RNAex: an RNA secondary structure prediction server enhanced by high-throughput structure-probing data

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INTRODUCTION

Several high-throughput technologies have been developed to probe RNA base pairs and loops at the transcriptome level in multiple species. However, to obtain the final RNA secondary structure, extensive effort and considerable expertise is required to statistically process the probing data and combine them with free energy models. Therefore, we developed an RNA secondary structure prediction server that is enhanced by experimental data (RNAex). RNAex is a web interface that enables non-specialists to easily access cutting-edge structure-probing data and predict RNA secondary structures enhanced by in vivo and in vitro data. RNAex annotates the RNA editing, RNA modification and SNP sites on the predicted structures. It provides four structure-folding methods, restrained MaxExpect, SeqFold, RNAstructure (Fold) and RNAfold that can be selected by the user. The performance of these four folding methods has been verified by previous publications on known structures. We re-mapped the raw sequencing data of the probing experiments to the whole genome for each species. RNAex thus enables users to predict secondary structures for both known and novel RNA transcripts in human, mouse, yeast and Arabidopsis. The RNAex web server is available at http://RNAex.ncrnalab.org/.

ABSTRACT

Several high-throughput technologies have been developed to probe RNA base pairs and loops at the transcriptome level in multiple species. However, to obtain the final RNA secondary structure, extensive effort and considerable expertise is required to statistically process the probing data and combine them with free energy models. Therefore, we developed an RNA secondary structure prediction server that is enhanced by experimental data (RNAex). RNAex is a web interface that enables non-specialists to easily access cutting-edge structure-probing data and predict RNA secondary structures enhanced by in vivo and in vitro data. RNAex annotates the RNA editing, RNA modification and SNP sites on the predicted structures. It provides four structure-folding methods, restrained MaxExpect, SeqFold, RNAstructure (Fold) and RNAfold that can be selected by the user. The performance of these four folding methods has been verified by previous publications on known structures. We re-mapped the raw sequencing data of the probing experiments to the whole genome for each species. RNAex thus enables users to predict secondary structures for both known and novel RNA transcripts in human, mouse, yeast and Arabidopsis. The RNAex web server is available at http://RNAex.ncrnalab.org/.
Four major steps are generally required to process the raw probing data and predict the data-enhanced RNA secondary structure. The first step is to map the reads, which is time-consuming and largely variable as the adaptor sequences specific to each probing experiment need to be trimmed. The second step is to calculate the RT stop read counts. Because enzyme cleavage truncates the RNA transcripts and chemical modification halts reverse transcription before the modification sites, each mapped read only gives information for the base immediately 5’ of the first mapping position. The third step is to derive the structural reactivity from the RT stop counts. Usually, there are two libraries in each probing experiment for controlling the background noise. A commonly used method is to control the effect of transcript abundance and length in each library, and then subtract the normalized counts in the control library from the normalized counts in the treatment library. The fourth step is to incorporate the experimental restraints into the final structure prediction. The structural reactivity derived from the third step needs to be transformed into the proper inputs (e.g., probabilities based on a statistical model) for a given structure-folding algorithm.

Several programs and platforms have been developed to accomplish the four steps that are required to use the latest structure-probing data. StructureFold provides an integrated solution to analyze the structure-probing data via the Galaxy platform. RSF provides a Perl framework for data processing and structure inference based on structure-probing data. Although these packages are incredibly helpful, intensive computational efforts and expertise in analyzing the probing data are required to use them, which generate large bottlenecks for wet lab biologists. The SAVoR web server was developed for structure prediction with experimental restraints and data visualization along the predicted structures, but users need to create and input the read alignment files. SAVoR does not provide optional folding methods other than RNAfold.

We designed the novel web server RNAex to bridge the gap between accumulating structure-probing data and improved RNA secondary structure prediction. RNAex enables non-specialists to easily access state-of-art high-throughput probing data in multiple species (human, mouse, yeast and Arabidopsis), and predict RNA secondary structures enhanced in vivo and in vitro experimental data. We have re-mapped the raw sequencing reads of published structure-probing experiments and transformed the read counts into corresponding inputs for four representative structure-folding methods (RME, SeqFold, RNAstructure (Fold), and RNAfold). RNAex predicts RNA secondary structure with many control options and provides an interactive visualization interface for users to explore the probing data, the processed structure profile and the predicted secondary structures. RNAex also displays the post-transcriptional regulation and mutation information for each predicted RNA secondary structure, including RNA modification sites (human and mouse) and RNA editing (human and mouse) and SNP sites (human). This enables the predicted RNA structure to be effectively annotated and linked with function, disease and post-transcriptional regulation events.

