## Sequence-structure relationships in RNA-loops:

## establishing the basis for loop homology modeling

## Supplementary Material

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## RLML – The RLooM Modeling Language

Modeling loops using the RLooM application is performed using a simple XML-like script language – RLML. Three parameters can be adjusted: the template data set that should be used, the maximum distance between the anchors of a loop and a target structure such that the inserted loop gives a valid model, and the threshold distance defining when a clash occurs between the new loop and the target molecule.

A single command is enclosed between tags specifying the loop-type of the query.

$$\langle x \rangle ... \langle /x \rangle$$
, with x = hairpin|segment|internal|multiloop

Each command has a number of anchors (hairpins/segments:2, internal loops:4, multiloop:6+):

The anchor-tag has an optional parameter id, which can be used for specifying the sequence of the anchors. By default, <anchor>-tags are processed in order of appearance.

Finally, each command requires a query:

with SEQUENCE being a nucleotide sequence, wildcards are allowed

The <query>-tag has three optional parameters: k, force, and mcsearch. The parameter k

specifies the tolerated number of mismatches, force denotes whether suitable candidate loops with a different sequence than the query shall be artificially mutated to match the query sequence. The parameter mcsearch, if set to true, allows a valid MC-Search script (for details see e.g. http://major.iric.ca) to be submitted instead of the query sequence. By default, k is set to 0, force to false, and mcsearch to true.

The optional <remodel>- tag specifies, whether loop candidates should be mutated into its enclosed sequence.

where SEQUENCE has to be a non-wildcard nucleotide sequence.

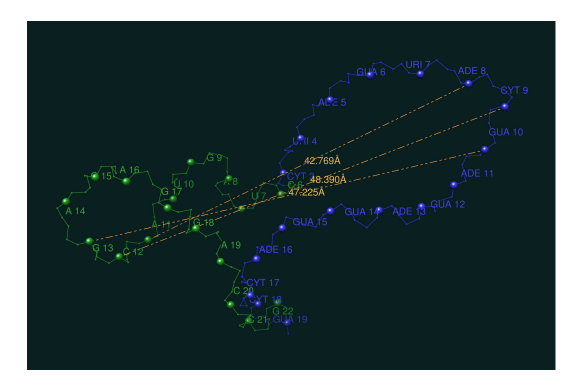


Figure 1. Backbone divergence between iFoldRNA model and native structure

blue: iFoldRNA model green: PDB structure 1Q9A, A6-A22. Structures are superposed at their anchors.

