CITS4008 Assignment 2

Max Ward

April 4, 2014

Abstract

RNA is the messenger molecule for DNA. It also plays a vital role in cellular metabolism. As a result, it is valuable to predict the secondary structure of RNA molecules. I have found that the RNA folding problem is equivalent to finding the maximum-weight independent set of a circle graph. In addition the Nussinov algorithm, which is used to fold RNA, can be used to solve the maximum-weight independent set problem, and is competitive with similar algorithms.

Contents

1	Dyr	namic Programming	2
	1.1	The Nussinov Algorithm	2
	1.2	The Zuker Algorithm	3
		1.2.1 Thermodynamic Model Improvements	5
	1.3	Parallel Implementations	5
2	Sto	chastic Context Free Grammars	5
3	Uni	fication and Comparison	5
4	Con	nclusion	5

Introduction

Ribonucleic Acid (RNA) is a biologically active molecule similar in structure to DNA. Recent research has found myriad functions for RNA. For example, RNA can act as a catalyst for mRNA splicing, peptide bond formation, and can alter the regulation of genes [19]. Because of its inherently single stranded nature, RNA forms bonds with itself, folding into secondary and tertiary structures [6].

It is axiomatic that chemical structure is tantamount to biological function, and RNA is no exception. For this reason, there has and continues to be an intense interest in predicting the secondary structure of RNA molecules in silico. This is because it will allow the detection and classification of unknown RNAs, and assist the design of new RNA based drugs [4]. The secondary structure of RNA is also highly conserved during evolution, indicating its importance [9].

I endeavour to give the reader a brisk but incisive review of RNA secondary structure prediction algorithms in this paper. For the sake of succinctness, my focus shall be pseudoknot-free RNA prediction methods. Particularly those able to predict RNA structures ex nihilo—that is with no information other than the primary RNA sequence itself.

1 Dynamic Programming

1.1 The Nussinov Algorithm

According to the thermodynamic hypothesis, biologically active molecules should form structures that have minimal free energy and thus maximum stability [ref]. For RNA molecules, maximising bonds is a crude but nonetheless accurate measure of energetic stability, as every bond increases the stability of a structure [14]. In the late 1970s, when the first large RNA molecules were being successfully sequenced, Nussinov et al. [14] introduced an algorithm to find a single structure with the maximal number of bonds using dynamic programming. This is possible only with the restriction that all bonding pairs are nested, and hence contains no pseudoknots.

Because of its dynamic programming nature, this algorithm performs recursive decompositions of an RNA and builds larger structures out of repeated substructures. A natural representation of this is depicted in Figure 1. Part A of Figure 1 shows bonds as arcs across a circular graph. Once can also see the nested nature of the structures being explored by the Nussinov algorithm. Part B shows how these structures translate to actual RNAs, and how these appear in vivo.

$$M(i,j) = \max \{A, B, C, D\}$$

$$A = M(i,j-1)$$

$$B = M(i+1,j)$$

$$C = M(i+1,j-1) + W(i,j)$$
(1)

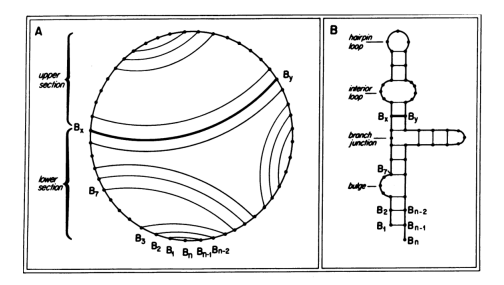


Figure 1: RNA secondary structure as described in the Nussinov algorithm. Taken from a publication by Nussinov & Jacobson [13].

$$D = \max \{ M(i,k) + M(k+1,j) \} \ when \ i < k < j$$

Equation 1 describes the recurrence relation for the Nussinov algorithm. The first two cases (A and B) find the score associated with not allowing the positions i and j to bond. The case C conversely determines the score given that positions i and j are bonded. The final case D computes the score associated with a bifurcation. A bifurcation here means decomposition of the RNA into two separate structures.

