Marc Vidal

List of Publications by Year in descending order

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4658 5988 41,535 160 85 160 citations h-index g-index papers 175 175 175 39618 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	The human disease network. Proceedings of the National Academy of Sciences of the United States of America, 2007, 104, 8685-8690.	7.1	2,924
2	Towards a proteome-scale map of the human protein–protein interaction network. Nature, 2005, 437, 1173-1178.	27.8	2,676
3	Evidence for dynamically organized modularity in the yeast protein–protein interaction network. Nature, 2004, 430, 88-93.	27.8	1,683
4	A Map of the Interactome Network of the Metazoan <i>C. elegans</i> . Science, 2004, 303, 540-543.	12.6	1,587
5	Drug—target network. Nature Biotechnology, 2007, 25, 1119-1126.	17.5	1,584
6	Interactome Networks and Human Disease. Cell, 2011, 144, 986-998.	28.9	1,543
7	High-Quality Binary Protein Interaction Map of the Yeast Interactome Network. Science, 2008, 322, 104-110.	12.6	1,297
8	Uncovering disease-disease relationships through the incomplete interactome. Science, 2015, 347, 1257601.	12.6	1,219
9	A Proteome-Scale Map of the Human Interactome Network. Cell, 2014, 159, 1212-1226.	28.9	1,199
10	Evidence for Network Evolution in an <i>Arabidopsis</i> Interactome Map. Science, 2011, 333, 601-607.	12.6	838
11	An empirical framework for binary interactome mapping. Nature Methods, 2009, 6, 83-90.	19.0	800
12	Independently Evolved Virulence Effectors Converge onto Hubs in a Plant Immune System Network. Science, 2011, 333, 596-601.	12.6	776
13	Protein Interaction Mapping in C. elegans Using Proteins Involved in Vulval Development. Science, 2000, 287, 116-122.	12.6	766
14	A reference map of the human binary protein interactome. Nature, 2020, 580, 402-408.	27.8	724
15	A Protein–Protein Interaction Network for Human Inherited Ataxias and Disorders of Purkinje Cell Degeneration. Cell, 2006, 125, 801-814.	28.9	714
16	Network modeling links breast cancer susceptibility and centrosome dysfunction. Nature Genetics, 2007, 39, 1338-1349.	21.4	602
17	How many human proteoforms are there?. Nature Chemical Biology, 2018, 14, 206-214.	8.0	580
18	[34] GATEWAY recombinational cloning: Application to the cloning of large numbers of open reading frames or ORFeomes. Methods in Enzymology, 2000, 328, 575-IN7.	1.0	570

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19	Correlation between transcriptome and interactome mapping data from Saccharomyces cerevisiae. Nature Genetics, 2001, 29, 482-486.	21.4	570
20	Proto-genes and de novo gene birth. Nature, 2012, 487, 370-374.	27.8	555
21	Annotation Transfer Between Genomes: Protein-Protein Interologs and Protein-DNA Regulogs. Genome Research, 2004, 14, 1107-1118.	5. 5	516
22	Intrinsic Disorder Is a Common Feature of Hub Proteins from Four Eukaryotic Interactomes. PLoS Computational Biology, 2006, 2, e100.	3.2	512
23	Widespread Macromolecular Interaction Perturbations in Human Genetic Disorders. Cell, 2015, 161, 647-660.	28.9	482
24	Widespread Expansion of Protein Interaction Capabilities by Alternative Splicing. Cell, 2016, 164, 805-817.	28.9	479
25	A public genome-scale lentiviral expression library of human ORFs. Nature Methods, 2011, 8, 659-661.	19.0	477
26	A Chaperome Subnetwork Safeguards Proteostasis in Aging and Neurodegenerative Disease. Cell Reports, 2014, 9, 1135-1150.	6.4	476
27	Identification of Potential Interaction Networks Using Sequence-Based Searches for Conserved Protein-Protein Interactions or "Interologs". Genome Research, 2001, 11, 2120-2126.	5. 5	431
