## Kevin M Weeks

## List of Publications by Citations

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153	10,795	53	102
papers	citations	h-index	g-index
161	12,735 ext. citations	11.8	6.69
ext. papers		avg, IF	L-index

#	Paper	IF	Citations
153	Architecture and secondary structure of an entire HIV-1 RNA genome. <i>Nature</i> , <b>2009</b> , 460, 711-6	50.4	607
152	RNA structure analysis at single nucleotide resolution by selective 2Shydroxyl acylation and primer extension (SHAPE). <i>Journal of the American Chemical Society</i> , <b>2005</b> , 127, 4223-31	16.4	573
151	Selective 2Shydroxyl acylation analyzed by primer extension (SHAPE): quantitative RNA structure analysis at single nucleotide resolution. <i>Nature Protocols</i> , <b>2006</b> , 1, 1610-6	18.8	563
150	Accurate SHAPE-directed RNA structure determination. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2009</b> , 106, 97-102	11.5	493
149	RNA motif discovery by SHAPE and mutational profiling (SHAPE-MaP). <i>Nature Methods</i> , <b>2014</b> , 11, 959-65	521.6	333
148	RNA recognition by Tat-derived peptides: interaction in the major groove?. <i>Cell</i> , <b>1991</b> , 66, 577-88	56.2	332
147	High-throughput SHAPE analysis reveals structures in HIV-1 genomic RNA strongly conserved across distinct biological states. <i>PLoS Biology</i> , <b>2008</b> , 6, e96	9.7	313
146	mRNA structure determines specificity of a polyQ-driven phase separation. <i>Science</i> , <b>2018</b> , 360, 922-927	33.3	284
145	A fast-acting reagent for accurate analysis of RNA secondary and tertiary structure by SHAPE chemistry. <i>Journal of the American Chemical Society</i> , <b>2007</b> , 129, 4144-5	16.4	270
144	Principles for targeting RNA with drug-like small molecules. <i>Nature Reviews Drug Discovery</i> , <b>2018</b> , 17, 547-558	64.1	246
143	Accurate SHAPE-directed RNA secondary structure modeling, including pseudoknots. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2013</b> , 110, 5498-503	11.5	210
142	Advances in RNA structure analysis by chemical probing. <i>Current Opinion in Structural Biology</i> , <b>2010</b> , 20, 295-304	8.1	210
141	SHAPE-directed RNA secondary structure prediction. <i>Methods</i> , <b>2010</b> , 52, 150-8	4.6	201
140	Major groove accessibility of RNA. <i>Science</i> , <b>1993</b> , 261, 1574-7	33.3	193
139	Catalysts from synthetic genetic polymers. <i>Nature</i> , <b>2015</b> , 518, 427-30	50.4	187
138	ShapeFinder: a software system for high-throughput quantitative analysis of nucleic acid reactivity information resolved by capillary electrophoresis. <i>Rna</i> , <b>2008</b> , 14, 1979-90	5.8	185
137	Selective 2Shydroxyl acylation analyzed by primer extension and mutational profiling (SHAPE-MaP) for direct, versatile and accurate RNA structure analysis. <i>Nature Protocols</i> , <b>2015</b> , 10, 1643-69	18.8	182

