

A Simple Protocol for the Inference of RNA Global Pairwise Alignments

Felix Karg

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University of Freiburg



Content

needle

Sequence Alignment

Needleman-Wunsch

LocARNA

Tree-based sequence
alignment

gardenia

RNA StrAT

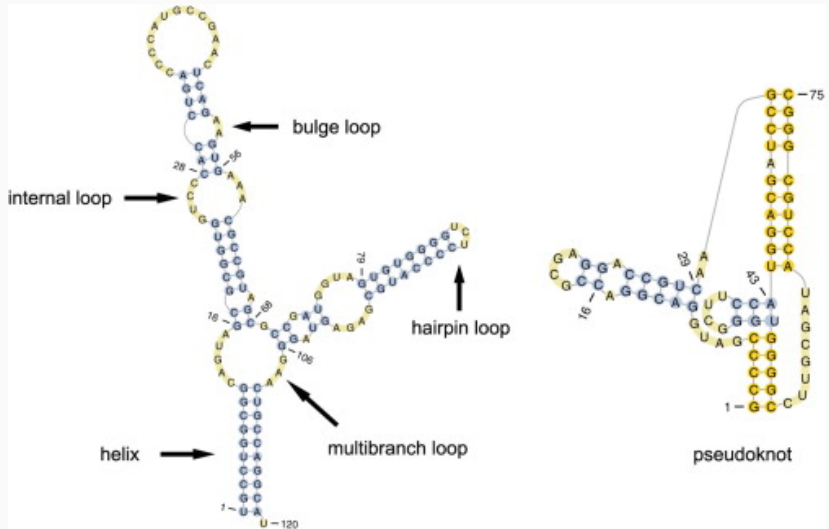
RNAdistance

RNAforester

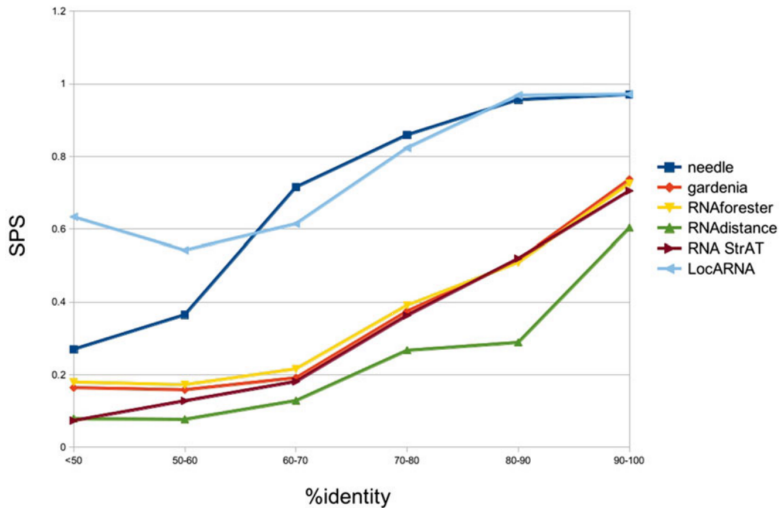
Comparison

Sources

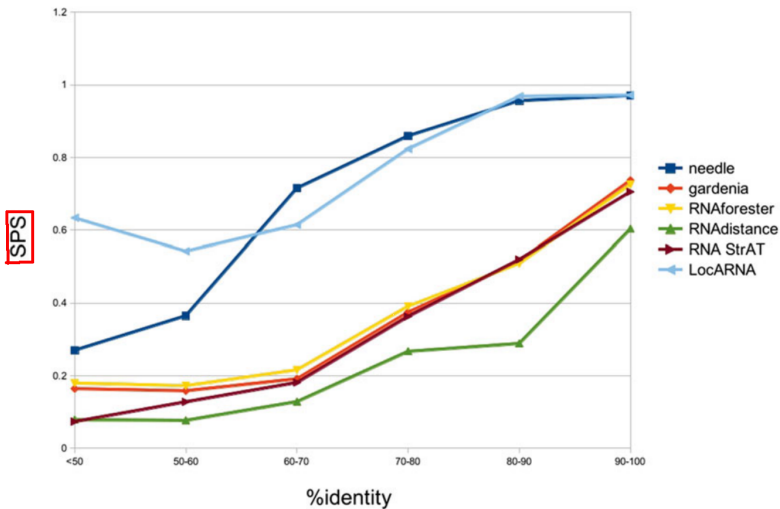
RNA secondary structure



Predicted Secondary Structures



Predicted Secondary Structures



Sum of Pairs Score

SPS - introduction

Sum of Pairs Score

Used to measure the alignment of two RNA sequences

SPS - introduction

Sum of Pairs Score

Used to measure the similarity of two RNA sequences

Sequence Similarity - Example

A: AAGGCTT

B: AAGGC

C: AAGGCAT

Similarity:

$1 - (\text{edit distance} / \text{unaligned length of shorter sequence})$

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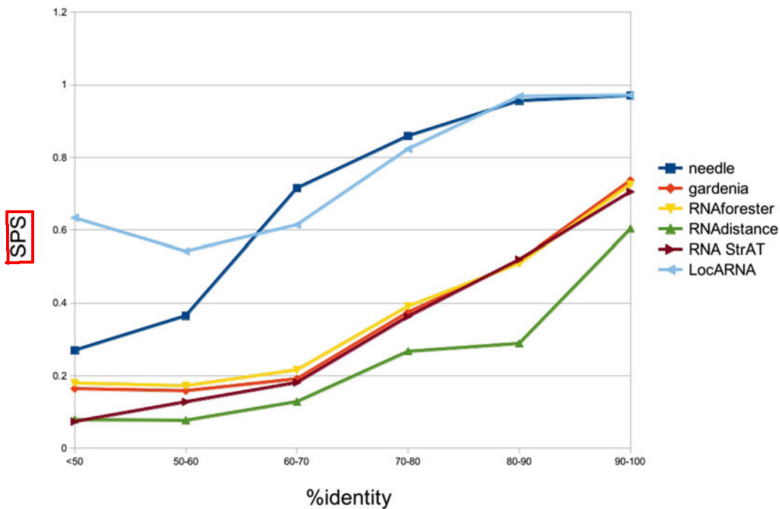
B: AAGGC

C: AAGGCAT

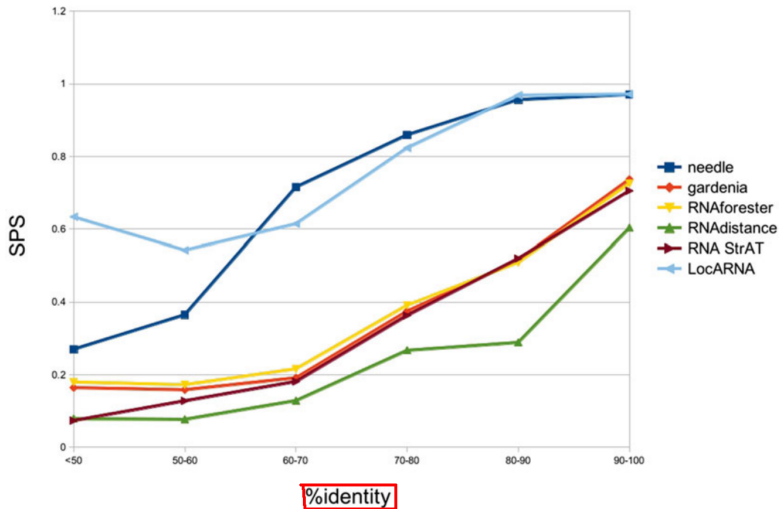
Similarity: $86\% = 1 - (1 / 7)$

$1 - (\text{edit distance} / \text{unaligned length of shorter sequence})$

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Sequence Identity - Example

A: AAGGCTT

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Identity:

Identical nucleotides / shorter sequence length

Sequence Identity - Example

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Sequence Identity - Example

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Identity: 100%

Identical nucleotides / shorter sequence length

Sequence Identity - Example

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Identity:

Identical nucleotides / shorter sequence length

Sequence Identity - Example

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Sequence Identity - Example

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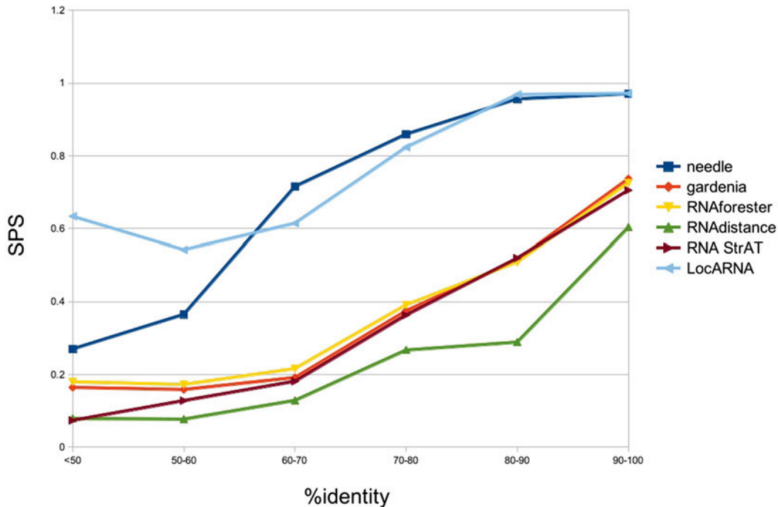
C: AAGGCAT

Identity: $85\% = 6 / 7$

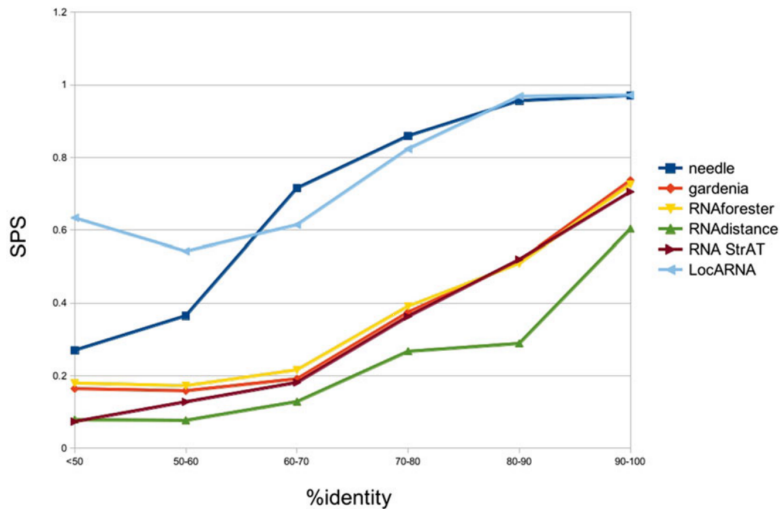
Identical nucleotides / shorter sequence length

Explain: predicted (RNAfold), test setup

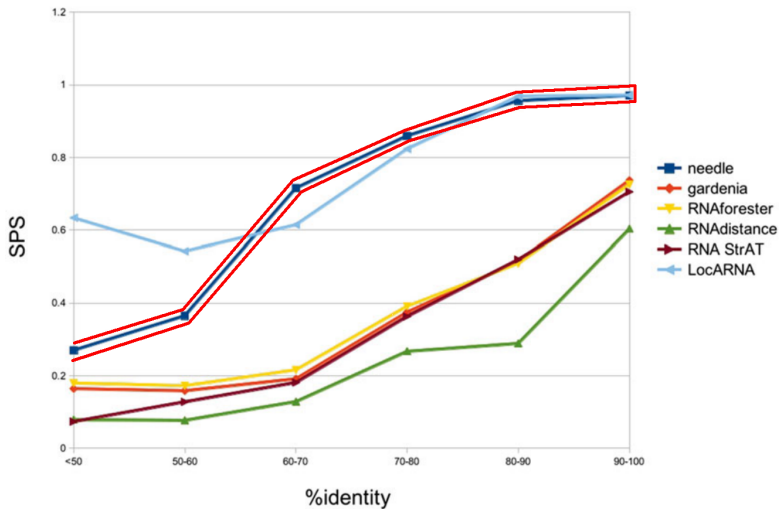
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Needleman-Wunsch

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- Sequence based Algorithm

- Sequence based Algorithm
- part of the EMBOSS package

- Sequence based Algorithm
- part of the EMBOSS package
- Implementation of the Needleman-Wunsch algorithm

Needleman-Wunsch Algorithm

Sometimes also called:

Needleman-Wunsch Algorithm

Sometimes also called:

- Optimal matching algorithm

Needleman-Wunsch Algorithm

Sometimes also called:

- Optimal matching algorithm
- Global alignment technique

needle

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Sequence Alignment

- local sequence alignment

Sequence Alignment

- local sequence alignment
- global sequence alignment

Sequence Alignment

- local sequence alignment
- global sequence alignment
- glocal sequence alignment

Sequence Alignment - local

RNA sequence A



RNA sequence B

Sequence Alignment - global

RNA sequence A



RNA sequence B

Sequence Alignment - glocal

RNA sequence A



RNA sequence B

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Needleman-Wunsch Algorithm - introduction

We need a scoring system.

