Loop Grammars to Identify RNA Structural Patterns

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Abstract: The biological functions of an RNA molecule are largely determined by molecular configuration. Understanding the link between the structure and the biological functions has been considered one of the challenges in biology. In this study, we face the problem of identifying a given structural pattern into an RNA pseudoknot-free secondary structure. We introduce a context-free grammar, Loop Grammar, that formalizes the primary and secondary structure of an RNA molecule as a composition of loops. Such composition is expressed as to concatenation or nesting of the simplest structural elements, hairpins, generated during the folding process when a bond between two nonconsecutive nucleotides is established. Then, we formalize the concatenation and nesting on Fatgraphs, oriented surfaces with boundary, and we define a Surface Loop Grammar, whose algebraic expressions uniquely identify such surfaces associated with given RNA structures. The terms of the Loop Grammar allow us to face the problems of identifying substructures considering both the primary and secondary structures, while the strings generated by Surface Loop Grammar permit to identify a given structural pattern in a secondary structure in terms of relations among hairpins. Both use the string pattern matching.

1 INTRODUCTION

RNA is a single strand polymer, named primary structure, that consists of four different nucleotides, Adenine (A), Guanine (G), Cytosine (C) and Uracil (U), linked together by phosphodiester bonds, referred to as strong bonds. RNA folds back on itself determining complex three-dimensional shapes known as secondary and tertiary structures (Dill, 1990; Ferré-D’Amaré and Doudna, 1999). During the folding process, each nucleotide can interact with another one by establishing a hydrogen bond, referred to as weak bond, mainly Watson-Crick (G-C and A-U) and wobble (G-U) base pairs. RNA molecules play numerous roles in cellular, and they are classified according to the functions that perform in the cell. Main classes of RNA are messenger RNA (mRNA), transfer RNA (tRNA) and ribosomal RNA (rRNA), each of which performs different but cooperative functions in protein synthesis. Functional RNA families such as tRNA and rRNA exhibit a highly conserved shape of the secondary structure, but little sequence similarity (Höchsmann et al., 2004). Therefore, it is of great interest the possibility of comparing and identifying RNA secondary structures directly, i.e., without relying on sequence similarity (Jiang et al., 2002), while the identification of common primary and secondary structures is useful to study the consequences of the RNA secondary structures changes in the RNA-RNA interactions. Moreover, searching for sequence motifs has been a powerful tool for analysis of DNA and proteins; but this approach does not work as effectively with RNA because conserved RNA structures may have no detectable sequence similarity (Li et al., 2008). In the literature, several approaches have been studied over the years for finding common patterns. Wang et al. proposed an algorithm for finding the largest approximately common substructures between two trees (Wang et al., 1998). Höchsmann et al. gives a method for finding local patterns in a tree-representation of RNAs (Höchsmann et al., 2003). Algorithms based on tree data structures were also proposed in (Mauri and Pavesi, 2005). Backofen and Siebert introduced an approach for computing common sequential and structural patterns based on dynamic programming (Backofen and Siebert, 2007).

According to Waterman (Waterman and Smith, 1978), an RNA secondary structure is composed of five basic structural elements namely hairpins, internal loops, bulges, helixes (or stacks) and multi-loops, illustrated in Figure 3 of Section 2. Each of them, characterized by strong and weak bonds, is a loop. They are generated when at least one base pair is formed. Disregarding the spatial configuration of the
molecule and reducing nucleotides to dots, an RNA secondary structure can be schematically represented by a squiggle-plot representation like the one in Figure 1-A, where solid and zigzag lines represent strong and weak bonds, respectively. Another way is the arc diagram, where the nucleotides are represented by vertices on a straight line (backbone) and the base pairs are depicted by arcs in the upper half-plane, as illustrated in Figure 1-B.

Figure 1: The secondary structure of homo sapiens miR-516a-3p predicted by Mfold (Zuker, 2003). (A) A squiggle-plot representation and (B) the arc diagram of the molecule is illustrated.

