snRNAs, which appear in multiple, often genomically clustered, copies. Lineage-specific examples of functional repeat-derived ncRNAs include, e.g., the mouse B2-element (Allen et al., 2004b). Since the RNAz classification values are very high for most members of these classes, we speculate that these groups contain functional ncRNAs that are associated with an ascidian-specific repeat family.

A similar pattern was observed in *C. elegans* (Missal et al., 2006). With slightly different blastclust setting, 148 clusters containing a total of 916 RNAz signals as well as 2,756 non-clustered sequences were found. In contrast to the urochordate data, however, all large clusters could be annotated. It is not clear at this point whether this

Size	Annotation	Size	Annotation	Size	Annotation
197	(1)	13	tRNA Arg:CCT	8	tRNA Thr:AGT
160	(2)	13	(16)	8	tRNA Val:AAC
144	(3)	12	tRNA Gly:GCC	8	
32	5S RNA	11	tRNA Gly:TCC	8	
26	tRNA Ile:YAT	11	(19)		
22	(6)	11	(20)	Size	Frequency
20	(7)	9		7	7
18	tRNA Pro:HGG	9	U5	6	1
17	(9)	9		5	10
17	(10)	9		4	13
14	tRNA Leu:WAG	9		3	22
13	U3	8	tRNA Ala:WGC/Ser:GCT	2	83
13	tRNA Arg:ACG	8		1	2065
13	tRNA Leu:TAA	8	tRNA Asn:GTT		

TABLE 5. Sequence-based clustering of Ciona intestinalis ncRNA candidates

Here we used blastclust requiring a sequence overlap of \geq 50%, 80% identity in the overlap region, and word size of 20, i.e., much less stringent settings than the defaults. Numbers in parentheses refer to sequence families for which consensus sequences are provided in the electronic supplement.

difference is biologically meaningful, or whether sequences with high copy numbers have been excluded more effectively from the nematode screen as a consequence of more complete exclusion of repetitive DNA. Not surprisingly, no large sequence-based clusters were found in the mammalian screen (Washietl et al., 2005a) since in this case the input alignments were already devoid of multi-copy genes including tRNAs and snRNAs.

Structure-based clustering

A more general approach to assign ncRNAs to families is based purely on structural similarity. Given a set of predicted ncRNAs, one may use a structural alignment method to compute all pairwise alignments, and subsequently cluster all ncRNAs by similarity. In principle, this should allow not only to assign predicted ncRNAs to known families, but even to define complete new ncRNA families. For the pairwise alignment step one would ideally use a variant of the Sankoff algorithm which simultaneously computes sequence alignment and consensus structure, but is computationally expensive (Sankoff, 1985). Performing structural alignments for all pairs of ncRNA candidates in a set of several ten thousand is therefore still problematic. Moreover, most existing implementations can use only two sequences (no profile alignments) and compute global instead of local similarity. A local variant is described by Hull Havgaard et al. (2005).

The goal of annotation tools that classify family membership in results of other surveys is different from the direct search for RNA family members in genomic data. In the latter case one is interested in a "short list" of candidates that contains as few false positives as possible (e.g., for use in experimental verification). In post-processing data such as those from RNAz we are interested in a more balanced trade-off between sensitivity and specificity similar to that of annotating protein motifs in known predicted protein coding genes.

For the purpose of a structural clustering of ncRNA candidates, we suggest a pipeline consisting of the following three major steps:

- generate all pairwise local sequence/structure alignments,
- based on this information, hierarchically cluster the ncRNAs using WPGMA (or any other suited hierarchical clustering method) into a tree,
- finally, extract relevant clusters and construct multiple alignments of the ncRNA candidates in each cluster.

Recent developments in pairwise sequencestructure alignment allow us to get very close to the ideal of using Sankoff's algorithm and in the same time increase the efficiency dramatically. Hofacker et al., (2004a) proposed a (global) scoring scheme that is based on all base pair probabilities (in the structure ensemble) of the two RNAs. Such probabilities can be reasonably predicted using McCaskill's pair probability algorithm (McCaskill, 1990). Since probabilities reflect thermodynamical properties of the RNAs, the new scoring scheme factors in thermodynamics without the need of computing a full energy model during alignment. It turns out that this idea can be used to design an even more time- and space-efficient algorithm that can also be extended to local alignment (Will et al., 2006). The resulting new algorithm LocARNA is ready to manage the envisioned ten thousands of RNAs. As a test case we consider the 3.332 C. intestinalis ncRNA candidates from Missal et al. (2005). In contrast to the previous section. we used here a very stringent sequence-homologybased pre-processing set that identifies sequences with more than 90% identity. Pairwise structural alignments of the resulting 2,804 distinct sequences can be computed in about 2 days on 10 dual core CPUs (Fig. 7).

