

Computational Prediction of RNA and DNA Secondary Structure

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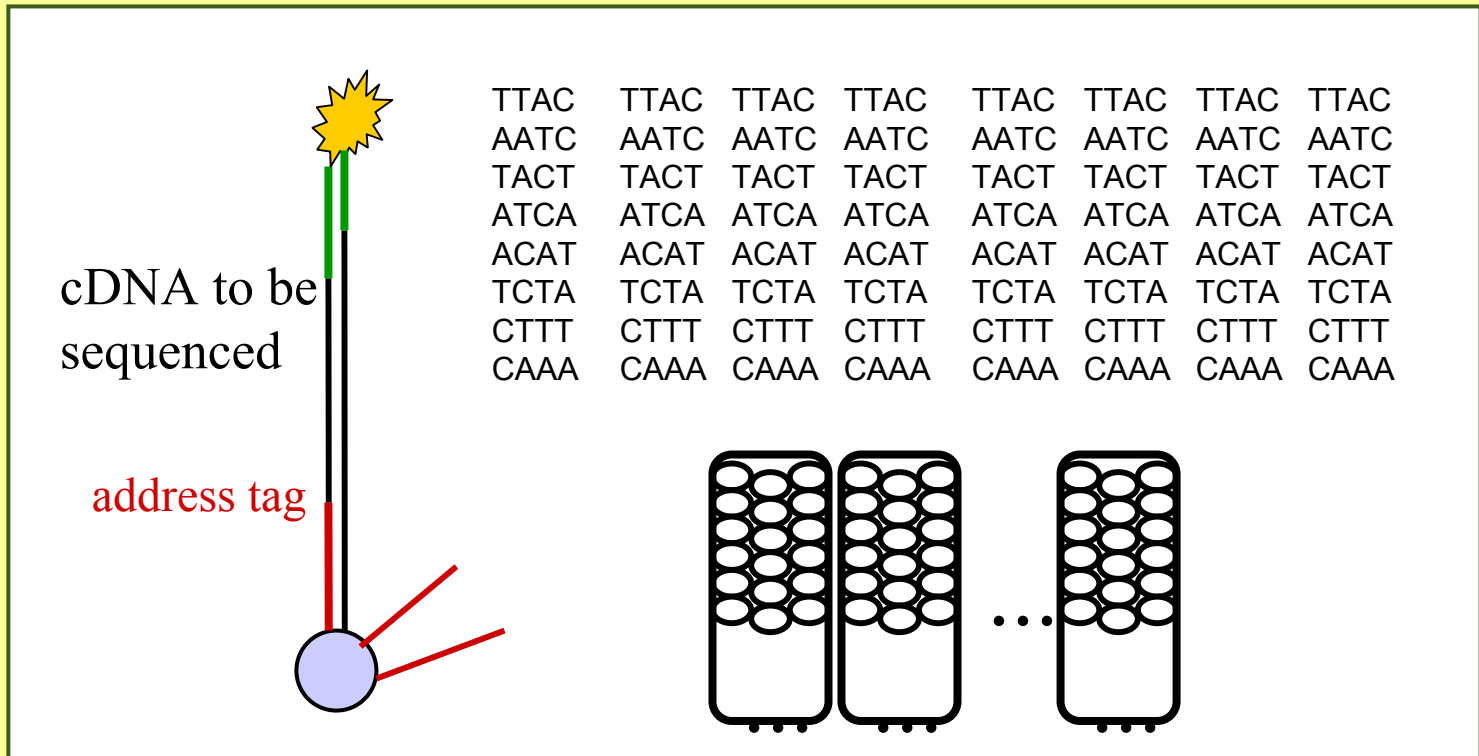
The Department of Computer Science, UBC

RNA plays varied roles in the cell...



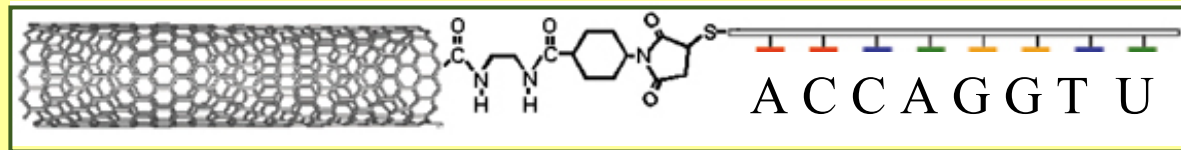
“... a familiar performer has turned up in a stunning variety of guises. RNA, long upstaged by its more glamorous sibling, is turning out to have star qualities of its own” – J. Couzin, Science, 2003

... in sequence analysis,



“...signature sequencing [permits] application of powerful statistical techniques for discovery of functional relationships among genes” – S. Brenner et al.

...nanotechnology,

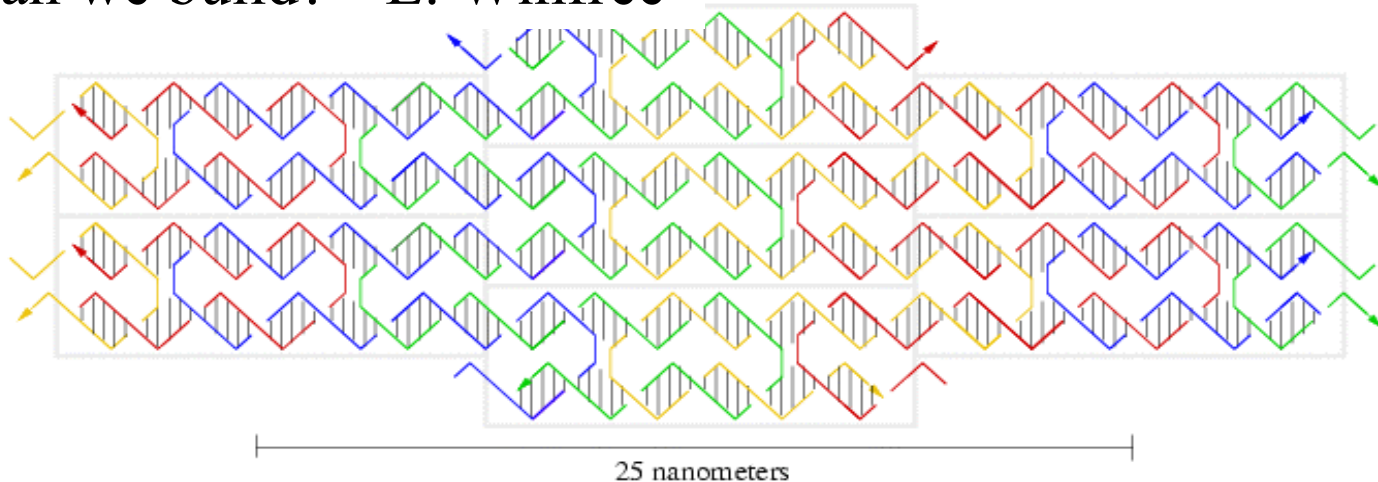
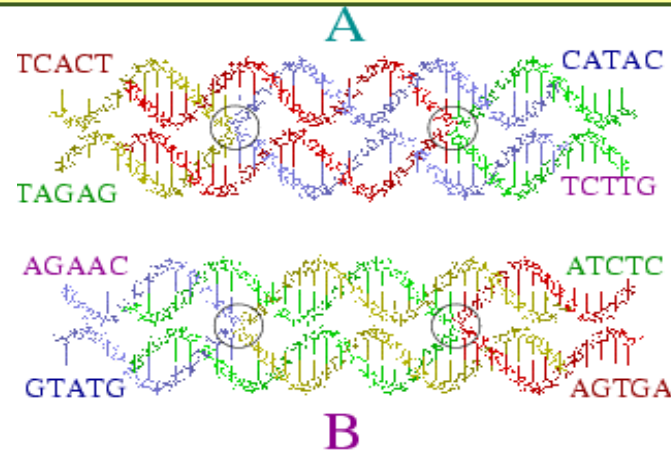