Taking advantage of the arc diagram representation, it is possible to observe that given two loops there are only two possible cases: a loop follows the other, referred to as loop concatenation, or it is nested into the other, referred to as loop nesting, as shown in Figure 2, respectively. In this work, we do not consider crossing between loops since we face the problem of identifying structural patterns in RNA pseudoknot-free secondary structures.

In this paper, we define a context-free grammar, called Loop Grammar. Each term of the grammar represents both primary and secondary structure of an RNA molecule as a composition of loops. The proposed composition is expressed as concatenation and nesting of hairpins, considered as base loops. Both concatenation and nesting are also formalized on surfaces. Such formalization permits to introduce another grammar, called Surface Loop Grammar, whose algebraic expressions uniquely identify fatgraphs, oriented surfaces with boundary, associated with given RNA structures (Penner et al., 2010). In other words, a string obtained by Loop Grammar models an RNA structure, while the corresponding term of Loop Surface Grammar identifies the surface associated with the given structure. The terms of the grammars allows us to identify RNA structural pattern in terms of the strings matching, which consists of finding all the matching strings occurrences of a pattern string in other string, and string pattern matching, that tries to find a place where one or several strings are found within a larger string or text.

The paper is organized as follows. In Section 2, we introduce some preliminary concepts. Firstly, the concept of the loop and the corresponding RNA decomposition are presented. Secondly, some mathematical definitions related to RNA secondary structures and the corresponding topological concepts are recalled. In Section 3, we introduce a context-free grammar, Loop Grammar. In Section 4, we define two topological operators over fatgraphs and formulate the Surface Loop Grammar taking advantage of the topological operators. In Section 5, we discuss the obtained results. The paper ends with some conclusions and future perspective, Section 6.

2 BASIC CONCEPTS

In this section, we introduce preliminary notions. The concept of the loop and the corresponding RNA decomposition are presented in Section 2.1, whereas some mathematical representations of RNA secondary structures and some topological concepts are recalled in Section 2.2.

2.1 RNA Secondary Structure: Loops as Structural Elements

Each RNA secondary structure is characterised by strong and weak bonds. Strong bonds link two consecutive nucleotides, whereas weak bonds connect two non-consecutive nucleotides. According to Waterman (Waterman and Smith, 1978), each RNA secondary structure can be uniquely decomposed into five basic structural elements, namely hairpin, internal loop, bulge, helix, and multi loop, illustrated in
Figure 3: Basic structural elements of RNA secondary structures: hairpin (A), internal loop (B), bulge (C), helix (D), and multi-loop (E). A strong bond is depicted with a line, a weak bond is drawn with a zigzag line and several consecutive strong bonds are represented by a dashed line.

Each of them is a loop made of a set of nucleotides linked by strong and weak bonds. A hairpin, depicted in Figure 3-A, is a loop characterised by one weak bond enclosing a sequence of nucleotides linked by strong bonds. An internal loop, represented in Figure 3-B, is defined by two weak bonds alternating with two non-empty sequences of nucleotides linked by strong bonds. A bulge, shown in Figure 3-C, is a special case of internal loop in which one of the two sequences of nucleotides is empty. A helix, illustrated in Figure 3-D, is also a special case of internal loop in which both sequences are empty. Finally, a multi-loop, depicted in Figure 3-E, consists of more than two weak bonds separated by non-empty sequences of nucleotides linked by strong bonds.

2.2 Representations and Topology of RNA Secondary Structures

An RNA secondary structure can be schematically represented by a squiggle-plot representation like the one in Figure 1-A, where solid and zigzag lines represent strong and weak bonds, respectively. A special case of this representation is the arc diagram, which is obtained from the mentioned above depicting each vertex on a straight line and connecting two non-consecutive vertices by an arc, which corresponds to weak bound, in the upper half-plane.

Definition 1 (Arc Diagram). An Arc Diagram is a labeled graph over the vertex set $[\ell] = \{1, \ldots, \ell\}$, in which each vertex has degree $\leq 3$, and the edges are all the segments $[i, i+1]$ for $i = 1, \ldots, \ell-1$ and some semi-circular arcs $(i, j)$ in the upper half-plane, with $1 \leq i < j \leq \ell$.

