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The Circular Secondary Structure Uncertainty Plot (CS²-UPlot) - Visualizing RNA Secondary Structure with Base Pair Binding Probabilities

Dan Tulpan

Abstract—The Circular Secondary Structure Uncertainty Plot (CS²-UPlot) is an intuitive visual representation of an RNA secondary structure that includes the uncertainty of all possible base pairings. The CS²-UPlot uses a chord diagram layout and is comprised of 3 concentric graphical layers representing the three main information components of an RNA secondary structure required by the BioVis 2015 Design challenge: (i) the RNA sequence (outer layer), (ii) uncertainty and free energy (*mfe*) scatter plots for each base (middle layer) and, (iii) uncertainty and minimum free energy (*mfe*) base pairings (inner layer). The CS²-UPlot challenges the classical ways of representing RNA secondary structures and combines base pairings with dot-plot values in a single graphical representation capable of assisting biologists in quickly spotting similarities and differences among a large number of secondary structures and their corresponding RNA sequences. **Availability:** Figures can be downloaded from: <http://www.nrcbioinformatics.ca/cs2uplot/>

1 THE CHALLENGES

The BioVis 2015 Design Competition includes two challenges: uncertainty visualization and sequence evolution visualization. My solution is designed to mainly address the first challenge, while it can be part of a solution for the second challenge, too, as it will be described in the following sections.

First, I will extrapolate lists of requirements that capture the main desired features for solutions to each challenge. I will use these requirements to design a solution and to determine to what extent the solution addresses each challenge.

The required features for *Challenge 1* are: (1.1) display the RNA sequence, (1.2) visualization of base pairing probabilities represented in the top-right triangle of the table associated with the dot-plot, (1.3) visualization of base-pairings corresponding to the *mfe* structure represented in the bottom-left triangle of the table associated with the dot-plot (optional since the *mfe* secondary structure already presents this), (1.4) must be a static picture.

The required features for *Challenge 2* are loosely defined as follows: (2.1) supports comparison of predicted RNA structures, (2.2) ability to identify changes in the RNA sequence that influence its structural stability, (2.3) must be one or more static pictures.

2 THE SOLUTION

Challenge 1 mainly requires an intuitive mean to visualize sparse tabular information, with the constraint that the (x,y) coordinates for each uncertainty value in the table corresponds to a base pair in an RNA sequence, whose *mfe* secondary structure is known and typically represented as a graph. Based on these constraints, my choice is to open-up and map the typical *mfe* secondary structure representation of the RNA sequence on a circular plot much in the same fashion as Circos [1] represents linear genomes and their interactions, inter-dependencies and features. In the interest of time the Circos graphical library was employed to implement an automatic process that produces the plots presented in this manuscript. Alternatively, the *D3.js* JavaScript library can also be used for the same purpose. I also acknowledge that simplified versions of circular plots were used for RNA structure representations as early as 1978 [2], nevertheless they represented only one type of information such as *mfe* secondary structures.

Figures 1, 2, 3 and 4 provide a first glimpse of the more advanced visualization method proposed in this manuscript. For a lack

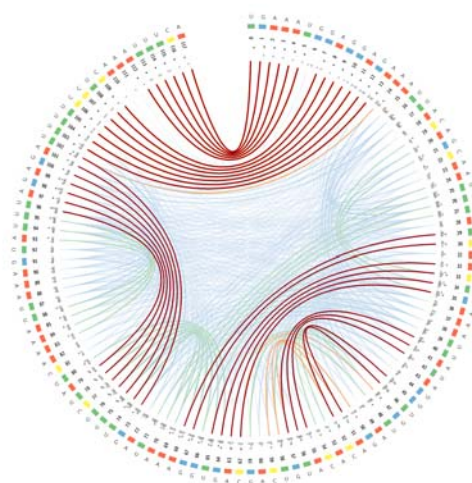


Fig. 1. The CS²-UPlot of the HAR1 ncRNA in ancestral chimp.

of better words, I called this plot type a Circular Secondary Structure Uncertainty Plot (CS²-UPlot).

A CS²-UPlot consists of 3 concentric layers of information: (outer layer) the RNA sequence, (middle layer) the uncertainty and minimum free energy (*mfe*) mini scatter plots for each base and, (inner layer) the uncertainty and *mfe* base pairings.

The outer layer (RNA sequence) consists of 4 types of equally spaced blocks colored corresponding to their base content (A, C, G and U). Each base and its corresponding position in the sequence (counting starts at 0) is displayed around each block. This addresses requirement 1.1.

The middle layer represents a mini scatter plot for each RNA base 4. The height and color intensity (gray scale) of each dot on the plot mark its pairing probability, while its x-location within a narrow segment that signifies the whole sequence suggests the relative location of the corresponding base pair on the RNA sequence. The density of the dots in a scatter plot is tightly related to the interaction capacity of each base in a given structural conformation. The higher the number of dots in the scatter lot, the more potential base pairings that base can form. This partially addresses requirement 1.2, which will be fully addressed by the inner layer.

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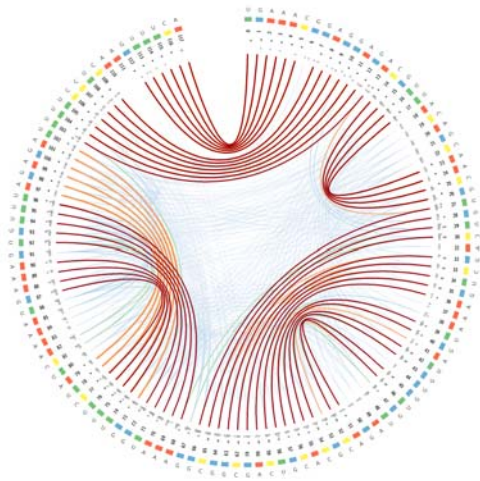


Fig. 2. The CS²-UPlot of the HAR1 ncRNA in Denisovan.

