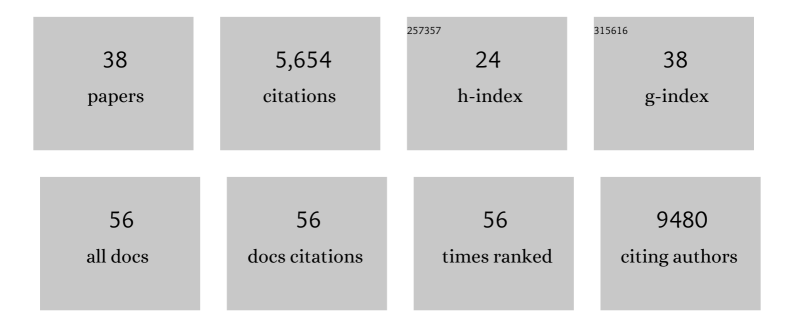
Gytis Dudas

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/2503531/publications.pdf Version: 2024-02-01



Οντις Πιιρλς

#	Article	IF	CITATIONS
1	Antibody escape and global spread of SARS-CoV-2 lineage A.27. Nature Communications, 2022, 13, 1152.	5.8	20
2	Lower respiratory tract infections in children requiring mechanical ventilation: a multicentre prospective surveillance study incorporating airway metagenomics. Lancet Microbe, The, 2022, 3, e284-e293.	3.4	24
3	Polymorphism of genetic ambigrams. Virus Evolution, 2021, 7, veab038.	2.2	5
4	Single mosquito metatranscriptomics identifies vectors, emerging pathogens and reservoirs in one assay. ELife, 2021, 10, .	2.8	69
5	Variant Analysis of SARS-CoV-2 Genomes from Belgian Military Personnel Engaged in Overseas Missions and Operations. Viruses, 2021, 13, 1359.	1.5	6
6	Global genome analysis reveals a vast and dynamic anellovirus landscape within the human virome. Cell Host and Microbe, 2021, 29, 1305-1315.e6.	5.1	59
7	Bulk Milk Tank Samples Are Suitable to Assess Circulation of Tick-Borne Encephalitis Virus in High Endemic Areas. Viruses, 2021, 13, 1772.	1.5	5
8	Emergence of an early SARS-CoV-2 epidemic in the United States. Cell, 2021, 184, 4939-4952.e15.	13.5	31
9	Conserved ancestral tropical niche but different continental histories explain the latitudinal diversity gradient in brush-footed butterflies. Nature Communications, 2021, 12, 5717.	5.8	33
10	Emergence and spread of SARS-CoV-2 lineage B.1.620 with variant of concern-like mutations and deletions. Nature Communications, 2021, 12, 5769.	5.8	51
11	Bayesian inference of reassortment networks reveals fitness benefits of reassortment in human influenza viruses. Proceedings of the National Academy of Sciences of the United States of America, 2020, 117, 17104-17111.	3.3	35
12	Estimating effective population size changes from preferentially sampled genetic sequences. PLoS Computational Biology, 2020, 16, e1007774.	1.5	14
13	Inferring time-dependent migration and coalescence patterns from genetic sequence and predictor data in structured populations. Virus Evolution, 2019, 5, vez030.	2.2	20
14	The ability of single genes vs full genomes to resolve time and space in outbreak analysis. BMC Evolutionary Biology, 2019, 19, 232.	3.2	35
15	MERS coronaviruses from camels in Africa exhibit region-dependent genetic diversity. Proceedings of the United States of America, 2018, 115, 3144-3149.	3.3	142
16	Active Ebola Virus Replication and Heterogeneous Evolutionary Rates in EVD Survivors. Cell Reports, 2018, 22, 1159-1168.	2.9	37
17	MERS-CoV spillover at the camel-human interface. ELife, 2018, 7, .	2.8	172
18	Phylodynamic assessment of intervention strategies for the West African Ebola virus outbreak. Nature Communications, 2018, 9, 2222.	5.8	59

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19	E-104 Virus genomes reveal factors that spread and sustained the Ebola epidemic. Journal of Acquired Immune Deficiency Syndromes (1999), 2018, 77, 43-43.	0.9	0
20	Virus genomes reveal factors that spread and sustained the Ebola epidemic. Nature, 2017, 544, 309-315.	13.7	346
21	Genomic epidemiology reveals multiple introductions of Zika virus into the United States. Nature, 2017, 546, 401-405.	13.7	298
22	Genome-wide evolutionary dynamics of influenza B viruses on a global scale. PLoS Pathogens, 2017, 13, e1006749.	2.1	78
23	The evolution of Ebola virus: Insights from the 2013–2016 epidemic. Nature, 2016, 538, 193-200.	13.7	264
24	Preliminary Evaluation of the Effect of Investigational Ebola Virus Disease Treatments on Viral Genome Sequences. Journal of Infectious Diseases, 2016, 214, S333-S341.	1.9	11
25	Comment on "Mutation rate and genotype variation of Ebola virus from Mali case sequences― Science, 2016, 353, 658-658.	6.0	6
26	Rapid outbreak sequencing of Ebola virus in Sierra Leone identifies transmission chains linked to sporadic cases. Virus Evolution, 2016, 2, vew016.	2.2	105
27	MERS-CoV recombination: implications about the reservoir and potential for adaptation. Virus Evolution, 2016, 2, vev023.	2.2	60
28	Real-time, portable genome sequencing for Ebola surveillance. Nature, 2016, 530, 228-232.	13.7	1,179
29	The global antigenic diversity of swine influenza A viruses. ELife, 2016, 5, e12217.	2.8	146
30	Are arthropods at the heart of virus evolution?. ELife, 2015, 4, .	2.8	26
31	Reassortment between Influenza B Lineages and the Emergence of a Coadapted PB1–PB2–HA Gene Complex. Molecular Biology and Evolution, 2015, 32, 162-172.	3.5	63
32	Evolution and Spread of Ebola Virus in Liberia, 2014–2015. Cell Host and Microbe, 2015, 18, 659-669.	5.1	87
33	Ebola Virus Epidemiology, Transmission, and Evolution during Seven Months in Sierra Leone. Cell, 2015, 161, 1516-1526.	13.5	275
34	Temporal and spatial analysis of the 2014–2015 Ebola virus outbreak in West Africa. Nature, 2015, 524, 97-101.	13.7	272
35	Integrating influenza antigenic dynamics with molecular evolution. ELife, 2014, 3, e01914.	2.8	299
36	Genomic surveillance elucidates Ebola virus origin and transmission during the 2014 outbreak. Science, 2014, 345, 1369-1372.	6.0	1,083

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37	The genetics of host–virus coevolution in invertebrates. Current Opinion in Virology, 2014, 8, 73-78.	2.6	35
38	Phylogenetic Analysis of Guinea 2014 EBOV Ebolavirus Outbreak. PLOS Currents, 2014, 6, .	1.4	62