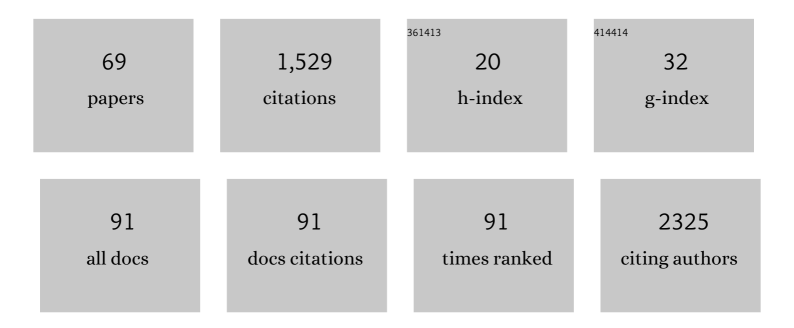
## Pieter Meysman

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/432484/publications.pdf Version: 2024-02-01



#	Article	IF	CITATIONS
1	MoMAC: Multi-objective optimization to combine multiple association rules into an interpretable classification. Applied Intelligence, 2022, 52, 3090-3102.	5.3	10
2	Preexisting memory CD4 T cells in na $\tilde{A}$ ve individuals confer robust immunity upon hepatitis B vaccination. ELife, 2022, 11, .	6.0	11
3	Recent advances in T-cell receptor repertoire analysis: Bridging the gap with multimodal single-cell RNA sequencing. ImmunoInformatics, 2022, 5, 100009.	2.2	27
4	Viral Evolution and Immunology of SARS-CoV-2 in a Persistent Infection after Treatment with Rituximab. Viruses, 2022, 14, 752.	3.3	10
5	Current challenges for unseen-epitope TCR interaction prediction and a new perspective derived from image classification. Briefings in Bioinformatics, 2021, 22, .	6.5	78
6	Preparing students for the data-driven life science era through a real-world viral infection case. Journal of Biological Education, 2021, 55, 178-187.	1.5	2
7	HLA-DRB1 Alleles Associated with Lower Leishmaniasis Susceptibility Share Common Amino Acid Polymorphisms and Epitope Binding Repertoires. Vaccines, 2021, 9, 270.	4.4	5
8	Sorted B cell transcriptomes point towards actively regulated B cell responses during ongoing chronic hepatitis B infections. Cellular Immunology, 2021, 362, 104283.	3.0	9
9	Constrained Standardization of Count Data from Massive Parallel Sequencing. Journal of Molecular Biology, 2021, 433, 166966.	4.2	1
10	ClusTCR: a python interface for rapid clustering of large sets of CDR3 sequences with unknown antigen specificity. Bioinformatics, 2021, 37, 4865-4867.	4.1	30
11	The use of plasma donor-derived, cell-free DNA to monitor acute rejection after kidney transplantation. Nephrology Dialysis Transplantation, 2020, 35, 714-721.	0.7	61
12	The Absence of C-5 DNA Methylation in Leishmania donovani Allows DNA Enrichment from Complex Samples. Microorganisms, 2020, 8, 1252.	3.6	9
13	Transcriptome profiling of liver biopsies before antiviral treatment start can predict HCC development 8.3 years before clinical diagnosis in chronic hepatitis B and C patients. Journal of Hepatology, 2020, 73, S392.	3.7	0
14	Transcriptomic profiling of different responder types in adults after a Priorix® vaccination. Vaccine, 2020, 38, 3218-3226.	3.8	1
15	Clustering association rules to build beliefs and discover unexpected patterns. Applied Intelligence, 2020, 50, 1943-1954.	5.3	10
16	Oocyte maturation under lipotoxic conditions induces carryover transcriptomic and functional alterations during post-hatching development of good-quality blastocysts: novel insights from a bovine embryo-transfer model. Human Reproduction, 2020, 35, 293-307.	0.9	17
17	MESSAR: Automated recommendation of metabolite substructures from tandem mass spectra. PLoS ONE, 2020, 15, e0226770.	2.5	24
18	Identification of Epitope-Specific T Cells in T-Cell Receptor Repertoires. Methods in Molecular Biology, 2020, 2120, 183-195	0.9	3

