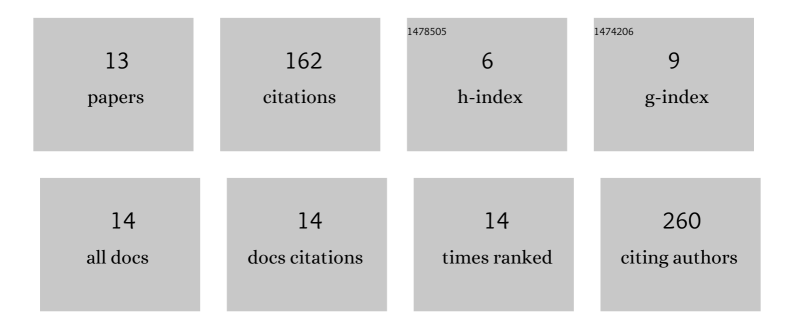
## Shintaro Minami

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/2070331/publications.pdf Version: 2024-02-01



#	Article	IF	CITATIONS
1	A prospective compound screening contest identified broader inhibitors for Sirtuin 1. Scientific Reports, 2019, 9, 19585.	3.3	15
2	Large-scale aggregation analysis of eukaryotic proteins reveals an involvement of intrinsically disordered regions in protein folding. Scientific Reports, 2018, 8, 678.	3.3	26
3	MICAN-SQ: a sequential protein structure alignment program that is applicable to monomers and all types of oligomers. Bioinformatics, 2018, 34, 3324-3331.	4.1	13
4	Characteristics of interactions at protein segments without non-local intramolecular contacts in the Protein Data Bank. PLoS ONE, 2018, 13, e0205052.	2.5	2
5	Rules for connectivity of secondary structure elements in protein: Two–layer αβ sandwiches. Protein Science, 2017, 26, 2257-2267.	7.6	2
6	An iterative compound screening contest method for identifying target protein inhibitors using the tyrosine-protein kinase Yes. Scientific Reports, 2017, 7, 12038.	3.3	28
7	Importance of consensus region of multiple-ligand templates in a virtual screening method. Biophysics and Physicobiology, 2016, 13, 149-156.	1.0	3
8	Non-sequential Structure Similarity in Proteins. Seibutsu Butsuri, 2016, 56, 027-029.	0.1	0
9	How a Spatial Arrangement of Secondary Structure Elements Is Dispersed in the Universe of Protein Folds. PLoS ONE, 2014, 9, e107959.	2.5	14
10	MICAN : a protein structure alignment algorithm that can handle Multiple-chains, Inverse alignments, C α only models, Alternative alignments, and Non-sequential alignments. BMC Bioinformatics, 2013, 14, 24.	2.6	55
11	2PT003 Modeling structure of large insersion regions in homology modeling bu utilizing non-sequential structure relationship(The 50th Annual Meeting of the Biophysical Society of Japan). Seibutsu Butsuri, 2012, 52, S120.	0.1	0
12	1E1448 Development of a novel protein structure predicion method emphasizing non-local interaction(Genome biology, Bioinformatics,The 49th Annual Meeting of the Biophysical Society of) Tj ETQq0 0 0	rgBT /Ove	erl <b>o</b> ck 10 Tf 5

13	3P292 A generalized ensemble Monte Carlo simulation for MSA to optimize a consistency-based score(Genome biology: Genome analysis,The 48th Annual Meeting of the Biophysical Society of Japan). Seibutsu Butsuri, 2010, 50, S197.	0.1	C	)	
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