R. Hamers

“Linking biological molecules with nanotubes and nano wires [is] a method for biological sensing and for controlling nanoscale assembly” – R. Hamers, Chemistry, U. Wisconsin

and beyond...

“..rather than examining in detail what occurs in nature (biological organisms), we take the engineering approach of asking, "what can we build?" E. Winfree



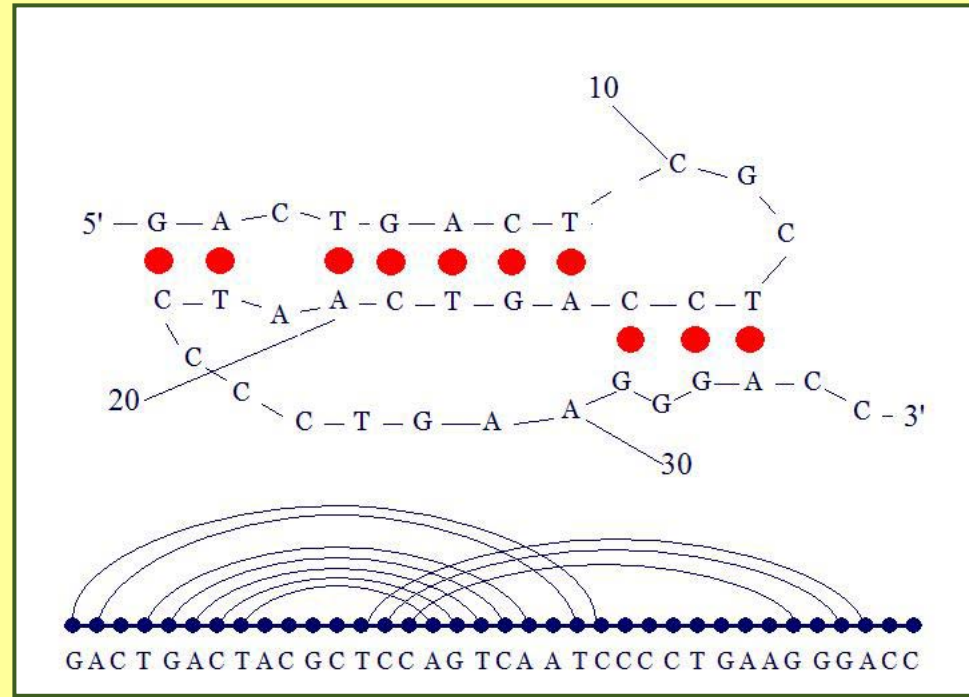
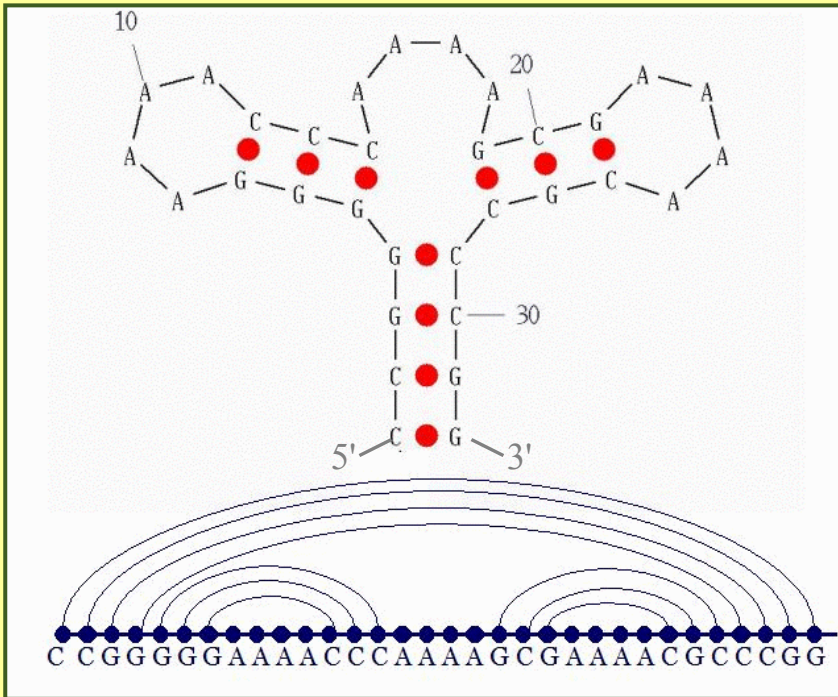
... hold promise in therapeutics,

and hold clues to our understanding of primitive life

“there are strong reasons to conclude that DNA and protein based life was preceded by a simpler life form based primarily on RNA” – G. F. Joyce

To understand the function of RNA molecules, *we need to understand their structure.*

DNA, RNA secondary structure



our goals

- ***predict*** from base sequence the secondary structure of
 - a single molecule
 - a small group of molecules
 - the most stable molecule in a combinatorial set of molecules
- ***design*** a molecule or molecules with a certain secondary structure

overview

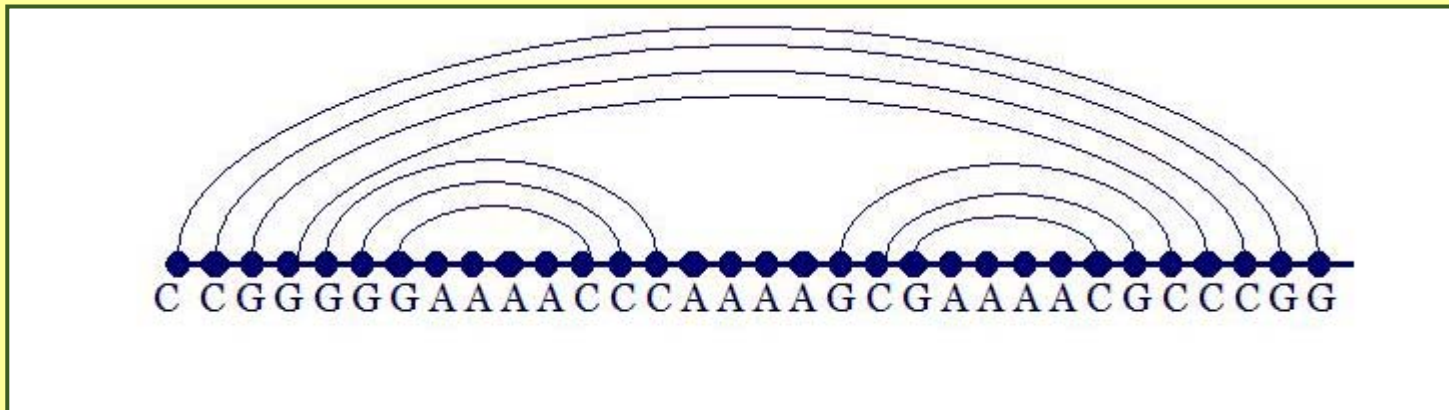
- background
 - *how is RNA secondary structure prediction currently done?*
- challenges
 - *how might it be done better?*
- projects
 - *what is the BETA-lab doing about it?*

