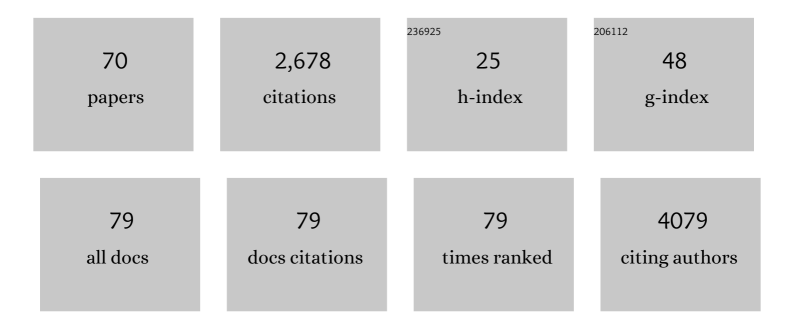
Giovanni Minervini

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	DisProt in 2022: improved quality and accessibility of protein intrinsic disorder annotation. Nucleic Acids Research, 2022, 50, D480-D487.	14.5	117
2	SARSâ€CoVâ€2 variants preferentially emerge at intrinsically disordered protein sites helping immune evasion. FEBS Journal, 2022, 289, 4240-4250.	4.7	25
3	Characterization of the pVHL Interactome in Human Testis Using High-Throughput Library Screening. Cancers, 2022, 14, 1009.	3.7	1
4	Expanding the clinical-pathological and genetic spectrum of RYR1-related congenital myopathies with cores and minicores: an Italian population study. Acta Neuropathologica Communications, 2022, 10, 54.	5.2	3
5	RING 3.0: fast generation of probabilistic residue interaction networks from structural ensembles. Nucleic Acids Research, 2022, 50, W651-W656.	14.5	75
6	Molecular Effects of Mutations in Human Genetic Diseases. International Journal of Molecular Sciences, 2022, 23, 6408.	4.1	0
7	Neurocognitive assessment and DNA sequencing expand the phenotype and genotype spectrum of Alström syndrome. American Journal of Medical Genetics, Part A, 2021, 185, 732-742.	1.2	5
8	Critical assessment of protein intrinsic disorder prediction. Nature Methods, 2021, 18, 472-481.	19.0	187
9	The f subunit of human ATP synthase is essential for normal mitochondrial morphology and permeability transition. Cell Reports, 2021, 35, 109111.	6.4	22
10	HIF1α-dependent induction of the mitochondrial chaperone TRAP1 regulates bioenergetic adaptations to hypoxia. Cell Death and Disease, 2021, 12, 434.	6.3	17
11	PED in 2021: a major update of the protein ensemble database for intrinsically disordered proteins. Nucleic Acids Research, 2021, 49, D404-D411.	14.5	95
12	Identification of Rare LRP5 Variants in a Cohort of Males with Impaired Bone Mass. International Journal of Molecular Sciences, 2021, 22, 10834.	4.1	5
13	DisProt: intrinsic protein disorder annotation in 2020. Nucleic Acids Research, 2020, 48, D269-D276.	14.5	141
14	The E3 ubiquitin-protein ligase MDM2 is a novel interactor of the von Hippel–Lindau tumor suppressor. Scientific Reports, 2020, 10, 15850.	3.3	2
15	The pVHL neglected functions, a tale of hypoxia-dependent and -independent regulations in cancer. Open Biology, 2020, 10, 200109.	3.6	14
16	Assessing predictors for new post translational modification sites: AÂcase study on hydroxylation. PLoS Computational Biology, 2020, 16, e1007967.	3.2	10
17	Novel Missense Variant in <i>MYL2</i> Gene Associated With Hypertrophic Cardiomyopathy Showing High Incidence of Restrictive Physiology. Circulation Genomic and Precision Medicine, 2020, 13, e002824.	3.6	6
18	The role of mitochondrial ATP synthase in cancer. Biological Chemistry, 2020, 401, 1199-1214.	2.5	29

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19	Insights into the molecular features of the von Hippel–Lindau-like protein. Amino Acids, 2019, 51, 1461-1474.	2.7	4
20	Arg-8 of yeast subunit e contributes to the stability of F-ATP synthase dimers and to the generation of the full-conductance mitochondrial megachannel. Journal of Biological Chemistry, 2019, 294, 10987-10997.	3.4	32
21	Genotype-phenotype relations of the von Hippel-Lindau tumor suppressor inferred from a large-scale analysis of disease mutations and interactors. PLoS Computational Biology, 2019, 15, e1006478.	3.2	24
22	A targeted next-generation gene panel reveals a novel heterozygous nonsense variant in the TP63 gene in patients with arrhythmogenic cardiomyopathy. Heart Rhythm, 2019, 16, 773-780.	0.7	15