An appropriate distance measure that is based on both sequence and structure information is necessary for applying the weighted pair group method (WPGMA) or any comparable method for tree construction and cluster extraction. Since the pairwise alignments are computed together with their similarity scores, one might naively attempt to use these scores also for clustering. This is not appropriate, however, since the local scores reflect the quality of local structure prediction, not the similarity of the different alignments.

We therefore used the following normalized sequence and structure similarity measures of different alignments of RNAs. On the sequence level, we use the average sequence identity between the RNA sequences of the alignment. For the scoring of the structural similarity, we use the SCI, which is the ratio between the mean single minimum free energy (mfe) and the consensus mfe. The similarities are then transformed into distances and WPGMA is applied onto the resulting distance matrix to produce the final tree. Figure 8 shows a subtree that contains about half of the known tRNA precursors. With very few exceptions, the tRNAs are clustered according to their amino acid and anticodon, demonstrating that the procedure indeed yields plausible results.

The resulting tree is then cut at a specific threshold to generate the clusters, from which we can then extract a common motif using an appropriate multiple alignment method. Sequence identity in the identified clusters can be rather low due to the structure influence in the clustering (often below 60%). The approach is thus capable of identifying families of structured RNAs in a range where global multiple sequence alignment already yields very poor results.

Interactions with mRNAs

Regulatory RNAs more often than not function by means of direct RNA-RNA binding via complementary base pairing. This mechanism underthe post-transcriptional gene silencing lies pathways of microRNAs and siRNAs (reviewed e.g. by Nelson et al., 2003) as well as RNAi (Elbashir and Tuschl, 2001), it is crucial for snoRNA-directed RNA editing (Gott and Emeson, 2000), and it is used in the gRNA-directed mRNA editing in kinetoplastids (Stuart et al., 1997). A wide range of ncRNA regulation in bacteria is based upon RNA duplex formation (Gottesman, 2004). Synthetic "modifier RNAs" have been used as experimental techniques for changing the gene expression patterns independent of the RNAi pathway (see e.g. Childs et al., 2002; Meisner et al., 2004; Nulf and Corey, 2004; Paulus et al., 2004). Recent studies of the transcriptome of various organisms have uncovered ample evidence for wide-spread anti-sense transcription



Fig. 7. Pipeline for clustering a set of ncRNAs A,B,C,D and E. Starting from the RNAs with pair probability matrices, all pairwise alignments are computed, clusters determined and the RNAs of each cluster multiply aligned.



Fig. 8. Structure-based clustering of the 3,332 *Ciona intestinalis* ncRNAs candidates predicted by RNAz (Missal et al., 2005) yields (among others) this cluster comprising 157 of the 301 detected tRNAs that code for 17 different amino acids. Some nodes represent groups of almost identical sequences (>90% identity) of which only one representative has been used for clustering.

(Shendure and Church, 2002; Yelin et al., 2003; Chen et al., 2005a; Katayama et al., 2005; Steigele and Nieselt, 2005; David et al., 2006). These transcripts might at least in part be involved in RNA-RNA interactions.

MicroRNA-mRNA interaction is a rather special type of interaction mediated by the RISC complex, which at least in metazoa appears to be governed by rules that are only partially derived from the thermodynamics of RNA–RNA interactions, see e.g. (Brennecke et al., 2005; Isaac, 2005). Several well publicized tools are available for this task, an non-exhaustive list is given in Table 6. Since miRNA target prediction is not primarily concerned with the specific functions of the miRNAs rather than with the primary annotation tasks, we will not expand on this topic.