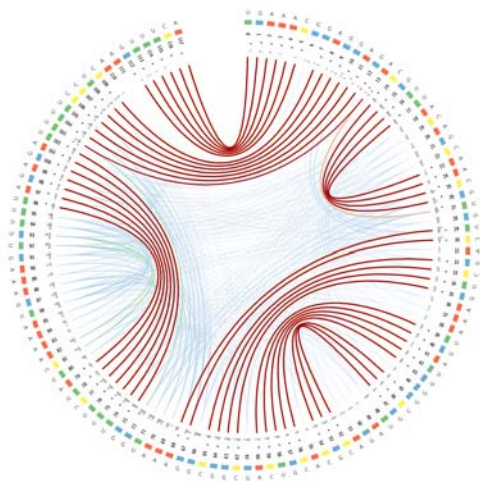


Fig. 3. The CS²-UPlot of the HAR1 ncRNA in human.

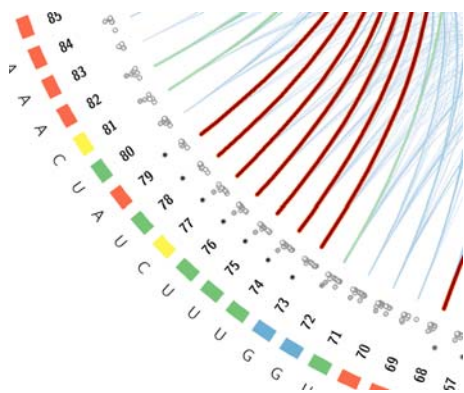


Fig. 4. A close-up view of the CS²-UPlot of the HAR1 ncRNA in human.

The inner layer consists of a set of colored arches that connect base pairs on the RNA sequence, which signify hydrogen bonds depicted by short segments in a typical graph-based RNA secondary structure representation. The arch colors (currently using a 5-color palette as-

signed to uncertainty values within 0.2 unit intervals span the interval [0,1]) are assigned based on the corresponding uncertainty probabilities, ranging from blue (less stable) to dark orange (more stable) with green representing medium stable base pairs. The thickness of each arch is also proportional with the uncertainty probabilities using 5 incremental sizes ranging from 1 (less stable) to 9 (more stable). Complemented by scatter plots, this representation addresses requirement 1.2. The dark red arches represent the *mfe* base pairing corresponding to the most stable interactions. Thus requirement 1.3 is also addressed.

The next section will explain how to interpret the information depicted in the CS²-UPlots and will shed light on how the plots can be used to address the requirements for Challenge 2.

3 THE INTERPRETATION

The information represented in Figures 1, 2 and 3 can be interpreted based on predominance and localization of various visual cues related to color and position. For example, the CS²-UPlot of the HAR1 ncRNA ancestral chimp sequence shows a fairly large number of low and medium base-pair probabilities (blue and green arches) localized in two areas of the RNA sequence between base pairs 19 and 35, and between base pairs 47 and 95. Subsequently, the plot also displays a fairly large subsequence with no base pairings between positions 14 and 28. By comparison, the corresponding subsequences in Denisovan and human HAR1 ncRNAs, contain a solid 4 base-pair stem formed due to 2 consecutive base pair mutations at positions 14-15 and 25-26 replacing the AA/UU base pairs in chimp with the more stable CG/GC in Denisovan and human.

A total of 17 single and consecutive base pair mutations (positions 5, 14-15, 25-26, 32, 40, 43, 53, 56, 63, 65, 72, 87, 93 and 112) can be identified between the ancestral chimp and Denisovan HAR1 ncRNAs, while only one mutation (U replaced by C) at position 46 occurred between the Denisovan and human HAR1 ncRNAs. The large number of mutations that distinguish the Denisovan from the chimp RNA sequences apparently caused not only the apparition of a new 4 base pair stem between positions 14-17 and 23-26 in Denisovan, but caused also the breakdown of an existing 7 base pair stem between positions 73-79 and 96-102 in ancestral chimp and the creation of 2 neighbouring 4 and 5 base pair stems in Denisovan bordered by positions 68-72/92-96 and 74-77/87-90. The orange arches in the Denisovan HAR1 CS²-UPlot suggest the existence of a powerful energetic pressure to recreate the ancestral chimp stem between positions 73-79 and 96-102. This stem re-appears in the human HAR1 ncRNA secondary structure due to changes in stem structures between positions 27-46 and 51-66.

We can also notice in the CS²-UPlot that the human secondary structure is by far the most stable out of all three HAR1 structures given the sparsity of stronger base pairing probabilities (less orange and green arches).

4 FUTURE IMPROVEMENTS AND USABILITY

The comparison of multiple RNA secondary structures using the CS²-UPlot can be also achieved by combining on a single plot all concentric layers representing the corresponding sequences and the uncertainty base pairings. The only modification that is required is to use different color palettes (one for each RNA sequence) for arch coloring, which in turn might limit the total number of sequences that can be represented and compared in this fashion to 6 or 7. The superposition of concentric rings representing the base sequences will also allow a fast identification of base pair mutations as opposed to looking at separate plots.

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