Zsuzsanna Dosztanyi

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/4131183/publications.pdf

Version: 2024-02-01

65 papers 11,448 citations

38 h-index 62 g-index

71 all docs

71 docs citations

times ranked

71

15239 citing authors

#	Article	IF	CITATIONS
1	IUPred: web server for the prediction of intrinsically unstructured regions of proteins based on estimated energy content. Bioinformatics, 2005, 21, 3433-3434.	4.1	1,832
2	InterPro in 2017â€"beyond protein family and domain annotations. Nucleic Acids Research, 2017, 45, D190-D199.	14.5	1,358
3	IUPred2A: context-dependent prediction of protein disorder as a function of redox state and protein binding. Nucleic Acids Research, 2018, 46, W329-W337.	14.5	1,080
4	The Pairwise Energy Content Estimated from Amino Acid Composition Discriminates between Folded and Intrinsically Unstructured Proteins. Journal of Molecular Biology, 2005, 347, 827-839.	4.2	911
5	D2P2: database of disordered protein predictions. Nucleic Acids Research, 2012, 41, D508-D516.	14.5	570
6	ANCHOR: web server for predicting protein binding regions in disordered proteins. Bioinformatics, 2009, 25, 2745-2746.	4.1	527
7	Prediction of Protein Binding Regions in Disordered Proteins. PLoS Computational Biology, 2009, 5, e1000376.	3.2	523
8	Disorder and Sequence Repeats in Hub Proteins and Their Implications for Network Evolution. Journal of Proteome Research, 2006, 5, 2985-2995.	3.7	312
9	IUPred3: prediction of protein disorder enhanced with unambiguous experimental annotation and visualization of evolutionary conservation. Nucleic Acids Research, 2021, 49, W297-W303.	14.5	258
10	PDB_TM: selection and membrane localization of transmembrane proteins in the protein data bank. Nucleic Acids Research, 2004, 33, D275-D278.	14.5	245
11	Molecular Principles of the Interactions of Disordered Proteins. Journal of Molecular Biology, 2007, 372, 549-561.	4.2	242
12	DisProt 7.0: a major update of the database of disordered proteins. Nucleic Acids Research, 2017, 45, D219-D227.	14.5	242
13	Analyzing Protein Disorder with IUPred2A. Current Protocols in Bioinformatics, 2020, 70, e99.	25.8	233
14	Transmembrane proteins in the Protein Data Bank: identification and classification. Bioinformatics, 2004, 20, 2964-2972.	4.1	216
15	MobiDB 3.0: more annotations for intrinsic disorder, conformational diversity and interactions in proteins. Nucleic Acids Research, 2018, 46, D471-D476.	14.5	190
16	MobiDB: intrinsically disordered proteins in 2021. Nucleic Acids Research, 2021, 49, D361-D367.	14.5	183
17	Prediction of protein disorder based on IUPred. Protein Science, 2018, 27, 331-340.	7.6	164
18	MobiDB-lite: fast and highly specific consensus prediction of intrinsic disorder in proteins. Bioinformatics, 2017, 33, 1402-1404.	4.1	161

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19	Stabilization centers in proteins:Identification, characterization and predictions. Journal of Molecular Biology, 1997, 272, 597-612.	4.2	144
20	DisProt: intrinsic protein disorder annotation in 2020. Nucleic Acids Research, 2020, 48, D269-D276.	14.5	141
21	TMDET: web server for detecting transmembrane regions of proteins by using their 3D coordinates. Bioinformatics, 2005, 21, 1276-1277.	4.1	132
22	Prevalent Structural Disorder in E. coliand S. cerevisia e Proteomes. Journal of Proteome Research, 2006, 5, 1996-2000.	3.7	119
23	DisProt in 2022: improved quality and accessibility of protein intrinsic disorder annotation. Nucleic Acids Research, 2022, 50, D480-D487.	14.5	117
24	Simultaneous quantification of protein order and disorder. Nature Chemical Biology, 2017, 13, 339-342.	8.0	113
25	Bioinformatical approaches to characterize intrinsically disordered/unstructured proteins. Briefings in Bioinformatics, 2010, 11, 225-243.	6.5	107
26	Degrons in cancer. Science Signaling, 2017, 10, .	3.6	100
27	PhaSePro: the database of proteins driving liquid–liquid phase separation. Nucleic Acids Research, 2020, 48, D360-D367.	14.5	100
28	SCide: identification of stabilization centers in proteins. Bioinformatics, 2003, 19, 899-900.	4.1	99
29	DIBS: a repository of disordered binding sites mediating interactions with ordered proteins. Bioinformatics, 2018, 34, 535-537.	4.1	72
30	Prediction of Protein Disorder at the Domain Level. Current Protein and Peptide Science, 2007, 8, 161-171.	1.4	71
31	PlaToLoCo: the first web meta-server for visualization and annotation of low complexity regions in proteins. Nucleic Acids Research, 2020, 48, W77-W84.	14.5	71
32	Disentangling the complexity of low complexity proteins. Briefings in Bioinformatics, 2020, 21, 458-472.	6.5	70
33	Disordered Binding Regions and Linear Motifs—Bridging the Gap between Two Models of Molecular Recognition. PLoS ONE, 2012, 7, e46829.	2.5	60
34	Systematic discovery of linear binding motifs targeting an ancient protein interaction surface on <scp>MAP</scp> kinases. Molecular Systems Biology, 2015, 11, 837.	7.2	60
35	Prediction of Protein Disorder. Methods in Molecular Biology, 2008, 426, 103-115.	0.9	56
36	The expanding view of protein–protein interactions: complexes involving intrinsically disordered proteins. Physical Biology, 2011, 8, 035003.	1.8	55