28	Reverse two-hybrid and one-hybrid systems to detect dissociation of protein-protein and DNA-protein interactions Proceedings of the National Academy of Sciences of the United States of America, 1996, 93, 10315-10320.	7.1	416
29	Integrating â€~omic' information: a bridge between genomics and systems biology. Trends in Genetics, 2003, 19, 551-560.	6.7	407
30	An experimentally derived confidence score for binary protein-protein interactions. Nature Methods, 2009, 6, 91-97.	19.0	397
31	Network-based in silico drug efficacy screening. Nature Communications, 2016, 7, 10331.	12.8	394
32	Convergent Targeting of a Common Host Protein-Network by Pathogen Effectors from Three Kingdoms of Life. Cell Host and Microbe, 2014, 16, 364-375.	11.0	367
33	Interpreting cancer genomes using systematic host network perturbations by tumour virus proteins. Nature, 2012, 487, 491-495.	27.8	349
34	Epsteinâ€"Barr virus and virus human protein interaction maps. Proceedings of the National Academy of Sciences of the United States of America, 2007, 104, 7606-7611.	7.1	348
35	C. elegans ORFeome version 1.1: experimental verification of the genome annotation and resource for proteome-scale protein expression. Nature Genetics, 2003, 34, 35-41.	21.4	347
36	E2F-4 and E2F-5, two members of the E2F family, are expressed in the early phases of the cell cycle Proceedings of the National Academy of Sciences of the United States of America, 1995, 92, 2403-2407.	7.1	342

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37	Interactome: gateway into systems biology. Human Molecular Genetics, 2005, 14, R171-R181.	2.9	329
38	Edgetic perturbation models of human inherited disorders. Molecular Systems Biology, 2009, 5, 321.	7.2	326
39	Effect of sampling on topology predictions of protein-protein interaction networks. Nature Biotechnology, 2005, 23, 839-844.	17.5	302
40	Network-based prediction of protein interactions. Nature Communications, 2019, 10, 1240.	12.8	293
41	Literature-curated protein interaction datasets. Nature Methods, 2009, 6, 39-46.	19.0	288
42	Combined Functional Genomic Maps of the C. elegans DNA Damage Response. Science, 2002, 295, 127-131.	12.6	270
43	A novel member of the RING finger family, KRIP-1, associates with the KRAB-A transcriptional repressor domain of zinc finger proteins. Proceedings of the National Academy of Sciences of the United States of America, 1996, 93, 15299-15304.	7.1	260
44	Predictive models of molecular machines involved in Caenorhabditis elegans early embryogenesis. Nature, 2005, 436, 861-865.	27.8	260
45	Empirically controlled mapping of the Caenorhabditis elegans protein-protein interactome network. Nature Methods, 2009, 6, 47-54.	19.0	260
46	High-Throughput Yeast Two-Hybrid Assays for Large-Scale Protein Interaction Mapping. Methods, 2001, 24, 297-306.	3.8	258
47	Next-generation sequencing to generate interactome datasets. Nature Methods, 2011, 8, 478-480.	19.0	258
48	hORFeome v3.1: A resource of human open reading frames representing over 10,000 human genes. Genomics, 2007, 89, 307-315.	2.9	248
49	Screening patients for heterozygous p53 mutations using a functional assay in yeast. Nature Genetics, 1993, 5, 124-129.	21.4	243
50	Combining biological networks to predict genetic interactions. Proceedings of the National Academy of Sciences of the United States of America, 2004, 101, 15682-15687.	7.1	225
51	Human ORFeome Version 1.1: A Platform for Reverse Proteomics. Genome Research, 2004, 14, 2128-2135.	5.5	208
52	A Protein Domain-Based Interactome Network for C. elegans Early Embryogenesis. Cell, 2008, 134, 534-545.	28.9	196
53	A Gateway-Compatible Yeast One-Hybrid System. Genome Research, 2004, 14, 2093-2101.	5.5	189
54	A Biological Atlas of Functional Maps. Cell, 2001, 104, 333-339.	28.9	188

