Alessandro Pedretti

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/438477/publications.pdf

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38 papers 2,373 citations

394421 19 h-index 302126 39 g-index

39 all docs 39 docs citations

39 times ranked

3728 citing authors

#	Article	IF	CITATIONS
1	A multiscale approach to predict the binding mode of metallo betaâ€lactamase inhibitors. Proteins: Structure, Function and Bioinformatics, 2022, 90, 372-384.	2.6	8
2	Extensive Sampling of Molecular Dynamics Simulations to Identify Reliable Protein Structures for Optimized Virtual Screening Studies: The Case of the hTRPM8 Channel. International Journal of Molecular Sciences, 2022, 23, 7558.	4.1	1
3	The VEGA suite of programs: an versatile platform for cheminformatics and drug design projects. Bioinformatics, 2021, 37, 1174-1175.	4.1	90
4	Combining Different Docking Engines and Consensus Strategies to Design and Validate Optimized Virtual Screening Protocols for the SARS-CoV-2 3CL Protease. Molecules, 2021, 26, 797.	3.8	14
5	In-Depth AGE and ALE Profiling of Human Albumin in Heart Failure: Ex Vivo Studies. Antioxidants, 2021, 10, 358.	5.1	4
6	MetaTREE, a Novel Database Focused on Metabolic Trees, Predicts an Important Detoxification Mechanism: The Glutathione Conjugation. Molecules, 2021, 26, 2098.	3.8	2
7	MetaClass, a Comprehensive Classification System for Predicting the Occurrence of Metabolic Reactions Based on the MetaQSAR Database. Molecules, 2021, 26, 5857.	3.8	4
8	Tree2C: A Flexible Tool for Enabling Model Deployment with Special Focus on Cheminformatics Applications. Applied Sciences (Switzerland), 2020, 10, 7704.	2.5	3
9	Towards the Inhibition of Protein–Protein Interactions (PPIs) in STAT3: Insights into a New Class of Benzothiadiazole Derivatives. Molecules, 2020, 25, 3509.	3.8	1
10	A Comprehensive Mapping of the Druggable Cavities within the SARS-CoV-2 Therapeutically Relevant Proteins by Combining Pocket and Docking Searches as Implemented in Pockets 2.0. International Journal of Molecular Sciences, 2020, 21, 5152.	4.1	34
11	Repositioning Dequalinium as Potent Muscarinic Allosteric Ligand by Combining Virtual Screening Campaigns and Experimental Binding Assays. International Journal of Molecular Sciences, 2020, 21, 5961.	4.1	9
12	Homology Modeling of Leishmanolysin (gp63) from <i>Leishmania panamensis</i> and Molecular Docking of Flavonoids. ACS Omega, 2020, 5, 14741-14749.	3.5	26
13	Type 2 Diabetes Mellitus: A Review of Multi-Target Drugs. Molecules, 2020, 25, 1987.	3.8	208
14	Combining Molecular Dynamics and Docking Simulations to Develop Targeted Protocols for Performing Optimized Virtual Screening Campaigns on the hTRPM8 Channel. International Journal of Molecular Sciences, 2020, 21, 2265.	4.1	15
15	FAME 3: Predicting the Sites of Metabolism in Synthetic Compounds and Natural Products for Phase 1 and Phase 2 Metabolic Enzymes. Journal of Chemical Information and Modeling, 2019, 59, 3400-3412.	5.4	60
16	Rescoring and Linearly Combining: A Highly Effective Consensus Strategy for Virtual Screening Campaigns. International Journal of Molecular Sciences, 2019, 20, 2060.	4.1	17
17	Prediction of UGT-mediated Metabolism Using the Manually Curated MetaQSAR Database. ACS Medicinal Chemistry Letters, 2019, 10, 633-638.	2.8	10
18	MetaQSAR: An Integrated Database Engine to Manage and Analyze Metabolic Data. Journal of Medicinal Chemistry, 2018, 61, 1019-1030.	6.4	18

