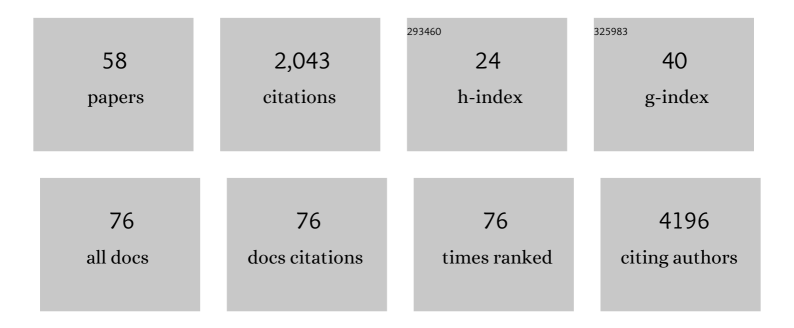
## **Stefan Prost**

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/5377029/publications.pdf Version: 2024-02-01



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#	Article	IF	CITATIONS
1	Genome-wide diversity loss in reintroduced Eurasian lynx populations urges immediate conservation management. Biological Conservation, 2022, 266, 109442.	1.9	18
2	Rapid in situ identification of biological specimens via DNA amplicon sequencing using miniaturized laboratory equipment. Nature Protocols, 2022, 17, 1415-1443.	5.5	23
3	Climate change leads to higher NPP at the end of the century in the Antarctic Tundra: Response patterns through the lens of lichens. Science of the Total Environment, 2022, 835, 155495.	3.9	6
4	Genomic analyses show extremely perilous conservation status of African and Asiatic cheetahs ( <i>Acinonyx jubatus</i> ). Molecular Ecology, 2022, 31, 4208-4223.	2.0	21
5	Unlocking the potential of a validated single nucleotide polymorphism array for genomic monitoring of trade in cheetahs (Acinonyx jubatus). Molecular Biology Reports, 2021, 48, 171-181.	1.0	6
6	NGSpeciesID: DNA barcode and amplicon consensus generation from longâ€read sequencing data. Ecology and Evolution, 2021, 11, 1392-1398.	0.8	30
7	Recent Evolutionary History of Tigers Highlights Contrasting Roles of Genetic Drift and Selection. Molecular Biology and Evolution, 2021, 38, 2366-2379.	3.5	35
8	Ecological Specialization and Evolutionary Reticulation in Extant Hyaenidae. Molecular Biology and Evolution, 2021, 38, 3884-3897.	3.5	15
9	Genomic insights into the conservation status of the world's last remaining Sumatran rhinoceros populations. Nature Communications, 2021, 12, 2393.	5.8	39
10	Nanopore sequencing in non-human forensic genetics. Emerging Topics in Life Sciences, 2021, 5, 465-473.	1.1	5
11	Developmental validation of Oxford Nanopore Technology MinION sequence data and the NGSpeciesID bioinformatic pipeline for forensic genetic species identification. Forensic Science International: Genetics, 2021, 53, 102493.	1.6	31
12	Chromosome-level reference genome of the European wasp spider <i>Argiope bruennichi</i> : a resource for studies on range expansion and evolutionary adaptation. GigaScience, 2021, 10, .	3.3	35
13	Evidence of two deeply divergent co-existing mitochondrial genomes in the Tuatara reveals an extremely complex genomic organization. Communications Biology, 2021, 4, 116.	2.0	16
14	Rhinoceros genomes uncover family secrets. Nature, 2021, 599, 209-210.	13.7	0
15	The <i>Tetragnatha kauaiensis</i> Genome Sheds Light on the Origins of Genomic Novelty in Spiders. Genome Biology and Evolution, 2021, 13, .	1.1	16
16	Functional analysis of natural <scp>PCSK</scp> 9 mutants in modern and archaic humans. FEBS Journal, 2020, 287, 515-528.	2.2	8
17	Tissue- and Population-Level Microbiome Analysis of the Wasp Spider Argiope bruennichi Identified a Novel Dominant Bacterial Symbiont. Microorganisms, 2020, 8, 8.	1.6	26
18	Chromosome-level de novo assembly of the pig-tailed macaque genome using linked-read sequencing and HiC proximity scaffolding. GigaScience, 2020, 9, .	3.3	6