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55	A protein–protein interaction map of the Caenorhabditis elegans 26S proteasome. EMBO Reports, 2001, 2, 821-828.	4.5	173
56	Integrating Interactome, Phenome, and Transcriptome Mapping Data for the C. elegans Germline. Current Biology, 2002, 12, 1952-1958.	3.9	170
57	Mycobacterium tuberculosis Type VII Secreted Effector EsxH Targets Host ESCRT to Impair Trafficking. PLoS Pathogens, 2013, 9, e1003734.	4.7	167
58	A disease module in the interactome explains disease heterogeneity, drug response and captures novel pathways and genes in asthma. Human Molecular Genetics, 2015, 24, 3005-3020.	2.9	162
59	Open-reading-frame sequence tags (OSTs) support the existence of at least 17,300 genes in C. elegans. Nature Genetics, 2001, 27, 332-336.	21.4	159
60	Dominant-negative p53 mutations selected in yeast hit cancer hot spots Proceedings of the National Academy of Sciences of the United States of America, 1996, 93, 4091-4095.	7.1	158
61	A First Version of the Caenorhabditis elegans Promoterome. Genome Research, 2004, 14, 2169-2175.	5.5	155
62	A framework for exhaustively mapping functional missense variants. Molecular Systems Biology, 2017, 13, 957.	7.2	146
63	Protein interaction maps for model organisms. Nature Reviews Molecular Cell Biology, 2001, 2, 55-63.	37.0	139
64	Spatiotemporal 16p11.2 Protein Network Implicates Cortical Late Mid-Fetal Brain Development and KCTD13-Cul3-RhoA Pathway in Psychiatric Diseases. Neuron, 2015, 85, 742-754.	8.1	139
65	Systematic Interactome Mapping and Genetic Perturbation Analysis of a C. elegans TGF- \hat{l}^2 Signaling Network. Molecular Cell, 2004, 13, 469-482.	9.7	136
66	Increasing specificity in high-throughput yeast two-hybrid experiments. Methods, 2004, 32, 363-370.	3.8	135
67	Analysis of the human E2 ubiquitin conjugating enzyme protein interaction network. Genome Research, 2009, 19, 1905-1911.	5.5	134
68	A genome-wide positioning systems network algorithm for in silico drug repurposing. Nature Communications, 2019, 10, 3476.	12.8	134
69	Protein interaction network of alternatively spliced isoforms from brain links genetic risk factors for autism. Nature Communications, 2014, 5, 3650.	12.8	131
70	Edgotype: a fundamental link between genotype and phenotype. Current Opinion in Genetics and Development, 2013, 23, 649-657.	3.3	129
71	Proteome-Scale Human Interactomics. Trends in Biochemical Sciences, 2017, 42, 342-354.	7.5	129
72	High-Quality Binary Interactome Mapping. Methods in Enzymology, 2010, 470, 281-315.	1.0	126

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73	Yeast Two-Hybrid Systems and Protein Interaction Mapping Projects for Yeast and Worm. Yeast, 2000, 1, 88-94.	1.7	121
74	Insight into transcription factor gene duplication from Caenorhabditis elegans Promoterome-driven expression patterns. BMC Genomics, 2007, 8, 27.	2.8	120
75	HSP90 Shapes the Consequences of Human Genetic Variation. Cell, 2017, 168, 856-866.e12.	28.9	117
76	An extended set of yeast-based functional assays accurately identifies human disease mutations. Genome Research, 2016, 26, 670-680.	5.5	116
77	Survey of variation in human transcription factors reveals prevalent DNA binding changes. Science, 2016, 351, 1450-1454.	12.6	114
78	APID database: redefining protein–protein interaction experimental evidences and binary interactomes. Database: the Journal of Biological Databases and Curation, 2019, 2019, .	3.0	113
79	Human Gene-Centered Transcription Factor Networks for Enhancers and Disease Variants. Cell, 2015, 161, 661-673.	28.9	111
80	A comprehensive library of human transcription factors for cell fate engineering. Nature Biotechnology, 2021, 39, 510-519.	17.5	110
