Woody Sherman

List of Publications by Citations

Source: https://exaly.com/author-pdf/1954400/woody-sherman-publications-by-citations.pdf

Version: 2024-04-17

This document has been generated based on the publications and citations recorded by exaly.com. For the latest version of this publication list, visit the link given above.

The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

12,456 103 91 47 h-index g-index citations papers 6.53 15,041 103 5.9 L-index avg, IF ext. citations ext. papers

#	Paper	IF	Citations
91	Protein and ligand preparation: parameters, protocols, and influence on virtual screening enrichments. <i>Journal of Computer-Aided Molecular Design</i> , 2013 , 27, 221-34	4.2	2521
90	Novel procedure for modeling ligand/receptor induced fit effects. <i>Journal of Medicinal Chemistry</i> , 2006 , 49, 534-53	8.3	1320
89	Prediction of Absolute Solvation Free Energies using Molecular Dynamics Free Energy Perturbation and the OPLS Force Field. <i>Journal of Chemical Theory and Computation</i> , 2010 , 6, 1509-19	6.4	959
88	Improved docking of polypeptides with Glide. <i>Journal of Chemical Information and Modeling</i> , 2013 , 53, 1689-99	6.1	815
87	Accurate and reliable prediction of relative ligand binding potency in prospective drug discovery by way of a modern free-energy calculation protocol and force field. <i>Journal of the American Chemical Society</i> , 2015 , 137, 2695-703	16.4	633
86	Use of an induced fit receptor structure in virtual screening. <i>Chemical Biology and Drug Design</i> , 2006 , 67, 83-4	2.9	376
85	Probing the alpha-helical structural stability of stapled p53 peptides: molecular dynamics simulations and analysis. <i>Chemical Biology and Drug Design</i> , 2010 , 75, 348-59	2.9	309
84	ConfGen: a conformational search method for efficient generation of bioactive conformers. <i>Journal of Chemical Information and Modeling</i> , 2010 , 50, 534-46	6.1	303
83	Analysis and comparison of 2D fingerprints: insights into database screening performance using eight fingerprint methods. <i>Journal of Molecular Graphics and Modelling</i> , 2010 , 29, 157-70	2.8	275
82	Mechanism of the hydrophobic effect in the biomolecular recognition of arylsulfonamides by carbonic anhydrase. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2011 , 108, 17889-94	11.5	268
81	Relative Binding Free Energy Calculations in Drug Discovery: Recent Advances and Practical Considerations. <i>Journal of Chemical Information and Modeling</i> , 2017 , 57, 2911-2937	6.1	254
80	Water networks contribute to enthalpy/entropy compensation in protein-ligand binding. <i>Journal of the American Chemical Society</i> , 2013 , 135, 15579-84	16.4	230
79	Large-scale systematic analysis of 2D fingerprint methods and parameters to improve virtual screening enrichments. <i>Journal of Chemical Information and Modeling</i> , 2010 , 50, 771-84	6.1	227
78	Novel method for generating structure-based pharmacophores using energetic analysis. <i>Journal of Chemical Information and Modeling</i> , 2009 , 49, 2356-68	6.1	224
77	Rational approaches to improving selectivity in drug design. <i>Journal of Medicinal Chemistry</i> , 2012 , 55, 1424-44	8.3	189
76	Improving the Prediction of Absolute Solvation Free Energies Using the Next Generation OPLS Force Field. <i>Journal of Chemical Theory and Computation</i> , 2012 , 8, 2553-8	6.4	159
75	Rapid shape-based ligand alignment and virtual screening method based on atom/feature-pair similarities and volume overlap scoring. <i>Journal of Chemical Information and Modeling</i> , 2011 , 51, 2455-6	56 ^{6.1}	147

(2014-2006)

