



**HAL**  
open science

## RNA-SequenLens for Visualizing RNA Secondary Structures

Florence Ying Wang, Arnaud Sallaberry, Mathieu Roche

► **To cite this version:**

Florence Ying Wang, Arnaud Sallaberry, Mathieu Roche. RNA-SequenLens for Visualizing RNA Secondary Structures. BioVis: Biological Data Visualization, Jul 2015, Dublin, Ireland. lirmm-01184562

**HAL Id: lirmm-01184562**

**<https://hal-lirmm.ccsd.cnrs.fr/lirmm-01184562>**

Submitted on 16 Aug 2015

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

# RNA-SequenLens for Visualizing RNA Secondary Structures

Florence Ying Wang

Arnaud Sallaberry

Mathieu Roche

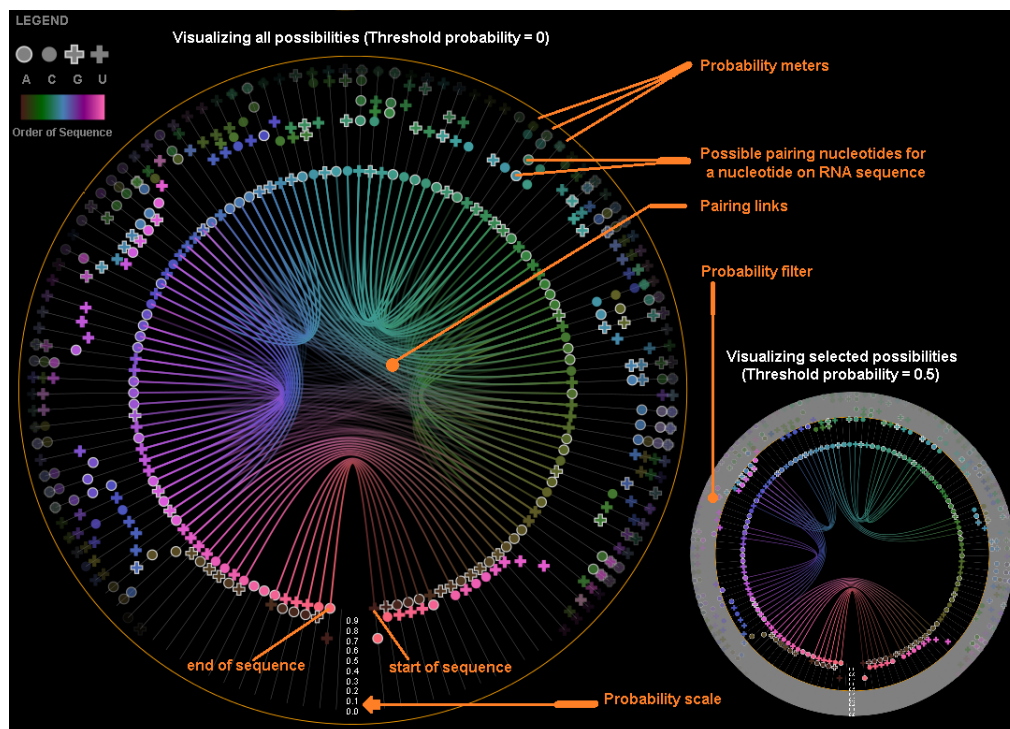


Fig. 1. Visualizing all the possible base pairing probabilities of ancestral(chimp) RNA with RNA-SequenLens. A threshold probability can be set interactively by the *probability filter* to determine the amount of pairing links shown in the center of SequenLens.

**Abstract**—In this paper, we present RNA-SequenLens to facilitate the visualization and comparison of RNA secondary structures. With RNA-SequenLens, all possible base pairings of a RNA sequence can be visualized at the desired probability threshold. Different RNA secondary structures can be easily compared. The interactive demo is available at <https://youtu.be/C6EDC8LZJXw>

## 1 INTRODUCTION

Traditionally, computational biologists use dot-plot to visualize the base pairing probabilities, where each grid on  $N \times N$  grids encodes the base pair binding probability [3]. Furthermore, the predicted secondary structure is often represented as a graph or arc diagram [1]. Although these visualizations have been popular, they have limitations such as: (1) with dot-plot and arc diagram, possible pairing nucleotides are not easily perceived, therefore it is difficult to compare RNA structures; (2) rainbow colors are often adopted, which can not reveal the characteristics of data that has implied ordering [2]. In this paper, we introduce a visual design called “RNA-SequenLens” to address above problems and the visualization tasks raised in the re-design contest.