Beyond the realm of microRNAs, RNA–RNA interactions might be more direct and hence more directly related to the thermodynamics of RNA– RNA hybridization. In order to gather evidence for a possible function of the ncRNAs candidates in ncRNA-mRNA interactions, we investigated whether the RNAz predictions show an increased propensity of interacting with known mRNAs. In detail, we used the following procedure:

True and shuffled RNAz hits from both genomic strands were aligned to human mRNA sequences from RefSeq (NCBI FTP server, March 7) using NCBI blast (version 2.2.10 with standard parameters for blastn except an *E*-value cutoff of $E \leq 0.1$ and filtering set to false). The resulting alignment was filtered by removing all blast alignments with a length of less than 20 nucleotides or less than 75% sequence identity. Furthermore, alignments were retained only if the query matched the antisense strand of an mRNA. Of the 71,970 RNAz hits, this yields 11,112 (15.4%) true and 1,319 (1.8%) shuffled predicted antisense interactions. In the control consisting of 71,968 conserved non-coding DNA sequences that were part of the input set in the vertebrate RNAz screen,

Program	Source	Reference
PicTar	http://pictar.bio.nyu.edu/	Grün et al. (2005), Lall et al. (2006), Krek et al. (2005)
MiRanda	http://cbio.mskcc.org/mirnaviewer/	Enright et al. (2003)
TargetScan	http://genes.mit.edu/targetscan/index.html	Lewis et al. (2003)
RNAhybrid	http://bibiserv.techfak.uni-bielefeld.de/rnahybrid/	Rehmsmeier et al. (2004)
EMBL	http://www.russell.embl.de/miRNAs/	Stark et al. (2003), Brennecke et al. (2005)
DIANA-microT	http://diana.pcbi.upenn.edu/cgi-bin/micro_t.cgi/	Kiriakidou et al. (2004)
MicroInspector	http://mirna.imbb.forth.gr/microinspector/	Rusinov et al. (2005)
miRU	http://bioinfo3.noble.org/miRNA/miRU.htm	Zhang (2005)
MovingTargets	available on DVD by request	Burgler and Macdonald (2005)

TABLE 6. Tools for microRNA target prediction

we find 9,055 (12.6%) predicted interactions. This corresponds to an enrichment between true and random fractions of 1.22.

After blast search, 1,396 of true, none of the shuffled, and 1,108 of the control hits were removed because they overlapped the mRNA sequence that they matched. This step eliminates potential false positives, but might also exclude true *cis*-antisense transcripts. Interestingly, this step does not affect the enrichment factor of 1.22.

For each RNAz hit, the longest alignment was kept for further analysis. For these interacting pairs of RNAz hits and mRNAs, a coarse grained estimate of the interaction free energy (IFE) was computed using RNAduplex. This component of the Vienna RNA Package computes a simplified hybridization of two RNAs which allows only intermolecular base pairs (see also Rehmsmeier et al., 2004; Dimitrov and Zuker, 2004).

IFE distributions of the true and shuffled RNAz hits were tested against the null hypothesis of a common distribution using the Kolmogorov–Smirnov test from the statistical package R. The null hypothesis that both IFE distributions of shuffled and true RNAz sequence originate from a common distribution was rejected with $P < 10^{-16}$. The remaining RNAz sequences were co-folded with their potential target mRNAs using RNAduplex to determine IFEs. Density plots of the IFE distributions are shown in Figure 9.

Interactions were classified according to their calculated IFEs. Class I interactors have an IFE lower or equal to the empirical 0.05 quantile of the shuffled IFE distribution, class II interactors lower or equal to the 0.1 empirical quantile, class III interactors lower or equal to the 0.25 empirical quantile and class IV interactions lower or equal to the empirical median IFE of the shuffled RNAz hits. Results of this classification are given in Table 7. Only 13 of the 11,112 RNAz sequences retained after Blast search are known miRNAs, and nine of these are classified as interacting based on IFE. Four of these sequences have an entry in Tarbase (Sethupathy et al., 2006), listing experimentally verified miRNA target mRNAs and in all four cases our approach would have predicted the correct target. Four of the predicted interactors are snoRNAs, which is in line with other reports that snoRNAs may play a role in mRNA modification (Kishore and Sham, 2006).

Consistent with published data, we identify e.g. the interaction between *mir-196a* and its target mRNA HOXB8. This miRNA has an exceptionally high complementarity to its target mRNAs compared to other miRNAs (Yekta et al., 2004) and one would therefore expect to find a particularly strong interaction. Nevertheless, we classify this interaction as not significantly more stable than random. This may indicate that microRNA function is not governed by RNA-RNA interaction energy but is dominated by structural constraints imposed by the RISC complex. This view is consistent with the observation that most miR-mRNA interactions are far from exact complementarity (Du and Zamore, 2005). At this point we cannot rule out, however, that the interaction energy model used here is too crude to properly describe individual binding patterns.