For each arc diagram, it is possible to associate the linear chord diagram deleting the unpaired nucleotides, i.e. nucleotides that do no any crossing.

Definition 2 (Linear Chord Diagram). A linear chord diagram consists of a line segment, called its backbones, to which are attached a number $n_0$ of chords with distinct endpoints.

As an example, the linear chord diagram associated with the miR-516a-3p secondary structure illustrated in Figure 1-B is shown in Figure 4-A.

Each linear chord diagram admits a fattening. A fatgraph $F$ is a graph equipped with a cyclic order on the edges incident on each vertex, as shown in Figure 4-B. It uniquely determines an oriented surface with boundary. For more details regarding the concept of fatgraph, interested readers can refer to (Penner et al., 2010).

3 LOOP GRAMMAR

Several context-free grammars have been proposed in the literature. Knudsen and Hein proposed a very simple grammar (Knudsen and Hein, 1999), which has been implemented into the secondary structure prediction software Pfold (Knudsen and Hein, 2003). Other proposed grammars are (Dowell and Eddy, 2004; Sakakibara et al., 1994). Each term of these grammars is the primary structure of an RNA molecule, while the parse tree has a natural correspondence with its secondary structure. Other grammars, such as RNAFeatures, are designed to explicitly designate the different structural features (Giegerich, 2014), while others describe RNA structures in dot-parenthesis notation such as (Anderson et al., 2012).

We introduce a context-free grammar that models the RNA secondary structure in terms of loops. Differently from RNAFeatures, the Loop Grammar does not explicitly identify structural components since it represents each RNA secondary structure as a composition of hairpins. As a consequence, the Loop grammar is characterized by a set of 5 productions, while Grammar RNAFeatures is defined using more of 20 rules. The Loop grammar is unambiguous and imposes a particular order to add a new weak bond.

Definition 3. Let $\Sigma_{RNA} = \{A, U, G, C\}$ be the alphabet of RNA nucleotides, and let $\Sigma_{RNA} =
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\{ (A, U), (U, A), (G, C), (C, G), (G, U), (U, G) \} be the alphabet of weak bonds, whose elements represent pairs of nucleotides. The Loop Grammar is \( \mathcal{L}_{RNA} = (V_N, V_T, P, S) \) where \( V_N = \{ S, S', H \} \), \( V_T = \Sigma_{RNA} \cup \Sigma_{\text{RNA'}} \cup \{ [ ] \} \), and the set of productions \( P \) is

\[
S ::= \epsilon \quad \text{empty structure} \\
S' ::= sS \quad \text{non-empty structure} \\
H ::= HS \quad \text{secondary structure} \\
H ::= x[S'] \quad \text{base loop}
\]

where \( x \in \Sigma_{RNA} \) and \( s \in \Sigma_{RNA} \).

The start symbol \( S \) formalizes empty or non-empty RNA structure, whereas non-terminal symbol \( S' \) represents any RNA primary and secondary structure. A primary structure, a sequence of unpaired nucleotides, can be uniquely represented applying production \( S \rightarrow S' \) followed by \( S' \rightarrow sS \). The secondary structure, \( HS \), is composed by a loop \( H \) followed by a structure \( S \). Each loop \( H \) is formalized by production \( H \rightarrow x[S'] \), where \( S' \) could be both primary and secondary structure. If \( S' \) is a sequence of unpaired nucleotides the grammar generates a hairpin, otherwise one of the other four loops (internal loop, helix, multiloop, bulge) is formalized as a composition of base loops. Note that this representation permits to naturally associate the Loop Energy Model, where the total energy \( E \) of a structure \( S \) is the sum over the energy contributions of each constituent loop \( H \) (Zucker and Stiegler, 1981). As a consequence, associating a probability distribution over the production rules we can also use this formalization to predict the RNA secondary structures based on this energy model.