## 2 VISUALIZATION REQUIREMENTS FOR CHALLENGE 1 AND 2

The overall purpose of challenge 1 is to design an intuitive visual representation of RNA secondary structure to encode the uncertainty for all the possible base pairing possibilities. Detailed visualization requirements for challenge 1 are summarized as follows:

[T1-1] Visualizing a RNA sequence.

[T1-2] For each nucleotide (i.e. A, C, G, U) on a RNA sequence, visualizing its possible pairing nucleotides.

[T1-3] For a RNA sequence, visualizing all base pairing possibilities.

The overall purpose of challenge 2 is to design a visual representation that supports comparison of the predicted RNA structures. Detailed visualization requirements for challenge 2 are summarized as follows:

[T2-1] Comparing different RNA sequences.

[T2-2] Comparing different predicted RNA secondary structures.

## 3 VISUAL MAPPING

The features of RNA-SequenLens are shown in Figure 1.

(I) For [T1-1], a RNA sequence is displayed circularly in the inner most circle, then color is used to encode the order of nucleotides on the RNA sequence. As shown by the legend, RNA sequence start with brown and ends with pink (i.e. counterclockwise). For visualizing different nucleotides, we use “circle” to present A (with outline) and C (without outline), “cross” to represent G (with outline) and U (without outline). This is due to the fact that A is likely to bind with U and C is

• Florence Ying Wang, LIRMM & Universite de Montpellier, France. E-mail: [ying.wang@lirmm.fr](mailto:ying.wang@lirmm.fr).

• Arnaud Sallaberry, LIRMM & Universite Paul Valery Montpellier, France. E-mail: [arnaud.sallaberry@lirmm.fr](mailto:arnaud.sallaberry@lirmm.fr).

• Mathieu Roche, TETIS & LIRMM & CIRAD, France. E-mail: [mathieu.roche@cirad.fr](mailto:mathieu.roche@cirad.fr)