136	RNA-Puzzles: a CASP-like evaluation of RNA three-dimensional structure prediction. Rna, 2012, 18, 610-	<b>25</b> 8	181
135	SHAPE reveals transcript-wide interactions, complex structural domains, and protein interactions across the Xist lncRNA in living cells. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2016</b> , 113, 10322-7	11.5	149
134	QuShape: rapid, accurate, and best-practices quantification of nucleic acid probing information, resolved by capillary electrophoresis. <i>Rna</i> , <b>2013</b> , 19, 63-73	5.8	147
133	Pervasive Regulatory Functions of mRNA Structure Revealed by High-Resolution SHAPE Probing. <i>Cell</i> , <b>2018</b> , 173, 181-195.e18	56.2	139
132	RNA SHAPE chemistry reveals nonhierarchical interactions dominate equilibrium structural transitions in tRNA(Asp) transcripts. <i>Journal of the American Chemical Society</i> , <b>2005</b> , 127, 4659-67	16.4	133
131	RNA-Puzzles Round III: 3D RNA structure prediction of five riboswitches and one ribozyme. <i>Rna</i> , <b>2017</b> , 23, 655-672	5.8	118
130	The mechanisms of RNA SHAPE chemistry. Journal of the American Chemical Society, 2012, 134, 6617-24	16.4	116
129	Exploring RNA structural codes with SHAPE chemistry. <i>Accounts of Chemical Research</i> , <b>2011</b> , 44, 1280-9	124.3	112
128	Protein-facilitated RNA folding. Current Opinion in Structural Biology, 1997, 7, 336-42	8.1	110
127	Detection of RNA-Protein Interactions in Living Cells with SHAPE. <i>Biochemistry</i> , <b>2015</b> , 54, 6867-75	3.2	109
126	Protein facilitation of group I intron splicing by assembly of the catalytic core and the 5Ssplice site domain. <i>Cell</i> , <b>1995</b> , 82, 221-30	56.2	108
125	Genomic RNA Elements Drive Phase Separation of the SARS-CoV-2 Nucleocapsid. <i>Molecular Cell</i> , <b>2020</b> , 80, 1078-1091.e6	17.6	98
124	Single-molecule correlated chemical probing of RNA. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2014</b> , 111, 13858-63	11.5	95
123	A collapsed non-native RNA folding state. <i>Nature Structural Biology</i> , <b>2000</b> , 7, 362-6		92
122	van <b>\$</b> Hoff enthalpies without baselines. <i>Protein Science</i> , <b>2000</b> , 9, 1416-9	6.3	91
121	Functionally conserved architecture of hepatitis C virus RNA genomes. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2015</b> , 112, 3692-7	11.5	89
120	Time-resolved RNA SHAPE chemistry: quantitative RNA structure analysis in one-second snapshots and at single-nucleotide resolution. <i>Nature Protocols</i> , <b>2009</b> , 4, 1413-21	18.8	87
119	The cellular environment stabilizes adenine riboswitch RNA structure. <i>Biochemistry</i> , <b>2013</b> , 52, 8777-85	3.2	86