As an example, we use Loop Grammar to represent the molecule illustrated in Figure 1. The first step is to formalize that the structure is not empty by rule \( S \rightarrow S' \) and to represent the head composed of the unpaired nucleotide \( U \) by \( S' \rightarrow sS \). This unpaired sequence is followed by loops formalized by rule \( S' \rightarrow HS \). Such loop, generated by the weak bond between the second and the last nucleotide, is formalized by production \( H \rightarrow x[S'] \). In this case, the substructure is determined by the weak bond that involves the third and the second to the last nucleotide and it is formalized by the only possible sequence of rules \( S' \rightarrow HS \), \( S \rightarrow \epsilon \) and \( H \rightarrow x[S'] \). The same sequence must be used to model the weak bond between the fourth and the third to the last nucleotide. Instead, to represent the sub-motif that involves the nucleotides from the fifth to the fifteenth one, it is necessary to include at the beginning and at the ending of the previous sequence the production \( S' \rightarrow sS \) for representing the two unpaired sequences of nucleotides, \( UC \) and \( AG \). Lastly, the unpaired sequence nested to the innermost weak bond is formalized using the production \( S' \rightarrow sS \). The string associated to the considered molecule is \( U(G,U)(C,G)(U,C,G)U(U,U,C,A)[A,G] \). Such scheme works in general to give a unique algebraic expression of each motif of RNA secondary structure. This observation yields the following:

**Theorem 1.** The Loop Grammar, \( \mathcal{L}_{RNA} \) generates uniquely all RNA structures.

It is equivalent to prove that the grammar \( \mathcal{L}_{RNA} \) is not ambiguous. A technique for proving it is by induction over nucleotides and loops or proving that the grammar is \( LR(1) \), but it is omitted since it is essentially the same as the proof of Theorem 2.

**Corollary 1.** Each derivation path of the Loop Grammar, \( \mathcal{L}_{RNA} \), corresponds uniquely to an RNA secondary structure.

Each term obtained by grammar \( \mathcal{L}_{RNA} \) uniquely represents a particular molecule. It is a word over \( \Sigma = \Sigma_{RNA} \cup \Sigma_{\text{RNA'}} \cup \{ [ ] \} \), where \( \Sigma_{RNA} \) and \( \Sigma_{\text{RNA'}} \) are the alphabets of unpaired nucleotides and weak bonds, respectively. In this work, we assumed that only Watson-Crick and wobble base pairs characterize the RNA molecule; if we want to add also the non-canonical weak bonds, it is enough to add the corresponding pairs in \( \Sigma_{\text{RNA'}} \). Instead, the extra symbols, \(" [ ] \)\”, are introduced to model the fact that a weak bond is contained into another one. In other words, a loop is nested into another one. By abuse of notation, we do not introduce another type of extra symbols to model that a weak bond is followed by is another one. The abuse consists in the fact that the usual concatenation has been used to concatenate both nucleotides and loops.

### 4 Topological Formalization

In this section, we will focus on the relations, nesting or concatenation, among hairpins. We define two topological operators, nesting and concatenation, in Section 4.1. They permit to define the Surface Loop Grammar in Section 4.2.

#### 4.1 Topological Operators

Each RNA secondary structure can be uniquely decomposed into loops. Each of them can be expressed as a composition of hairpins, as defined by Loop Grammar in Section 3. The proposed composition is based on nesting and concatenation. Briefly, nesting corresponds to the insertion of a structure into
another one, as shown in Figure 2-A for the simple case in which a hairpin is nested into another one; while concatenation is used to represent a motif in which a structure is followed by another one, as illustrated in Figure 2-B for two simple hairpins. The two operators are described over fatgraph, which is a two-dimensional topological object that uniquely determines an oriented surface with boundary as introduced in Section 2. In topology, cutting and gluing are two methods for analysing surfaces. Taking advantage of these two methods, we define the following two operations: cutting backbone and gluing backbone. The former cuts the backbone of a fatgraph in two parts, as shown on top of Figure 5; the latter glues two parts of backbone, as illustrated in the bottom of Figure 5.

The two topological operators, nesting and concatenation, are defined as follows.

**Definition 4 (Concatenation).** Given two fatgraphs, $F_1$ and $F_2$, the concatenation, $F_1 \circ F_2$, is a fatgraph defined as $F_1$ followed by $F_2$, whose backbones are glued.

As an example, we consider the two fatgraphs, $F_1$ and $F_2$, illustrated in Figures 6-A and 6-B, respectively, whose concatenation, $F_1 \circ F_2$, is shown in Figure 6-C.