74	Affinity enhancement of an in vivo matured therapeutic antibody using structure-based computational design. <i>Protein Science</i> , 2006 , 15, 949-60	6.3	137
73	Modeling Local Structural Rearrangements Using FEP/REST: Application to Relative Binding Affinity Predictions of CDK2 Inhibitors. <i>Journal of Chemical Theory and Computation</i> , 2013 , 9, 1282-93	6.4	128
72	Energetic analysis of fragment docking and application to structure-based pharmacophore hypothesis generation. <i>Journal of Computer-Aided Molecular Design</i> , 2009 , 23, 541-54	4.2	125
71	Applying physics-based scoring to calculate free energies of binding for single amino acid mutations in protein-protein complexes. <i>PLoS ONE</i> , 2013 , 8, e82849	3.7	116
70	Thermodynamic analysis of water molecules at the surface of proteins and applications to binding site prediction and characterization. <i>Proteins: Structure, Function and Bioinformatics</i> , 2012 , 80, 871-83	4.2	105
69	Contribution of explicit solvent effects to the binding affinity of small-molecule inhibitors in blood coagulation factor serine proteases. <i>ChemMedChem</i> , 2011 , 6, 1049-66	3.7	103
68	Understanding kinase selectivity through energetic analysis of binding site waters. <i>ChemMedChem</i> , 2010 , 5, 618-27	3.7	99
67	High-energy water sites determine peptide binding affinity and specificity of PDZ domains. <i>Protein Science</i> , 2009 , 18, 1609-19	6.3	94
66	Hydration Site Thermodynamics Explain SARs for Triazolylpurines Analogues Binding to the A2A Receptor. <i>ACS Medicinal Chemistry Letters</i> , 2010 , 1, 160-4	4.3	91
65	Computational approaches for fragment-based and de novo design. <i>Current Topics in Medicinal Chemistry</i> , 2010 , 10, 14-32	3	87
64	Accurate Binding Free Energy Predictions in Fragment Optimization. <i>Journal of Chemical Information and Modeling</i> , 2015 , 55, 2411-20	6.1	83
63	Predicting Binding Affinities for GPCR Ligands Using Free-Energy Perturbation. <i>ACS Omega</i> , 2016 , 1, 293-304	3.9	81
62	Impact of binding site waters on inhibitor design: contemplating a novel inverse binding mode of indirubin derivatives in DYRK kinases. <i>Journal of Cheminformatics</i> , 2014 , 6,	8.6	78
61	Interactions between Hofmeister anions and the binding pocket of a protein. <i>Journal of the American Chemical Society</i> , 2015 , 137, 3859-66	16.4	76
60	Current assessment of docking into GPCR crystal structures and homology models: successes, challenges, and guidelines. <i>Journal of Chemical Information and Modeling</i> , 2012 , 52, 3263-77	6.1	74
59	Exploring protein flexibility: incorporating structural ensembles from crystal structures and simulation into virtual screening protocols. <i>Journal of Physical Chemistry B</i> , 2012 , 116, 6952-9	3.4	71
58	Allosteric inhibition of the NS2B-NS3 protease from dengue virus. ACS Chemical Biology, 2013, 8, 2744-5	5 4 .9	66
57	A computational approach to enzyme design: predicting faminotransferase catalytic activity using docking and MM-GBSA scoring. <i>Journal of Chemical Information and Modeling</i> , 2014 , 54, 2334-46	6.1	62