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19	Memory CD4+ T cell receptor repertoire data mining as a tool for identifying cytomegalovirus serostatus. Genes and Immunity, 2019, 20, 255-260.	4.1	19
20	Detection of Enriched T Cell Epitope Specificity in Full T Cell Receptor Sequence Repertoires. Frontiers in Immunology, 2019, 10, 2820.	4.8	108
21	Diagnosing enterovirus meningitis via blood transcriptomics: an alternative for lumbar puncture?. Journal of Translational Medicine, 2019, 17, 282.	4.4	10
22	Using Expert Driven Machine Learning to Enhance Dynamic Metabolomics Data Analysis. Metabolites, 2019, 9, 54.	2.9	15
23	MILES: a Java tool to extract node-specific enriched subgraphs in biomolecular networks. Bioinformatics, 2019, , .	4.1	0
24	On the viability of unsupervised T-cell receptor sequence clustering for epitope preference. Bioinformatics, 2019, 35, 1461-1468.	4.1	52
25	Mining the Enriched Subgraphs for Specific Vertices in a Biological Graph. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2019, 16, 1496-1507.	3.0	5
26	Integrated genomic and metabolomic profiling of ISC1, an emerging Leishmania donovani population in the Indian subcontinent. Infection, Genetics and Evolution, 2018, 62, 170-178.	2.3	32
27	Increased herpes zoster risk associated with poor HLA-A immediate early 62 protein (IE62) affinity. Immunogenetics, 2018, 70, 363-372.	2.4	8
28	On the feasibility of mining CD8+ T cell receptor patterns underlying immunogenic peptide recognition. Immunogenetics, 2018, 70, 159-168.	2.4	64
29	Plasma donor-derived cell-free DNA kinetics after kidney transplantation using a single tube multiplex PCR assay. PLoS ONE, 2018, 13, e0208207.	2.5	50
30	Fast Open Modification Spectral Library Searching through Approximate Nearest Neighbor Indexing. Journal of Proteome Research, 2018, 17, 3463-3474.	3.7	69
31	Grasping frequent subgraph mining for bioinformatics applications. BioData Mining, 2018, 11, 20.	4.0	30
32	Transcriptome profiling in blood before and after hepatitis B vaccination shows significant differences in gene expression between responders and non-responders. Vaccine, 2018, 36, 6282-6289.	3.8	47
33	FO017UNILATERAL NEPHRECTOMY SUPPRESSES IMMUNE RESPONSE PATHWAYS IN ACUTE KIDNEY INJURY AND OVERCOMES PROGRESSION TO CHRONIC KIDNEY DISEASE IN MICE. Nephrology Dialysis Transplantation, 2018, 33, i8-i8.	0.7	Ο
34	speaq 2.0: A complete workflow for high-throughput 1D NMR spectra processing and quantification. PLoS Computational Biology, 2018, 14, e1006018.	3.2	42
35	Protein complex analysis: From raw protein lists to protein interaction networks. Mass Spectrometry Reviews, 2017, 36, 600-614.	5.4	22
36	sfinx: an R package for the elimination of false positives from affinity purification–mass spectrometry datasets. Bioinformatics, 2017, 33, 1902-1904.	4.1	3