23	The lipoprotein <scp>HP1454</scp> of <i>Helicobacter pylori</i> regulates <scp>T</scp> â€cell response by shaping <scp>T</scp> â€cell receptor signalling. Cellular Microbiology, 2019, 21, e13006.	2.1	27
24	Pore formation by yeast mitochondrial ATP synthase involves subunits e, g and b. Biochimica Et Biophysica Acta - Bioenergetics, 2018, 1859, e16-e17.	1.0	0
25	Role of F-ATP synthase f subunit in dimer formation and PTP modulation. Biochimica Et Biophysica Acta - Bioenergetics, 2018, 1859, e110.	1.0	0
26	High-Conductance Channel Formation in Yeast Mitochondria is Mediated by F-ATP Synthase e and g Subunits. Cellular Physiology and Biochemistry, 2018, 50, 1840-1855.	1.6	57
27	The clinical spectrum of CASQ1-related myopathy. Neurology, 2018, 91, e1629-e1641.	1.1	14
28	Arginine 107 of yeast ATP synthase subunit g mediates sensitivity of the mitochondrial permeability transition to phenylglyoxal. Journal of Biological Chemistry, 2018, 293, 14632-14645.	3.4	40
29	Calmodulin Enhances Cryptochrome Binding to INAD in Drosophila Photoreceptors. Frontiers in Molecular Neuroscience, 2018, 11, 280.	2.9	15
30	Whole-Exome Sequencing Identifies Pathogenic Variants in <i>TJP1</i> Gene Associated With Arrhythmogenic Cardiomyopathy. Circulation Genomic and Precision Medicine, 2018, 11, e002123.	3.6	38
31	Novel interactions of the von Hippel-Lindau (pVHL) tumor suppressor with the CDKN1 family of cell cycle inhibitors. Scientific Reports, 2017, 7, 46562.	3.3	6
32	Performance of in silico tools for the evaluation of p16INK4a (CDKN2A) variants in CAGI. Human Mutation, 2017, 38, 1042-1050.	2.5	13
33	FELLS: fast estimator of latent local structure. Bioinformatics, 2017, 33, 1889-1891.	4.1	72
34	DisProt 7.0: a major update of the database of disordered proteins. Nucleic Acids Research, 2017, 45, D219-D227.	14.5	242
35	Ca ²⁺ binding to Fâ€ATP synthase β subunit triggers the mitochondrial permeability transition. EMBO Reports, 2017, 18, 1065-1076.	4.5	170
36	<i>In silico</i> study of the structure and function of <i>Streptococcus mutans</i> plasmidic proteins. Bio-Algorithms and Med-Systems, 2017, 13, 51-61.	2.4	0

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37	Mapping pathogenic mutations suggests an innovative structural model for the pendrin (SLC26A4) transmembrane domain. Biochimie, 2017, 132, 109-120.	2.6	19
38	VHLdb: A database of von Hippel-Lindau protein interactors and mutations. Scientific Reports, 2016, 6, 31128.	3.3	36
39	Computational analysis of prolyl hydroxylase domain-containing protein 2 (PHD2) mutations promoting polycythemia insurgence in humans. Scientific Reports, 2016, 6, 18716.	3.3	8
40	NaÃ ⁻ ve Bayes ant colony optimization for designing high dimensional experiments. Applied Soft Computing Journal, 2016, 49, 259-268.	7.2	6
41	The Ca 2+ regulatory site of the permeability transition pore is within the catalytic core of F-ATP synthase. Biochimica Et Biophysica Acta - Bioenergetics, 2016, 1857, e65-e66.	1.0	1
42	The RING 2.0 web server for high quality residue interaction networks. Nucleic Acids Research, 2016, 44, W367-W374.	14.5	369
43	Assessment of phenolic herbicide toxicity and mode of action by different assays. Environmental Science and Pollution Research, 2016, 23, 7398-7408.	5.3	27
44	lsoform-specific interactions of the von Hippel-Lindau tumor suppressor protein. Scientific Reports, 2015, 5, 12605.	3.3	26
45	Structural in silico dissection of the collagen V interactome to identify genotype–phenotype correlations in classic Ehlers–Danlos Syndrome (EDS). FEBS Letters, 2015, 589, 3871-3878.	2.8	11
46	Heterozygous Reelin Mutations Cause Autosomal-Dominant Lateral Temporal Epilepsy. American Journal of Human Genetics, 2015, 96, 992-1000.	6.2	94
