An assessment of bacterial small RNA target prediction programs

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Keywords: bacteria, sRNA, sRNA target prediction

Most bacterial regulatory RNAs exert their function through base-pairing with target RNAs. Computational prediction of targets is a busy research field that offers biologists a variety of web sites and software. However, it is difficult for a non-expert to evaluate how reliable those programs are. Here, we provide a simple benchmark for bacterial sRNA target prediction based on trusted E. coli sRNA/target pairs. We use this benchmark to assess the most recent RNA target predictors as well as earlier programs for RNA-RNA hybrid prediction. Moreover, we consider how the definition of mRNA boundaries can impact overall predictions. Recent algorithms that exploit both conservation of targets and accessibility information offer improved accuracy over previous software. However, even with the best predictors, the number of true biological targets with low scores and non-targets with high scores remains puzzling.

Introduction

Identification of regulatory small RNAs (sRNAs) in bacterial species is rapidly advancing owing to recent improvement of bioinformatics and sequencing technologies.¹ Bacterial sRNAs act usually through sequence-specific binding to target mRNAs, leading to altered stability or translational repression/activation of their targets.² In the order of 100 to 200 sRNAs are present in a typical bacterial genome and each sRNA may regulate a dozen or more targets, making the sRNA/mRNA interaction network a significant part of the overall gene regulation network. Biological functions controlled by sRNAs are diverse, related to stress responses as well as carbon and amino acid metabolism, virulence and cellular development.³

As sRNA collections build up, the main bottleneck in understanding bacterial sRNA function has now shifted to the question of target prediction. Experimental validation of specific sRNA/mRNA pairs using reporter gene assays or compensatory mutations of binding sites is labor intensive,⁴ and high throughput in vitro screens are subject to high rates of false positives.⁵ On the other hand, computational target prediction is challenging since sRNAs and their target often form imperfect and relatively short RNA-RNA hybrids that are not easily distinguishable from many other hybrids formed by random pairs of transcripts. Yet, computational methods have recently benefited from new algorithms and the introduction of RNA accessibility and conservation information that improved their performances.⁶⁻⁸

Here, we evaluate computational target prediction methods using a collection of trusted, experimentally validated sRNA/mRNA pairs in E. coli. We designed a benchmark system so that wet lab scientists can easily interpret it for practical purposes. We also evaluated program speeds and abilities to locate the actual duplex region in each sRNA/mRNA pair.

Results

Target predictors

Programs for the identification of RNA-RNA interactions can be grouped in 3 main categories that we term “alignment-like,” “inter-RNA” and “independent fold.” The “alignment-like” category includes programs such as Guugle⁹ and RIsearch¹⁰ that can quickly scan large sequence files for reverse complements of a given RNA sequence. While Guugle looks only for stretches of Watson-Crick or G:U matches, RIsearch implements a Turner-like energy model¹¹ and allows gaps in the base paired segments. The “inter-RNA” category includes predictors that use a nearest neighbor thermodynamic model restricted to interactions between sRNA and mRNA, neglecting intramolecular base-pairs. Representative programs in this class are Pairfold,¹²,¹³ RNAcofold,¹⁴ TargetRNA,¹⁵ RNAtarget,¹⁶,¹⁷ RNAhybrid,¹⁶,¹⁷ RNAplex,¹⁸ and RNAduplex.¹⁸ These programs identify intermolecular contacts and compute a binding free energy for the interaction. RNAcofold
and Pairfold achieve this by considering the joint structure predicted after concatenating the 2 RNA sequences. The “independent fold” approach analyzes the secondary structure folding landscape of each RNA independently and models the total binding energy as a sum of 2 contributions. RNAup\(^{19}\) and IntaRNA\(^{7}\) are the major programs in this class. Finally, a recent generation of programs including CopraRNA\(^{20,21}\) and TargetRNA2\(^{22}\) combine the benefits of the “inter-RNA” class of programs to the use of conservation information. While TargetRNA2 requires conservation of the sRNA region involved in target binding, based on observations by Peer and Margalit,\(^{23}\) CopraRNA uses the conservation of the interaction itself.