![Figure 5](image)

Figure 5: The cutting and the gluing backbone operations are illustrated on top and in the bottom, respectively.

![Figure 6](image)

Figure 6: (A) The fatgraph $F_1$; (B) the fatgraph $F_2$; (C) the concatenation of $F_1$ and $F_2$; (D) a nesting of $F_1$ and $F_2$.

**Definition 5 (Nesting).** Given two fatgraphs, $F_1$ and $F_2$, the nesting, $F_1 \sqcap F_2$, is a fatgraph defined by cutting the backbone under an arch of $F_1$ once and by gluing $F_2$, where the backbone of $F_1$ has been cut.

Differently from the concatenation, the resulting of a nesting, $F_1 \sqcap F_2$, is not unique fatgraph, since it depends on where the backbone of $F_1$ is cut. As an example, we again consider the two surfaces of Figures 6-A and 6-B, and a possible nesting is shown in Figure 6-D. Moreover, another resulting structure can also be obtained cutting the backbone in the last components and gluing the backbone of fatgraph $F_2$ that corresponds to the concatenation. In fact, from the topological point of view, both operators correspond to the connected sum (Gilbert and Porter, 1994).

### 4.2 Loop Surface Grammar

The two topological operators defined above allows us to formalize the surface associated with an RNA secondary structure using a term of a language of expressions whose grammar would look like the following one:

$$
F ::= \varepsilon \quad \text{empty structure}
| \ L \quad \text{base loop}
| F \circ F \quad \text{concatenation of fatgraphs}
| F \sqcap F \quad \text{nesting of two fatgraphs}
$$

where $F$ is a generic fatgraph and $L$ is the loop base. Each fatgraph $F$ can be defined in terms of nesting and concatenation of other fatgraphs. Such grammar can generate different derivation trees representing the same fatgraph, but this problem has been solved by the ordered given by the backbone.

**Definition 6.** The Surface Loop Grammar is $S_{RNA} = (V_N, V_T, P, S)$, where $V_N = \{S, S', L\}$, $V_T = \{(x, \bar{x})\}$, and the set of productions $P$ is

$$
S ::= \varepsilon \quad \text{empty structure}
| S' \quad \text{non-empty structure}
S' ::= L \circ S \quad \text{concatenation}
L ::= L \sqcap S' \quad \text{nesting}
| (x, \bar{x}) \quad \text{base loop}
$$

The start symbol $S$ represents any surfaces associated with RNA secondary structures, empty or non-empty. If a secondary structure is not empty, the associated fatgraph is composed of the surface associated a base loop, $L := (x, \bar{x})$, concatenated to another structure, eventually empty, or the considered surface contains another structure not empty.

**Theorem 2.** The Surface Loop Grammar $S_{RNA}$ is context-free and it generates uniquely the surface associated to the RNA structures.

The proof is reported in Appendix A.

**Corollary 2.** Each derivation path of the Surface Loop Grammar, $S_{RNA}$, corresponds uniquely to the surface associated to an RNA secondary structure.
5 RESULTS AND DISCUSSION

In this study, we have defined two context-free grammars, Loop Grammar and Surface Loop Grammar. The former models both RNAs primary and secondary structure in terms of a string as a composition of loops. The proposed composition is based on concatenation or nesting of hairpins, considered as base loops. These two operators have been also described over fatgraphs, two-dimensional topological objects that uniquely determine oriented surfaces with boundary. The operators have permitted to define a Surface Loop grammar, whose algebraic expressions uniquely identify fatgraphs. Each term of the grammar represents an RNA secondary structure without relying on sequence similarity in terms of hairpins and relations among them. To prove that the grammar is unambiguous, we have associated with it the generating function, which determines the Catalan numbers able to enumerate all possible linear chord diagrams without any crossings. The terms obtained by the two grammars allow us to face the problem of identifying substructures considering both the primary and secondary structures in terms of both strings matching and string pattern matching. In the literature, the problem has been widely addressed using trees as data structures (Cserkuti et al., 2006). Considering only pseudoknot-free structures represents a limitation of our approach due to the use of context-free grammar. This formalism is inadequate to model RNA pseudoknotted structures (Harrison, 1978).

