Sarel J Fleishman

List of Publications by Citations

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88
papers

5,713
citations

36
h-index

9-index

75
g-index

7,041
ext. papers

2,041
ext. citations

3,041
ext. citations

4,041
ext. citations

2,041
ext. citations

#	Paper	IF	Citations
88	ROSETTA3: an object-oriented software suite for the simulation and design of macromolecules. <i>Methods in Enzymology</i> , 2011 , 487, 545-74	1.7	1216
87	Computational design of proteins targeting the conserved stem region of influenza hemagglutinin. <i>Science</i> , 2011 , 332, 816-21	33.3	441
86	High-resolution mapping of protein sequence-function relationships. <i>Nature Methods</i> , 2010 , 7, 741-6	21.6	350
85	RosettaScripts: a scripting language interface to the Rosetta macromolecular modeling suite. <i>PLoS ONE</i> , 2011 , 6, e20161	3.7	311
84	Optimization of affinity, specificity and function of designed influenza inhibitors using deep sequencing. <i>Nature Biotechnology</i> , 2012 , 30, 543-8	44.5	279
83	A putative molecular-activation switch in the transmembrane domain of erbB2. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2002 , 99, 15937-40	11.5	220
82	Automated Structure- and Sequence-Based Design of Proteins for High Bacterial Expression and Stability. <i>Molecular Cell</i> , 2016 , 63, 337-346	17.6	204
81	Dominant Mutations in the Autoimmune Regulator AIRE Are Associated with Common Organ-Specific Autoimmune Diseases. <i>Immunity</i> , 2015 , 42, 1185-96	32.3	156
80	Community-wide assessment of protein-interface modeling suggests improvements to design methodology. <i>Journal of Molecular Biology</i> , 2011 , 414, 289-302	6.5	114
79	Principles of Protein Stability and Their Application in Computational Design. <i>Annual Review of Biochemistry</i> , 2018 , 87, 105-129	29.1	106
78	Quasi-symmetry in the cryo-EM structure of EmrE provides the key to modeling its transmembrane domain. <i>Journal of Molecular Biology</i> , 2006 , 364, 54-67	6.5	102
77	The structural and energetic basis for high selectivity in a high-affinity protein-protein interaction. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2010 , 107, 10080-5	11.5	101
76	A Calpha model for the transmembrane alpha helices of gap junction intercellular channels. <i>Molecular Cell</i> , 2004 , 15, 879-88	17.6	100
75	Role of the biomolecular energy gap in protein design, structure, and evolution. <i>Cell</i> , 2012 , 149, 262-73	56.2	83
74	Free diffusion of steroid hormones across biomembranes: a simplex search with implicit solvent model calculations. <i>Biophysical Journal</i> , 2004 , 87, 768-79	2.9	81
73	Automated Design of Efficient and Functionally Diverse Enzyme Repertoires. <i>Molecular Cell</i> , 2018 , 72, 178-186.e5	17.6	79
72	Community-wide evaluation of methods for predicting the effect of mutations on protein-protein interactions. <i>Proteins: Structure, Function and Bioinformatics</i> , 2013 , 81, 1980-7	4.2	78

(2013-2017)

71	Principles for computational design of binding antibodies. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2017 , 114, 10900-10905	11.5	73	
70	Computational design of a protein-based enzyme inhibitor. <i>Journal of Molecular Biology</i> , 2013 , 425, 35	636755	71	
69	Transmembrane protein structures without X-rays. <i>Trends in Biochemical Sciences</i> , 2006 , 31, 106-13	10.3	71	
68	A novel scoring function for predicting the conformations of tightly packed pairs of transmembrane alpha-helices. <i>Journal of Molecular Biology</i> , 2002 , 321, 363-78	6.5	68	
67	Design and in vitro realization of carbon-conserving photorespiration. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2018 , 115, E11455-E11464	11.5	66	
66	Computational design of a pH-sensitive IgG binding protein. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2014 , 111, 675-80	11.5	57	
65	Co-evolving residues in membrane proteins. <i>Bioinformatics</i> , 2007 , 23, 3312-9	7.2	57	
64	A putative mechanism for downregulation of the catalytic activity of the EGF receptor via direct contact between its kinase and C-terminal domains. <i>Structure</i> , 2004 , 12, 2265-75	5.2	57	
63	AbDesign: An algorithm for combinatorial backbone design guided by natural conformations and sequences. <i>Proteins: Structure, Function and Bioinformatics</i> , 2015 , 83, 1385-406	4.2	55	
62	Progress in structure prediction of alpha-helical membrane proteins. <i>Current Opinion in Structural Biology</i> , 2006 , 16, 496-504	8.1	55	
61	An evolutionarily conserved network of amino acids mediates gating in voltage-dependent potassium channels. <i>Journal of Molecular Biology</i> , 2004 , 340, 307-18	6.5	52	
60	An automatic method for predicting transmembrane protein structures using cryo-EM and evolutionary data. <i>Biophysical Journal</i> , 2004 , 87, 3448-59	2.9	48	
59	Computational design of protein-protein interactions. <i>Current Opinion in Structural Biology</i> , 2013 , 23, 903-10	8.1	46	
58	Structure of the ultra-high-affinity colicin E2 DNaseIm2 complex. <i>Journal of Molecular Biology</i> , 2012 , 417, 79-94	6.5	45	
57	One-step design of a stable variant of the malaria invasion protein RH5 for use as a vaccine immunogen. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2017 , 114, 998-1002	11.5	44	
56	Mutational scanning reveals the determinants of protein insertion and association energetics in the plasma membrane. <i>ELife</i> , 2016 , 5,	8.9	43	
55	Overcoming an optimization plateau in the directed evolution of highly efficient nerve agent bioscavengers. <i>Protein Engineering, Design and Selection</i> , 2017 , 30, 333-345	1.9	41	
54	Emerging themes in the computational design of novel enzymes and protein-protein interfaces. <i>FEBS Letters</i> , 2013 , 587, 1147-54	3.8	41	