Another, more practical, categorization of RNA target identification software has on one hand actual target predictors, \textit{i.e.} programs that identify and score the most favorable local interaction (CopraRNA, IntaRNA, RNAplex, RNAup, TargetRNA2) and on the other hand programs that model the full hybrid formed between the small RNA and the longer RNA (RNAhybrid, RNAduplex, RNAcofold, Pairfold). While the latter category can be suitable to microRNA/mRNA binding analysis, where the 21 nt RNA can be bound almost entirely to its target, results are more difficult to interpret when dealing with bacterial sRNAs that are generally over 100 nt long and are bound to the target over a small fraction of their size.

The pros and cons of sRNA target prediction algorithms have been reviewed.\(^{24}\) The goal we set here is to evaluate the ability of programs to differentiate a \textit{bona fide} sRNA/mRNA regulatory pair from a random duplex that may form between any transcripts in a cell. To this aim, we need to rank RNA-RNA duplexes and thus to rely on scores or free energy values provided by the programs. This requirement excludes programs whose energy model is too simplistic, such as Gugule or RiSearch. Although RiSearch does compute free energy values, the authors recommend the program to be used for fast screening of potential pairs, but not to evaluate specific interactions.

We included in our comparison programs from the major categories above, including the most recent bacterial sRNA predictors CopraRNA, TargetRNA (in its last implementation TargetRNA2\(^{22}\)) and IntaRNA, programs for local hybrid prediction that can deal with both bacterial sRNAs and miRNAs (RNAup, RNAplex also known as its web server implementation RNApredator\(^{25}\)), and programs predicting full RNA-RNA hybrids, including Pairfold, RNAcofold, RNAhybrid and RNAduplex.

We ran IntaRNA, RNAplex, RNAup, RNAhybrid, RNAduplex, RNAcofold and Pairfold using Unix command line versions and CopraRNA and TargetRNA2 from their respective web servers (see Suppl. Methods).

**Benchmark sets and search domains**

\textit{Escherichia coli} is the organism with the highest number of experimentally validated sRNA/target pairs. We compiled a list of 102 “trusted” sRNA/target pairs involving 22 different sRNAs (Table S1), each one bing supported by at least one published experimental evidence from the list described in Methods. This includes \textit{E. coli} pairs from the datasets of Peer and Margalit\(^{23}\) and Wright et al.\(^{20}\) plus a few additions from the literature.\(^{26,27}\)

Bacterial small RNAs are known to interact preferentially with mRNA 5′ regions with few exceptions to this rule (\textit{e.g.}, see ref.\(^{28,29}\)). Within our data set, all pairs, except one, are found to occur in the mRNA region located −150/+100 nucleotides (nt) around the start codon (Fig. S1). A common practice that consists in restricting sRNA target searches to the −200/+100 region (\textit{e.g.}, see ref.\(^{20}\)) around the start codon of all mRNAs is thus justified at least for enteric bacteria. Indeed, restricting the search space to the 5′ region efficiently reduces false positive occurrences and computing time. However, bacterial mRNAs often have short 5′ UTRs that do not extend much upstream of the Shine-Dalgarno sequence. Therefore, since the long 200 nt upstream region may be a source of false positives, this artifact can be avoided when exact transcription starts are known. Such information requires massive, strand-specific sequencing of transcripts and is still rarely available. However, this situation is quickly changing and, for certain bacteria including \textit{E. coli}, accurate UTR information is already available.\(^{30}\) Thus, we asked whether a precise delineation of UTRs could improve target identification. When possible, we performed predictions on 2 target data sets: one made of arbitrary −200/+100 nt regions around each start codon (hereafter named “default UTRs”) and one made of regions predicted from publicly available strand-specific RNA-seq data (hereafter named “real UTRs”). Note that we also included in our analysis 5′ regions of open reading frames located in polycistronic genes, as these regions are also subject to regulation as exemplified by the \textit{malXYZ} operon mRNA targeted at 2 locations by the sRNA SgrS,\(^{31}\) or the \textit{Staphylococcus aureus} opp3 operon mRNA containing at least 2 RsaE-binding sites.\(^{32}\) Details of the 5′ UTR definition procedures are provided in Methods and Suppl. Methods.