1.2 The Zuker Algorithm

Soon after the work of Nussinov & Jacobson, Zuker & Stiegler [?] described an altered version of the same algorithm which, instead of maximising base pair weights, minimized the free energy of the secondary structure. This was done by introducing a number of thermodynamic rules for canonical structures like hairpin loops, internal bulges, multiloops, unbonded base pairs, and stacked base pairs. The algorithm is similar to the Nussinov algorithm but adds another mutually recursive dynamic programming recurrence to inject a complex and relatively comprehensive energy system. The original energy system used is borrowed from the work of Studnicka et al. [?] who presented a complex but theoretically similar algorithm, albeit with much worse asymptotic and implementation complexities.

First I shall introduce some useful terminology which should clarify aspects of Zuker & Stieglers algorithm. The bases of an RNA molecule can be thought of as vertices in a planar graph. Edges between such vertices are then represented as chords on a semicircular diagram (Figures 1 and 2). These chords are not allowed to touch. A chord is admissible if it represents a chemically valid bond, and an admissible structure is a structure whose graph contains only admissible bonds. Thence, one can define a face of such a graph as any planar region bounded on all sides—either

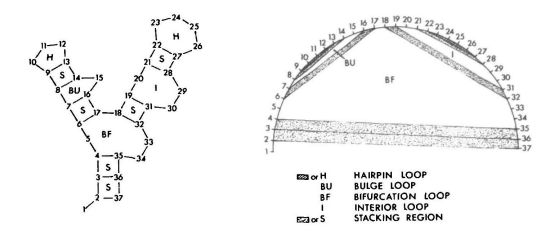


Figure 2: Diagram of faces used in the Zuker algorithm. Taken from original publication [?].

by chords, or the edge of the graph. The folding algorithm of Zuker & Stiegler considers such faces as the basic contributing factor to a molecule's stability, unlike the algorithm of Nussinov & Jacobson which considers only individual bonds.

Let E(F) represent the energy of a face F; impossible structures are given an energy value of infinity, for example hairpin loops with less than three bases in the intervening loop region. In addition let V(i,j) be defined as the minimum free energy of all structures in which bases i and j are bonded, and let W(i,j) represent the minimum free energy of all structures contained within bases i and j inclusive. Note that for W(i,j) there may or may not be a bond between bases i and j. Also, if i and j cannot bond then $V(i,j) = \infty$. Finally note that FH(i,j) represents a hairpin loop structure from i to j, and that FL(i,j,i',j') is defined as the region bounded by the bonds i,j and i',j'. Examples of these decompositions are shown diagrammatically in the right half of Figure 2. The labelled regions show faces in a semicircular graph representing a strand of RNA. In the accompanying left half of the figure, the same RNA structure is shown as it would appear in a real RNA rather than in a purely diagrammatic depiction.

$$V(i,j) = \min \{E1, E2, E3\}$$

$$E1 = E(FH(i,j))$$

$$E2 = \min \{E(FL(i,j,i',j')) + V(i',j')\} \text{ where } i < i' < j' < j$$

$$E3 = \min \{W(i+1,i') + W(i'+1,j1)\} \text{ where } i+1 < i' < j-2$$

As shown by the definition provided in Equation 2, V(i, j) is computed by minimizing three cases. The first case considers the bond between i and j closing off a hairpin loop (H in Figure 2). The second accounts for situations in which i and j are bonded. This can result in a bulge (BU in Figure 2), internal loop (I in Figure 2), or the continuation of a stacking region with the interior bond i', j' (S in Figure 2). The third and final case considers bifurcations (BF in Figure 2).

$$W(i,j) = \min \{W(i+1,j), W(i,j-1), V(i,j), E4\}$$

$$E4 = \min \{W(i,i') + W(i'+1,j)\} \text{ where } i < i' < j-1$$
(3)

Equation 3 is the recurrence for W(i,j) as described by Zuker & Stiegler. Again there are three cases. The first two cases W(i+1,j) and W(i,j-1) should be thought of as a single case which consider situations in which there is no bond between i and j. This is similar to cases A and B from the Nussinov algorithm (Equation 1). The final case considers taking the bond from i to j. This final case allows for bifurcations in which two bonding pairs split the structure into two sections. The final minimum free energy of the best structure is defined by W(1,n), where n is the length of the RNA molecule. It should be noted that the free energy for small molecules (fewer than 6 nucleotides in length) can easily be precomputed, and forms the base case of the given recurrence relations. Because of its efficiency $(O(N^3))$ time and $O(N^2)$ space), robustness, and extensibility, this method is, even today, still the most popular available. The most widely used packages for RNA secondary structure prediction all contain implementations of the Zuker algorithm [11, ?].