	1EVV:A	1evv:b	13vv:c	1L2x	1Q8N	1Q $9$ A	1 <sub>RMN</sub>	2cky	2F88
Bases	13-22	30-40	53-61	7-14	14-19	6-22	16-21	26-36	23-27
Best Hits									
RLooM	1EHZ*	1мј1	2к4c*	1L3D*	1м90*	2D3O*	3BBN*	3D2x*	1JZX*
$\mathrm{RMSD}_{\mathbf{a}}$	0.14	0.00	0.15	0.90	0.48	0.16	0.20	0.25	0.44
$\mathrm{RMSD}_{\mathrm{b}}$	1.04	0.78	1.11	1.72	4.12	2.76	3.12	0.53	1.58
$\mathrm{RMSD_s}$	0.98	0.69	0.90	0.75	2.05	1.05	1.71	0.38	1.34
iFoldRNA									
$\mathrm{RMSD}_{\mathbf{a}}$	1.08	0.30	1.21	0.50	0.48	2.58	0.60	1.39	1.01
$\mathrm{RMSD}_{\mathrm{b}}$	10.88	3.90	4.72	4.40	1.26	23.19	2.51	9.21	1.45
$\mathrm{RMSD}_{\mathrm{s}}$	6.19	6.42	6.90	5.43	5.31	8.75	5.68	6.31	1.18
All Hits									
RLooM									
#models	14	11	16	3	25	23	58	2	13
$\mathrm{RMSD}_{\mathbf{a}}$									
Range	[0.00,0.96]	[0.00,0.52]	[0.00,0.95]	[0.11,0.99]	[0.00,1.02]	[0.03,1.13]	[0.00,1.23]	[0.22,0.25]	[0.00,2.63]
Mean	$0.48 \pm 0.27$	$0.29 \pm 0.17$	$0.48 \pm 0.25$	$0.67 \pm 0.48$	$0.70 \pm 0.20$	$0.44 \pm 0.28$	$0.52 \pm 0.23$	$0.24 \pm 0.02$	$1.16 \pm 0.80$
Median	0.52	0.31	0.50	0.90	0.70	0.33	0.47	0.24	0.97
$\mathrm{RMSD}_{\mathrm{b}}$									
Range	[0.00,2.98]	[0.78, 1.74]	[0.00,3.81]	[0.25,2.19]	[0.00,2.31]	[0.15,8.66]	[0.00,1.26]	[0.41,0.53]	[0.00,3.51]
Mean	$2.73 \pm 2.20$	$1.10 \pm 0.45$	$1.93 \pm 1.27$	$1.39 \pm 1.01$	$1.69 \pm 0.86$	$4.35 \pm 2.09$	$1.18 \pm 0.35$	$0.47 \pm 0.08$	$2.30 \pm 0.97$
Median	2.26	1.15	1.70	1.72	1.52	3.99	1.10	0.47	2.19
$\mathrm{RMSD}_{\mathrm{s}}$									
Range	[0.00,2.26]	[0.69,1.1]	[0.00,3.15]	[0.10, 0.79]	[0.00,1.67]	[0.14,2.05]	[0.00,1.44]	[0.38,0.49]	[0.00,2.89]
Mean	$1.43 \pm 0.82$	$0.73 \pm 0.28$	$1.29 \pm 0.91$	$0.55 \pm 0.39$	$1.24 \pm 0.42$	$1.45 \pm 0.45$	$1.74 \pm 0.73$	$0.44 \pm 0.08$	$1.84 \pm 0.75$
Median	1.43	0.72	0.93	0.75	1.20	1.40	1.54	0.44	1.76
iFoldRNA									
#models	10	10	10	10	10	10	10	10	10
$\mathrm{RMSD}_{\mathbf{a}}$									
Range	[1.08,4.12]	[0.30,3.04]	[1.21,3.51]	[0.50,1.22]	[0.48,4.78]	[2.58, 3.99]	[0.60,3.42]	[1.39,7.28]	[1.01,5.92]
Mean	$2.42 \pm 0.92$	$0.85 \pm 0.89$	$2.19 \pm 0.79$	$0.76 \pm 0.24$	$1.36 \pm 1.36$	$3.18 \pm 0.52$	$1.82 \pm 0.89$	$5.04 \pm 1.82$	$2.91 \pm 1.46$
Median	2.48	0.48	2.15	0.76	0.76	3.01	1.97	5.24	2.93
$\mathrm{RMSD}_{\mathrm{b}}$									
Range	[9.60,21.67]	[2.44,16.45]	[4.56,17.46]	[3.18,7.55]	[1.26,10.98]	[19.05,28.28]	[2.34,9.69]	[9.21,18.62]	[1.45,5.75]
Mean	$14.68 \pm 4.24$	$8.18 \pm 5.40$	$10.02 \pm 5.08$	$4.94 \pm 1.34$	$3.48 \pm 2.83$	$24.11 \pm 2.86$	$4.10 \pm 2.23$	$13.57 \pm 2.91$	$3.41 \pm 1.39$
Median	14.51	6.82	9.39	4.65	2.45	23.95	3.08	13.37	3.27
$\mathrm{RMSD}_{\mathrm{s}}$									
Range	[4.77,6.83]	[6.37,9.77]	[5.53,7.83]	[4.94,5.89]	[5.31,6.25]	[7.61,9.50]	[5.04,8.46]	[6.31,8.86]	[1.18,4.35]
Mean	$6.10 \pm 0.65$	$7.23 \pm 1.26$	$6.61 \pm 0.63$	$5.30 \pm 0.29$	$5.72 \pm 0.33$	$8.55 \pm 0.53$	$6.09 \pm 1.03$	$7.55 \pm 0.84$	$2.48 \pm 0.91$
Median	6.26	6.59	6.50	5.33	5.68	8.62	5.80	7.42	2.34
Table 1									

Table 1

Loop modeling results Modeled structures are indicated by their PDB-identifier and are located in chain A.  $RMSD_a$ : RMSD between anchors,  $RMSD_b$ : RMSD between reduced backbones given anchor superposition,  $RMSD_s$ : structural similarity – RMSD between reduced backbones given optimal superposition, values are given in Å, \*: second best template (cf. text)