Needleman-Wunsch Algorithm - introduction

We need a scoring system.

Example:

Needleman-Wunsch Algorithm - introduction

We need a scoring system.

Example:

- Match: 1
- Mismatch: -1
- Indel: -1

Needleman-Wunsch Algorithm

Two sequences to compare:

Needleman-Wunsch Algorithm

Two sequences to compare:

GCATGCU

GATTACA

Needleman-Wunsch Algorithm - Example

		G	C	A	T	G	C	U
G								
A								
T								
T								
A								
C								
A								

Needleman-Wunsch Algorithm - Example

		G	C	A	T	G	C	U
	0							
G								
A								
T								
T								
A								
C								
A								

Needleman-Wunsch Algorithm - Example

		G	C	A	T	G	C	U
	0	-1	-2	-3	-4	-5	-6	-7
G	-1							
A	-2							
T	-3							
T	-4							
A	-5							
C	-6							
A	-7							

Needleman-Wunsch Algorithm - Example

		G	C	A	T	G	C	U
	0	-1	-2	-3	-4	-5	-6	-7
G	-1	X						
A	-2							
T	-3							
T	-4							
A	-5							
C	-6							
A	-7							

Needleman-Wunsch Algorithm - Example

		G	C	A	T	G	C	U
	0	-1	-2	-3	-4	-5	-6	-7
G	-1	1						
A	-2							
T	-3							
T	-4							
A	-5							
C	-6							
A	-7							

Needleman-Wunsch Algorithm - Example

		G	C	A	T	G	C	U
	0	-1	-2	-3	-4	-5	-6	-7
G	-1	1	X					
A	-2	Y						
T	-3							
T	-4							
A	-5							
C	-6							
A	-7							

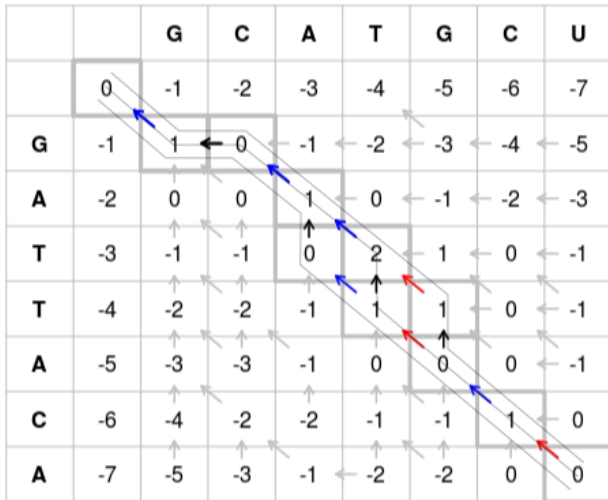
Needleman-Wunsch Algorithm - Example

		G	C	A	T	G	C	U
	0	-1	-2	-3	-4	-5	-6	-7
G	-1	1	0					
A	-2	0						
T	-3							
T	-4							
A	-5							
C	-6							
A	-7							

Needleman-Wunsch Algorithm - Example

		G	C	A	T	G	C	U
	0	-1	-2	-3	-4	-5	-6	-7
G	-1	1	0	-1	-2	-3	-4	-5
A	-2	0	0	1	0	-1	-2	-3
T	-3	-1	-1	0	2	1	0	-1
T	-4	-2	-2	-1	1	1	0	-1
A	-5	-3	-3	-1	0	0	0	-1
C	-6	-4	-2	-2	-1	-1	1	0
A	-7	-5	-3	-1	-2	-2	0	0

Needleman-Wunsch Algorithm - Best Matches



Needleman-Wunsch Algorithm - Results

Sequences	Best alignments		
GCATGCU	GCATG-CU	GCA-TGCU	GCAT-GCU
GATTACA	G-ATTACA	G-ATTACA	G-ATTACA

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Two steps:

Two steps:

- create base pair probability matrix using RNAfold

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- using these as guide for optimal alignment

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- create base pair probability matrix using RNAfold
- using these as guide for optimal alignment

(folding and aligning)