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19	Prediction of the Formation of Reactive Metabolites by A Novel Classifier Approach Based on Enrichment Factor Optimization (EFO) as Implemented in the VEGA Program. Molecules, 2018, 23, 2955.	3.8	9
20	WarpEngine, a Flexible Platform for Distributed Computing Implemented in the VEGA Program and Specially Targeted for Virtual Screening Studies. Journal of Chemical Information and Modeling, 2018, 58, 1154-1160.	5.4	3
21	Binding Space Concept: A New Approach To Enhance the Reliability of Docking Scores and Its Application to Predicting Butyrylcholinesterase Hydrolytic Activity. Journal of Chemical Information and Modeling, 2017, 57, 1691-1702.	5.4	31
22	Key factors regulating protein carbonylation by $\hat{l}\pm,\hat{l}^2$ unsaturated carbonyls: A structural study based on a retrospective meta-analysis. Biophysical Chemistry, 2017, 230, 20-26.	2.8	3
23	Novel selective, potent naphthyl TRPM8 antagonists identified through a combined ligand- and structure-based virtual screening approach. Scientific Reports, 2017, 7, 10999.	3.3	23
24	Structural Effects of Some Relevant Missense Mutations on the MECP2â€DNA Binding: A MD Study Analyzed by Rescore+, a Versatile Rescoring Tool of the VEGA ZZ Program. Molecular Informatics, 2016, 35, 424-433.	2.5	18
25	Computational approaches in the rational design of improved carbonyl quenchers: focus on histidine containing dipeptides. Future Medicinal Chemistry, 2016, 8, 1721-1737.	2.3	21
26	Enhancing the Reliability of GPCR Models by Accounting for Flexibility of Their Pro ontaining Helices: the Case of the Human mAChR1 Receptor. Molecular Informatics, 2015, 34, 216-227.	2.5	2
27	Synthesis, structure–activity relationships and stereochemical investigations of new tricyclic pyridazinone derivatives as potential STAT3 inhibitors. MedChemComm, 2013, 4, 1181.	3.4	16
28	Biological and computational evaluation of an oxadiazole derivative (MD77) as a new lead for direct STAT3 inhibitors. MedChemComm, 2012, 3, 592.	3.4	26
29	Reactions and enzymes in the metabolism of drugs and other xenobiotics. Drug Discovery Today, 2012, 17, 549-560.	6.4	183
30	Homology modeling and metabolism prediction of human carboxylesterase-2 using docking analyses by GriDock: a parallelized tool based on AutoDock 4.0. Journal of Computer-Aided Molecular Design, 2010, 24, 771-787.	2.9	40
31	In silico prediction of human carboxylesterase-1 (hCES1) metabolism combining docking analyses and MD simulations. Bioorganic and Medicinal Chemistry, 2010, 18, 320-329.	3.0	42
32	Synthesis, modeling, and crystallographic study of 3,4-disubstituted-1,2,5-oxadiazoles and evaluation of their ability to decrease STAT3 activity. MedChemComm, 2010, 1, 156.	3.4	20
33	Design, Synthesis, and Evaluation of Carnosine Derivatives as Selective and Efficient Sequestering Agents of Cytotoxic Reactive Carbonyl Species. ChemMedChem, 2009, 4, 967-975.	3.2	55
34	Comparative modeling of the quaternary structure for the human TRPM8 channel and analysis of its binding features. Biochimica Et Biophysica Acta - Biomembranes, 2009, 1788, 973-982.	2.6	45
35	Assessing drug-likeness – what are we missing?. Drug Discovery Today, 2008, 13, 285-294.	6.4	235
36	VEGA – An open platform to develop chemo-bio-informatics applications, using plug-in architecture and script programming. Journal of Computer-Aided Molecular Design, 2004, 18, 167-173.	2.9	635

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37	Modeling of Binding Modes and Inhibition Mechanism of Some Natural Ligands of Farnesyl Transferase Using Molecular Docking. Journal of Medicinal Chemistry, 2002, 45, 1460-1465.	6.4	37
38	VEGA: a versatile program to convert, handle and visualize molecular structure on Windows-based PCs. Journal of Molecular Graphics and Modelling, 2002, 21, 47-49.	2.4	404