To test the approach, we have considered the Vertebrate Telomerase RNAs. Telomerase is a ribonucleoprotein enzyme that maintains telomere length by adding telomeric sequence repeats onto chromosome ends. The most remarkable feature of this molecule is the evolutionary conservation of four structural domains: the pseudoknot domain, the CR4-CR5 domain, the Box H/ACA domain, and the CR7 domain (Chen et al., 2000). In this test, we have considered the CR4-CR5 domain from human, quoll, Xenopus, and Typhlonectes telomerase RNAs. We have represented them as strings using Loop Grammar and Surface Loop Grammar. The terms obtained with the former are reported in Tables 1, while terms obtained with Surface Loop Grammar are the same for each species. It is 

\[ S = L \sqcap L \sqcap L \sqcap L \sqcap L \sqcap L \sqcap L \sqcap L \sqcap L \sqcap L \sqcap L \sqcap L \sqcap L \sqcap L \sqcap L. \]

The term of the human CR4-CR5 domain obtained by Surface Loop Grammar matches with the strings that identify the ones of other considered species. Such string matching has been done using Notepad++ 7.6 as a text editor. Moreover, using the strings obtained with the Loop Grammar, we have identified the pattern \( \langle C, G \rangle \langle C, G \rangle \langle C, G \rangle \langle C, G \rangle \langle G, C \rangle \langle S_x \rangle \), which determines the Catalan numbers only in human and in Quoll CR4-CR5 domain. To identify these string patterns, we have developed a prototype Java tool based on regular expressions and Java Regular Expression Tester (Expression Tester, 2018). The regular expressions allow us to recognize the corresponding closed bracket of the weak interaction.

Table 1: The terms of CR4-CR5 domain of Human, Quoll, Xenopus and Typhlonectes obtained with Loop Grammar. \( S_{h}, S_{q}, S_{x}, S_{t} \) are the substructure omitted due to lack of space.

<table>
<thead>
<tr>
<th>Species</th>
<th>Term</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human</td>
<td>( S_{h} = \langle C, G \rangle \langle C, G \rangle \langle C, G \rangle \langle C, G \rangle \langle G, C \rangle \langle S_{x} \rangle )</td>
</tr>
<tr>
<td>Quoll</td>
<td>( S_{q} = \langle C, G \rangle \langle C, G \rangle \langle C, G \rangle \langle G, C \rangle \langle S_{x} \rangle )</td>
</tr>
<tr>
<td>Xenopus</td>
<td>( S_{x} = \langle C, G \rangle \langle C, G \rangle \langle C, G \rangle \langle C, G \rangle \langle C, G \rangle \langle S_{x} \rangle )</td>
</tr>
<tr>
<td>Typhlonectes</td>
<td>( S_{t} = \langle C, G \rangle \langle C, G \rangle \langle A, U \rangle \langle C, G \rangle \langle S_{x} \rangle )</td>
</tr>
</tbody>
</table>

6 FUTURE PERSPECTIVES

The biological functions of RNAs are largely determined by molecular configuration. In this work, we have faced the problem of the identification of a given structural pattern into an RNA pseudoknot-free secondary structure in terms of string introducing two context-free grammars able to represent both primary and secondary structure of an RNA molecule. We are working on the development of a toolchain that implements the presented methodology and takes as input dot-bracket notation, an output of RNAStrand database (Andronescu et al., 2008). We have planned to compare our tool with the existing ones using an appropriate benchmark to study technical features such as scalability. Moreover, the tool will be tested on real RNAs. It will be carried out in collaboration with experts of the biological domain in order to test the impact of our approach on the creation of new biological knowledge.