**Rank-based prediction performance**

Our first performance measure aimed at addressing the following question: how many putative targets for a given sRNA should be experimentally tested (for instance using a reporter assay) before finding the first true target? Programs were run using the best options suggested by authors. RNA accessibility information, which is known to improve target prediction significantly, was selected when possible.\(^{19,33}\) Programs from websites (\textit{i.e.}, CopraRNA, TargetRNA2) were run using preselected mRNA target sequences that correspond to “default UTRs” as defined above. These servers cannot take arbitrary lists of target sequences as input. Therefore tests with “real UTR” targets were limited to programs ran from the command line. Parameter details are presented in Suppl. Methods.

Putative targets were ranked by scores or free energy values provided by each program. The median rank is the median position of the highest ranking validated target among predicted targets for all sRNAs in the dataset (Fig. 1A–B). Median ranks range from 4 (CopraRNA) to 1340 (RNAcofold). Expectedly, programs designed for evaluating local interactions (CopraRNA, intaRNA, RNAplex, RNAup, TargetRNA2) perform far better than programs computing complete hybrids. In this latter category, RNAhybrid, RNAduplex, perform significantly better due to their free energy scores restricted to intermolecular base-pairs.
The least performing programs RNAcofold and Pairfold are both “concatenation” algorithms that are not intended to evaluate the binding energy of the complex and therefore cannot perform any better than a random ranking.

Our test probably underestimates the performance of programs since some of the pairs ranking before the first confirmed target are also possibly true targets. Nevertheless, some sRNA/target pairs rank very poorly, even by the best performing predictors (Fig. 1A), which questions their presence in our benchmark set. Indeed, some sRNA/target pairs are part of the trusted pair list because target expression is affected by the amount of the corresponding sRNA. However, this effect may occur indirectly in the absence of any contact between the 2 RNAs. To test whether putative indirect pairs may have impacted our benchmark, we considered only pairs for which we had direct evidence of sRNA/target interaction (82 pairs, see Table S1 and Methods). There was no improvement of prediction accuracy over the full data set (Fig. S2A), suggesting the presence of indirect pairs in our dataset is not a reason for the low ranking of some actual targets. Further inspection of poorly ranked trusted sRNA/target pairs shows that the same pairs tend to be missed by all predictors. Typical mispredictions involve pairs with target sites located 5' of upstream genes such as in RyhB/fur mRNA\textsuperscript{34} and hybrid structures formed by double kissing complexes such as in OxyS/fhlA mRNA\textsuperscript{35}.

We tested the effect of using the “real” UTR sequences (RNA-seq-based UTRs) of mRNAs instead of arbitrary upstream regions (−200 nt before AUG codon) on prediction performance (Table 1). Using “real” UTRs improved results moderately (AUC gains of 4.9% to 17.5%) albeit not significantly for most predictors. We observed a significant AUC improvement (Wilcoxon P = 0.03) only for concatenation methods RNAcofold and Pairfold, however this should be put into perspective as these methods had a very small AUC to start with. Yet, we may hypothesize that more accurate mRNA boundaries can improve free energy estimations of the hybrid in some cases.

The best prediction accuracy overall was obtained using CopraRNA, suggesting an advantage to a method using both RNA accessibility and conservation information, in line with the authors own estimation\textsuperscript{20}. Although CopraRNA performed best overall, it should be noted that its use is limited compared to other programs since it requires an homolog set for each sRNA of interest and targets are evaluated only in a predefined list of genomes.

