## Lucianna Helene Santos

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/5702755/publications.pdf

Version: 2024-02-01

30 papers 536 citations

8 h-index 752698 20 g-index

32 all docs 32 docs citations

times ranked

32

692 citing authors

#	Article	IF	CITATIONS
1	Propedia: a database for protein–peptide identification based on a hybrid clustering algorithm. BMC Bioinformatics, 2021, 22, 1.	2.6	261
2	Integrating Molecular Docking and Molecular Dynamics Simulations. Methods in Molecular Biology, 2019, 2053, 13-34.	0.9	87
3	nAPOLI: a graph-based strategy to detect and visualize conserved protein-ligand interactions in large-scale. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2019, 17, 1-1.	3.0	25
4	Computational drug design strategies applied to the modelling of human immunodeficiency virus-1 reverse transcriptase inhibitors. Memorias Do Instituto Oswaldo Cruz, 2015, 110, 847-864.	1.6	23
5	Characterization of glucose-tolerant $\hat{l}^2$ -glucosidases used in biofuel production under the bioinformatics perspective: a systematic review. Genetics and Molecular Research, 2017, 16, .	0.2	19
6	Understanding Structure–Activity Relationships for Trypanosomal Cysteine Protease Inhibitors by Simulations and Free Energy Calculations. Journal of Chemical Information and Modeling, 2019, 59, 137-148.	5.4	17
7	Introducing Programming Skills for Life Science Students. Biochemistry and Molecular Biology Education, 2019, 47, 288-295.	1.2	16
8	A Computational Method to Propose Mutations in Enzymes Based on Structural Signature Variation (SSV). International Journal of Molecular Sciences, 2019, 20, 333.	4.1	11
9	Proteingo: Motivation, user experience, and learning of molecular interactions in biological complexes. Entertainment Computing, 2019, 29, 31-42.	2.9	10
10	Glutantl̂²ase: a database for improving the rational design of glucose-tolerant l̂²-glucosidases. BMC Molecular and Cell Biology, 2020, 21, 50.	2.0	9
11	Structural insights into NS5B protein of novel equine hepaciviruses and pegiviruses complexed with polymerase inhibitors. Virus Research, 2020, 278, 197867.	2.2	8
12	Benzimidazole inhibitors of the major cysteine protease of <i>Trypanosoma brucei</i> . Future Medicinal Chemistry, 2019, 11, 1537-1551.	2.3	7
13	Antigenic and Substrate Preference Differences between Scorpion and Spider Dermonecrotic Toxins, a Comparative Investigation. Toxins, 2020, 12, 631.	3.4	7
14	Proteus: An algorithm for proposing stabilizing mutation pairs based on interactions observed in known protein 3D structures. BMC Bioinformatics, 2020, 21, 275.	2.6	6
15	Studying effects of different protonation states of His11 and His102 in ribose-5-phosphate isomerase of Trypanosoma cruzi: an example of cooperative behavior. Journal of Biomolecular Structure and Dynamics, 2020, 38, 2047-2056.	3.5	4
16	Insights into the Mechanism of Ethionamide Resistance in Mycobacterium tuberculosis through an in silico Structural Evaluation of EthA and Mutants Identified in Clinical Isolates. Catalysts, 2020, 10, 543.	3.5	4
17	Characterization of an Allosteric Pocket in Zika Virus NS2B-NS3 Protease. Journal of Chemical Information and Modeling, 2022, 62, 945-957.	5.4	4
18	Thiophenacetamide as a potential modulator to NF-κB: structure and dynamics study using in silico and molecular biology assays. Journal of Biomolecular Structure and Dynamics, 2019, 37, 4395-4406.	3.5	3

#	Article	IF	CITATIONS
19	Shared Binding Mode of Perrottetinene and Tetrahydrocannabinol Diastereomers inside the CB1 Receptor May Incentivize Novel Medicinal Drug Design: Findings from an in Silico Assay. ACS Chemical Neuroscience, 2020, 11, 4289-4300.	3.5	3
20	pH and non-covalent ligand binding modulate Zika virus NS2B/NS3 protease binding site residues: Discoveries from MD and constant pH MD simulations. Journal of Biomolecular Structure and Dynamics, 2022, 40, 10359-10372.	3.5	3
21	Bacterial 2′-Deoxyguanosine Riboswitch Classes as Potential Targets for Antibiotics: A Structure and Dynamics Study. International Journal of Molecular Sciences, 2022, 23, 1925.	4.1	3
22	Modelagem computacional de proteÃnas. , 0, , .		2
23	Profiling selectivity of chagasin mutants towards cysteine proteases cruzain or cathepsin L through molecular dynamics simulations. Journal of Biomolecular Structure and Dynamics, 2020, 39, 1-13.	3.5	1
24	Docagem molecular: em busca do encaixe perfeito e acessÃvel., 0, , .		1
25	ToxAnalyzer: A user-friendly web tool for interactive data analysis and visualization of chemical compounds from the Comparative Toxicogenomics Database (CTD)â,,¢. Computational Toxicology, 2021, 19, 100170.	3.3	1
26	Global and Local Controllability. Boletim Da Sociedade Paranaense De Matematica, 2014, 32, 27.	0.4	0
27	A Brief View of Molecular Modeling Approaches to P2 Receptors. , 0, , .		O
28	A Brief History of Bioinformatics Told by Data Visualization. Lecture Notes in Computer Science, 2020, , 235-246.	1.3	0
29	Rad5 HIRAN domain: Structural insights into its interaction with ssDNA through molecular modeling approaches. Journal of Biomolecular Structure and Dynamics, 2023, 41, 3062-3075.	3.5	O
30	Using Computers to Improve Biofuel Production. Frontiers for Young Minds, 0, 10, .	0.8	0