Yu-Yen Ou

List of Publications by Year in Descending Order

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The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

54	1,066	19	32
papers	citations	h-index	g-index
59	1,372 ext. citations	4.3	5.06
ext. papers		avg, IF	L-index

#	Paper	IF	Citations
54	Using k-mer embeddings learned from a Skip-gram based neural network for building a cross-species DNA N6-methyladenine site prediction model. <i>Plant Molecular Biology</i> , 2021 , 107, 533-542	₂ 4.6	1
53	Identification of efflux proteins based on contextual representations with deep bidirectional transformer encoders. <i>Analytical Biochemistry</i> , 2021 , 633, 114416	3.1	
52	Incorporating a transfer learning technique with amino acid embeddings to efficiently predict N-linked glycosylation sites in ion channels. <i>Computers in Biology and Medicine</i> , 2021 , 130, 104212	7	2
51	FAD-BERT: Improved prediction of FAD binding sites using pre-training of deep bidirectional transformers. <i>Computers in Biology and Medicine</i> , 2021 , 131, 104258	7	8
50	GT-Finder: Classify the family of glucose transporters with pre-trained BERT language models. <i>Computers in Biology and Medicine</i> , 2021 , 131, 104259	7	5
49	Addressing data imbalance problems in ligand-binding site prediction using a variational autoencoder and a convolutional neural network. <i>Briefings in Bioinformatics</i> , 2021 , 22,	13.4	2
48	A transformer architecture based on BERT and 2D convolutional neural network to identify DNA enhancers from sequence information. <i>Briefings in Bioinformatics</i> , 2021 , 22,	13.4	28
47	DeepSIRT: A deep neural network for identification of sirtuin targets and their subcellular localizations. <i>Computational Biology and Chemistry</i> , 2021 , 93, 107514	3.6	2
46	ActTRANS: Functional classification in active transport proteins based on transfer learning and contextual representations. <i>Computational Biology and Chemistry</i> , 2021 , 93, 107537	3.6	2
45	mCNN-ETC: identifying electron transporters and their functional families by using multiple windows scanning techniques in convolutional neural networks with evolutionary information of protein sequences. <i>Briefings in Bioinformatics</i> , 2021,	13.4	1
44	TRP-BERT: Discrimination of transient receptor potential (TRP) channels using contextual representations from deep bidirectional transformer based on BERT. <i>Computers in Biology and Medicine</i> , 2021 , 137, 104821	7	3
43	TNFPred: identifying tumor necrosis factors using hybrid features based on word embeddings. <i>BMC Medical Genomics</i> , 2020 , 13, 155	3.7	3
42	Advantages and Constraints of a Hybrid Model K-12 E-Learning Assistant Chatbot. <i>IEEE Access</i> , 2020 , 8, 77788-77801	3.5	26
41	Using Language Representation Learning Approach to Efficiently Identify Protein Complex Categories in Electron Transport Chain. <i>Molecular Informatics</i> , 2020 , 39, e2000033	3.8	1
40	A Self-Relevant CNN-SVM Model for Problem Classification in K-12 Question-Driven Learning. <i>IEEE Access</i> , 2020 , 8, 225822-225830	3.5	O
39	DeepETC: A deep convolutional neural network architecture for investigating and classifying electron transport chain's complexes. <i>Neurocomputing</i> , 2020 , 375, 71-79	5.4	27
38	Use Chou's 5-steps rule with different word embedding types to boost performance of electron transport protein prediction model. <i>IEEE/ACM Transactions on Computational Biology and Bioinformatics</i> , 2020 , PP,	3	7