Assessing other predictor qualities

While the above comparison used the ranks of the best scoring targets of each sRNA, we found interesting to analyze ranks of all trusted targets combined. When all 102 trusted targets are considered, median ranks for all programs increase mathematically (Fig. S3). However, considering that our test set contains only 4–5 targets per sRNA, these new median ranks ranging from 50 to

Table 1. Area under ROC-like curve (AUC) for target prediction with 9 programs, using default UTR or “real” RNA-seq-derived UTRs

| Program       | AUC default UTR | AUC real UTR | AUC gain (%) |
|---------------|-----------------|--------------|--------------|
| CopraRNA      | 46.41           | na\textsuperscript{a} | na\textsuperscript{a} |
| IntaRNA       | 27.02           | 28.83        | 6.3          |
| RNAplex       | 20.81           | 21.88        | 4.9          |
| RNAup         | 22.75           | 27.59        | 17.5         |
| TargetRNA2    | 18.52           | na\textsuperscript{a} | na\textsuperscript{a} |
| RNAhybrid     | 4.14            | 4.38         | 5.5          |
| RNAduplex     | 3.86            | 3.55         | –8.7         |
| RNAcofold     | 0.57            | 1.42         | 59.9         |
| Pairfold      | 0.25            | 1.22         | 79.5         |

\*Programs CopraRNA and TargetRNA2 can be run only using default UTRs.

Table 2. Recall of experimentally demonstrated base pairs (%)

| Program                  | Recall (%) |
|--------------------------|------------|
| IntaRNA/CopraRNA\textsuperscript{a} | 76.7       |
| RNAplex                  | 73.6       |
| TargetRNA2               | 55.9       |
| RNAup                    | 78.9       |

\*IntaRNA and CopraRNA predict the same hybrid region.
600 for the best predictors are a good indication of how predicted target sets remain populated mostly by unconfirmed targets.

We also tested how accurately each program was able to determine the correct hybrid regions assuming a correct sRNA/mRNA pair was detected. Since experimental validations of sRNA/target pairs do not always specify the duplex structure, we focused on the set of 55 pairs supported by compensatory mutations (see Methods). We excluded from this test all programs that predict a hybrid structure involving the full sRNA. Table 2 shows the fraction of compensatory mutations that are indeed found within the predicted hybrid regions. Interestingly, this ratio is far from perfect (56% to 79%) even though we only considered correct RNA/target pairs.

Finally, we compared the speed of the different programs (Table 3). Program Texte vary by 2 orders of magnitude, the fastest programs are able to match a sRNA to the complete set of E. coli mRNAs in about one minute while some require one to 2 hours to complete the same task. The best performing predictors are generally the slowest, with the exception of RNAPlex, which achieves both high prediction accuracy and fast running time.

### Conclusion

RNA target prediction programs have improved continuously over the past 10 years owing to algorithmic innovation, as well as the exploitation of structural and evolutionary constraints. We confirm here that CopraRNA, a predictor that uses both RNA accessibility and evolutionary conservation of the interaction outperforms other programs in terms of ranking actual sRNA/mRNA pairs. However, conservation data is not often available and programs that predict sRNA targets based on sequence alone are still much needed. IntaRNA, RNAPlex and RNAup are the best predictors in this class. The presence of RNAPlex in this shortlist is surprising as this program is up to 100 times faster than competitors owing to a simplified structural model. Furthermore, our tests do not show significant accuracy gains from using RNA-seq derived UTRs instead of fixed-length 5’ regions, supporting the common practice for defining mRNA target regions.