53	Restricted sidechain plasticity in the structures of native proteins and complexes. <i>Protein Science</i> , 2011 , 20, 753-7	6.3	37
52	Highly active enzymes by automated combinatorial backbone assembly and sequence design. <i>Nature Communications</i> , 2018 , 9, 2780	17.4	32
51	Hotspot-centric de novo design of protein binders. Journal of Molecular Biology, 2011, 413, 1047-62	6.5	32
50	Intrinsically disordered C-terminal segments of voltage-activated potassium channels: a possible fishing rod-like mechanism for channel binding to scaffold proteins. <i>Bioinformatics</i> , 2006 , 22, 1546-50	7.2	32
49	Ultrahigh specificity in a network of computationally designed protein-interaction pairs. <i>Nature Communications</i> , 2018 , 9, 5286	17.4	32
48	Optimizing antibody affinity and stability by the automated design of the variable light-heavy chain interfaces. <i>PLoS Computational Biology</i> , 2019 , 15, e1007207	5	31
47	Computational design of novel protein binders and experimental affinity maturation. <i>Methods in Enzymology</i> , 2013 , 523, 1-19	1.7	30
46	Why reinvent the wheel? Building new proteins based on ready-made parts. <i>Protein Science</i> , 2016 , 25, 1179-87	6.3	28
45	Interplay between hydrophobicity and the positive-inside rule in determining membrane-protein topology. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2016 , 113, 10340-5	11.5	26
44	RosettaDock in CAPRI rounds 6-12. <i>Proteins: Structure, Function and Bioinformatics</i> , 2007 , 69, 758-63	4.2	26
43	A new twist in TCR diversity revealed by a forbidden alphabeta TCR. <i>Journal of Molecular Biology</i> , 2008 , 375, 1306-19	6.5	21
42	Local energetic frustration affects the dependence of green fluorescent protein folding on the chaperonin GroEL. <i>Journal of Biological Chemistry</i> , 2017 , 292, 20583-20591	5.4	19
41	Rosetta in CAPRI rounds 13-19. Proteins: Structure, Function and Bioinformatics, 2010, 78, 3212-8	4.2	17
40	Has the code for protein translocation been broken?. <i>Trends in Biochemical Sciences</i> , 2006 , 31, 192-6	10.3	17
39	Prediction and simulation of motion in pairs of transmembrane alpha-helices. <i>Bioinformatics</i> , 2007 , 23, e212-8	7.2	17
38	A lipophilicity-based energy function for membrane-protein modelling and design. <i>PLoS Computational Biology</i> , 2019 , 15, e1007318	5	16
37	Practically useful protein-design methods combining phylogenetic and atomistic calculations. <i>Current Opinion in Structural Biology</i> , 2020 , 63, 58-64	8.1	16
36	A "fuzzy"-logic language for encoding multiple physical traits in biomolecules. <i>Journal of Molecular Biology</i> , 2014 , 426, 4125-4138	6.5	16

(2021-2017)