37	Using two-dimensional convolutional neural networks for identifying GTP binding sites in Rab proteins. <i>Journal of Bioinformatics and Computational Biology</i> , 2019 , 17, 1950005	1	7
36	Using word embedding technique to efficiently represent protein sequences for identifying substrate specificities of transporters. <i>Analytical Biochemistry</i> , 2019 , 577, 73-81	3.1	15
35	Deeplon: Deep learning approach for classifying ion transporters and ion channels from membrane proteins. <i>Journal of Computational Chemistry</i> , 2019 , 40, 1521-1529	3.5	7
34	iMotor-CNN: Identifying molecular functions of cytoskeleton motor proteins using 2D convolutional neural network via Chou & 5-step rule. <i>Analytical Biochemistry</i> , 2019 , 575, 17-26	3.1	43
33	Prediction of ATP-binding sites in membrane proteins using a two-dimensional convolutional neural network. <i>Journal of Molecular Graphics and Modelling</i> , 2019 , 92, 86-93	2.8	10
32	iEnhancer-5Step: Identifying enhancers using hidden information of DNA sequences via Chou ∜ 5-step rule and word embedding. <i>Analytical Biochemistry</i> , 2019 , 571, 53-61	3.1	76
31	DeepEfflux: a 2D convolutional neural network model for identifying families of efflux proteins in transporters. <i>Bioinformatics</i> , 2018 , 34, 3111-3117	7.2	24
30	Classifying the molecular functions of Rab GTPases in membrane trafficking using deep convolutional neural networks. <i>Analytical Biochemistry</i> , 2018 , 555, 33-41	3.1	46
29	Incorporating post translational modification information for enhancing the predictive performance of membrane transport proteins. <i>Computational Biology and Chemistry</i> , 2018 , 77, 251-260	3.6	10
28	Identifying the molecular functions of electron transport proteins using radial basis function networks and biochemical properties. <i>Journal of Molecular Graphics and Modelling</i> , 2017 , 73, 166-178	2.8	26
27	Incorporating deep learning with convolutional neural networks and position specific scoring matrices for identifying electron transport proteins. <i>Journal of Computational Chemistry</i> , 2017 , 38, 2000) - 2006	65
26	Incorporating efficient radial basis function networks and significant amino acid pairs for predicting GTP binding sites in transport proteins. <i>BMC Bioinformatics</i> , 2016 , 17, 501	3.6	18
25	Using Deep Learning with Position Specific Scoring Matrices to Identify Efflux Proteins in Membrane and Transport Proteins 2016 ,		1
24	Prediction of FAD binding sites in electron transport proteins according to efficient radial basis function networks and significant amino acid pairs. <i>BMC Bioinformatics</i> , 2016 , 17, 298	3.6	41
23	Bioinformatics approaches for functional annotation of membrane proteins. <i>Briefings in Bioinformatics</i> , 2014 , 15, 155-68	13.4	29
22	Identification of efflux proteins using efficient radial basis function networks with position-specific scoring matrices and biochemical properties. <i>Proteins: Structure, Function and Bioinformatics</i> , 2013 , 81, 1634-43	4.2	9
21	ETMB-RBF: discrimination of metal-binding sites in electron transporters based on RBF networks with PSSM profiles and significant amino acid pairs. <i>PLoS ONE</i> , 2013 , 8, e46572	3.7	7
20	Predicting Protein Metal Binding Sites with RBF Networks based on PSSM Profiles and Additional Properties. <i>Current Bioinformatics</i> , 2012 , 7, 180-186	4.7	2

19	Prediction of transporter targets using efficient RBF networks with PSSM profiles and biochemical properties. <i>Bioinformatics</i> , 2011 , 27, 2062-7	7.2	61
18	Incorporating distant sequence features and radial basis function networks to identify ubiquitin conjugation sites. <i>PLoS ONE</i> , 2011 , 6, e17331	3.7	58
17	Structural and Functional Discrimination of Membrane Proteins 2011 , 1-32		
16	Incorporating significant amino acid pairs to identify O-linked glycosylation sites on transmembrane proteins and non-transmembrane proteins. <i>BMC Bioinformatics</i> , 2010 , 11, 536	3.6	29
15	Prediction of membrane spanning segments and topology in beta-barrel membrane proteins at better accuracy. <i>Journal of Computational Chemistry</i> , 2010 , 31, 217-23	3.5	36
14	Classification of transporters using efficient radial basis function networks with position-specific scoring matrices and biochemical properties. <i>Proteins: Structure, Function and Bioinformatics</i> , 2010 , 78, 1789-97	4.2	49
13	Topology Prediction of EHelical and EBarrel Transmembrane Proteins Using RBF Networks. <i>Lecture Notes in Computer Science</i> , 2010 , 642-649	0.9	2
12	Using Efficient RBF Networks to Classify Transport Proteins Based on PSSM Profiles and Biochemical Properties. <i>Lecture Notes in Computer Science</i> , 2009 , 869-876	0.9	2
11	Prediction of protein secondary structures with a novel kernel density estimation based classifier. <i>BMC Research Notes</i> , 2008 , 1, 51	2.3	4
10	TMBETADISC-RBF: Discrimination of beta-barrel membrane proteins using RBF networks and PSSM profiles. <i>Computational Biology and Chemistry</i> , 2008 , 32, 227-31	3.6	67
9	Using Efficient RBF Network to Identify Interface Residues Based on PSSM Profiles and Biochemical Properties. <i>Lecture Notes in Computer Science</i> , 2007 , 132-141	0.9	1
8	Enhancing Protein Disorder Detection by Refined Secondary Structure Prediction 2007 , 395-409		1
7	Protein disorder prediction by condensed PSSM considering propensity for order or disorder. <i>BMC Bioinformatics</i> , 2006 , 7, 319	3.6	78
6	Expediting Model Selection for Support Vector Machines Based on an Advanced Data Reduction Algorithm. <i>Lecture Notes in Computer Science</i> , 2006 , 1017-1021	0.9	2
5	Expediting Model Selection for Support Vector Machines Based on an Advanced Data Reduction Algorithm 2006 , 1017-1021		
4	Data classification with radial basis function networks based on a novel kernel density estimation algorithm. <i>IEEE Transactions on Neural Networks</i> , 2005 , 16, 225-36		115
3	A novel radial basis function network classifier with centers set by hierarchical clustering		2
2	Expediting model selection for Support Vector Machines based on data reduction		2

TNFPred: Identifying tumor necrosis factors using hybrid features based on word embeddings

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