approaches to RNA secondary structure prediction

- comparative sequence analysis
- prediction from base sequence
 - find *minimum free energy (mfe)* structure

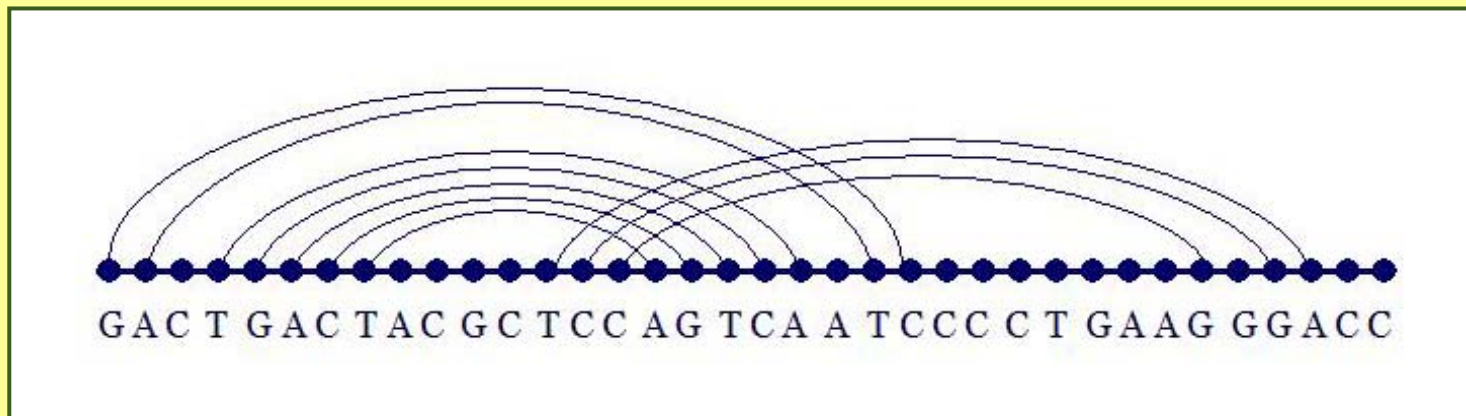
free energy model

- free energy of structure (at fixed temperature, ionic concentration) = sum of loop energies
- **standard model** uses experimentally determined thermodynamic parameters where available; extrapolations for long loops



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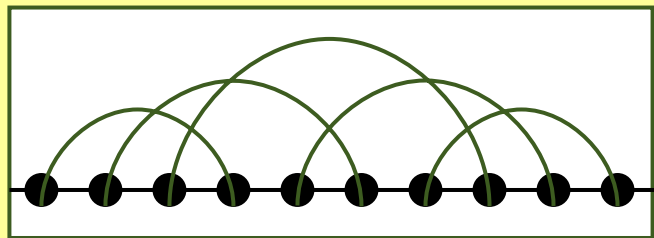


on the mfe approach

- mfe approach ignores folding pathway, metal ions, nonstandard bonds
- “some species can remain kinetically trapped in nonequilibrium states... we expect that most RNA’s exist naturally in their thermodynamically most stable configurations” —Tinoco and Bustamante, J. Mol. Biol. 1999.

why is mfe secondary structure prediction hard?

- mfe structure can be found by calculating free energy of all possible structures
- but, number of potential structures grows exponentially with the number, n , of bases
- structures can be arbitrarily complex



- *success for restricted classes of structures*

predicting mfe pseudoknot free structures

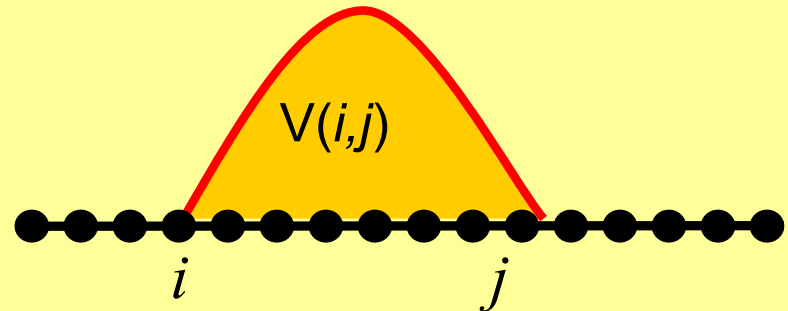
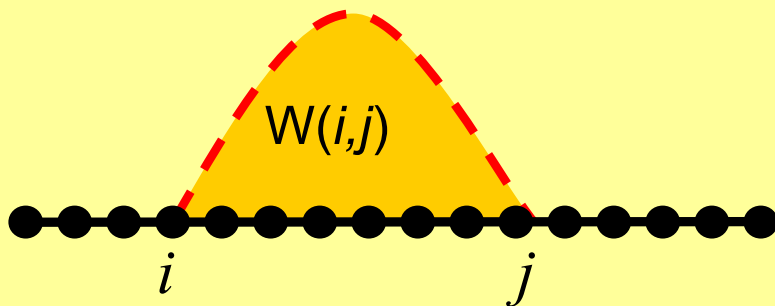
- dynamic programming avoids explicit enumeration of all pseudoknot free structures (Zuker & Stiegler 1981)
- suboptimal folds, probabilities of base pairings can also be calculated
- software: mfold, Vienna package

dynamic programming (Zuker and Steigler)

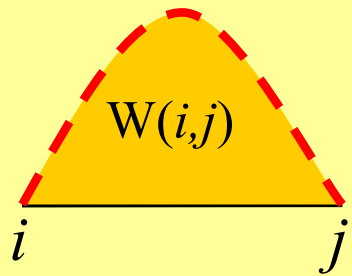
- based on the “more is less” principle: by calculating more than you need, less work is needed overall
- construct mfe structure for whole strand from mfe structures for substrands
- running time is $O(n^3)$

dynamic programming (Zuker and Steigler)

