

Helices of RNAs

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Outline

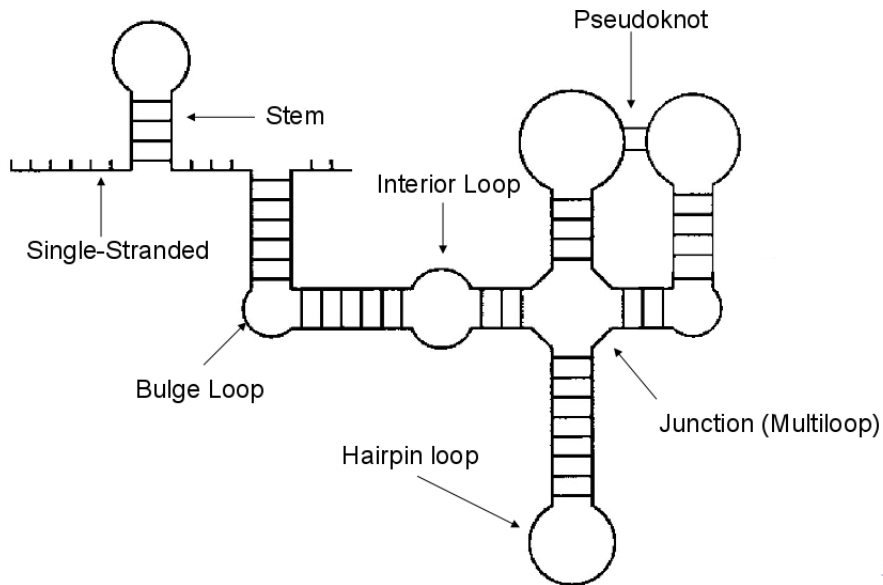
- Introducing abstract shapes
- Development of a new structure abstraction
- Implementation of an algorithm based on the new abstraction
- Possible problems

Structural components of RNA

RNA has different structural components:

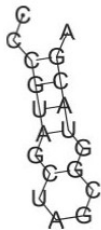
- single-stranded regions (SS)
- hairpin loops (HL)
- stacking regions (SR)
- bulges on 5' side (BL) or 3' side (BR)
- internal loops (IL)
- multiloops (ML).

Structural components of RNA

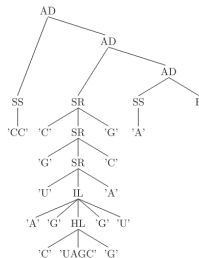


Visualisation of structures

CCCGUAGCUAGCGGUACGA



squiggle plot



tree

Suboptimal structures

But the "true" structure is not always the one with the lowest predicted free energy. So what to do?

- Enumerate suboptimal structures within a given energy range R .
- Hope to find a structure fulfilling your expectation or coming close to experimental results.

But the number of suboptimal structures grows exponentially with the energy range considered.

Introducing abstract shapes

Solution: Use abstract shapes to describe a set of structures.

- An abstract shape represents a class of similar structures sharing a common pattern of helix nesting and adjacency.
- "Abstract" since we do not care about all details of the structures.
- Each shape class has a representative structure called shrep (with minimum folding energy).

Introducing abstract shapes

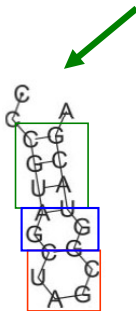
In the domain of shapes, we care only about

- open structures (OP)
- closed structure (CP)
- branching ("fork", FK)
- adjacency of structures (AD)

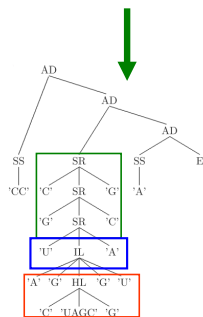
Level of detail is defined by an abstraction function π . The abstraction function can be defined on the level of the structural components.

Example

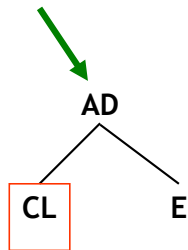
CCCGUAGCUAGCGGUACGA



squiggle plot



tree



abstract shape

Abstract shape, Energy range: 5 kcal/mol

UCGCGCACAGGACAUCCUAGGUACAAGGCCGCCGU

-6.30	.(((.(..(((....))).(((.....)))))).	[[] []]
-4.90(((....))).(((.....))).....	[] []
-3.90((..(((....)))...)).....	[]

Drawback of abstract shape

- The major drawback of abstract shape analysis is the position independence of the abstraction

new algorithm should be:

- develop a new algorithm for RNA secondary structures which keeps track of the position of structural elements
- be implemented in the framework of "Abstract shapes of RNA"
- be evaluated and compared to existing algorithms
- The new algorithm will be used to design prediction strategies for various classes of RNAs

develop a new structure abstraction

The straightforward idea to overcome the position independence of the current available shape abstractions is

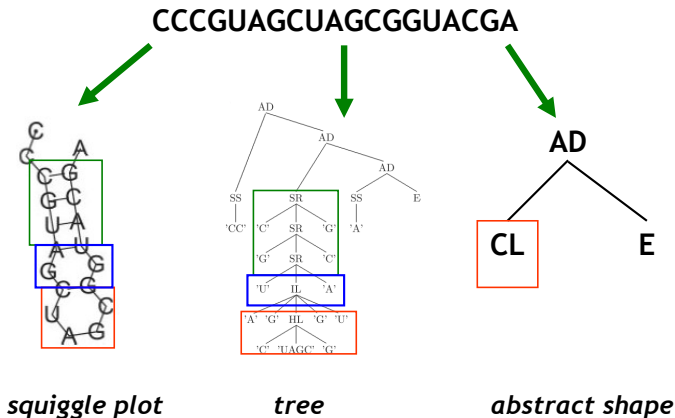
what keeps track of positions

- hairpin loop, because it is the least expensive one
- multiloops, as possible braching points, are structurally important
- stacked pairs, bulge and internal loops are the main structural contributors
- possible different helices type

which positions of the base pair will be tracked

- i
- j
- (i,j)
- $i+j/2$

Todo: capture of the pdf picture



Output from the first trial version of helices shape

Helices shape, Energy range: 5 kcal/mol

	UCGCGCACAGGACAUCCUAGGUACAAGGCCGCCGU	
-6.30	.((.(.((.(.((....)))((.(.((....)))))))).	[13.5,25]
-4.60((.(.((....)))((.(.((....))))..	[13.5,26.5]
-3.90((.(.((.(.((....)))..)).....	[13.5]
-3.60((.(.((....((.(.((....))))..))..	[22]
-3.40((.(.((.(.((....)))..))....((....))..	[13.5,30]
-3.20	..(((.(.((....((.(.((....))))..))....	[17]
-2.80((.(.((....((.(.((....))))..))..	[25]
-2.40((.(.((....((....))))..	[26.5]
-1.60((.(.((.(.((....)))..))....((....))	[13.5,31.5]

Possible problems

- do not preserve nesting of elements and might lead to abstract shapes where a helix index occurs more than once → solved with different representation form
- abstracting from bulge and internal loops might be too rigorous → refine the definition of the abstraction, such that all critical criteria are met
- records of helices is a lot more than the records of shapes → runtime problem

Outlook: Designing RNA class predictors

- specific classes of RNAs, such as riboswitches or miRNA precursors, show characteristic features within their folding space
- the energy landscape of riboswitches harbors two equally low, but well separated minima, but well separated minima
- miRNA precursors is governed by one deep and well-defined minimum
- develop specialized prediction algorithms for classes of RNAs

End

- Thanks a lot for your attention !
- Questions ?