**DATASETS**

We first collected various representative high-throughput structure-probing data in human, mouse, yeast and Arabidopsis. We analyzed the raw sequencing data and mapped the short reads to the whole genome for each species (see detailed mapping methods in Supplementary File 1). The following genomes were used: hg19 for human, mm10 for mouse, SacCer2 for yeast and TAIR10 for Arabidopsis. Then, we calculated the RT stop counts for each individual nucleotide following the same procedures that were described in the original papers.

Next, we collected the post-transcriptional regulation and mutation information for human and mouse to display in the predicted structures. The RNA modification sites for human and mouse were downloaded from the RMBase database. The RNA editing sites for human and mouse were obtained from the RADAR and DARNED databases. The human SNP information was downloaded from NCBI dbSNP.

**ALGORITHMS**

The RNAex server embedded the structure-probing data and the regulation and mutation information in a background database and implemented all computation steps through backward scripts. Users input their transcripts of interest (IDs or genomic locations) and RNAex performs four major steps. (i) RNAex extracts the sequence and structure-probing data for the given transcripts. (ii) RNAex processes the structure-probing datasets selected by the user. (iii) RNAex predicts the data-enhanced RNA secondary structures using any of the four well known folding methods (RME, SeqFold, RNAstructure (Fold) and RNAfold), which are selected by the user. (iv) RNAex visualizes the predicted structures, the processed structure-probing data and the post-transcriptional regulation and mutation information (Figure 1).

The first step is to extract the RNA sequence and the RT stop counts of the structure-probing experiments for the selected transcripts. RNAex displays the sequences in FASTA format as a check page for users to confirm. Then, it extracts the probing read counts along the transcript from the user-selected structure-probing datasets. RNAex checks if sufficient probing data were mapped on the selected transcripts. RNAex only proceeds to fold the data-enhanced structure for transcripts that have been mapped with sufficient probing data. Otherwise, RNAex predicts the structure without the enhanced data, which generates a structure prediction that is no different from those of other sequence-based RNA secondary structure prediction servers.
The next step is to process the structure-probing data, which differs according to the four different folding methods available [RME, SeqFold, RNAstructure (Fold) and RNAfold]. Usually, as mentioned in the above introduction section, all methods need a paired control data as the background to compare and normalize the signal data (Supplementary Table S1). For instance, we used a denatured sample as the control for the signal data of DMS-seq, as did the version of genome annotation is selected based on the original paper. aWe provide the whole genome data (hg19) by re-mapping the raw reads, and transcriptome only data (transcriptome) mapped in the original paper.

The last step is visualizing the predicted structures and all relevant information. For each selected input transcript, RNAex provides a module for structure folding without incorporating probing data, which is convenient for users to compare and evaluate the contributions of the structure-probing data. If users are interested in the raw data, RNAex provides a genome browser page that directly displays the RT stop counts. RNAex also provides post-transcriptional regulation and mutation information on the
Figure 1. Schematic of the RNAex web server. RNAex embeds the structure-probing data, post-transcriptional regulation data and mutation information. Users select specific probing datasets and a folding method and input their selected transcripts of interest. Then, RNAex performs four major steps in the background: extracting the sequence and structure-probing data, processing the structure-probing data, predicting the structures and visualizing the results.

RNAex WEB SERVER

The RNAex web server is free to access and is unrestricted (without a login procedure). We adapted several JavaScript libraries and the HTML framework Bootstrap to build the RNAex server. The genome browser is based on JBrowse (31). The server is accessible from http://RNAex.ncrnalab.org (redirected to http://lulab.life.tsinghua.edu.cn/RNAex) and is compatible with most web browsers (Mozilla Firefox 15+, Google Chrome version 38+, Internet Explorer 9+ and Safari 8+). The processing and predicting functions of the RNAex server also are available through the command line version of RME, SeqFold, RNAstructure (Fold) and RNAfold, which are open source and available for downloading.

Inputs

To use RNAex, users are required to first select two matched probing datasets and a folding method. Then, users need to input their transcripts of interest for structure prediction, by providing the transcript IDs or the genomic locations in GTF format. RNAex can run multiple transcripts in parallel.

RNAex accepts the following transcript IDs for known transcripts: TAIR (v10) ID (32) for Arabidopsis transcripts, SGD (SacCer2) ID (33) for yeast transcripts, GENCODE (v19 for DMS-seq and v12 for PARS data) ID (34) for human transcripts and GENCODE (v2) ID (35) for mouse transcripts. RNAex also accepts genomic locations in GTF format for novel transcripts.