81	Comprehensive characterization of protein–protein interactions perturbed by disease mutations. Nature Genetics, 2021, 53, 342-353.	21.4	109
82	C. elegans ORFeome Version 3.1: Increasing the Coverage of ORFeome Resources With Improved Gene Predictions. Genome Research, 2004, 14, 2064-2069.	5 . 5	107
83	A unifying view of 21st century systems biology. FEBS Letters, 2009, 583, 3891-3894.	2.8	105
84	A Novel Human DnaJ Protein, hTid-1, a Homolog of the Drosophila Tumor Suppressor Protein Tid56, Can Interact with the Human Papillomavirus Type 16 E7 Oncoprotein. Virology, 1998, 247, 74-85.	2.4	102
85	Viral Perturbations of Host Networks Reflect Disease Etiology. PLoS Computational Biology, 2012, 8, e1002531.	3.2	102
86	Pooledâ€matrix protein interaction screens using Barcode Fusion Genetics. Molecular Systems Biology, 2016, 12, 863.	7.2	102
87	Genome-Scale Networks Link Neurodegenerative Disease Genes to α-Synuclein through Specific Molecular Pathways. Cell Systems, 2017, 4, 157-170.e14.	6.2	102
88	Dynamic Role of trans Regulation of Gene Expression in Relation to Complex Traits. American Journal of Human Genetics, 2017, 100, 571-580.	6.2	101
89	Interactome modeling. FEBS Letters, 2005, 579, 1834-1838.	2.8	99
90	Selecting causal genes from genome-wide association studies via functionally coherent subnetworks. Nature Methods, 2015, 12, 154-159.	19.0	96

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91	In Situ Peroxidase Labeling and Mass-Spectrometry Connects Alpha-Synuclein Directly to Endocytic Trafficking and mRNA Metabolism in Neurons. Cell Systems, 2017, 4, 242-250.e4.	6.2	91
92	Prospects for drug screening using the reverse two-hybrid system. Trends in Biotechnology, 1999, 17, 374-381.	9.3	88
93	Systematic screening reveals a role for BRCA1 in the response to transcription-associated DNA damage. Genes and Development, 2014, 28, 1957-1975.	5.9	86
94	A Zebrafish Genetic Screen Identifies Neuromedin U as a Regulator of Sleep/Wake States. Neuron, 2016, 89, 842-856.	8.1	81
95	Knocking out multigene redundancies via cycles of sexual assortment and fluorescence selection. Nature Methods, 2011, 8, 159-164.	19.0	74
96	Human T cell leukemia virus type 1 Tax associates with a molecular chaperone complex containing hTid-1 and Hsp70. Current Biology, 2001, 11, 1771-1775.	3.9	73
97	Multiplex single-molecule interaction profiling of DNA-barcoded proteins. Nature, 2014, 515, 554-557.	27.8	72
98	'Edgetic' perturbation of a C. elegans BCL2 ortholog. Nature Methods, 2009, 6, 843-849.	19.0	71
99	ORFeome Cloning and Systems Biology: Standardized Mass Production of the Parts From the Parts-List. Genome Research, 2004, 14, 2001-2009.	5.5	69
100	The yeast two-hybrid assay: still finding connections after 25 years. Nature Methods, 2014, 11, 1203-1206.	19.0	68
101	Mapping transcription factor interactome networks using HaloTag protein arrays. Proceedings of the National Academy of Sciences of the United States of America, 2016, 113, E4238-47.	7.1	67
102	Host-pathogen interactome mapping for HTLV-1 and -2 retroviruses. Retrovirology, 2012, 9, 26.	2.0	64
103	SH3 interactome conserves general function over specific form. Molecular Systems Biology, 2013, 9, 652.	7.2	61
104	Comparative analysis of virus–host interactomes with a mammalian high-throughput protein complementation assay based on Gaussia princeps luciferase. Methods, 2012, 58, 349-359.	3.8	59
105	Protein Domain-Level Landscape of Cancer-Type-Specific Somatic Mutations. PLoS Computational Biology, 2015, 11, e1004147.	3.2	59
106	Identification of FAM111A as an SV40 Host Range Restriction and Adenovirus Helper Factor. PLoS Pathogens, 2012, 8, e1002949.	4.7	58