As a future work, we want to generalize the string pattern matching of RNA secondary structures with arbitrary pseudoknots. A promising approach is based on our preliminary results (Quadrini et al., 2017; Quadrini et al., 2018; Quadrini and Merelli, 2018) since it is able to model each kind of pseudoknots differently classical approaches, such as (Rivas and Eddy, 2000). Another direction of future work is to face the problem of RNAs classification using the Surface Loop Grammar. A suitable database of functional molecules is RNA Strand Database (Harrison, 1978). Moreover, the problem of folding of RNA without pseudoknots can be also addressed with the Loop Grammar defining an opportune probability distribution.
REFERENCES


APPENDIX A

Taking advantage of the Chomsky-Schützenberger enumeration theorem, that allows one to construct an
algebraic equation whose power series expansion provides the enumeration (Chomsky and Schützenberger, 1963), we can associate to the considered grammar the following generating function observing that operator concatenation and nesting over surface correspond to the connected sum from a topology point of view

\[ S(z) = 1 + zS(z)^2 \quad (1) \]

Equation 1 is just a quadratic equation in \( S(z) \) which we can solve using the quadratic formula. In a more familiar form, we can rewrite it as: \( zS(z)^2 - S(z) + 1 = 0 \) whose solution is

\[ S(z) = \frac{1 \pm \sqrt{1 - 4z}}{2z} \]

Since it is known that \( S(0) = 1 \) and for \( z \to 0^+ \), \( S(z) \to +\infty \), choosing the positive sign in the quadratic formula, the only possible solution is

\[ S(z) = \frac{1 - \sqrt{1 - 4z}}{2z} \]

whereas for \( z \to 0^+ \),

\[ \lim_{z \to 0^+} C(z) = \lim_{z \to 0^+} \frac{2(1 - 4z)^{-\frac{1}{2}}}{2} = 1 \]

since \( \lim_{z \to 0} C(z) \) is an indetermined form of \( 0/0 \) type. To expand \( S(z) \) we will just use the binomial formula on

\[ \sqrt{1 - 4z} = (1 - 4z)^{-\frac{1}{2}} \]

whence

\[ S(z) = \frac{1 - \sqrt{1 - 4z}}{2z} = \frac{2}{1 + \sqrt{1 - 4z}} \]

Using of the binomial formula with fractional exponents follows

\[ C(z) = \frac{1}{2z} (1 - \sqrt{1 - 4z}) = \frac{1}{2z} (1 - \sum_{n \geq 0} \binom{1/2}{n} (-4z)^n) \]

(2)

it follows that

\[ (-4)^n \binom{\alpha}{n} = \frac{1}{2} \cdot (\frac{1}{2} - 1) \cdots (\frac{1}{2} - n + 1) \cdot (-4)^n \]

\[ = \frac{1}{2} \cdot (\frac{1}{2} - (\frac{2n+3}{2}) \cdot (-1)^n \cdot (2 \cdot 2)^n \]

\[ = \frac{3 \cdot 5 \cdots (3 + 2n)}{n! n!} \cdot 2^n \cdot n \cdot (n - 1) \cdots 1 \]

\[ = \frac{3 \cdot 5 \cdots (3 - 2n)}{n! n!} \cdot 2n \cdot (2n - 2) \cdots 2 \]

\[ = \frac{1}{n! n! (2n - 1)} \]

\[ = \binom{2n}{n} \frac{1}{2n - 1} \]

(3)

Substituting 3 into equation 2, we obtain

\[ S(z) = \frac{1}{2z} \left( 1 + \sum_{n \geq 0} \binom{2n}{n} \cdot \frac{1}{2n - 1} \cdot z^n \right) \]

\[ = \frac{1}{2z} \left( 1 - 1 + \sum_{n \geq 1} \binom{2n}{n} \cdot \frac{1}{2n - 1} \cdot z^n \right) \]

\[ = \frac{1}{2z} \left( \sum_{n = 0} \binom{2(n + 1)}{n + 1} \cdot \frac{1}{2(n + 1) - 1} \cdot z^n \right) \]

(4)

Applying the definition of the binomial formula follows that

\[ C(z) = \sum_{n \geq 0} \binom{2n}{n} z^n \]

where

\[ s_n = \frac{2n!}{(n + 1)! n!} \]

with \( n \) is the number of base loops, and \( s_n \) is the Catalan numbers, i.e., the number of all possible linear chord diagrams without crossing. This proves the corresponding surface context-free language is unambiguous.