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37	A comprehensive assessment of long intrinsic protein disorder from the DisProt database. Bioinformatics, 2018, 34, 445-452.	4.1	53
38	MobiDB-lite 3.0: fast consensus annotation of intrinsic disorder flavors in proteins. Bioinformatics, 2021, 36, 5533-5534.	4.1	47
39	Is there a biological cost of protein disorder? Analysis of cancer-associated mutations. Molecular BioSystems, 2012, 8, 296-307.	2.9	43
40	Substrate Preference of Transglutaminase 2 Revealed by Logistic Regression Analysis and Intrinsic Disorder Examination. Journal of Molecular Biology, 2008, 383, 390-402.	4.2	35
41	Mutations of Intrinsically Disordered Protein Regions Can Drive Cancer but Lack Therapeutic Strategies. Biomolecules, 2021, 11, 381.	4.0	26
42	Amino acid similarity matrices based on force fields. Bioinformatics, 2001, 17, 686-699.	4.1	22
43	Proteins with Complex Architecture as Potential Targets for Drug Design: A Case Study of Mycobacterium tuberculosis. PLoS Computational Biology, 2011, 7, e1002118.	3.2	21
44	Integron-associated Mobile Gene Cassettes Code for Folded Proteins: The Structure of Bal32a, a New Member of the Adaptable $\hat{l}_{\pm}+\hat{l}^{2}$ Barrel Family. Journal of Molecular Biology, 2005, 346, 1229-1241.	4.2	20
45	Towards Proteomic Approaches for the Identification of Structural Disorder. Current Protein and Peptide Science, 2007, 8, 173-179.	1.4	20
46	Novel linear motif filtering protocol reveals the role of the LC8 dynein light chain in the Hippo pathway. PLoS Computational Biology, 2017, 13, e1005885.	3.2	20
47	Largeâ€Scale Analysis of Redoxâ€Sensitive Conditionally Disordered Protein Regions Reveals Their Widespread Nature and Key Roles in Highâ€Level Eukaryotic Processes. Proteomics, 2019, 19, e1800070.	2.2	17
48	Evolutionary Study of Disorder in Protein Sequences. Biomolecules, 2020, 10, 1413.	4.0	16
49	Systematic analysis of somatic mutations driving cancer: uncovering functional protein regions in disease development. Biology Direct, 2016, 11, 23.	4.6	15
50	Servers for sequence-structure relationship analysis and prediction. Nucleic Acids Research, 2003, 31, 3359-3363.	14.5	12
51	Sequential, Structural and FunctionalÂProperties of Protein Complexes Are Defined by How Folding and Binding Intertwine. Journal of Molecular Biology, 2019, 431, 4408-4428.	4.2	12
52	An intrinsically disordered proteins community for ELIXIR. F1000Research, 2019, 8, 1753.	1.6	12
53	Ancient Evolutionary Origin of Intrinsically Disordered Cancer Risk Regions. Biomolecules, 2020, 10, 1115.	4.0	11
54	Functions of intrinsically disordered proteins through evolutionary lenses. Progress in Molecular Biology and Translational Science, 2021, 183, 45-74.	1.7	10

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55	Prediction and Analysis of Intrinsically Disordered Proteins. Methods in Molecular Biology, 2015, 1261, 35-59.	0.9	9
56	Stabilization centers in various proteins. Theoretical Chemistry Accounts, 1999, 101, 27-32.	1.4	8
57	The N-Terminal Intrinsically Disordered Domain of Mgm101p Is Localized to the Mitochondrial Nucleoid. PLoS ONE, 2013, 8, e56465.	2.5	8
58	The MemMoRF database for recognizing disordered protein regions interacting with cellular membranes. Nucleic Acids Research, 2021, 49, D355-D360.	14.5	8
59	Noncovalent Cross-links in Context with Other Structural and Functional Elements of Proteinsâ€. Journal of Chemical Information and Computer Sciences, 2004, 44, 347-351.	2.8	7
60	The role of long-range interactions in defining the secondary structure of proteins is overestimated. Bioinformatics, 1997, 13, 297-301.	4.1	6
61	Bioinformatics Approaches to the Structure and Function of Intrinsically Disordered Proteins. , 2017, , 167-203.		5
62	Bioinformatical Approaches to Unstructured/Disordered Proteins and Their Interactions. Springer Series in Bio-/neuroinformatics, 2014, , 525-556.	0.1	1
63	Noncovalent Cross-Links in Context with Other Structural and Functional Elements of Proteins ChemInform, 2004, 35, no.	0.0	0
64	Bioinformatical Approaches to Unstructured/Disordered Proteins and Their Complexes. Springer Series on Bio- and Neurosystems, 2019, , 561-596.	0.2	0
65	Interresidue interactions in protein classes. Proteins: Structure, Function and Bioinformatics, 1997, 27, 360-366.	2.6	O