56	Generation of receptor structural ensembles for virtual screening using binding site shape analysis and clustering. <i>Chemical Biology and Drug Design</i> , 2012 , 80, 182-93	2.9	62
55	Improving docking results via reranking of ensembles of ligand poses in multiple X-ray protein conformations with MM-GBSA. <i>Journal of Chemical Information and Modeling</i> , 2014 , 54, 2697-717	6.1	60
54	Chemical basis for the recognition of trimethyllysine by epigenetic reader proteins. <i>Nature Communications</i> , 2015 , 6, 8911	17.4	57
53	Novel Inverse Binding Mode of Indirubin Derivatives Yields Improved Selectivity for DYRK Kinases. <i>ACS Medicinal Chemistry Letters</i> , 2013 , 4, 22-26	4.3	57
52	Improving database enrichment through ensemble docking. <i>Journal of Computer-Aided Molecular Design</i> , 2008 , 22, 621-7	4.2	56
51	Alchemical Binding Free Energy Calculations in AMBER20: Advances and Best Practices for Drug Discovery. <i>Journal of Chemical Information and Modeling</i> , 2020 , 60, 5595-5623	6.1	56
50	Predicting the Effect of Amino Acid Single-Point Mutations on Protein Stability-Large-Scale Validation of MD-Based Relative Free Energy Calculations. <i>Journal of Molecular Biology</i> , 2017 , 429, 948-	963	54
49	Rigorous Free Energy Simulations in Virtual Screening. <i>Journal of Chemical Information and Modeling</i> , 2020 , 60, 4153-4169	6.1	54
48	Consensus Induced Fit Docking (cIFD): methodology, validation, and application to the discovery of novel Crm1 inhibitors. <i>Journal of Computer-Aided Molecular Design</i> , 2012 , 26, 1217-28	4.2	54
47	Boosting virtual screening enrichments with data fusion: coalescing hits from two-dimensional fingerprints, shape, and docking. <i>Journal of Chemical Information and Modeling</i> , 2013 , 53, 1531-42	6.1	54
46	New hypotheses about the structure-function of proprotein convertase subtilisin/kexin type 9: analysis of the epidermal growth factor-like repeat A docking site using WaterMap. <i>Proteins: Structure, Function and Bioinformatics</i> , 2010 , 78, 2571-86	4.2	53
45	AutoQSAR: an automated machine learning tool for best-practice quantitative structure-activity relationship modeling. <i>Future Medicinal Chemistry</i> , 2016 , 8, 1825-1839	4.1	50
44	Kernel-based partial least squares: application to fingerprint-based QSAR with model visualization. Journal of Chemical Information and Modeling, 2013 , 53, 2312-21	6.1	46
43	Contributions of water transfer energy to protein-ligand association and dissociation barriers: Watermap analysis of a series of p38IMAP kinase inhibitors. <i>Proteins: Structure, Function and Bioinformatics</i> , 2013 , 81, 1509-26	4.2	46
42	Type II kinase inhibitors show an unexpected inhibition mode against Parkinson u disease-linked LRRK2 mutant G2019S. <i>Biochemistry</i> , 2013 , 52, 1725-36	3.2	44
41	Selecting an optimal number of binding site waters to improve virtual screening enrichments against the adenosine A2A receptor. <i>Journal of Chemical Information and Modeling</i> , 2014 , 54, 1737-46	6.1	43
40	The binding of benzoarylsulfonamide ligands to human carbonic anhydrase is insensitive to formal fluorination of the ligand. <i>Angewandte Chemie - International Edition</i> , 2013 , 52, 7714-7	16.4	43
39	Deciphering Cryptic Binding Sites on Proteins by Mixed-Solvent Molecular Dynamics. <i>Journal of Chemical Information and Modeling</i> , 2017 , 57, 1388-1401	6.1	41

(2020-2017)