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37	Multidisciplinary study of the secondary immune response in grandparents re-exposed to chickenpox. Scientific Reports, 2017, 7, 1077.	3.3	28
38	Multiplexed Spliced-Leader Sequencing: A high-throughput, selective method for RNA-seq in Trypanosomatids. Scientific Reports, 2017, 7, 3725.	3.3	24
39	Ten simple rules for forming a scientific professional society. PLoS Computational Biology, 2017, 13, e1005226.	3.2	7
40	Second ISCB Latin American Student Council Symposium (LA-SCS) 2016. F1000Research, 2017, 6, 1491.	1.6	5
41	Immunological evasion of immediate-early varicella zoster virus proteins. Immunogenetics, 2016, 68, 483-486.	2.4	8
42	Practical Approaches for Mining Frequent Patterns in Molecular Datasets. Bioinformatics and Biology Insights, 2016, 10, BBI.S38419.	2.0	2
43	Highlights from the 11th ISCB Student Council Symposium 2015. BMC Bioinformatics, 2016, 17, 95.	2.6	4
44	Unsupervised Quality Assessment of Mass Spectrometry Proteomics Experiments by Multivariate Quality Control Metrics. Journal of Proteome Research, 2016, 15, 1300-1307.	3.7	15
45	COLOMBOS v3.0: leveraging gene expression compendia for cross-species analyses: Table 1 Nucleic Acids Research, 2016, 44, D620-D623.	14.5	69
46	SFINX: Straightforward Filtering Index for Affinity Purification–Mass Spectrometry Data Analysis. Journal of Proteome Research, 2016, 15, 332-338.	3.7	20
47	Highlights from the ISCB Student Council Symposia in 2016. F1000Research, 2016, 5, 2852.	1.6	4
48	ISCB-Student Council Narratives: Strategical development of the ISCB-Regional Student Groups in 2016. F1000Research, 2016, 5, 2882.	1.6	6
49	Mining the human proteome for conserved mechanisms. BMC Bioinformatics, 2015, 16, .	2.6	1
50	Varicella-Zoster Virus-Derived Major Histocompatibility Complex Class I-Restricted Peptide Affinity Is a Determining Factor in the HLA Risk Profile for the Development of Postherpetic Neuralgia. Journal of Virology, 2015, 89, 962-969.	3.4	20
51	Mining the entire Protein DataBank for frequent spatially cohesive amino acid patterns. BioData Mining, 2015, 8, 4.	4.0	12
52	Mining Association Rules in Graphs Based on Frequent Cohesive Itemsets. Lecture Notes in Computer Science, 2015, , 637-648.	1.3	15
53	A primer to frequent itemset mining for bioinformatics. Briefings in Bioinformatics, 2015, 16, 216-231.	6.5	102
54	Comparative Analysis of Gene Expression: Uncovering Expression Conservation and Divergence Between Salmonella enterica Serovar Typhimurium Strains LT2 and 14028S. Methods in Molecular Biology, 2015, 1231, 125-135.	0.9	0

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55	Structural Properties of Prokaryotic Promoter Regions Correlate with Functional Features. PLoS ONE, 2014, 9, e88717.	2.5	22
56	Soft Skills: An Important Asset Acquired from Organizing Regional Student Group Activities. PLoS Computational Biology, 2014, 10, e1003708.	3.2	15
57	COLOMBOS v2.0: an ever expanding collection of bacterial expression compendia: Table 1 Nucleic Acids Research, 2014, 42, D649-D653.	14.5	38
58	MAGIC: access portal to a cross-platform gene expression compendium for maize. Bioinformatics, 2014, 30, 1316-1318.	4.1	2
59	Discovery of Spatially Cohesive Itemsets in Three-Dimensional Protein Structures. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2014, 11, 814-825.	3.0	5
60	Inferring gene expression from ribosomal promoter sequences, a crowdsourcing approach. Genome Research, 2013, 23, 1928-1937.	5.5	12
61	Expression Divergence between Escherichia coli and Salmonella enterica serovar Typhimurium Reflects Their Lifestyles. Molecular Biology and Evolution, 2013, 30, 1302-1314.	8.9	27
62	Mining spatially cohesive itemsets in protein molecular structures. , 2013, , .		2
63	Identifying Common Structural DNA Properties in Transcription Factor Binding Site Sets of the Lacl-GalR Family. Current Bioinformatics, 2013, 8, 483-488.	1.5	4
64	DNA Structural Properties in the Classification of Genomic Transcription Regulation Elements. Bioinformatics and Biology Insights, 2012, 6, BBI.S9426.	2.0	27
65	Directed Module Detection in a Large-Scale Expression Compendium. Methods in Molecular Biology, 2012, 804, 131-165.	0.9	3
66	COLOMBOS: Access Port for Cross-Platform Bacterial Expression Compendia. PLoS ONE, 2011, 6, e20938.	2.5	25
67	Query-based biclustering of gene expression data using Probabilistic Relational Models. BMC Bioinformatics, 2011, 12, S37.	2.6	14
68	Use of structural DNA properties for the prediction of transcription-factor binding sites in Escherichia coli. Nucleic Acids Research, 2011, 39, e6-e6.	14.5	42
69	Conditional Random Fields Feature Subset Selection Based on Genetic Algorithms for Phosphorylation Site Prediction. , 2009, , .		Ο