118	Functional classification of long non-coding RNAs by k-mer content. <i>Nature Genetics</i> , <b>2018</b> , 50, 1474-148 <b>3</b> 6.3	86
117	On the significance of an RNA tertiary structure prediction. <i>Rna</i> , <b>2010</b> , 16, 1340-9 5.8	82
116	Efficient protein-facilitated splicing of the yeast mitochondrial bI5 intron. <i>Biochemistry</i> , <b>1995</b> , 34, 7728-38,2	76
115	Strong correlation between SHAPE chemistry and the generalized NMR order parameter (S2) in RNA. <i>Journal of the American Chemical Society</i> , <b>2008</b> , 130, 12244-5	75
114	Accurate detection of chemical modifications in RNA by mutational profiling (MaP) with ShapeMapper 2. <i>Rna</i> , <b>2018</b> , 24, 143-148	74
113	Influence of nucleotide identity on ribose 2Shydroxyl reactivity in RNA. <i>Rna</i> , <b>2009</b> , 15, 1314-21 5.8	70
112	RNA secondary structure modeling at consistent high accuracy using differential SHAPE. <i>Rna</i> , <b>2014</b> , 20, 846-54	67
111	High-throughput SHAPE and hydroxyl radical analysis of RNA structure and ribonucleoprotein assembly. <i>Methods in Enzymology</i> , <b>2009</b> , 468, 67-89	67
110	Time-resolved RNA SHAPE chemistry. <i>Journal of the American Chemical Society</i> , <b>2008</b> , 130, 16178-80 16.4	66
109	Comparison of SIV and HIV-1 genomic RNA structures reveals impact of sequence evolution on conserved and non-conserved structural motifs. <i>PLoS Pathogens</i> , <b>2013</b> , 9, e1003294	65
108	Three-dimensional RNA structure refinement by hydroxyl radical probing. <i>Nature Methods</i> , <b>2012</b> , 9, 603- <b>&amp;</b> 1.6	64
107	Facile conversion of aptamers into sensors using a 2Sribose-linked fluorophore. <i>Journal of the American Chemical Society</i> , <b>2005</b> , 127, 12766-7	64
106	Fluorogenic resolution of ligand binding by a nucleic acid aptamer. <i>Journal of the American Chemical Society</i> , <b>2003</b> , 125, 12370-1	62
105	Fingerprinting noncanonical and tertiary RNA structures by differential SHAPE reactivity. <i>Journal of the American Chemical Society</i> , <b>2012</b> , 134, 13160-3	60
104	Native-like RNA tertiary structures using a sequence-encoded cleavage agent and refinement by discrete molecular dynamics. <i>Journal of the American Chemical Society</i> , <b>2009</b> , 131, 2541-6	60
103	RNA-tethered phenyl azide photocrosslinking via a short-lived indiscriminant electrophile. <i>Journal of the American Chemical Society</i> , <b>2003</b> , 125, 10850-61	57
102	Definition of a high-affinity Gag recognition structure mediating packaging of a retroviral RNA genome. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2010</b> , 107, 19248-53	56
101	Validating fragment-based drug discovery for biological RNAs: lead fragments bind and remodel the TPP riboswitch specifically. <i>Chemistry and Biology</i> , <b>2014</b> , 21, 591-5	55

## (2002-2015)

1	100	In-cell SHAPE reveals that free 30S ribosome subunits are in the inactive state. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2015</b> , 112, 2425-30	11.5	51	
Š	99	Two distinct binding modes of a protein cofactor with its target RNA. <i>Journal of Molecular Biology</i> , <b>2006</b> , 361, 771-84	6.5	50	
Ş	98	Pervasive tertiary structure in the dengue virus RNA genome. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2018</b> , 115, 11513-11518	11.5	50	
Ç	97	In-cell RNA structure probing with SHAPE-MaP. <i>Nature Protocols</i> , <b>2018</b> , 13, 1181-1195	18.8	49	
Ş	96	Protein-dependent transition states for ribonucleoprotein assembly. <i>Journal of Molecular Biology</i> , <b>2001</b> , 309, 1087-100	6.5	48	
Š	95	Guidelines for SHAPE Reagent Choice and Detection Strategy for RNA Structure Probing Studies. <i>Biochemistry</i> , <b>2019</b> , 58, 2655-2664	3.2	45	
9	94	Multiple conformations are a conserved and regulatory feature of the RB1 5SUTR. <i>Rna</i> , <b>2015</b> , 21, 1274-8	<b>35</b> .8	45	
ç	93	RNA structures facilitate recombination-mediated gene swapping in HIV-1. <i>Journal of Virology</i> , <b>2010</b> , 84, 12675-82	6.6	45	
Ş	92	Lack of secondary structure characterizes the 5Sends of mammalian mitochondrial mRNAs. <i>Rna</i> , <b>2008</b> , 14, 862-71	5.8	45	
٥	91	Recruitment of intron-encoded and co-opted proteins in splicing of the bI3 group I intron RNA. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2002</b> , 99, 128-33	11.5	45	
Ş	90	RNA base-pairing complexity in living cells visualized by correlated chemical probing. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2019</b> , 116, 24574-24582	11.5	45	
8	89	The genetic code as expressed through relationships between mRNA structure and protein function. <i>FEBS Letters</i> , <b>2013</b> , 587, 1180-1188	3.8	44	
8	88	Challenge of mimicking the influences of the cellular environment on RNA structure by PEG-induced macromolecular crowding. <i>Biochemistry</i> , <b>2015</b> , 54, 6447-53	3.2	42	
8	87	A guanosine-centric mechanism for RNA chaperone function. <i>Science</i> , <b>2013</b> , 340, 190-5	33.3	42	
8	86	Visualizing induced fit in early assembly of the human signal recognition particle. <i>Nature Structural Biology</i> , <b>2001</b> , 8, 515-20		42	
8	85	Slow conformational dynamics at C2Sendo nucleotides in RNA. <i>Journal of the American Chemical Society</i> , <b>2008</b> , 130, 8884-5	16.4	41	
8	84	Complex ligand-induced conformational changes in tRNA(Asp) revealed by single-nucleotide resolution SHAPE chemistry. <i>Biochemistry</i> , <b>2008</b> , 47, 3454-61	3.2	40	
8	83	Catalysis of amide synthesis by RNA phosphodiester and hydroxyl groups. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2002</b> , 99, 14688-93	11.5	39	