38	Calculating Water Thermodynamics in the Binding Site of Proteins - Applications of WaterMap to Drug Discovery. <i>Current Topics in Medicinal Chemistry</i> , 2017 , 17, 2586-2598	3	41	
37	Relative Binding Free Energy Calculations Applied to Protein Homology Models. <i>Journal of Chemical Information and Modeling</i> , 2016 , 56, 2388-2400	6.1	40	
36	Structure-based approach to the prediction of disulfide bonds in proteins. <i>Protein Engineering, Design and Selection</i> , 2014 , 27, 365-74	1.9	39	
35	The Hsp70 interdomain linker is a dynamic switch that enables allosteric communication between two structured domains. <i>Journal of Biological Chemistry</i> , 2017 , 292, 14765-14774	5.4	37	
34	Computer-aided drug design of falcipain inhibitors: virtual screening, structure-activity relationships, hydration site thermodynamics, and reactivity analysis. <i>Journal of Chemical Information and Modeling</i> , 2012 , 52, 696-710	6.1	37	
33	Improving the Resistance Profile of Hepatitis C NS3/4A Inhibitors: Dynamic Substrate Envelope Guided Design. <i>Journal of Chemical Theory and Computation</i> , 2013 , 9, 5693-5705	6.4	30	
32	Physics-based enzyme design: predicting binding affinity and catalytic activity. <i>Proteins: Structure, Function and Bioinformatics</i> , 2014 , 82, 3397-409	4.2	29	
31	Structure-based virtual screening of MT2 melatonin receptor: influence of template choice and structural refinement. <i>Journal of Chemical Information and Modeling</i> , 2013 , 53, 821-35	6.1	28	
30	Differential Water Thermodynamics Determine PI3K-Beta/Delta Selectivity for Solvent-Exposed Ligand Modifications. <i>Journal of Chemical Information and Modeling</i> , 2016 , 56, 886-94	6.1	26	
29	Exploring conformational search protocols for ligand-based virtual screening and 3-D QSAR modeling. <i>Journal of Computer-Aided Molecular Design</i> , 2015 , 29, 165-82	4.2	25	
28	Water-Restructuring Mutations Can Reverse the Thermodynamic Signature of Ligand Binding to Human Carbonic Anhydrase. <i>Angewandte Chemie - International Edition</i> , 2017 , 56, 3833-3837	16.4	24	
27	Molecular dynamics techniques for modeling G protein-coupled receptors. <i>Current Opinion in Pharmacology</i> , 2016 , 30, 69-75	5.1	24	
26	Novel method for probing the specificity binding profile of ligands: applications to HIV protease. <i>Chemical Biology and Drug Design</i> , 2008 , 71, 387-407	2.9	22	
25	On the Rational Design of Zeolite Clusters. ACS Catalysis, 2015 , 5, 2859-2865	13.1	21	
24	Close intramolecular sulfur-oxygen contacts: modified force field parameters for improved conformation generation. <i>Journal of Computer-Aided Molecular Design</i> , 2012 , 26, 1195-205	4.2	18	
23	Highly efficient implementation of pseudospectral time-dependent density-functional theory for the calculation of excitation energies of large molecules. <i>Journal of Computational Chemistry</i> , 2016 , 37, 1425-41	3.5	17	
22	Predicting GPCR promiscuity using binding site features. <i>Journal of Chemical Information and Modeling</i> , 2014 , 54, 184-94	6.1	17	
21	Improved Alchemical Free Energy Calculations with Optimized Smoothstep Softcore Potentials. Journal of Chemical Theory and Computation, 2020, 16, 5512-5525	6.4	17	