In spite of software improvements, this study is also a reminder that the list of highest scoring mRNAs for a given E. coli sRNA remains populated by a majority of unconfirmed targets. One possible explanation for this could be that many true targets have not been validated yet, which is unlikely in this well-studied bacterium. Alternatively, in vivo sRNA-mRNA interaction can be positively affected by RNA-chaperon proteins (e.g. Hfq), which may significantly confuse computed targets ranks. Another reason that may explain the relatively poor ranking of many true targets is related to the way programs score an RNA-RNA pair. Predictors score targets based on binding free energy values (as well as conservation for CopraRNA and targetRNA2). However, a high binding energy does not necessarily imply an efficient processing of the mRNA target by the Hfq–sRNA complex. The situation in bacteria could be similar to that of miRNA-target pairs in eukaryotes, where, due to miRNA turnover36 lower stability of the microRNA/target duplex may favor transient interactions and therefore an efficient down-regulation of multiple targets.37 The large accuracy gain of CopraRNA obtained by introducing phylogenetic conservation in its scoring system illustrates again that there is much more than thermodynamics in understanding sRNA-based regulation.

### Methods

Definition of UTR regions: «default» UTR regions were automatically extracted from MG1655 E. coli K12 genome, selecting the −200/+100 fragments around each start codon (4317 regions). «Real» UTR regions were extracted based on RNA-seq-derived transcriptional start site (TSS).30 For each TSS, the region from the TSS to 100 nt past the start codon was extracted. Further details are provided in Suppl. methods.

Experimentally validated pairs (Table S1) were classified as described based on supporting experimental evidence23: (a) sRNA affecting the level of a protein or a translational fusion; (b) sRNA gene affecting the level of a targeted mRNA; (c) sRNA and mRNA forming a stable duplex in vitro; (d) sRNA mutation affecting a regulation; (e) sRNA mutations affecting a regulation rescued by compensatory mutations; (f) in vitro chemical/ enzymatic probing of the sRNA/target duplex; (g) sRNA/target duplex cleaved by RNase III; (h) sRNA preventing or allowing ribosome binding demonstrated by toe-printing assay. We considered that the criteria c) to h) demonstrated direct or near-direct interactions for 82 sRNA/mRNA duplexes.

Performance measures (Fig. 1 and S2) were obtained by running target predictors with each sRNA against the complete set of E. coli mRNAs (using “default” or “real” UTRs). Ranks of trusted targets (Fig. 1A and S2A) were computed based on the best ranking target of each sRNA. For the “random” distribution, we performed for each sRNA a set of N random draws between one and the total number of E. coli mRNAs, where N is the number of known targets for this sRNA, and kept the lowest ranking value. Receiver operating characteristic (ROC)-like curves (Fig. 1B and S2B) were obtained by counting the number of true targets found among the best X candidates (shown on X axis). Program versions, options and command lines are provided in Suppl. methods. Accessibility information was computed prior

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**Table 3. Average run time for matching 1 sRNA to all 4317 E. coli mRNAs**

| Method                  | Time (minutes) |
|-------------------------|----------------|
| TargetRNA2 (web site)   | 1              |
| RNAPlex (incl. accessibility calculation) | 1 |
| RNAhybrid               | 2              |
| RNAcofold               | 23             |
| RNAup                   | 55             |
| IntaRNA                 | 66             |
| RNAduplex               | 75             |
| CopraRNA (website)      | 105            |
| Pairfold                | 120            |
to RNAplex runs using RNAplfold. CopraRNA requires homologous sRNA sequences from other species and we used those provided by the CopraRNA server whenever possible. For IstR, RseX, RydC we provided our own sets (see Suppl. Files).

Recall of experimentally validated base pairs: 55 sRNA-target pairs have experimental support in the form of compulsory mutations. For each pair, we measured the ratio of experimentally validated bps that were included in the predicted hybrid region. This ratio was then averaged over all 55 sRNA/target pairs to produce percentages in Table 2. As TargetRNA2 could predict only 49 of the 55 experimentally supported pairs, the average value was corrected accordingly.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

Funding
This work was funded by Agence Nationale pour la Recherche (grant 2010-BLAN-1602-01) ‘Duplexomics’.

Supplemental Material
Supplemental data for this article can be accessed on the publisher’s website.

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