82	IFNL3 mRNA structure is remodeled by a functional non-coding polymorphism associated with hepatitis C virus clearance. <i>Scientific Reports</i> , <b>2015</b> , 5, 16037	4.9	37
81	RNA flexibility in the dimerization domain of a gamma retrovirus. <i>Nature Chemical Biology</i> , <b>2005</b> , 1, 104	<b>-1:1</b> :.7	37
80	Model-Free RNA Sequence and Structure Alignment Informed by SHAPE Probing Reveals a Conserved Alternate Secondary Structure for 16S rRNA. <i>PLoS Computational Biology</i> , <b>2015</b> , 11, e100412	2 <i>§</i>	36
79	Structure of an RNA switch that enforces stringent retroviral genomic RNA dimerization.  Proceedings of the National Academy of Sciences of the United States of America, 2006, 103, 13640-5	11.5	36
78	Architecture of a gamma retroviral genomic RNA dimer. <i>Biochemistry</i> , <b>2006</b> , 45, 12664-72	3.2	36
77	Principles for understanding the accuracy of SHAPE-directed RNA structure modeling. <i>Biochemistry</i> , <b>2013</b> , 52, 588-95	3.2	35
76	C2Sendo nucleotides as molecular timers suggested by the folding of an RNA domain. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2009</b> , 106, 15622-7	11.5	35
75	A collapsed state functions to self-chaperone RNA folding into a native ribonucleoprotein complex. <i>Nature Structural Biology</i> , <b>2001</b> , 8, 135-40		35
74	Ribosome RNA assembly intermediates visualized in living cells. <i>Biochemistry</i> , <b>2014</b> , 53, 3237-47	3.2	34
73	Selective 2Shydroxyl acylation analyzed by protection from exoribonuclease. <i>Journal of the American Chemical Society</i> , <b>2010</b> , 132, 9940-3	16.4	34
72	SHAPE analysis of long-range interactions reveals extensive and thermodynamically preferred misfolding in a fragile group I intron RNA. <i>Biochemistry</i> , <b>2008</b> , 47, 8504-13	3.2	34
71	An RNA structure-mediated, posttranscriptional model of human E1-antitrypsin expression.  Proceedings of the National Academy of Sciences of the United States of America, 2017, 114, E10244-E102	2 <del>53</del> .5	33
70	SHAPE-directed discovery of potent shRNA inhibitors of HIV-1. <i>Molecular Therapy</i> , <b>2012</b> , 20, 820-8	11.7	31
69	Mapping Local Nucleotide Flexibility by Selective Acylation of 2FAmine Substituted RNA. <i>Journal of the American Chemical Society</i> , <b>2000</b> , 122, 216-224	16.4	31
68	Long-range architecture in a viral RNA genome. <i>Biochemistry</i> , <b>2013</b> , 52, 3182-90	3.2	29
67	Structure-Based Alignment and Consensus Secondary Structures for Three HIV-Related RNA Genomes. <i>PLoS Computational Biology</i> , <b>2015</b> , 11, e1004230	5	28
66	Specific viral RNA drives the SARS CoV-2 nucleocapsid to phase separate <b>2020</b> ,		28
65	Structure and dynamics of the HIV-1 frameshift element RNA. <i>Biochemistry</i> , <b>2014</b> , 53, 4282-91	3.2	27