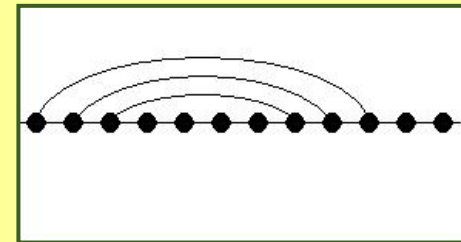
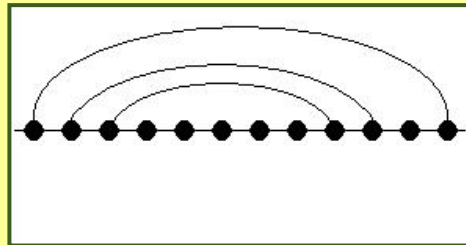
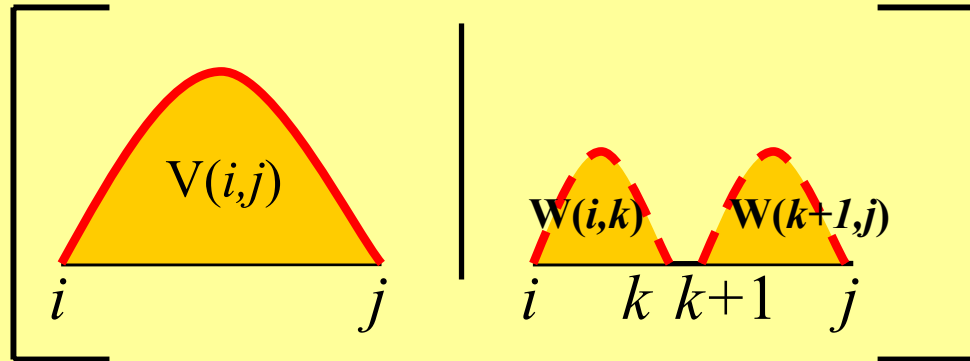
- **$W(i,j)$** : mfe structure of substrand from i to j
- **$V(i,j)$** : mfe structure of substrand from i to j , in which i th and j th bases are paired



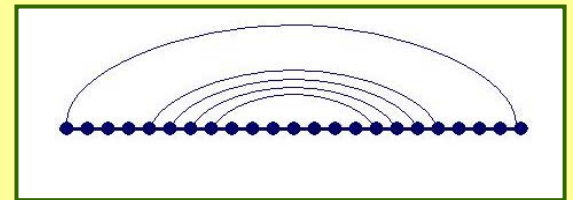
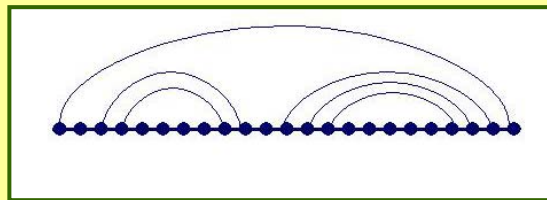
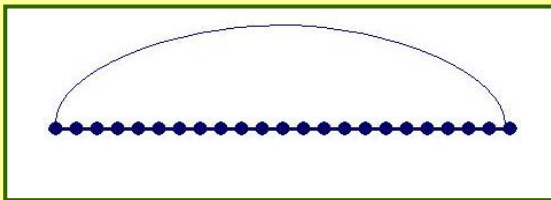
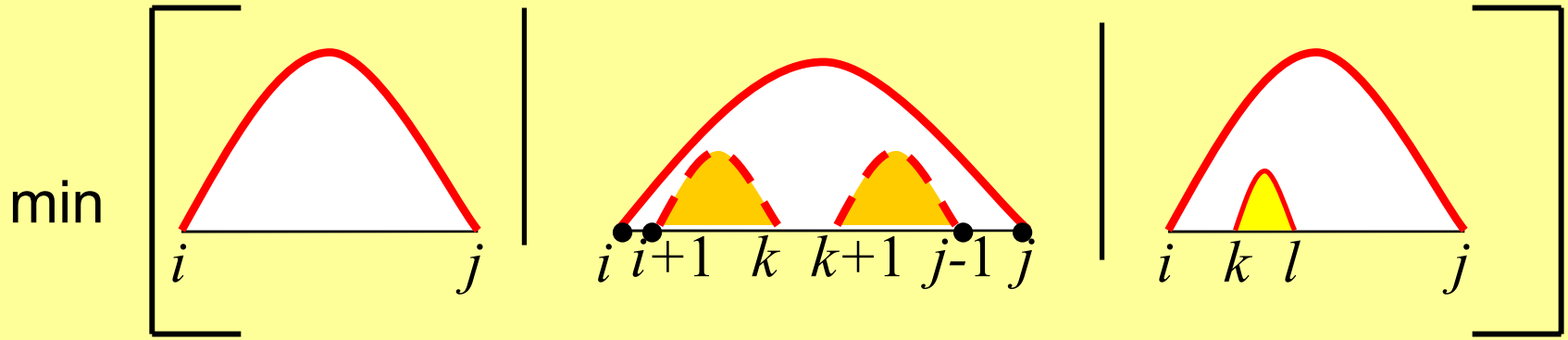
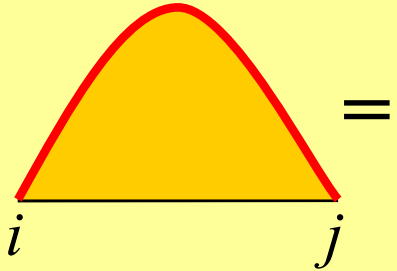
recurrences