We have identified here a large number of evolutionary conserved structured ncRNA candidate genes that interact with mRNAs significantly stronger than random sequences. Almost none of them belong to one of the established ncRNA families. This observation stimulates speculations on the functional role of these transcripts. Given the stable interactions, one might consider siRNAlike functions. Alternatively, it is conceivable that some of these genes act as "modifier RNAs" by influencing mRNA secondary structure



Fig. 9. Densities of interaction free energy distributions. The density of the interaction free energy distribution of true RNAz hit—mRNA interactions is shown in black, those of shuffled RNAz hits—mRNA interactions in brown. Dotted lines indicate the energy thresholds used for classification, at -215.82, -174.28, -126.91 and -91.80 kcal/mol, corresponding to the 0.05, 0.10, 0.25, 0.50 quantiles of the randomized distribution and defining classes I–IV respectively as interactions with energy lower or equal the threshold.

 TABLE 7. Interaction with mRNAs — relative to total number

 of RNAz hits

Interaction	True RNAz hits		Random RNAz hits		
class	Number	Fraction	Number	Fraction	Enr.
I	2,036	0.028	883	0.012	2.3
II	1,193	0.017	651	0.009	1.9
III	1,949	0.027	1,348	0.019	1.4
IV	2,087	0.029	1,861	0.026	1.1

Absolute numbers of RNAz hit sequences matching a particular interaction class and the respective fraction of total RNAz hit sequences (71,970 true, 71,968 random) are given. About 2,451 true and 3,204 randomized RNAz hits have an interaction energy smaller than the median of shuffled RNAz hit—mRNA interaction energies and are not mentioned in the table.

(Hackermüller et al., 2005). The fact that these ncRNAs are conserved in sequence and structure may suggest that other co-factors, such as proteins which recognize specific structured binding motifs, are involved in their function. It remains to be demonstrated whether these observed interactions are restricted to conserved structured RNAs or are also common among conserved nonstructured RNAs.

Structured RNAs are depleted in predicted TFBS

A recent study by Drake et al. (2005) demonstrated that evolutionarily conserved non-coding sequences are selectively constrained and thus can be expected to have discernible function(s). These sequences are most often interpreted as *cis*-acting DNA motifs. This class of functional sequence motifs consists in particular of binding sites for proteins involved in transcriptional regulation (Tagle et al., '88; Davidson, 2001; Butler and Kadonaga, 2002). In order to corroborate the fact that the RNAz predictions are indeed functional at the RNA level, we consider the distribution of known TFBSs within the RNAz candidates.

We consider a subset of 493 vertebrate TFBS patterns from the transfac database (Heinemeyer et al., 1998). These are mapped to the human sequence of every 10th alignment "slice" that scored as "structured RNA" in the mammalian RNAz screen by (Washietl et al., 2005a). For comparison, 10% of the negatively scored input alignments as well as shuffled datasets of both the positive and negative sets were used. The mapping was performed with pwmatch,⁵ a re-implementation of the scoring algorithm published by Kel et al. (2003), using a cut-off of 0:9. For simplicity, we will refer to these hits TFBSs in the following, irrespective of whether the detected sequence motif is a functional binding site in vivo or not.

We find that TFBSs are slightly enriched in true versus shuffled data sets. Furthermore, there is a small enrichment of predicted TFBSs in conserved non-coding DNA that is not classified as structured RNA (0.24 TFBS/nt) compared to the

⁵The pwmatch tool is available from www.bioinf.uni-leipzig. de/Software/pwmatch.

putative ncRNAs (0.20 TFBS/nt). Since randomized sequences have only a slightly smaller density of TFBS (0.18–0.19 TFBS/nt), we conclude this (high) background level is spurious, i.e., that most of the computationally predicted TFBS are not functional. The false discovery rate of the human RNAz screen was estimated on the order of 10% (Washietl et al., 2005a). The data are thus consistent with an increased frequency of TFBS in evolutionarily conserved non-coding DNA, while structured RNAs approximately behave like random background.