64	Sharing and archiving nucleic acid structure mapping data. <i>Rna</i> , <b>2011</b> , 17, 1204-12	5.8	27
63	Robust and generic RNA modeling using inferred constraints: a structure for the hepatitis C virus IRES pseudoknot domain. <i>Biochemistry</i> , <b>2010</b> , 49, 4931-3	3.2	26
62	Physical and Functional Analysis of Viral RNA Genomes by SHAPE. <i>Annual Review of Virology</i> , <b>2019</b> , 6, 93-117	14.6	25
61	Role of context in RNA structure: flanking sequences reconfigure CAG motif folding in huntingtin exon 1 transcripts. <i>Biochemistry</i> , <b>2013</b> , 52, 8219-25	3.2	25
60	Secondary structure of the mature ex virio Moloney murine leukemia virus genomic RNA dimerization domain. <i>Journal of Virology</i> , <b>2010</b> , 84, 898-906	6.6	25
59	Review toward all RNA structures, concisely. <i>Biopolymers</i> , <b>2015</b> , 103, 438-48	2.2	23
58	Selective 2Shydroxyl acylation analyzed by protection from exoribonuclease (RNase-detected SHAPE) for direct analysis of covalent adducts and of nucleotide flexibility in RNA. <i>Nature Protocols</i> , <b>2011</b> , 6, 1683-94	18.8	23
57	Messenger RNA Structure Regulates Translation Initiation: A Mechanism Exploited from Bacteria to Humans. <i>Biochemistry</i> , <b>2018</b> , 57, 3537-3539	3.2	22
56	Nonhierarchical ribonucleoprotein assembly suggests a strain-propagation model for protein-facilitated RNA folding. <i>Biochemistry</i> , <b>2010</b> , 49, 5418-25	3.2	22
55	Differential helix stabilities and sites pre-organized for tertiary interactions revealed by monitoring local nucleotide flexibility in the bI5 group I intron RNA. <i>Biochemistry</i> , <b>2003</b> , 42, 901-9	3.2	22
54	Structure-independent and quantitative ligation of single-stranded DNA. <i>Analytical Biochemistry</i> , <b>2006</b> , 349, 242-6	3.1	21
53	Evolution from DNA to RNA recognition by the bI3 LAGLIDADG maturase. <i>Nature Structural and Molecular Biology</i> , <b>2005</b> , 12, 779-87	17.6	21
52	Targeting the Oncogenic Long Non-coding RNA SLNCR1 by Blocking Its Sequence-Specific Binding to the Androgen Receptor. <i>Cell Reports</i> , <b>2020</b> , 30, 541-554.e5	10.6	20
51	Direct Duplex Detection: An Emerging Tool in the RNA Structure Analysis Toolbox. <i>Trends in Biochemical Sciences</i> , <b>2016</b> , 41, 734-736	10.3	20
50	Direct identification of base-paired RNA nucleotides by correlated chemical probing. <i>Rna</i> , <b>2017</b> , 23, 6-13	<b>3</b> 5.8	20
49	Structural basis for the self-chaperoning function of an RNA collapsed state. <i>Biochemistry</i> , <b>2004</b> , 43, 151	13 <u>9</u> -86	20
48	Visualization of RNA structure models within the Integrative Genomics Viewer. <i>Rna</i> , <b>2017</b> , 23, 1012-101	<b>8</b> 5.8	17
47	Kinetic and thermodynamic framework for assembly of the six-component bI3 group I intron ribonucleoprotein catalyst. <i>Biochemistry</i> , <b>2003</b> , 42, 9980-8	3.2	17