107	<i>MECP2</i> Is a Frequently Amplified Oncogene with a Novel Epigenetic Mechanism That Mimics the Role of Activated RAS in Malignancy. Cancer Discovery, 2016, 6, 45-58.	9.4	57
108	Maximizing binary interactome mapping with a minimal number of assays. Nature Communications, 2019, 10, 3907.	12.8	57

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109	InSite: a computational method for identifying protein-protein interaction binding sites on a proteome-wide scale. Genome Biology, 2007, 8, R192.	9.6	53
110	Array MAPPIT: High-Throughput Interactome Analysis in Mammalian Cells. Journal of Proteome Research, 2009, 8, 877-886.	3.7	53
111	A Genetic Strategy to Eliminate Self-Activator Baits Prior to High-Throughput Yeast Two-Hybrid Screens. Genome Research, 1999, 9, 1128-1134.	5.5	51
112	Interactome Mapping of the Phosphatidylinositol 3-Kinase-Mammalian Target of Rapamycin Pathway Identifies Deformed Epidermal Autoregulatory Factor-1 as a New Glycogen Synthase Kinase-3 Interactor. Molecular and Cellular Proteomics, 2010, 9, 1578-1593.	3.8	51
113	A proactive genotype-to-patient-phenotype map for cystathionine beta-synthase. Genome Medicine, 2020, 12, 13.	8.2	45
114	Protein interactions of the transcription factor Hoxa1. BMC Developmental Biology, 2012, 12, 29.	2.1	43
115	An interâ€species protein–protein interaction network across vast evolutionary distance. Molecular Systems Biology, 2016, 12, 865.	7.2	42
116	Shifted Transversal Design smart-pooling for high coverage interactome mapping. Genome Research, 2009, 19, 1262-1269.	5.5	36
117	WorfDB: the Caenorhabditis elegans ORFeome Database. Nucleic Acids Research, 2003, 31, 237-240.	14.5	35
118	The transcription factor ERG recruits CCR4–NOT to control mRNA decay and mitotic progression. Nature Structural and Molecular Biology, 2016, 23, 663-672.	8.2	35
119	Network Analysis of UBE3A/E6AP-Associated Proteins Provides Connections to Several Distinct Cellular Processes. Journal of Molecular Biology, 2018, 430, 1024-1050.	4.2	32
120	How much of the human protein interactome remains to be mapped?. Science Signaling, 2016, 9, eg7.	3.6	31
121	Isoform discovery by targeted cloning, 'deep-well' pooling and parallel sequencing. Nature Methods, 2008, 5, 597-600.	19.0	30
122	Identifying pathogenicity of human variants via paralog-based yeast complementation. PLoS Genetics, 2017, 13, e1006779.	3.5	30
123	Precision medicine â€" networks to the rescue. Current Opinion in Biotechnology, 2020, 63, 177-189.	6.6	30
124	Domain-based prediction of the human isoform interactome provides insights into the functional impact of alternative splicing. PLoS Computational Biology, 2017, 13, e1005717.	3.2	30
125	Global Edgetic Rewiring in Cancer Networks. Cell Systems, 2015, 1, 251-253.	6.2	28
126	Global Analysis of Intercellular Homeodomain Protein Transfer. Cell Reports, 2019, 28, 712-722.e3.	6.4	28

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127	Histone deacetylase-dependent transcriptional repression by pRB in yeast occurs independently of interaction through the LXCXE binding cleft. Proceedings of the National Academy of Sciences of the United States of America, 2001, 98, 8720-8725.	7.1	26
128	From genome to proteome: developing expression clone resources for the human genome. Human Molecular Genetics, 2006, 15, R31-R43.	2.9	26
129	Controllability in an islet specific regulatory network identifies the transcriptional factor NFATC4, which regulates Type 2 Diabetes associated genes. Npj Systems Biology and Applications, 2018, 4, 25.	3.0	25
130	Protein–Protein Interactions and Networks: Forward and Reverse Edgetics. Methods in Molecular Biology, 2011, 759, 197-213.	0.9	24
131	Proteome-scale Binary Interactomics in Human Cells. Molecular and Cellular Proteomics, 2016, 15, 3624-3639.	3.8	23