CONCLUDING REMARKS: UNSTRUCTURED RNAs

In prokaryotic genomes, the structure of genes, and in particular the promotor and terminator elements are sufficiently well understood that they help to detect non-coding genes independently of RNA structure or comparative sequence information. In eukaryotes, on the other hand, computational approaches to de novo ncRNA prediction are at present limited to structured RNAs.

A substantial number of mlncRNAs, including *Xist* and H19, appear to contain one or more domains with conserved RNA secondary structures, which can be detected (Washietl et al., 2005a). Without additional experimental information such as EST or cDNA data, however, it is not possible at present to reliably predict the structure of such genes from genomic sequence data. Interestingly, *Xist* has arisen from the protein-coding *Lnx3* during the formation of the mammalian X-chromosome (Duret et al., 2006). Such ancient "pseudo-genes" that acquired different

functions, and hence different patterns of sequence conservation, could at least conceivably be detected by means of specialized computational approaches.

As the example of the non-coding gene *Evf-1* (Faedo et al., 2004; Kohtz and Fishell, 2004) shows, not all well-defined non-coding RNAs have detectable evolutionarily conserved secondary structures, see Figure 10. This gene is one of the few representative of mlncRNAs that has been studied in some detail. The expression of Evf-1 depends both on the "Sonic hedgehog" (shh) and Dlx genes. The molecule exhibits splice variants of similar patterns in human, mouse and rat. The splice variant Evf-2 (Feng et al., 2006) has two 5' exons, one of which overlaps one of the two known ultra-conserved enhancer elements that interact with *Dlx-2* to activate transcription of the two adjacent Dlx genes. Feng et al. (2006) found that this ncRNA acts as co-activator by directly interacting with *Dlx-2*, indicating a novel mechanism whereby transcription is controlled by the cooperative actions of an ncRNA and a homeodomain protein. In a similar vain, the trithorax response element derived transcripts mentioned in the introduction also show no detectable conserved secondary structure. Despite the rather well-defined architecture of such transcripts and their evolutionary conservation at sequence level, currently available bioinformatics methods are insufficient to reliably detect such unstructured ncRNA genes.

Recent tiling array (Cawley et al., 2004) and cDNA data (Carninci et al., 2005) strongly suggest that ncRNAs genes of this type are the rule rather than the exception. Even in the presence of cDNA



Fig. 10. Chromosomal region of the Dlx-5/6 bigene cluster (modified from the UCSC genome browser, hg17 assembly). Two known ultra-conserved enhancer elements, ei and eii, which are targets of Dlx2 (Ghanem et al., 2003), are associated with the Evf-2 transcript, which in turn interacts directly with Dlx2 as a co-activator of Dlx-5 and Dlx-6 (Feng et al., 2006). None of the predicted RNA secondary motives (RNAz, EvoFold) in this region is located in an exon of the non-coding RNAs Evf-1, while EvoFold predicts conserved structure at the two ultra-conserved enhancer location ei and eii.

and/or EST data, it is not an easy task to distinguish genes with short ORFs that code for short peptides from bona fide ncRNAs.

An interesting observation in this context is that a small class of large ncRNAs (which includes Xist and Air) give rise to a substantial number of short unspliced cDNAs that appear to be internally primed at A-rich regions of a much longer fulllength transcript. Starting from this property, Furuno et al. (2006) searched the entire RIKEN mouse cDNA dataset and characterized 66 "long expressed non-coding regions" (ENORs) longer than 10 kb, many of which are involved in imprinting and/or anti-sense transcription. A substantial number of these long ncRNAs (42 of 66 ENORs), and notably Xist, contains small regions with conserved secondary structure as determined by comparison with the mammalian RNAz screen by Washietl et al. (2005a).

The distribution of TFBS discussed in the previous section might provide another starting point. Further investigations with selected sets of TF binding motifs will be needed to determine whether TFBS frequencies can be used to discern between *cis*-acting DNA elements and sequence elements that are functional at transcript level.

In this contribution, we have attempted to provide an overview of the state of the art in ncRNA annotation. In summary, both the detection of functional RNAs in genomic sequence data and the classification of the candidate sequences is a challenging problem, despite significant recent advances in RNA bioinformatics. Reliably automatic annotation that would be applicable routinely on newly sequenced genomes remains elusive beyond those cases that can be handled by sequences homology with known ncRNA genes.

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