20	Large-Scale Validation of Mixed-Solvent Simulations to Assess Hotspots at Protein-Protein Interaction Interfaces. <i>Journal of Chemical Information and Modeling</i> , 2018 , 58, 784-793	6.1	16
19	Docking and Virtual Screening Strategies for GPCR Drug Discovery. <i>Methods in Molecular Biology</i> , 2015 , 1335, 251-76	1.4	14
18	A Water-Bridged Cysteine-Cysteine Redox Regulation Mechanism in Bacterial Protein Tyrosine Phosphatases. <i>CheM</i> , 2017 , 3, 665-677	16.2	13
17	In search of novel ligands using a structure-based approach: a case study on the adenosine A receptor. <i>Journal of Computer-Aided Molecular Design</i> , 2016 , 30, 863-874	4.2	13
16	Synthesis, biological evaluation, hydration site thermodynamics, and chemical reactivity analysis of Eketo substituted peptidomimetics for the inhibition of Plasmodium falciparum. <i>Bioorganic and Medicinal Chemistry Letters</i> , 2014 , 24, 1274-9	2.9	13
15	Hole filling and library optimization: application to commercially available fragment libraries. <i>Bioorganic and Medicinal Chemistry</i> , 2012 , 20, 5379-87	3.4	12
14	The Binding of Benzoarylsulfonamide Ligands to Human Carbonic Anhydrase is Insensitive to Formal Fluorination of the Ligand. <i>Angewandte Chemie</i> , 2013 , 125, 7868-7871	3.6	11
13	Fully integrated FPGA molecular dynamics simulations 2019 ,		11
12	Mechanistic and computational studies of the reductive half-reaction of tyrosine to phenylalanine active site variants of D-arginine dehydrogenase. <i>Biochemistry</i> , 2014 , 53, 6574-83	3.2	10
11	Accounting for the Central Role of Interfacial Water in Protein-Ligand Binding Free Energy Calculations. <i>Journal of Chemical Theory and Computation</i> , 2020 , 16, 7883-7894	6.4	6
11	•	6.4	5
	Calculations. Journal of Chemical Theory and Computation, 2020 , 16, 7883-7894	6.4	
10	Calculations. Journal of Chemical Theory and Computation, 2020, 16, 7883-7894 2019, Mechanism of biomolecular recognition of trimethyllysine by the fluorinated aromatic cage of		
10	Calculations. Journal of Chemical Theory and Computation, 2020, 16, 7883-7894 2019, Mechanism of biomolecular recognition of trimethyllysine by the fluorinated aromatic cage of KDM5A PHD3 finger. Communications Chemistry, 2020, 3, Automated Protocol for Large-Scale Modeling of Gene Expression Data. Journal of Chemical	6.3	5
10 9 8	Calculations. Journal of Chemical Theory and Computation, 2020, 16, 7883-7894 2019, Mechanism of biomolecular recognition of trimethyllysine by the fluorinated aromatic cage of KDM5A PHD3 finger. Communications Chemistry, 2020, 3, Automated Protocol for Large-Scale Modeling of Gene Expression Data. Journal of Chemical Information and Modeling, 2016, 56, 2216-2224 On the Rational Design of Zeolite Clusters for Converging Reaction Barriers: Quantum Study of	6.3	5 4 3
10 9 8 7	Calculations. Journal of Chemical Theory and Computation, 2020, 16, 7883-7894 2019, Mechanism of biomolecular recognition of trimethyllysine by the fluorinated aromatic cage of KDM5A PHD3 finger. Communications Chemistry, 2020, 3, Automated Protocol for Large-Scale Modeling of Gene Expression Data. Journal of Chemical Information and Modeling, 2016, 56, 2216-2224 On the Rational Design of Zeolite Clusters for Converging Reaction Barriers: Quantum Study of Aldol Kinetics Confined in HZSM-5. Journal of Physical Chemistry C, 2018, 122, 23230-23241	6.3 6.1 3.8	5433
10 9 8 7 6	Calculations. Journal of Chemical Theory and Computation, 2020, 16, 7883-7894 2019, Mechanism of biomolecular recognition of trimethyllysine by the fluorinated aromatic cage of KDM5A PHD3 finger. Communications Chemistry, 2020, 3, Automated Protocol for Large-Scale Modeling of Gene Expression Data. Journal of Chemical Information and Modeling, 2016, 56, 2216-2224 On the Rational Design of Zeolite Clusters for Converging Reaction Barriers: Quantum Study of Aldol Kinetics Confined in HZSM-5. Journal of Physical Chemistry C, 2018, 122, 23230-23241 Free Energy Methods in Drug DiscoveryIntroduction. ACS Symposium Series, 1-38 Water-Restructuring Mutations Can Reverse the Thermodynamic Signature of Ligand Binding to	6.3 6.1 3.8 0.4	54333

- 2 Improving Force Field Parameters for Small-Molecule Conformation Generation **2016**, 57-85
- 11th German Conference on Chemoinformatics (GCC 2015): Fulda, Germany. 8-10 November 2015.

 Journal of Cheminformatics, **2016**, 8, 18

8.6