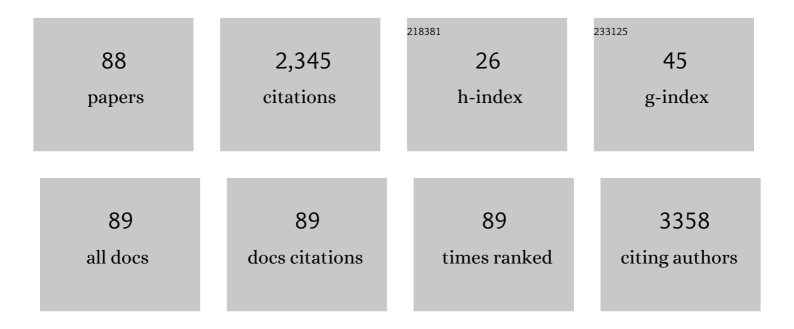
## Ting-Yi Sung

List of Publications by Year in descending order

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TINC-YI SUNC

#	Article	IF	CITATIONS
1	Calibr improves spectral library search for spectrum-centric analysis of data independent acquisition proteomics. Scientific Reports, 2022, 12, 2045.	1.6	7
2	Comparison of different variant sequence types coupled with decoy generation methods used in concatenated target-decoy database searches for proteogenomic research. Journal of Proteomics, 2021, 231, 104021.	1.2	2
3	Multi-Q 2 software facilitates isobaric labeling quantitation analysis with improved accuracy and coverage. Scientific Reports, 2021, 11, 2233.	1.6	3
4	A data-independent acquisition-based global phosphoproteomics system enables deep profiling. Nature Communications, 2021, 12, 2539.	5.8	44
5	Somatic mutation subtypes of lung adenocarcinoma in East Asian reveal divergent biological characteristics and therapeutic vulnerabilities. IScience, 2021, 24, 102522.	1.9	4
6	MinProtMaxVP: Generating a minimized number of protein variant sequences containing all possible variant peptides for proteogenomic analysis. Journal of Proteomics, 2020, 223, 103819.	1.2	6
7	Proteogenomics of Non-smoking Lung Cancer in East Asia Delineates Molecular Signatures of Pathogenesis and Progression. Cell, 2020, 182, 226-244.e17.	13.5	178
8	iHPDM: In Silico Human Proteome Digestion Map with Proteolytic Peptide Analysis and Graphical Visualizations. Journal of Proteome Research, 2019, 18, 4124-4132.	1.8	2
9	WinProphet: A User-Friendly Pipeline Management System for Proteomics Data Analysis Based on Trans-Proteomic Pipeline. Analytical Chemistry, 2019, 91, 9403-9406.	3.2	5
10	N-GlyDE: a two-stage N-linked glycosylation site prediction incorporating gapped dipeptides and pattern-based encoding. Scientific Reports, 2019, 9, 15975.	1.6	46
11	Subcellular Proteome Landscape of Human Embryonic Stem Cells Revealed Missing Membrane Proteins. Journal of Proteome Research, 2018, 17, 4138-4151.	1.8	19
12	Evaluating the Possibility of Detecting Variants in Shotgun Proteomics via LeTE-Fusion Analysis Pipeline. Journal of Proteome Research, 2018, 17, 2937-2952.	1.8	5
13	Phosphoproteomics Reveals HMGA1, a CK2 Substrate, as a Drug-Resistant Target in Non-Small Cell Lung Cancer. Scientific Reports, 2017, 7, 44021.	1.6	31
14	Decoding the Effect of Isobaric Substitutions on Identifying Missing Proteins and Variant Peptides in Human Proteome. Journal of Proteome Research, 2017, 16, 4415-4424.	1.8	8
15	iTop-Q: an Intelligent Tool for Top-down Proteomics Quantitation Using DYAMOND Algorithm. Analytical Chemistry, 2017, 89, 13128-13136.	3.2	2
16	MAGIC-web: a platform for untargeted and targeted N-linked glycoprotein identification. Nucleic Acids Research, 2016, 44, W575-W580.	6.5	13
17	iMet-Q: A User-Friendly Tool for Label-Free Metabolomics Quantitation Using Dynamic Peak-Width Determination. PLoS ONE, 2016, 11, e0146112.	1.1	17
18	Metabolite Identification for Mass Spectrometry-Based Metabolomics Using Multiple Types of Correlated Ion Information. Analytical Chemistry, 2015, 87, 2143-2151.	3.2	65