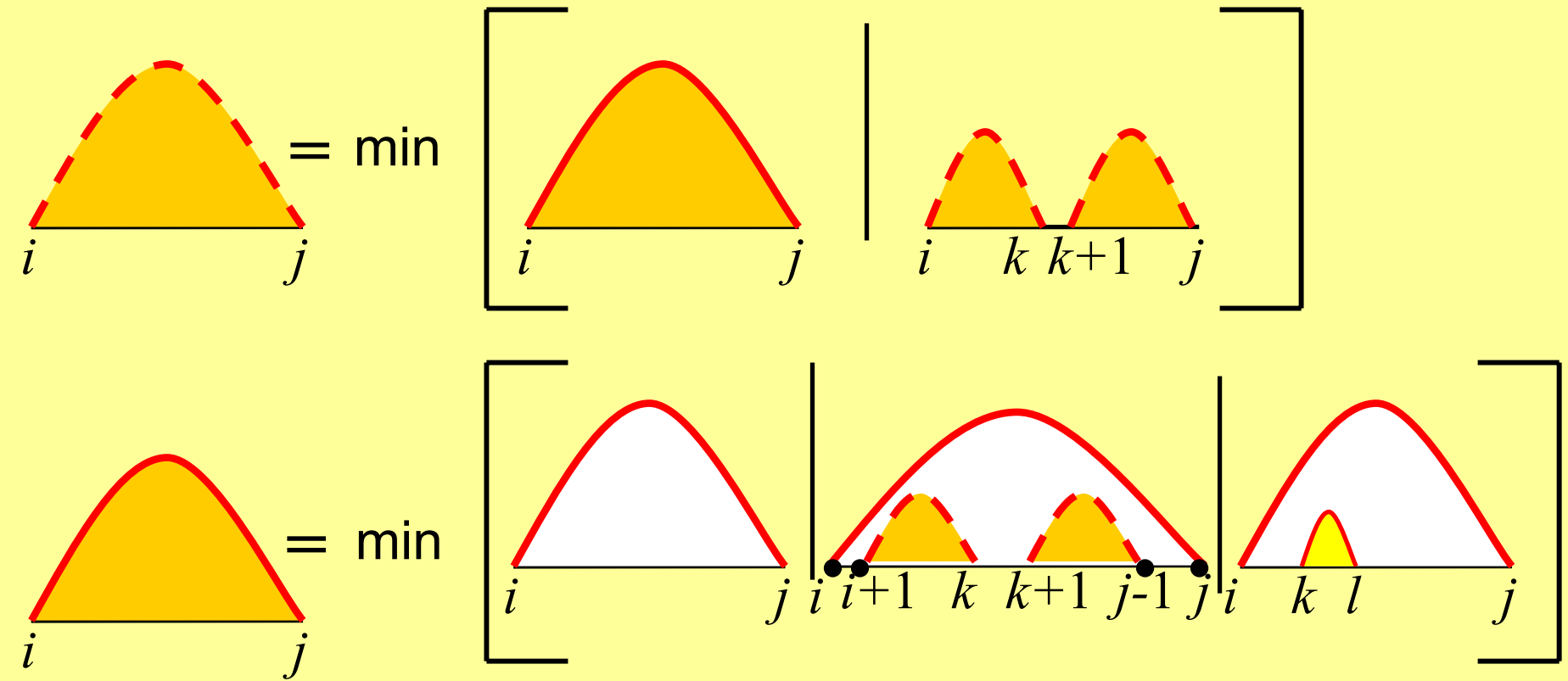
= min



recurrences



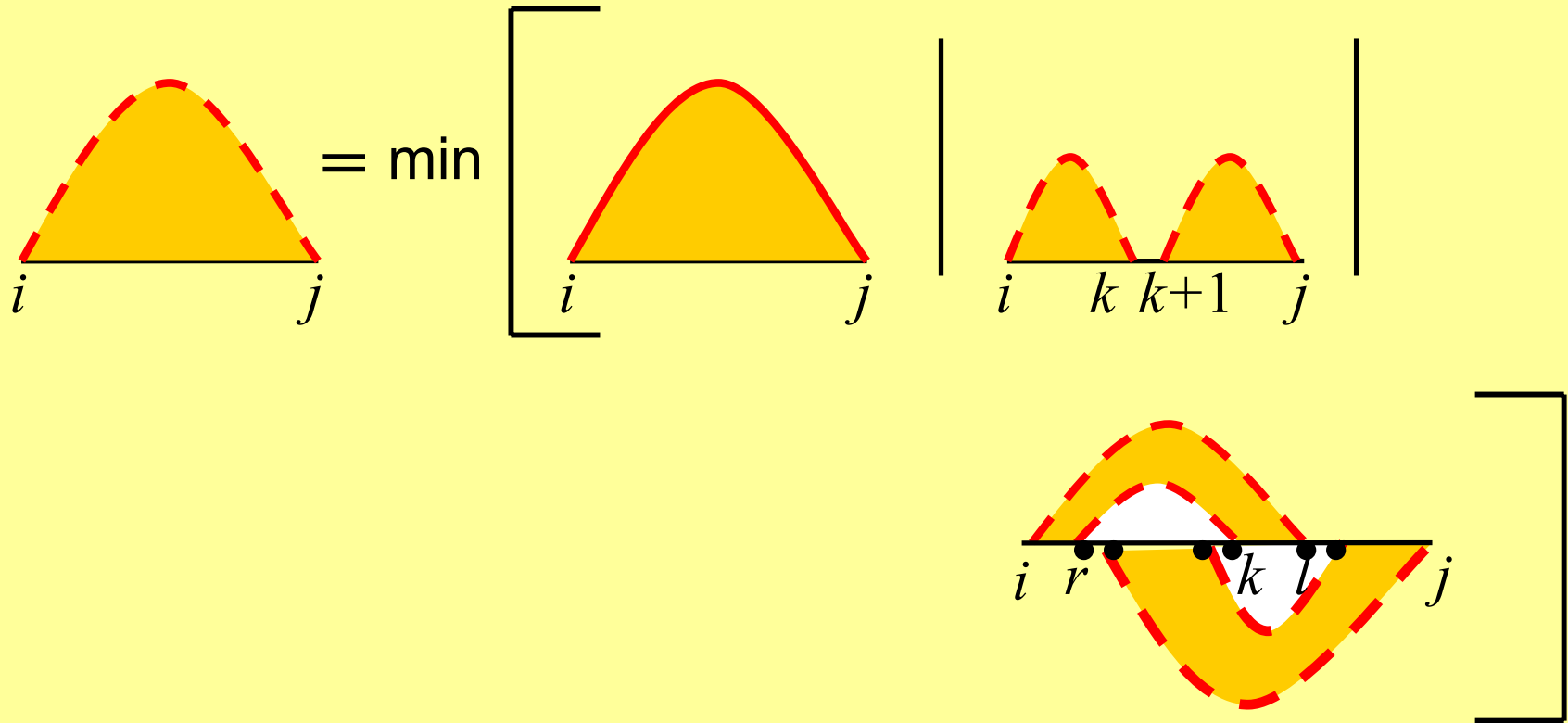
recurrences



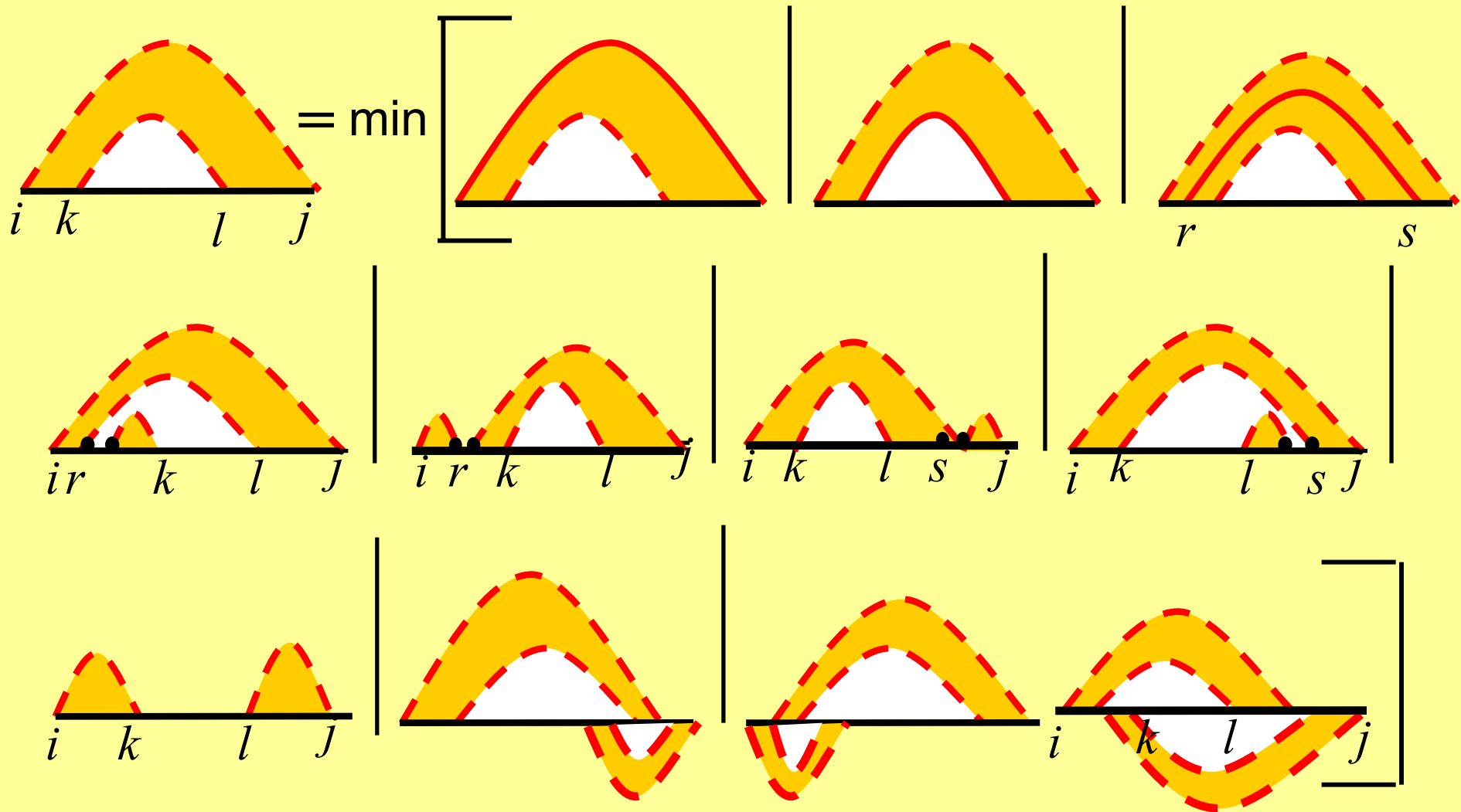
current approaches: pseudoknotted structures

- loop energies augmented to fit known data
- mfe algorithms have been extended to handle certain pseudoknotted structures (Akutsu, 2000; Rivas and Eddy, 1999)
- heuristic approaches also used (STAR software, van Batenburg et al.)

$W(i,j)$: Rivas and Eddy algorithm



recurrence for gapped structures



Rivas and Eddy algorithm

- running time is $O(n^6)$
- “*we lack a systematic a priori characterization of the class of configurations that this algorithm can solve*” (Rivas and Eddy, 1999)

heuristic approaches

- space of structures is explored, guided by random decisions as well as goal of minimizing free energy
- can incorporate better energy models, model folding pathways, and be substantially faster than mfe algorithms
- no guarantee that optimal solution is found

STAR algorithm (van Batenburg et al.)

- simulates RNA folding by stepwise addition and removal of stems to the structure formed at previous steps
- choice of stem to add/remove is random, biased by “fitness” of resulting structure
- process is carried out on large population of structures, not just one

prediction challenges

- predict structures formed from multiple strands
- predict which strand in a combinatorial set has minimum free energy structure
- predict pseudoknotted structures

BETA-Lab Projects

