## Yee Ling Wu

## List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/11336910/publications.pdf

Version: 2024-02-01

706676 843174 1,375 22 14 20 h-index citations g-index papers 22 22 22 2658 all docs docs citations times ranked citing authors

#	Article	IF	CITATIONS
1	Gene Copy-Number Variation and Associated Polymorphisms of Complement Component C4 in Human Systemic Lupus Erythematosus (SLE): Low Copy Number Is a Risk Factor for and High Copy Number Is a Protective Factor against SLE Susceptibility in European Americans. American Journal of Human Genetics, 2007, 80, 1037-1054.	2.6	411
2	DNA deaminases induce break-associated mutation showers with implication of APOBEC3B and 3A in breast cancer kataegis. ELife, 2013, 2, e00534.	2.8	322
3	Early Components of the Complement Classical Activation Pathway in Human Systemic Autoimmune Diseases. Frontiers in Immunology, 2016, 7, 36.	2.2	143
4	Sensitive and Specific Real-Time Polymerase Chain Reaction Assays to Accurately Determine Copy Number Variations (CNVs) of Human Complement <i>C4A</i> , <i>C4B</i> , <i>C4-Short</i> , and RCCX Modules: Elucidation of <i>C4</i> CNVs in 50 Consanguineous Subjects with Defined HLA Genotypes. Journal of Immunology, 2007, 179, 3012-3025.	0.4	77
5	Effects of Complement <i>C4</i> Gene Copy Number Variations, Size Dichotomy, and <i>C4A</i> Deficiency on Genetic Risk and Clinical Presentation of Systemic Lupus Erythematosus in East Asian Populations. Arthritis and Rheumatology, 2016, 68, 1442-1453.	2.9	58
6	Active RNAP pre-initiation sites are highly mutated by cytidine deaminases in yeast, with AID targeting small RNA genes. ELife, 2014, 3, e03553.	2.8	51
7	Intrinsic transcriptional heterogeneity in B cells controls early class switching to IgE. Journal of Experimental Medicine, 2017, 214, 183-196.	4.2	49
8	Great genotypic and phenotypic diversities associated with copy-number variations of complement C4 and RP-C4-CYP21-TNX (RCCX) modules: A comparison of Asian-Indian and European American populations. Molecular Immunology, 2009, 46, 1289-1303.	1.0	43
9	Gene copy-number variations (CNVs) of complement <i>C4</i> and <i>C4A</i> deficiency in genetic risk and pathogenesis of juvenile dermatomyositis. Annals of the Rheumatic Diseases, 2016, 75, 1599-1606.	0.5	36
10	Increased frequency of complement <i>C4B</i> deficiency in rheumatoid arthritis. Arthritis and Rheumatism, 2012, 64, 1338-1344.	6.7	28
11	Human Complement Components C4A and C4B Genetic Diversities: Complex Genotypes and Phenotypes. Current Protocols in Immunology, 2005, 68, Unit 13.8.	3.6	26
12	Four Systemic Lupus Erythematosus Subgroups, Defined by Autoantibodies Status, Differ Regarding <i>HLAâ€DRB1</i> Genotype Associations and Immunological and Clinical Manifestations. ACR Open Rheumatology, 2022, 4, 27-39.	0.9	25
13	Assessment of complement C4 gene copy number using the paralog ratio test. Human Mutation, 2010, 31, 866-874.	1.1	23
14	Opposite Profiles of Complement in Antiphospholipid Syndrome (APS) and Systemic Lupus Erythematosus (SLE) Among Patients With Antiphospholipid Antibodies (aPL). Frontiers in Immunology, 2019, 10, 885.	2.2	20
15	Gene CNVs and protein levels of complement C4A and C4B as novel biomarkers for partial disease remissions in new-onset type 1 diabetes patients. Pediatric Diabetes, 2012, 13, 408-418.	1.2	14
16	Elevated serum complement levels and higher gene copy number of complement <i>C4B</i> associated with hypertension and effective response to statin therapy in childhood-onset systemic lupus erythematosus (SLE). Lupus Science and Medicine, 2019, 6, e000333.	1.1	11
17	Human Complement C4B Allotypes and Deficiencies in Selected Cases With Autoimmune Diseases. Frontiers in Immunology, 2021, 12, 739430.	2.2	11
18	Strong Association of Combined Genetic Deficiencies in the Classical Complement Pathway With Risk of Systemic Lupus Erythematosus and Primary Sjögren's Syndrome. Arthritis and Rheumatology, 2022, 74, 1842-1850.	2.9	10

#	Article	IF	CITATIONS
19	Determination of the Loss of Function Complement C4 Exon 29 CT Insertion Using a Novel Paralog-Specific Assay in Healthy UK and Spanish Populations. PLoS ONE, 2011, 6, e22128.	1.1	6
20	Loop extrusion promotes an alternate pathway for isotype switching. Cell Reports, 2021, 37, 110059.	2.9	6
21	Complement Deficiencies in Human Systemic Lupus Erythematosus (SLE) and SLE Nephritis: Epidemiology and Pathogenesis., 2007,, 183-193.		4
22	Complement in Rheumatic Diseases. , 2014, , 286-302.		1