Users are free to adjust various parameters by selecting ‘Advanced options’. For example, users can define the criteria of sufficient probing data for transcripts by adjusting the options for ‘percentage of nucleotides with sufficient probing reads’ and ‘minimum number of reads to define sufficiency’. Users also can easily choose whether to compare the data-enhanced structure prediction with the structure prediction without data. Users also can adjust the contribution of probing data in the structure prediction by changing the parameters for each folding method, including the weight \(m\), gamma1 and gamma2 for RME (7), the \(P\)-value cutoff for SeqFold (20), the slope and intercept for RNAstructure (Fold) (21) and the algorithm, slope, intercept for RNAfold (19). Detailed descriptions of these parameters...
and the available options are provided in the manual on the RNAex server.

Selecting the ‘Submit’ button triggers RNAex to run the job. A user-provided E-mail address is optional but advisable when submitting computationally intensive jobs. A message containing the URL for the result page will be sent to the E-mail address when the job is finished.

### Computation

After submitting the job, RNAex provides a check page that users can browse to confirm the job. The check page displays the extracted RNA sequence for each input transcript in FASTA format. Users can check job accuracy in this page, and then go back to revise the job if they find input errors. RNAex also shows a warning or error panel if it detects errors; the warning message contains a brief description that helps users identify potential problems. Users should note that structure prediction for long transcripts is quite time-consuming and is not as accurate as predictions for short transcripts (36). RNAex allows the users to fold long transcript by dividing the sequence into small chunks (600 nt each). Then, the server will merge the chunks into one structure file (ct file) for the users to download. In the visualization page, considering the visualization effect might be largely affected if we incorporate too long sequence into one viewer, the server still makes these chunks being visualized and interacted separately. Moreover, because most of the transcripts have less than 3000 nt, we still have a limitation on a maximum length as 3000 nt. Therefore, we also provide download links for the command line version of each folding method, which can be used if users choose to fold an entire long transcript locally.

After submitting and checking the input transcripts, the folding jobs will run on the server. Structure-folding analyses are usually time consuming, and the results may not be available immediately. A waiting page is provided that displays a URL to the result page. Users can retrieve their results later using the URL. Users also can bookmark the waiting page, which will be directed to the same URL. Results remain on the server for up to one month. The real-time log information is provided on the waiting page to make the server more user-friendly; the log actively reports the current processing steps and the job status.

### Outputs

The final output of RNAex contains a summary page showing all the result files and the parameters used for the run (Figure 2A). The output displays four sections in the following order.

(i) Main parameters. General information is shown for the structure-probing data (species, build, data type and dataset) and the folding method [RME, SeqFold, RNAstructure (Fold) or RNAfold] selected by the user.

(ii) Advanced parameters. Three types of advanced parameters are listed in order. The first type is the user-defined or default criteria to determine whether there are sufficient experimental probing data mapped onto the selected transcripts, including percentage of nucleotides with sufficient probing reads and read cut (the minimum number of reads to define sufficiency). The second type of advance parameter is the option to perform comparisons between the data-enhanced structure prediction and the prediction without data enhancement. The default selection is ‘yes’. The third type of advance parameter includes the weighting parameters for each folding method, which are explained in the manual on the RNAex server.

(iii) Summary table of results (Figure 2A). Each row of the table lists various types of information for one selected transcript, including basic description, structure-probing data, predicted structures, post-transcriptional regulation and mutation information.

(a) The basic description includes the transcript ID, the genomic coordinate (chromosome, strand, range and length) and the downloadable sequence file in FASTA format for each transcript. The ‘range’ column records the detailed starts and ends for all exons of the transcript, which are separated by commas.

(b) The structure-probing data information includes a statistic and a downloadable file. The statistic represents the average ‘data coverage’, which is the percent of nucleotides associated with structure-probing data for the selected transcript in two different samples. The downloadable file is provided in the ‘processed data’ column, which has different meaning for each probing method. For RME and RNAfold, the raw data are processed into single-stranded probabilities on every nucleotide. For SeqFold, the raw data are processed into single-stranded probabilities, but the values are discrete (either 1 or 0). For RNAstructure (Fold), the processed data represent the probing reactivity. A nucleotide with higher reactivity value has higher probability to be single-stranded. These descriptions are included in the data file as comment lines.

(c) Structure information includes the predicted secondary structure enhanced by the probing data [structure (enhanced)], and the predicted secondary structure without restrained data [structure (no data)]. These can be downloaded in CT format (17). The predicted folding free energies for these two kinds of structures also are provided, and their difference (structure difference) is calculated by dividing the number of unique base pairs by the number of all base pairs in the two structures.