35	High-accuracy modeling of antibody structures by a search for minimum-energy recombination of backbone fragments. <i>Proteins: Structure, Function and Bioinformatics</i> , 2017 , 85, 30-38	4.2	14	
34	Improved antibody-based ricin neutralization by affinity maturation is correlated with slower off-rate values. <i>Protein Engineering, Design and Selection</i> , 2017 , 30, 611-617	1.9	13	
33	Assigning transmembrane segments to helices in intermediate-resolution structures. <i>Bioinformatics</i> , 2004 , 20 Suppl 1, i122-9	7.2	13	
32	Estimating Interprotein Pairwise Interaction Energies in Cell Lysates from a Single Native Mass Spectrum. <i>Analytical Chemistry</i> , 2018 , 90, 10090-10094	7.8	12	
31	Manipulating the Folding Landscape of a Multidomain Protein. <i>Journal of Physical Chemistry B</i> , 2018 , 122, 11030-11038	3.4	12	
30	The structural context of disease-causing mutations in gap junctions. <i>Journal of Biological Chemistry</i> , 2006 , 281, 28958-63	5.4	12	
29	Computationally Designed Armadillo Repeat Proteins for Modular Peptide Recognition. <i>Journal of Molecular Biology</i> , 2016 , 428, 4467-4489	6.5	12	
28	A combined computational-experimental approach to define the structural origin of antibody recognition of sialyl-Tn, a tumor-associated carbohydrate antigen. <i>Scientific Reports</i> , 2018 , 8, 10786	4.9	11	
27	Computationally designed pyocyanin demethylase acts synergistically with tobramycin to kill recalcitrant biofilms. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2021 , 118,	11.5	10	
26	AbPredict 2: a server for accurate and unstrained structure prediction of antibody variable domains. <i>Bioinformatics</i> , 2019 , 35, 1591-1593	7.2	10	
25	Incorporating an allosteric regulatory site in an antibody through backbone design. <i>Protein Science</i> , 2017 , 26, 807-813	6.3	9	
24	Combined Crystal Structure of a Type I Cohesin: MUTATION AND AFFINITY BINDING STUDIES REVEAL STRUCTURAL DETERMINANTS OF COHESIN-DOCKERIN SPECIFICITIES. <i>Journal of Biological Chemistry</i> , 2015 , 290, 16215-25	5.4	9	
23	Computational protein design suggests that human PCNA-partner interactions are not optimized for affinity. <i>Proteins: Structure, Function and Bioinformatics</i> , 2013 , 81, 341-8	4.2	8	
22	pANT: a method for the pairwise assessment of nonfunctionalization times of processed pseudogenes. <i>Molecular Biology and Evolution</i> , 2003 , 20, 1876-80	8.3	8	
21	Community-Wide Experimental Evaluation of the PROSS Stability-Design Method. <i>Journal of Molecular Biology</i> , 2021 , 433, 166964	6.5	8	
20	PROTEIN DESIGN. Inspired by nature. <i>Science</i> , 2016 , 352, 657-8	33.3	8	
19	Overcoming a species-specificity barrier in development of an inhibitory antibody targeting a modulator of tumor stroma. <i>Protein Engineering, Design and Selection</i> , 2016 , 29, 135-47	1.9	7	
18	The AbDesign computational pipeline for modular backbone assembly and design of binders and enzymes. <i>Protein Science</i> , 2021 , 30, 151-159	6.3	7	

17	Collective repacking reveals that the structures of protein cores are uniquely specified by steric repulsive interactions. <i>Protein Engineering, Design and Selection</i> , 2017 , 30, 387-394	1.9	6
16	One-step sequence and structure-guided optimization of HIV-1 envelope gp140. <i>Current Research in Structural Biology</i> , 2020 , 2, 45-55	2.8	6
15	Extending the New Generation of Structure Predictors to Account for Dynamics and Allostery. <i>Journal of Molecular Biology</i> , 2021 , 433, 167007	6.5	6
14	PROSS 2: a new server for the design of stable and highly expressed protein variants. <i>Bioinformatics</i> , 2020 ,	7.2	5
13	Local Mutations Can Serve as a Game Changer for Global Protein Solvent Interaction. <i>Jacs Au</i> , 2021 , 1, 1076-1085		5
12	Computationally designed dual-color MRI reporters for noninvasive imaging of transgene expression <i>Nature Biotechnology</i> , 2022 ,	44.5	4
11	Stable and Functionally Diverse Versatile Peroxidases Designed Directly from Sequences <i>Journal of the American Chemical Society</i> , 2022 , 144, 3564-3571	16.4	4
10	A computationally designed fluorescent biosensor for D-serine		3
9	Design of a basigin-mimicking inhibitor targeting the malaria invasion protein RH5. <i>Proteins:</i> Structure, Function and Bioinformatics, 2020 , 88, 187-195	4.2	3
8	The neutralization potency of anti-SARS-CoV-2 therapeutic human monoclonal antibodies is retained against viral variants. <i>Cell Reports</i> , 2021 , 36, 109679	10.6	3
7	What Have We Learned from Design of Function in Large Proteins?. <i>Biodesign Research</i> , 2022 , 2022, 1-1	13.1	2
6	De novodesigned receptor transmembrane domains enhance CAR-T cell cytotoxicity and attenuate cytokine release		1
5	A Rationally and Computationally Designed Fluorescent Biosensor for d-Serine. <i>ACS Sensors</i> , 2021 , 6, 4193-4205	9.2	1
4	Computational Enzyme Engineering Pipelines for Optimized Production of Renewable Chemicals. <i>Frontiers in Bioengineering and Biotechnology</i> , 2021 , 9, 673005	5.8	1
3	Biomolecular Recognition of the Glycan Neoantigen CA19-9 by Distinct Antibodies. <i>Journal of Molecular Biology</i> , 2021 , 433, 167099	6.5	1
2	Direct-MS analysis of antibody-antigen complexes. <i>Proteomics</i> , 2021 , 21, e2000300	4.8	1
1	Computer-aided engineering of staphylokinase toward enhanced affinity and selectivity for plasmin <i>Computational and Structural Biotechnology Journal</i> , 2022 , 20, 1366-1377	6.8	0