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19	MAGIC: An Automated N-Linked Glycoprotein Identification Tool Using a Y1-Ion Pattern Matching Algorithm and <i>in Silico</i> MS <sup>2</sup> Approach. Analytical Chemistry, 2015, 87, 2466-2473.	3.2	67
20	Mining Missing Membrane Proteins by High-pH Reverse-Phase StageTip Fractionation and Multiple Reaction Monitoring Mass Spectrometry. Journal of Proteome Research, 2015, 14, 3658-3669.	1.8	24
21	Quest for Missing Proteins: Update 2015 on Chromosome-Centric Human Proteome Project. Journal of Proteome Research, 2015, 14, 3415-3431.	1.8	53
22	Rapid High-pH Reverse Phase StageTip for Sensitive Small-Scale Membrane Proteomic Profiling. Analytical Chemistry, 2015, 87, 12016-12023.	3.2	47
23	Informatics View on the Challenges of Identifying Missing Proteins from Shotgun Proteomics. Journal of Proteome Research, 2015, 14, 5396-5407.	1.8	14
24	Sequential Phosphoproteomic Enrichment through Complementary Metal-Directed Immobilized Metal Ion Affinity Chromatography. Analytical Chemistry, 2014, 86, 685-693.	3.2	100
25	Chromosome-centric Human Proteome Project (C-HPP): Chromosome 12. Journal of Proteome Research, 2014, 13, 3160-3165.	1.8	4
26	Decoding the Disease-Associated Proteins Encoded in the Human Chromosome 4. Journal of Proteome Research, 2013, 12, 33-44.	1.8	9
27	Lipid exposure prediction enhances the inference of rotational angles of transmembrane helices. BMC Bioinformatics, 2013, 14, 304.	1.2	13
28	Spectrum-based Method to Generate Good Decoy Libraries for Spectral Library Searching in Peptide Identifications. Journal of Proteome Research, 2013, 12, 2305-2310.	1.8	24
29	Efficient and Interpretable Prediction of Protein Functional Classes by Correspondence Analysis and Compact Set Relations. PLoS ONE, 2013, 8, e75542.	1.1	7
30	Label-free Quantitative Proteomics and N-Glycoproteomics Analysis of KRAS-activated Human Bronchial Epithelial Cells. Molecular and Cellular Proteomics, 2012, 11, 901-915.	2.5	23
31	Phosphoproteomic Analysis of Human Mesenchymal Stromal Cells during Osteogenic Differentiation. Journal of Proteome Research, 2012, 11, 586-598.	1.8	14
32	Prediction of nuclear proteins using nuclear translocation signals proposed by probabilistic latent semantic indexing. BMC Bioinformatics, 2012, 13, S13.	1.2	10
33	Computational Comparative Study of Tuberculosis Proteomes Using a Model Learned from Signal Peptide Structures. PLoS ONE, 2012, 7, e35018.	1.1	9
34	TMPad: an integrated structural database for helix-packing folds in transmembrane proteins. Nucleic Acids Research, 2011, 39, D347-D355.	6.5	21
35	Phosphoproteomics Identifies Oncogenic Ras Signaling Targets and Their Involvement in Lung Adenocarcinomas. PLoS ONE, 2011, 6, e20199.	1.1	35
36	Improving the Alignment Quality of Consistency Based Aligners with an Evaluation Function Using Synonymous Protein Words. PLoS ONE, 2011, 6, e27872.	1.1	4

