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List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Computational Prediction of N- and O-Linked Glycosylation Sites for Human and Mouse Proteins. Methods in Molecular Biology, 2022, , 177-186.	0.9	1
2	SEMal: Accurate protein malonylation site predictor using structural and evolutionary information. Computers in Biology and Medicine, 2020, 125, 104022.	7.0	10
3	Accurately Predicting Glutarylation Sites Using Sequential Bi-Peptide-Based Evolutionary Features. Genes, 2020, 11, 1023.	2.4	17
4	Accurate prediction of RNA 5-hydroxymethylcytosine modification by utilizing novel position-specific gapped k-mer descriptors. Computational and Structural Biotechnology Journal, 2020, 18, 3528-3538.	4.1	11
5	Mal-Light: Enhancing Lysine Malonylation Sites Prediction Problem Using Evolutionary-based Features. IEEE Access, 2020, 8, 77888-77902.	4.2	12
6	Predicting protein-peptide binding sites with a deep convolutional neural network. Journal of Theoretical Biology, 2020, 496, 110278.	1.7	25
7	SPRINT-Gly: predicting <i>N-</i> and <i>O-</i> linked glycosylation sites of human and mouse proteins by using sequence and predicted structural properties. Bioinformatics, 2019, 35, 4140-4146.	4.1	48
8	Structure-based prediction of protein– peptide binding regions using Random Forest. Bioinformatics, 2018, 34, 477-484.	4.1	62
9	SumSec: Accurate Prediction of Sumoylation Sites Using Predicted Secondary Structure. Molecules, 2018, 23, 3260.	3.8	13
10	Predicting lysineâ€malonylation sites of proteins using sequence and predicted structural features. Journal of Computational Chemistry, 2018, 39, 1757-1763.	3.3	30
11	Improving succinylation prediction accuracy by incorporating the secondary structure via helix, strand and coil, and evolutionary information from profile bigrams. PLoS ONE, 2018, 13, e0191900.	2.5	51
12	PSSM-Suc: Accurately predicting succinylation using position specific scoring matrix into bigram for feature extraction. Journal of Theoretical Biology, 2017, 425, 97-102.	1.7	65
13	SucStruct: Prediction of succinylated lysine residues by using structural properties of amino acids. Analytical Biochemistry, 2017, 527, 24-32.	2.4	55
14	Sequenceâ€based prediction of protein–peptide binding sites using support vector machine. Journal of Computational Chemistry, 2016, 37, 1223-1229.	3.3	81
15	Sequence-Based Prediction of Protein–Carbohydrate Binding Sites Using Support Vector Machines. Journal of Chemical Information and Modeling, 2016, 56, 2115-2122.	5.4	59