46	Packaged and Free Satellite Tobacco Mosaic Virus (STMV) RNA Genomes Adopt Distinct Conformational States. <i>Biochemistry</i> , <b>2017</b> , 56, 2175-2183	3.2	16	
45	Quantitative analysis of RNA solvent accessibility by N-silylation of guanosine. <i>Biochemistry</i> , <b>2009</b> , 48, 2109-14	3.2	16	
44	The SL1-SL2 (stem-loop) domain is the primary determinant for stability of the gamma retroviral genomic RNA dimer. <i>Journal of Biological Chemistry</i> , <b>2006</b> , 281, 37952-61	5.4	16	
43	A threefold RNA-protein interface in the signal recognition particle gates native complex assembly. Journal of Molecular Biology, <b>2007</b> , 369, 512-24	6.5	16	
42	Compartmentalization directs assembly of the signal recognition particle. <i>Biochemistry</i> , <b>2006</b> , 45, 14955	5-56 <u>4</u>	15	
41	The RNA Domain Vc1 Regulates Downstream Gene Expression in Response to Cyclic Diguanylate in Vibrio cholerae. <i>PLoS ONE</i> , <b>2016</b> , 11, e0148478	3.7	14	
40	Structure modeling of RNA using sparse NMR constraints. <i>Nucleic Acids Research</i> , <b>2017</b> , 45, 12638-1264	<b>7</b> 20.1	13	
39	Analysis of RNA-protein networks with RNP-MaP defines functional hubs on RNA. <i>Nature Biotechnology</i> , <b>2021</b> , 39, 347-356	44.5	13	
38	RNA tertiary structure analysis by 2Shydroxyl molecular interference. <i>Biochemistry</i> , <b>2014</b> , 53, 6825-33	3.2	12	
37	The Mrs1 splicing factor binds the bI3 group I intron at each of two tetraloop-receptor motifs. <i>PLoS ONE</i> , <b>2010</b> , 5, e8983	3.7	12	
36	Femtomole SHAPE reveals regulatory structures in the authentic XMRV RNA genome. <i>Journal of the American Chemical Society</i> , <b>2011</b> , 133, 20326-34	16.4	12	
35	Crystal structures, reactivity and inferred acylation transition states for 2Samine substituted RNA. Journal of the American Chemical Society, <b>2005</b> , 127, 13622-8	16.4	12	
34	Tris-borate is a poor counterion for RNA: a cautionary tale for RNA folding studies. <i>Nucleic Acids Research</i> , <b>2004</b> , 32, e184	20.1	12	
33	Chemical interrogation of mismatches in DNA-DNA and DNA-RNA duplexes under nonstringent conditions by selective 2Samine acylation. <i>Biochemistry</i> , <b>2002</b> , 41, 6866-74	3.2	12	
32	SHAPE analysis of small RNAs and riboswitches. <i>Methods in Enzymology</i> , <b>2014</b> , 549, 165-87	1.7	11	
31	Tagging DNA mismatches by selective 2Samine acylation. <i>Chemistry and Biology</i> , <b>2000</b> , 7, 405-10		11	
30	SHAPE Directed Discovery of New Functions in Large RNAs. <i>Accounts of Chemical Research</i> , <b>2021</b> , 54, 2502-2517	24.3	11	
29	Single-molecule correlated chemical probing reveals large-scale structural communication in the ribosome and the mechanism of the antibiotic spectinomycin in living cells. <i>PLoS Biology</i> , <b>2019</b> , 17, e300	083793	10	