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#	Article	IF	CITATIONS
37	Improving protein secondary structure prediction based on short subsequences with local structure similarity. BMC Genomics, 2010, 11, S4.	1.2	29
38	IDEAL-Q, an Automated Tool for Label-free Quantitation Analysis Using an Efficient Peptide Alignment Approach and Spectral Data Validation. Molecular and Cellular Proteomics, 2010, 9, 131-144.	2.5	114
39	An Informatics-assisted Label-free Quantitation Strategy that Depicts Phosphoproteomic Profiles in Lung Cancer Cell Invasion. Journal of Proteome Research, 2010, 9, 5582-5597.	1.8	57
40	Automated Generic Analysis Tools for Protein Quantitation Using Stable Isotope Labeling. Methods in Molecular Biology, 2010, 604, 257-272.	0.4	0
41	Abstract 5565: Quantitative proteomics identifies phosphorylation targets, modulators and pathway signatures of oncogenic RAS-mediated MAPK signaling in lung adenocarcinoma. , 2010, , .		0
42	Predicting helix–helix interactions from residue contacts in membrane proteins. Bioinformatics, 2009, 25, 996-1003.	1.8	56
43	MaXIC-Q Web: a fully automated web service using statistical and computational methods for protein quantitation based on stable isotope labeling and LC–MS. Nucleic Acids Research, 2009, 37, W661-W669.	6.5	10
44	Protein subcellular localization prediction of eukaryotes using a knowledge-based approach. BMC Bioinformatics, 2009, 10, S8.	1.2	35
45	PSLDoc: Protein subcellular localization prediction based on gappedâ€dipeptides and probabilistic latent semantic analysis. Proteins: Structure, Function and Bioinformatics, 2008, 72, 693-710.	1.5	44
46	Diameter variability of cycles and tori. Information Sciences, 2008, 178, 2960-2967.	4.0	9
47	Predicting RNA-binding sites of proteins using support vector machines and evolutionary information. BMC Bioinformatics, 2008, 9, S6.	1.2	110
48	Enhanced Membrane Protein Topology Prediction Using a Hierarchical Classification Method and a New Scoring Function. Journal of Proteome Research, 2008, 7, 487-496.	1.8	35
49	The Multi-Q web server for multiplexed protein quantitation. Nucleic Acids Research, 2007, 35, W707-W712.	6.5	14
50	BIOSMILE: A semantic role labeling system for biomedical verbs using a maximum-entropy model with automatically generated template features. BMC Bioinformatics, 2007, 8, 325.	1.2	45
51	Protein subcellular localization prediction based on compartment-specific features and structure conservation. BMC Bioinformatics, 2007, 8, 330.	1.2	54
52	Detection of the inferred interaction network in hepatocellular carcinoma from EHCO (Encyclopedia) Tj ETQq0 0	0 rgBT /O	verlock 10 Tf
53	Multi-Q:Â A Fully Automated Tool for Multiplexed Protein Quantitation. Journal of Proteome Research, 2006, 5, 2328-2338.	1.8	107

54Integrating linguistic knowledge into a conditional random fieldframework to identify biomedical<br/>named entities. Expert Systems With Applications, 2006, 30, 117-128.4.439