- predict structures formed from multiple strands

➡ PairFold

- predict which strand in a combinatorial set has minimum free energy structure

➡ CombFold

- predict pseudoknotted structures

➡ PseudoSpotter

PairFold (Andronescu)

- predicts minimum energy pseudoknot free secondary structure of a *pair* of strands
- energy model: loop plus initiation energies from Turner, Santa Lucia labs

CombFold (Andronescu et al.)

- predicts which strand in a combinatorial set has minimum free energy structure

TTAC TTAC TTAC TTAC TTAC TTAC TTAC TTAC
AATC AATC AATC AATC AATC AATC AATC AATC
TACT TACT TACT TACT TACT TACT TACT TACT
ATCA ATCA ATCA ATCA ATCA ATCA ATCA ATCA
ACAT ACAT ACAT ACAT ACAT ACAT ACAT ACAT
TCTA TCTA TCTA TCTA TCTA TCTA TCTA TCTA
CTTT CTTT CTTT CTTT CTTT CTTT CTTT CTTT
CAAA CAAA CAAA CAAA CAAA CAAA CAAA CAAA

- uses dynamic programming to avoid examining all combinatorial possibilities

CombFold (Andronescu et al.)

- predicts which strand in a combinatorial set has minimum free energy structure

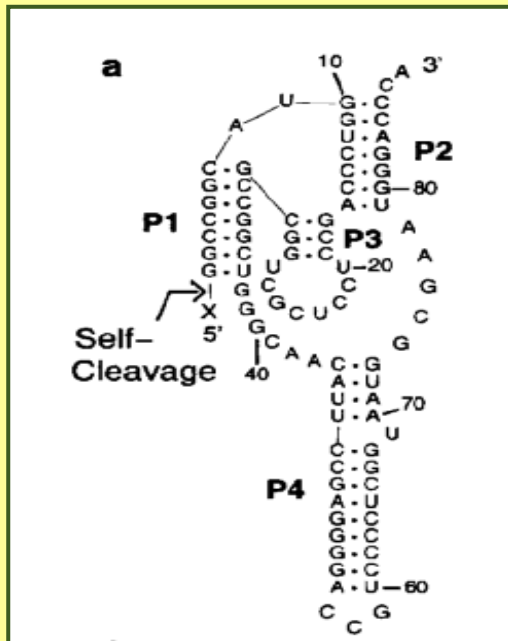
TTAC	TTAC	TTAC	TTAC	TTAC	TTAC	TTAC	TTAC
AATC	AATC	AATC	AATC	AATC	AATC	AATC	AATC
TACT	TACT	TACT	TACT	TACT	TACT	TACT	TACT
ATCA	ATCA	ATCA	ATCA	ATCA	ATCA	ATCA	ATCA
ACAT	ACAT	ACAT	ACAT	ACAT	ACAT	ACAT	ACAT
TCTA	TCTA	TCTA	TCTA	TCTA	TCTA	TCTA	TCTA
CTTT	CTTT	CTTT	CTTT	CTTT	CTTT	CTTT	CTTT
CAAA	CAAA	CAAA	CAAA	CAAA	CAAA	CAAA	CAAA

- TCTAATCATACTAATCCTTTTACTATCATTAC has a slightly stable secondary structure at 37°C

PseudoSpotter (Ren et al.)