(d) Post-transcriptional regulation and mutation sites are displayed on the RNA structures, including RNA modification sites (human and mouse) and RNA editing and SNP sites (human). If these data are missing, an ‘NA’ value is displayed.

(e) The link to visualize corresponding transcript’s genomic location in jbrowse is also provided.

(iv) Bulk download button. In addition to the summary table where files for each transcript can be downloaded separately, a bulk download button also is provided. An Excel file containing all information and the URL for each downloadable file can be obtained by clicking the button.
Figure 2. Visualization of structure-probing data, the predicted structures, regulation and mutation information. (A) The summary page contains the main parameters, advanced parameters, a summary table of results and a bulk download button. (B) The first visualization module displays various datasets in bar plots and the predicted structure in arc diagrams. The control options in the bottom-left corner are unfolded for illustration. (C) The second visualization module displays the secondary structures with control options unfolded. (D) Example visualization of post-transcriptional regulation and mutation information by selecting the ‘SNP site’, ‘editing site’ and ‘modification site’ options in the ‘datasets’ control panel.

Visualization

RNAex also implements interactive visualization tools for users to view the probing data and the predicted structures. The result page for each run contains \( n + 1 \) tabs for an easy visualization, where \( n \) is the number of input transcripts or long transcript’ fragments (each fragment is \(< 600 \) nt and counts as one tab). Users are free to navigate these results through the tabs at the top (Figure 2A). The first tab shown as default is the summary page. The other \( n \) tabs are visualization pages for each selected transcript. In each visualization page, RNAex provides two modules to view the probing data and predicted secondary structure. The first module is illustrated in Figure 2B, which is a combination of two plots. The upper plot shows the probing data as a bar plot, where nucleotides with higher values have higher probability to be single-stranded. Users can see the exact value for each base by hovering the mouse over the base. The lower plot shows the predicted secondary structure in arc diagrams. RNAex provides another interactive module adapted from ViennaRNA forna (37), which allows users to view the secondary structure (Figure 2C). The structure can be fine-tuned and color-coded according to various datasets, which can be controlled by users.

These two visualization modules can be configured in many ways using the buttons available in the bottom-left corner of the screen. The star at the left of the screen is used to reset the figure to default parameters. The ‘datasets’ button is used to color-code the plots according to various datasets. For example, clicking the ‘SNP site’ button prompts the upper panel of the first module to display several red bars, with each one showing a nucleotide with SNP
annotation (Figure 2D). If the transcript does not contain any post-transcriptional regulation information, then only one choice is provided (i.e. ‘probing data’). The next button shows many configuration options. For example, users can edit the y-axis labels through the input boxes for ‘y-axis label for the upper panel’ and ‘y-axis label for the lower panel’. If the transcripts are too long, users can view only a segment of the transcript by sliding the two icons in the ‘show local region (range)’ option. Users also can control the color of the bars shown in the upper panel by sliding the icons appearing in the ‘data cutoff for single-stranded bases’ and ‘data cutoff for paired bases’ options. Finally, a full-screen button is provided to enlarge the figure to the entire display region.

By default, two kinds of secondary structures are predicted for each transcript. One is restrained by the structure-probing data (data-enhanced), whereas the other is not restrained by any probing data (without data). Thus, four visualization panels are available for each input transcript. A fold/unfold option (the ‘+’ button before the panel title) is provided for users to conveniently compare the four different plots. All these figures can be downloaded in SVG or PNG format by clicking the download button in the bottom-right corner. The background color can be set to be transparent or white by changing the option of ‘background’.

DISCUSSION

We present RNAex as a web-based server that enables non-specialists to easily predict RNA secondary structures using cutting-edge high-throughput structure-probing data. Users can easily access and incorporate these data into the RNAex server to enhance the RNA secondary structure prediction using four different folding methods. RNAex results allow users to clearly view, identify and understand SNP and post-transcriptional regulation sites on the RNA structures. Overall, the server is an early web-based platform for the emerging new field of high-throughput RNA structure-probing analysis. We will continue enriching RNAex as additional data become available, by including more species, diverse types of structure-probing data and other types of information related to RNA structure dynamics. RNAex only uses the folding tools to predict a single secondary structure (e.g. the structure with the lowest folding free energy change). However, RNA secondary structure at non-zero temperatures is an ensemble of configurations, not a single structure. We will improve this in the future. Moreover, we also will continue to improve the user experience of the website.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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