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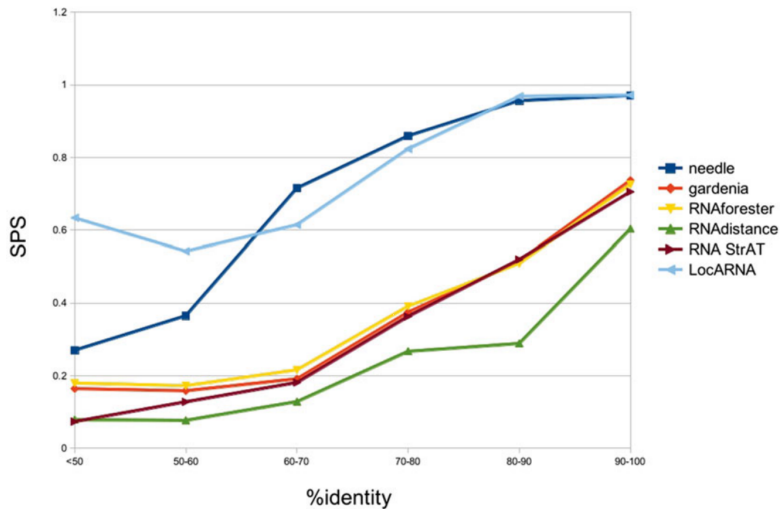
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Run time Comparison

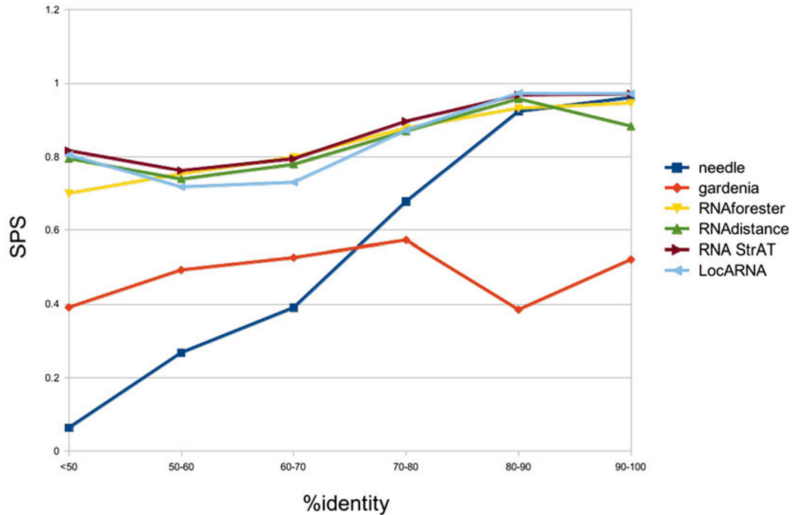
Mean running time (in seconds) per sequence

	bralibase	RNAspa	RNAstrand
needle	0.01	0.01	0.01
gardenia	0.01	0.01	0.02
RNA StrAT	0.02	0.32	0.46
LocARNA	0.02	0.08	0.02
RNAdistance	0.01	0.01	0.01
RNAforester	0.03	0.73	0.81

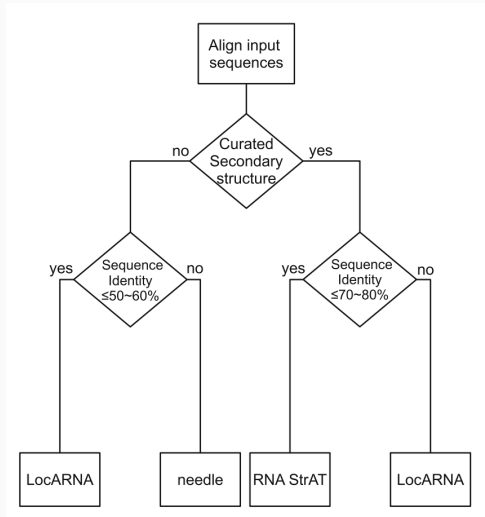
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Curated Secondary Structures



Proposed Workflow



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The slides can be found at:



Github

`https://github.com/fkarg/things-to-talk-about/
tree/master/proseminar`



Image

https://upload.wikimedia.org/wikipedia/commons/3/3f/Needleman-Wunsch_pairwise_sequence_alignment.png



Secondary structures Image

<https://www.sciencedirect.com/science/article/pii/B9780124200371000014>