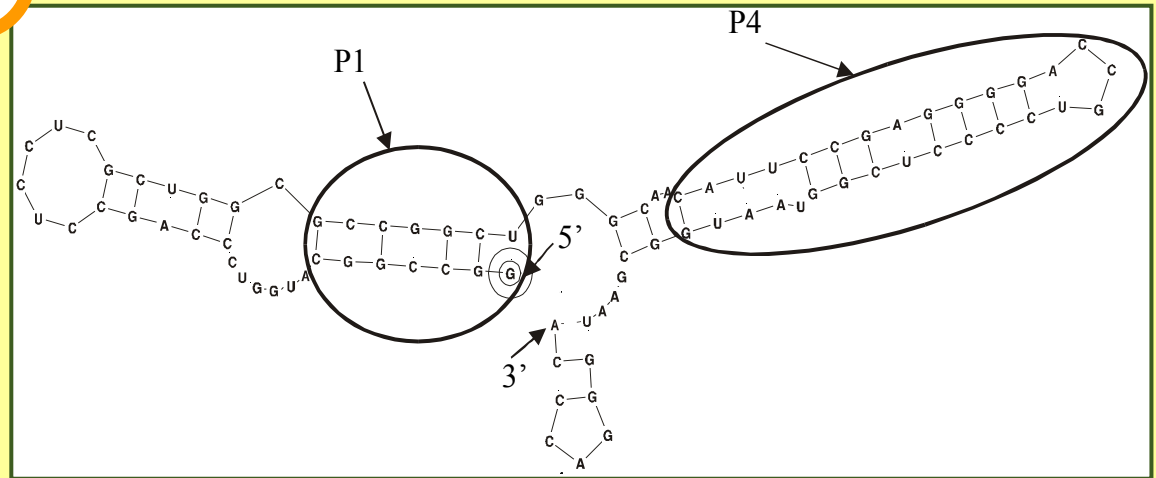
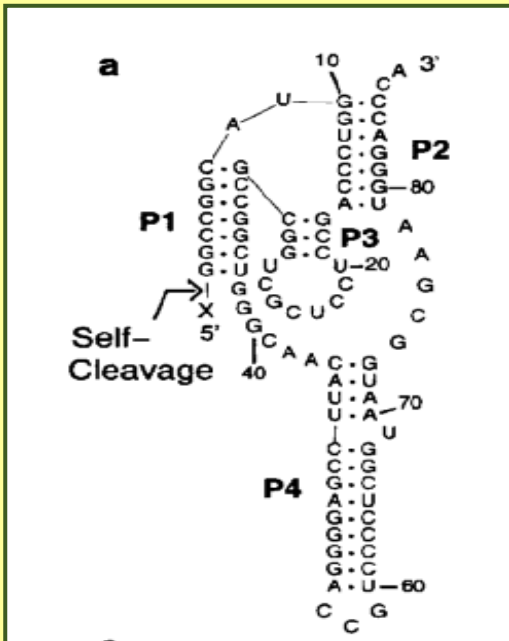
- heuristic algorithm for predicting pseudoknotted structures

PseudoSpotter (Ren et al.)



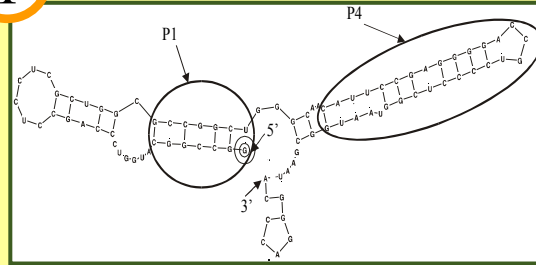
PseudoSpotter (Ren et al.)

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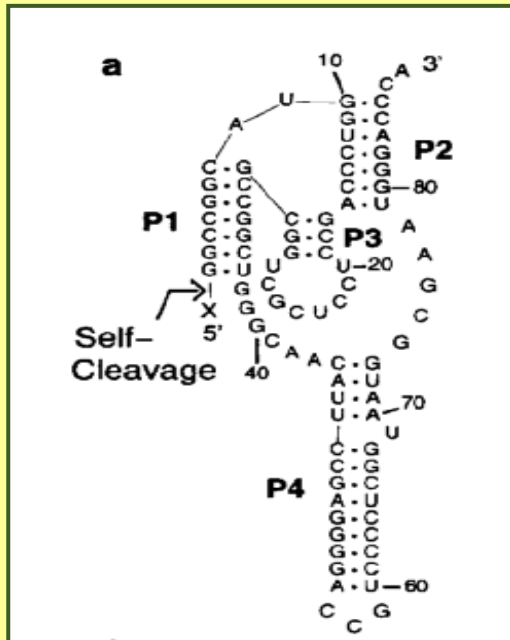
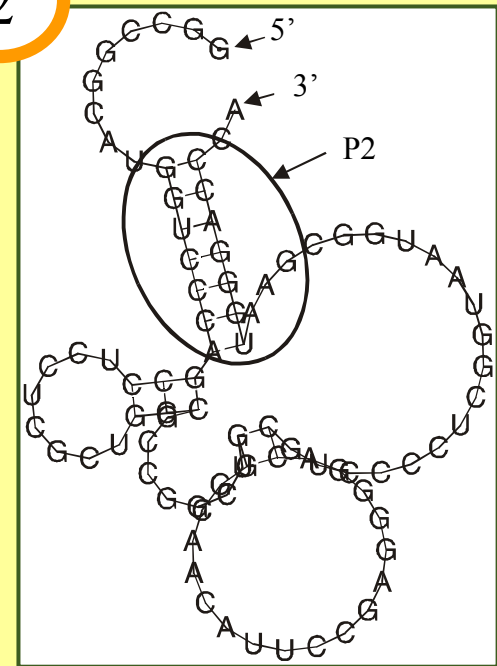


PseudoSpotter (Ren et al.)