1.2.1 Thermodynamic Model Improvements

- 1.3 Parallel Implementations
- 2 Stochastic Context Free Grammars
- 3 Unification and Comparison
- 4 Conclusion

References

- [1] B Alberts, D Bray, K Hopkin, A Johnson, J Lewis, M Raff, K Roberts, and P Walter. Essential cell biology; garland science: New york, 2009. pages 427–452.
- [2] Michaël Bon and Henri Orland. Tt2ne: a novel algorithm to predict rna secondary structures with pseudoknots. *Nucleic acids research*, 39(14):e93–e93, 2011.
- [3] Paul Bonsma and Felix Breuer. Counting hexagonal patches and independent sets in circle graphs. *Algorithmica*, 63(3):645–671, 2012.
- [4] Anne Condon. Problems on rna secondary structure prediction and design. In *Automata, Languages and Programming*, pages 22–32. Springer, 2003.

- [5] Jason Cong and CL Liu. Over-the-cell channel routing. Computer-Aided Design of Integrated Circuits and Systems, IEEE Transactions on, 9(4):408–418, 1990.
- [6] Graeme L Conn and David E Draper. Rna structure. Current opinion in structural biology, 8(3):278–285, 1998.
- [7] Dominique de Werra, Ch Eisenbeis, Sylvain Lelait, and Bruno Marmol. On a graph-theoretical model for cyclic register allocation. *Discrete Applied Mathematics*, 93(2):191–203, 1999.
- [8] Fanica Gavril. Algorithms for a maximum clique and a maximum independent set of a circle graph. *Networks*, 3(3):261–273, 1973.
- [9] Ivo L Hofacker. Rna consensus structure prediction with rnaalifold. In *Comparative Genomics*, pages 527–543. Springer, 2008.
- [10] Junjie Li, Sanjay Ranka, and Sartaj Sahni. Multicore and gpu algorithms for nussinov rna folding. In Computational Advances in Bio and Medical Sciences (ICCABS), 2013 IEEE 3rd International Conference on, pages 1–2. IEEE, 2013.
- [11] Ronny Lorenz, Stephan HF Bernhart, Christian Hoener Zu Siederdissen, Hakim Tafer, Christoph Flamm, Peter F Stadler, Ivo L Hofacker, et al. Viennarna package 2.0. Algorithms for Molecular Biology, 6(1):26, 2011.
- [12] Nicholas Nash and David Gregg. New algorithms for maximum independent sets of circle graphs. 2013.
- [13] Ruth Nussinov and Ann B Jacobson. Fast algorithm for predicting the secondary structure of single-stranded rna. *Proceedings of the National Academy of Sciences*, 77(11):6309–6313, 1980.
- [14] Ruth Nussinov, George Pieczenik, Jerrold R Griggs, and Daniel J Kleitman. Algorithms for loop matchings. SIAM Journal on Applied mathematics, 35(1):68–82, 1978.
- [15] Jana Sperschneider and Amitava Datta. Knotseeker: Heuristic pseudoknot detection in long rna sequences. RNA, 14(4):630–640, 2008.
- [16] Krister M Swenson, Yokuki To, Jijun Tang, and Bernard ME Moret. Maximum independent sets of commuting and noninterfering inversions. BMC bioinformatics, 10(Suppl 1):S6, 2009.
- [17] Ignacio Tinoco Jr and Carlos Bustamante. How rna folds. *Journal of molecular biology*, 293(2):271–281, 1999.
- [18] Gabriel Valiente. A new simple algorithm for the maximum-weight independent set problem on circle graphs. In *Algorithms and Computation*, pages 129–137. Springer, 2003.
- [19] Zhenjiang Xu, Anthony Almudevar, and David H Mathews. Statistical evaluation of improvement in rna secondary structure prediction. *Nucleic acids* research, 40(4):e26–e26, 2012.