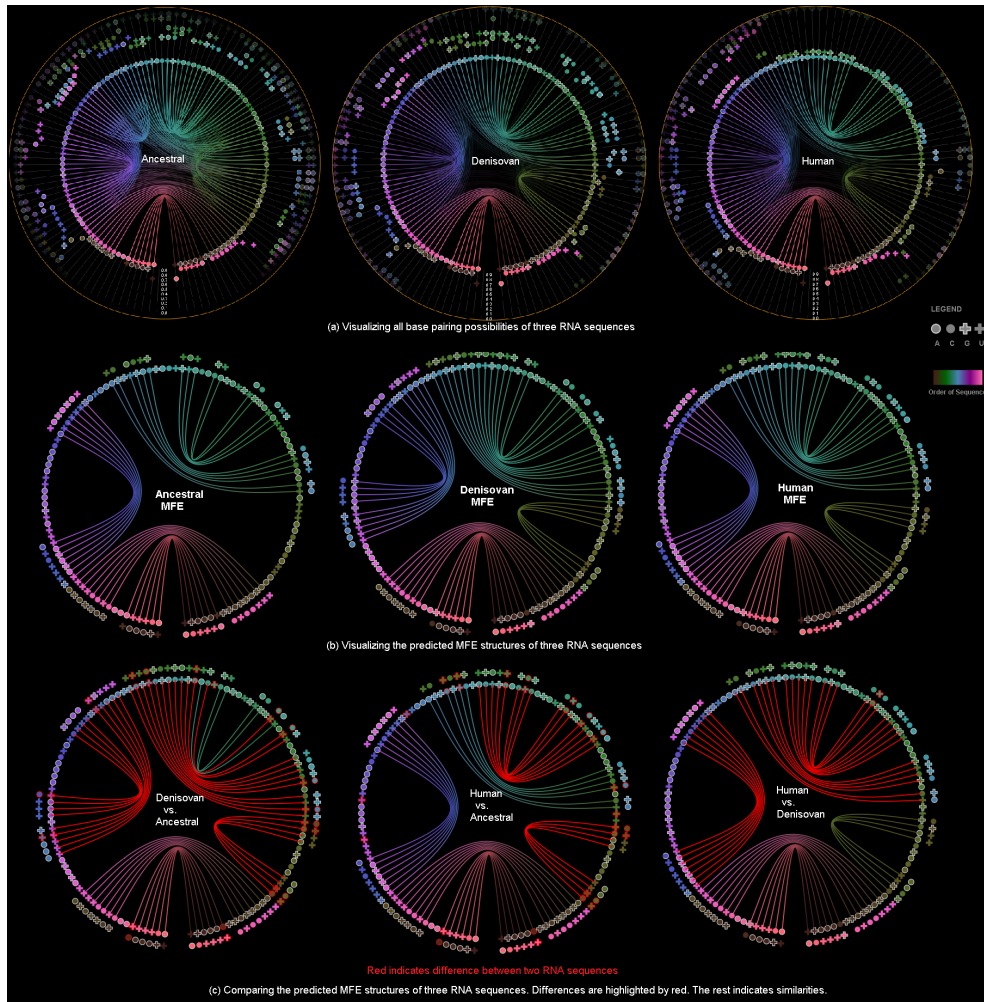


Fig. 2. Visualizing the changes of RNA primary and secondary structures of Ancestral(chimp), Denisovan and Human with RNA-SequenLens.

likely to bind with G. Hence, in SequenLens, circles often bind with crosses.

(2) For [T1-2], we plot one *probability meter* for each nucleotide on a RNA sequence. On the *probability meter* of a nucleotide, all its possible pairing nucleotides are displayed sequentially based on probabilities. The closer the pairing nucleotides to the center, the higher the probability. *Pairing links* are used to link possible binding nucleotide pairs. The color of a *Pairing link* is gradient color that gradually changes from one nucleotide color to its pair's color. The opacity of both *pairing links* and pairing nucleotides on *probability meters* indicate probabilities. The higher the opacity, the higher the probability.

(3) For [T1-3], we introduce a *probability filter* (gray mask in Figure 1 right corner) to allow the interactive visualization of pairing possibilities at the desired threshold. By interactively changing the size of the *probability filter*, *pairing links* of different probabilities will be displayed in the center of SequenLens.

(4) The visualization of predicted RNA secondary structures (e.g. MFE structure) can be considered as a special case of uncertainty visualization where the threshold probability is fixed (Figure 2(b)). Then, for [T2-1][T2-2], we can highlight the differences between two RNA sequences and their predicted secondary structures (Figure 2(c)).

#### 4 CASE STUDIES

In this section, we use two case studies to further illustrate our visual design in solving problems raised in the re-design contest.

**Case Study 1: Visualizing Uncertainty of Ancestral RNA** In Figure 1, circles often pair with crosses, symbols with white outlines often

pair with symbols without outlines, which validate the binding properties of A, C, G, U. Also, there is a strong likelihood of pairing between the start and end segments of the sequence. By changing the probability threshold to 0.5 (Figure 1 right corner), we can see that there are 3 major groups of base pairings indicated by 3 major *pairing link* colors: green, purple and pink. This finding also aligns with the 3 groups of *pairing links* in the MFE structure (1st image Figure 2 (b)).

**Case Study 2: Visualizing Sequence Evolution** In Figure 2(c), the first image shows that between the MFE structures of denisovan and ancestral, segments with the most variations in primary structure also have the most difference in secondary structure. The middle image shows that human and ancestral have less differences in the MFE structure than human and denisovan, whereas the last image indicates that human and denisovan have the least differences in terms of RNA primary structure (only 1 nucleotide is different).

#### 5 CONCLUSION

In this paper, we have introduced RNA-SequenLens for visualizing RNA secondary structures. Our case studies show the effectiveness of our design in visualizing uncertainty and sequence evolution.

#### REFERENCES

- [1] D. P. Aalberts and W. K. Jannen. Visualizing rna base-pairing probabilities with rnabow diagrams. *RNA*, 19(4):475–478, 2013.
- [2] D. Borland and R. M. Taylor II. Rainbow color map (still) considered harmful. *IEEE computer graphics and applications*, 27(2):14–17, 2007.
- [3] I. L. Hofacker. Vienna rna secondary structure server. *Nucleic acids research*, 31(13):3429–3431, 2003.