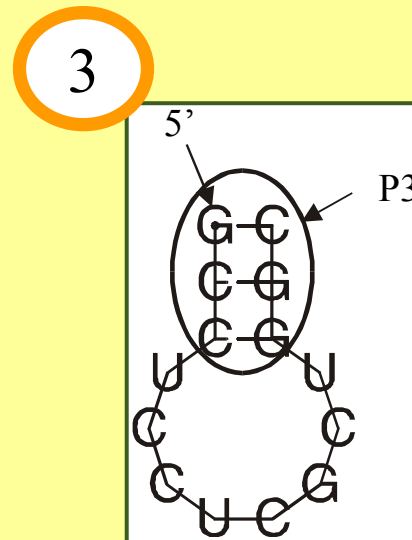
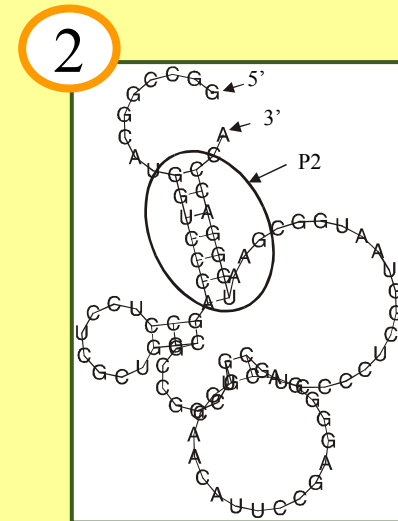
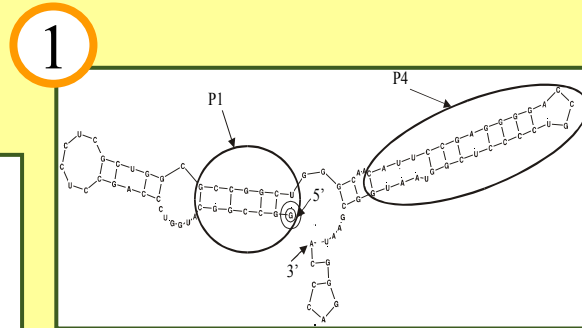
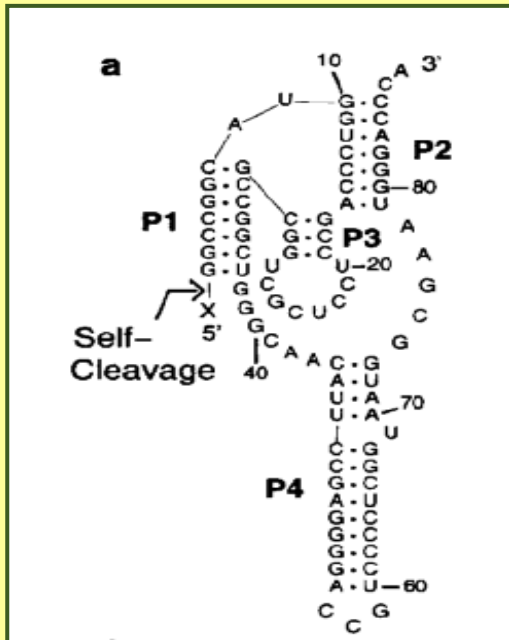
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PseudoSpotter (Ren et al.)



PseudoSpotter algorithm

- initial structure pool contains the empty structure
- repeat
 - for each structure in the pool, identify promising stems (hotspots) that can be added
 - augment each structure with each of its hotspots, thereby expanding the pooluntil all structures are saturated
- several structures are output

comparison of pseudoknot algorithms

	Optimal	Subopt, Close	Fail
R&E	18	5	9
STAR	19		13
Pseudo-spotter	21	7	4

- Quality of solutions found over 32 sample strands taken from RDB, Pseudobase

RNASoft - Software for RNA/DNA secondary structure prediction and design

[[Conditions of use](#) | [About RNASoft](#)]

Quick Launch [[PairFold](#) | [CombFold](#) | [RNA Designer](#)]

PairFold

[[Run](#)] [[About](#)]

PairFold predicts the minimum free secondary structure formed by two input molecules. **PairFold** can be used, for example, to predict interactions between a probe and target RNA molecule, or between pairs of strands in molecular-based nanostructures.

CombFold

[[Run](#)] [[About](#)]

CombFold predicts which strand, out of a combinatorial set formed from input strands, folds to a secondary structure with the lowest free energy. **CombFold** can, for example, efficiently test that no strand in a large tag library forms unwanted secondary structure.

RNA Designer

[[Run](#)] [[About](#)]

RNA Designer designs an RNA sequence that folds to a given input secondary structure. The tool is intended for designers of RNA molecules with particular structural or functional properties.

next steps

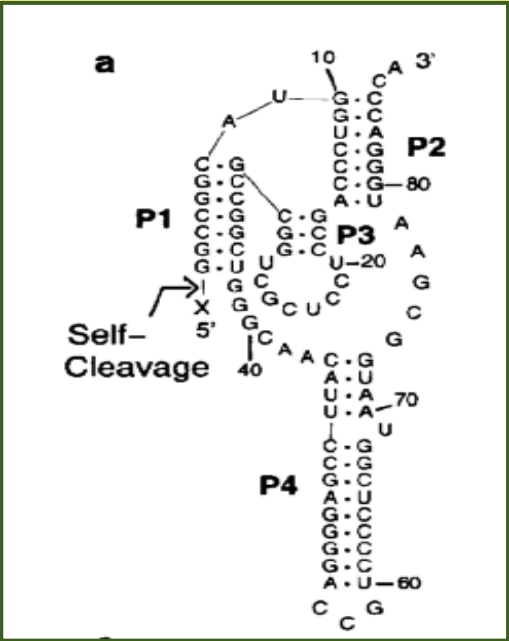
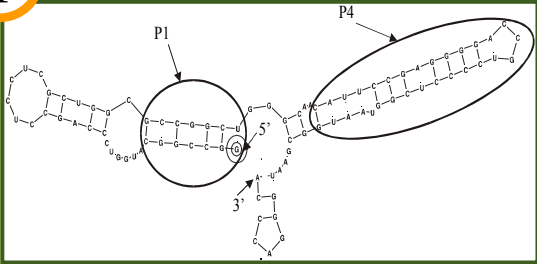
- better energy parameters for model, model evaluation and refinement
- develop better heuristic algorithms for predicting pseudoknotted structures
- improved mfe algorithms for natural pseudoknotted structures
- suboptimal foldings, partition function

how to get better energy models?

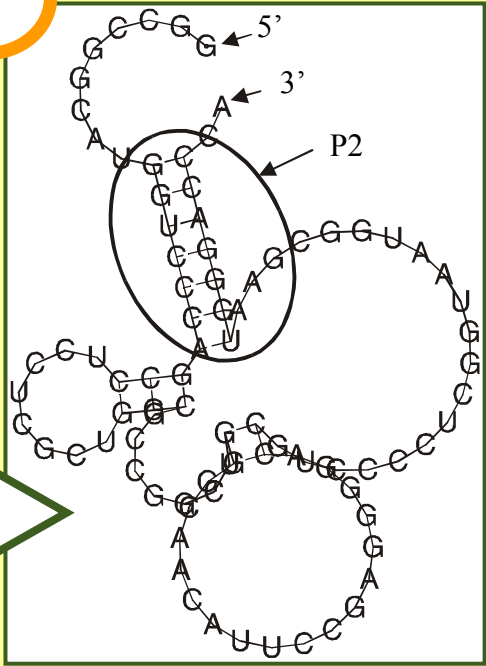
- compare variations of standard free energy model in terms of prediction quality
- perform lab experiments to find energy parameters
- incorporate tertiary structural elements (nonstandard base pairs or motifs, folding pathway, metal ions)

how to improve heuristic algorithm?

1



2



maybe here?

how to improve mfe algorithm?

- we have a simple characterization of the class of structures that the Rivas and Eddy algorithm can handle
- perhaps an $O(n^5)$ algorithm for natural structures?

acknowledgements

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Pseudospotter

Sohrab Shah

structural motif finding

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DNA word design

Shelly Zhao

Combfold, RNA structure analysis

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www.cs.ubc.ca/labs/beta