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#	Article	IF	CITATIONS
55	Various criteria in the evaluation of biomedical named entity recognition. BMC Bioinformatics, 2006, 7, 92.	1.2	80
56	NERBio: using selected word conjunctions, term normalization, and global patterns to improve biomedical named entity recognition. BMC Bioinformatics, 2006, 7, S11.	1.2	98
57	HYPLOSP: A KNOWLEDGE-BASED APPROACH TO PROTEIN LOCAL STRUCTURE PREDICTION. Journal of Bioinformatics and Computational Biology, 2006, 04, 1287-1307.	0.3	4
58	RIBRA—An Error-Tolerant Algorithm for the NMR Backbone Assignment Problem. Journal of Computational Biology, 2006, 13, 229-244.	0.8	18
59	TRANSMEMBRANE HELIX AND TOPOLOGY PREDICTION USING HIERARCHICAL SVM CLASSIFIERS AND AN ALTERNATING GEOMETRIC SCORING FUNCTION. , 2006, , .		5
60	PROTEIN SUBCELLULAR LOCALIZATION PREDICTION BASED ON COMPARTMENT-SPECIFIC BIOLOGICAL FEATURES. , 2006, , .		3
61	A semi-automatic method for annotating a biomedical proposition bank. , 2006, , .		21
62	Protein subcellular localization prediction based on compartment-specific biological features. Computational Systems Bioinformatics / Life Sciences Society Computational Systems Bioinformatics Conference, 2006, , 325-30.	0.4	3
63	A note on edge fault tolerance with respect to hypercubes. Applied Mathematics Letters, 2005, 18, 1125-1128.	1.5	4
64	HYPROSP II-A knowledge-based hybrid method for protein secondary structure prediction based on local prediction confidence. Bioinformatics, 2005, 21, 3227-3233.	1.8	50
65	GANAa genetic algorithm for NMR backbone resonance assignment. Nucleic Acids Research, 2005, 33, 4593-4601.	6.5	22
66	A KNOWLEDGE-BASED APPROACH TO PROTEIN LOCAL STRUCTURE PREDICTION. , 2005, , .		1
67	HYPROSP: a hybrid protein secondary structure prediction algorithma knowledge-based approach. Nucleic Acids Research, 2004, 32, 5059-5065.	6.5	24
68	On the construction of combinedk-fault-tolerant Hamiltonian graphs. Networks, 2001, 37, 165-170.	1.6	10
69	The shuffle-cubes and their generalization. Information Processing Letters, 2001, 77, 35-41.	0.4	27
70	Construction schemes for fault-tolerant Hamiltonian graphs. Networks, 2000, 35, 233-245.	1.6	6
71	On the isomorphism between cyclic-cubes and wrapped butterfly networks. IEEE Transactions on Parallel and Distributed Systems, 2000, 11, 864.	4.0	4
72	OPTIMUM CONGESTED ROUTING STRATEGY ON TWISTED CUBES. Journal of Interconnection Networks, 2000, 01, 115-134.	0.6	4

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#	Article	IF	CITATIONS
73	Edge congestion and topological properties of crossed cubes. IEEE Transactions on Parallel and Distributed Systems, 2000, 11, 64-80.	4.0	126
74	Christmas tree: A versatile 1-fault-tolerant design for token rings. Information Processing Letters, 1999, 72, 55-63.	0.4	17
75	Fault tolerant token ring embedding in double loop networks. Information Processing Letters, 1998, 66, 201-207.	0.4	8
76	The Recognition of Double Euler Trails in Series-Parallel Networks. Journal of Algorithms, 1998, 28, 216-257.	0.9	3
77	Multiple-edge-fault tolerance with respect to hypercubes. IEEE Transactions on Parallel and Distributed Systems, 1997, 8, 187-192.	4.0	14
78	Faithful 1-edge fault tolerant graphs. Information Processing Letters, 1997, 61, 173-181.	0.4	2
79	Routing and transmitting problems in de Bruijn networks. IEEE Transactions on Computers, 1996, 45, 1056-1062.	2.4	16
80	Transmitting on various network topologies. Networks, 1996, 27, 145-157.	1.6	2
81	A response to Volgenant's Addendum on the most vital edges. Networks, 1996, 27, 255-255.	1.6	0
82	Finding the most vital edges with respect to the number of spanning trees. IEEE Transactions on Reliability, 1994, 43, 600-603.	3.5	23
83	The most vital edges of matching in a bipartite graph. Networks, 1993, 23, 309-313.	1.6	4
84	A new shortest path routing algorithm and embedding cycles of crossed cube. , 0, , .		3
85	Christmas tree: a 1-fault-tolerant network for token rings. , 0, , .		0
86	GANAA Genetic Algorithm for NMR Backbone Resonance Assignment. , 0, , .		0
87	A Two-stage Classifier for Protein B-turn Prediction Using Support Vector Machines. , 0, , .		1
88	Multiaspect Examinations of Possible Alternative Mappings of Identified Variant Peptides: A Case Study on the HEK293 Cell Line. ACS Omega, 0, , .	1.6	1