132	A Model of Elegance. American Journal of Human Genetics, 1998, 63, 955-961.	6.2	22
133	Mutational analysis of the J recombination signal sequence binding protein (RBP-J)/Epstein-Barr virus nuclear antigen 2 (EBNA2) and RBP-J/Notch interaction. FEBS Journal, 2001, 268, 4639-4646.	0.2	21
134	[6] A green fluorescent protein-based reverse two-hybrid system: Application to the characterization of large numbers of potential protein-protein interactions. Methods in Enzymology, 2000, 328, 74-IN1.	1.0	19
135	Challenges and recommendations for epigenomics in precision health. Nature Biotechnology, 2017, 35, 1128-1132.	17.5	19
136	ORF Capture-Seq as a versatile method for targeted identification of full-length isoforms. Nature Communications, 2020, 11, 2326.	12.8	19
137	The HTLV-1 viral oncoproteins Tax and HBZ reprogram the cellular mRNA splicing landscape. PLoS Pathogens, 2021, 17, e1009919.	4.7	19
138	A Genome-Wide Gene Function Prediction Resource for Drosophila melanogaster. PLoS ONE, 2010, 5, e12139.	2.5	17
139	A massively parallel barcoded sequencing pipeline enables generation of the first ORFeome and interactome map for rice. Proceedings of the National Academy of Sciences of the United States of America, 2020, 117, 11836-11842.	7.1	16
140	Yeast Two-Hybrid Systems and Protein Interaction Mapping Projects for Yeast and Worm. Yeast, 2000, 1, 88-94.	1.7	16
141	New Views of Old Proteins: Clarifying the Enigmatic Proteome. Molecular and Cellular Proteomics, 2022, 21, 100254.	3.8	16
142	Yeast genetic interaction screen of human genes associated with amyotrophic lateral sclerosis: identification of MAP2K5 kinase as a potential drug target. Genome Research, 2017, 27, 1487-1500.	5.5	12
143	Use of protein-interaction maps to formulate biological questions. Current Opinion in Chemical Biology, 2001, 5, 57-62.	6.1	11
144	Integrated version of reverse two-hybrid system for the postproteomic era. Methods in Enzymology, 2002, 350, 525-545.	1.0	10

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145	Alternative glycosylation controls endoplasmic reticulum dynamics and tubular extension in mammalian cells. Science Advances, 2021, 7, .	10.3	8
146	Reply to "Does mapping reveal correlation between gene expression and protein–protein interaction?". Nature Genetics, 2003, 33, 16-17.	21.4	7
147	Perturbing interactions. Nature Methods, 2005, 2, 412-414.	19.0	6
148	Protein Interactomics by Two-Hybrid Methods. Methods in Molecular Biology, 2018, 1794, 1-14.	0.9	6
149	QUICKstep and GS-TAP: new moves for protein-interaction analysis. Nature Methods, 2006, 3, 975-976.	19.0	5
150	Interactome Networks., 2013,, 45-63.		5
151	Yeast-Based Genetic Interaction Analysis of Human Kinome. Cells, 2020, 9, 1156.	4.1	5
152	A Protein Domain-Based Interactome Network for C.Âelegans Early Embryogenesis. Cell, 2012, 151, 1633.	28.9	4
153	Interrogation of kinase genetic interactions provides a global view of PAK1-mediated signal transduction pathways. Journal of Biological Chemistry, 2020, 295, 16906-16919.	3.4	4
154	Reply to "Exhaustive benchmarking of the yeast two-hybrid system". Nature Methods, 2010, 7, 668-668.	19.0	3
155	OpenPIP: An Open-source Platform for Hosting, Visualizing and Analyzing Protein Interaction Data. Journal of Molecular Biology, 2022, 434, 167603.	4.2	3
156	Reply to Toward the complete interactome. Nature Biotechnology, 2006, 24, 615-615.	17.5	1
157	From genome to proteome: developing expression clone resources for the human genome. Human Molecular Genetics, 2006, 15, 2184-2184.	2.9	1
158	TheC.elegansinteractome project. , 2005, , .		0
159	Forward and Reverse Proteomics. , 2003, , 255-276.		0
160	Interactome Networks. FASEB Journal, 2008, 22, 262.1.	0.5	0