47	Structural protein reorganization and fold emergence investigated through amino acid sequence permutations. Amino Acids, 2015, 47, 147-152.	2.7	3
48	BOOGIE: Predicting Blood Groups from High Throughput Sequencing Data. PLoS ONE, 2015, 10, e0124579.	2.5	31
49	Insights into the proline hydroxylase (PHD) family, molecular evolution and its impact on human health. Biochimie, 2015, 116, 114-124.	2.6	17
50	Design and Analysis of a Petri Net Model of the Von Hippel-Lindau (VHL) Tumor Suppressor Interaction Network. PLoS ONE, 2014, 9, e96986.	2.5	18
51	Evaluation of the steric impact of flavin adenine dinucleotide in Drosophila melanogaster cryptochrome function. Biochemical and Biophysical Research Communications, 2014, 450, 1606-1611.	2.1	13
52	<i>CDKN2A</i> Unclassified Variants in Familial Malignant Melanoma: Combining Functional and Computational Approaches for Their Assessment. Human Mutation, 2014, 35, 828-840.	2.5	17
53	RepeatsDB: a database of tandem repeat protein structures. Nucleic Acids Research, 2014, 42, D352-D357.	14.5	53
54	In silico investigation of PHDâ€3 specific HIF1â€Î± proline 567 hydroxylation: A new player in the VHL/HIFâ€1α interaction pathway?. FEBS Letters, 2013, 587, 2996-3001.	2.8	11

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55	RAPHAEL: recognition, periodicity and insertion assignment of solenoid protein structures. Bioinformatics, 2012, 28, 3257-3264.	4.1	27
56	Bluues server: electrostatic properties of wild-type and mutated protein structures. Bioinformatics, 2012, 28, 2189-2190.	4.1	72
57	Do Natural Proteins Differ from Random Sequences Polypeptides? Natural vs. Random Proteins Classification Using an Evolutionary Neural Network. PLoS ONE, 2012, 7, e36634.	2.5	25
58	Selection Dynamic of <i>Escherichia coli</i> Host in M13 Combinatorial Peptide Phage Display Libraries. Bioscience, Biotechnology and Biochemistry, 2011, 75, 812-815.	1.3	8
59	The invasive Manila clam Ruditapes philippinarum (Adams and Reeve, 1850) in Northern Adriatic Sea: Population genetics assessed by an integrated molecular approach. Fisheries Research, 2011, 110, 259-267.	1.7	25
60	Design and dynamic simulation of minimal metallo-proteins. Journal of Molecular Modeling, 2011, 17, 2919-2925.	1.8	1
61	Probing mammalian spermine oxidase enzyme–substrate complex through molecular modeling, site-directed mutagenesis and biochemical characterization. Amino Acids, 2011, 40, 1115-1126.	2.7	35
62	Validation of a new tool for seafood safety and traceability: the case of Manila clam Ruditapes philippinarum. Economia Agro-Alimentare, 2011, , 503-507.	0.5	3
63	Health state of mollusc productive sites: Biochemical, physiological and molecular markers. Comparative Biochemistry and Physiology Part A, Molecular & Integrative Physiology, 2010, 157, S38.	1.8	0
64	A Model Based Ant Colony Design for the Protein Engineering Problem. Lecture Notes in Computer Science, 2010, , 352-359.	1.3	2
65	Massive non-natural proteins structure prediction using grid technologies. BMC Bioinformatics, 2009, 10, S22.	2.6	13
66	<i>In silico</i> Structural Study of Random Amino Acid Sequence Proteins Not Present in Nature. Chemistry and Biodiversity, 2009, 6, 2311-2336.	2.1	10
67	Porting Biological Applications in Grid: An Experience within the EUChinaGRID Framework. , 2009, , 67-71.		0
68	Human haptoglobin structure and function – a molecular modelling study. FEBS Journal, 2008, 275, 5648-5656.	4.7	78
69	Never born proteins as a test case for ab initio protein structures prediction. Bioinformation, 2008, 3, 177-179.	0.5	6
70	Lys300 Plays a Major Role in the Catalytic Mechanism of Maize Polyamine Oxidaseâ€. Biochemistry, 2005, 44, 16108-16120.	2.5	48