## (2021-2014)

28	An immature retroviral RNA genome resembles a kinetically trapped intermediate state. <i>Journal of Virology</i> , <b>2014</b> , 88, 6061-8	6.6	9
27	Direct detection of RNA modifications and structure using single-molecule nanopore sequencing <i>Cell Genomics</i> , <b>2022</b> , 2, 100097-100097		9
26	Direct detection of RNA modifications and structure using single molecule nanopore sequencing		9
25	RNA structure probing dash seq. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2011</b> , 108, 10933-4	11.5	8
24	Anti-cooperative assembly of the SRP19 and SRP68/72 components of the signal recognition particle. <i>Biochemical Journal</i> , <b>2008</b> , 415, 429-37	3.8	7
23	Mechanics of DNA flexibility visualized by selective 2Samine acylation at nucleotide bulges. <i>Journal of Molecular Biology</i> , <b>2004</b> , 337, 611-9	6.5	6
22	The roles of five conserved lentiviral RNA structures in HIV-1 replication. Virology, 2018, 514, 1-8	3.6	6
21	Effects of Refolding on Large-Scale RNA Structure. <i>Biochemistry</i> , <b>2019</b> , 58, 3069-3077	3.2	5
20	SHAPE Probing Reveals Human rRNAs Are Largely Unfolded in Solution. <i>Biochemistry</i> , <b>2019</b> , 58, 3377-33	3852	5
19	Time-Resolved, Single-Molecule, Correlated Chemical Probing of RNA. <i>Journal of the American Chemical Society</i> , <b>2020</b> , 142, 18735-18740	16.4	5
18	RNA targeting by the HIV-1 Tat protein. Journal of Computer - Aided Molecular Design, 1993, 1, 225-234		4
17	Subsite Ligand Recognition and Cooperativity in the TPP Riboswitch: Implications for Fragment-Linking in RNA Ligand Discovery ACS Chemical Biology, 2022,	4.9	3
16	Direct Mapping of Higher-Order RNA Interactions by SHAPE-JuMP. <i>Biochemistry</i> , <b>2021</b> , 60, 1971-1982	3.2	3
15	Discovery of a large-scale, cell-state-responsive allosteric switch in the 7SK RNA using DANCE-MaP <i>Molecular Cell</i> , <b>2022</b> ,	17.6	3
14	Small structural costs for evolution from RNA to RNP-based catalysis. <i>Journal of Molecular Biology</i> , <b>2003</b> , 331, 57-73	6.5	2
13	RNP-MaP: In-cell analysis of protein interaction networks defines functional hubs in RNA		2
12	Double-stranded RNA drives SARS-CoV-2 nucleocapsid protein to undergo phase separation at specific temperatures <b>2021</b> ,		2
11	Piercing the fog of the RNA structure-ome. <i>Science</i> , <b>2021</b> , 373, 964-965	33.3	2

10	Visualization of lncRNA and mRNA Structure Models Within the Integrative Genomics Viewer. <i>Methods in Molecular Biology</i> , <b>2021</b> , 2254, 15-25	1.4	2
9	Identifying proximal RNA interactions from cDNA-encoded crosslinks with ShapeJumper		1
8	Distinct MUNC lncRNA structural domains regulate transcription of different promyogenic factors <i>Cell Reports</i> , <b>2022</b> , 38, 110361	10.6	1
7	SHAPE-enabled fragment-based ligand discovery for RNA <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2022</b> , 119, e2122660119	11.5	1
6	Identifying proximal RNA interactions from cDNA-encoded crosslinks with ShapeJumper <i>PLoS Computational Biology</i> , <b>2021</b> , 17, e1009632	5	О
5	RNA clubs. <i>Rna</i> , <b>2015</b> , 21, 760-1	5.8	
4	High-Throughput Analysis of RNA Structure by SHAPE Chemistry. <i>FASEB Journal</i> , <b>2009</b> , 23, 326.3	0.9	
4	High-Throughput Analysis of RNA Structure by SHAPE Chemistry. <i>FASEB Journal</i> , <b>2009</b> , 23, 326.3  Accurate SHAPE-directed RNA structure prediction. <i>FASEB Journal</i> , <b>2009</b> , 23, 843.2	0.9	
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