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19	The tuatara genome reveals ancient features of amniote evolution. Nature, 2020, 584, 403-409.	13.7	105
20	Pre-extinction Demographic Stability and Genomic Signatures of Adaptation in the Woolly Rhinoceros. Current Biology, 2020, 30, 3871-3879.e7.	1.8	41
21	Improving the Chromosome-Level Genome Assembly of the Siamese Fighting Fish ( <i>Betta) Tj ETQq1 1 0.78431</i>	4 rgBT /O	verlock 10 Tf
22	Education in the genomics era: Generating high-quality genome assemblies in university courses. GigaScience, 2020, 9, .	3.3	9
23	Interspecific Gene Flow and the Evolution of Specialization in Black and White Rhinoceros. Molecular Biology and Evolution, 2020, 37, 3105-3117.	3.5	20
24	Rigorous wildlife disease surveillance. Science, 2020, 369, 145-147.	6.0	78
25	High-throughput sequencing for community analysis: the promise of DNA barcoding to uncover diversity, relatedness, abundances and interactions in spider communities. Development Genes and Evolution, 2020, 230, 185-201.	0.4	39
26	MinION-Based DNA Barcoding of Preserved and Non-Invasively Collected Wildlife Samples. Genes, 2020, 11, 445.	1.0	30
27	Portable sequencing as a teaching tool in conservation and biodiversity research. PLoS Biology, 2020, 18, e3000667.	2.6	37
28	Host and geography together drive early adaptive radiation of Hawaiian planthoppers. Molecular Ecology, 2019, 28, 4513-4528.	2.0	6
29	Genetic Biomonitoring and Biodiversity Assessment Using Portable Sequencing Technologies: Current Uses and Future Directions. Genes, 2019, 10, 858.	1.0	69
30	Comparative analyses identify genomic features potentially involved in the evolution of birds-of-paradise. GigaScience, 2019, 8, .	3.3	22
31	Nanopore sequencing of long ribosomal DNA amplicons enables portable and simple biodiversity assessments with high phylogenetic resolution across broad taxonomic scale. GigaScience, 2019, 8, .	3.3	126
32	The Genome and mRNA Transcriptome of the Cosmopolitan Calanoid Copepod Acartia tonsa Dana Improve the Understanding of Copepod Genome Size Evolution. Genome Biology and Evolution, 2019, 11, 1440-1450.	1.1	26
33	Dynamic evolutionary history and gene content of sex chromosomes across diverse songbirds. Nature Ecology and Evolution, 2019, 3, 834-844.	3.4	68
34	Mitogenomic evidence of close relationships between New Zealand's extinct giant raptors and small-sized Australian sister-taxa. Molecular Phylogenetics and Evolution, 2019, 134, 122-128.	1.2	17
35	Annotated Draft Genomes of Two Caddisfly Species Plectrocnemia conspersa CURTIS and Hydropsyche tenuis NAVAS (Insecta: Trichoptera). Genome Biology and Evolution, 2019, 11, 3445-3451.	1.1	21
36	Cost-effective assembly of the African wild dog ( <i>Lycaon pictus</i> ) genome using linked reads. GigaScience, 2019, 8, .	3.3	22

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37	Real-time DNA barcoding in a rainforest using nanopore sequencing: opportunities for rapid biodiversity assessments and local capacity building. GigaScience, 2018, 7, .	3.3	176
38	Rapid divergence of mussel populations despite incomplete barriers to dispersal. Molecular Ecology, 2018, 27, 1556-1571.	2.0	29
39	Draft Genome Sequence and Annotation of the Lichen-Forming Fungus Arthonia radiata. Genome Announcements, 2018, 6, .	0.8	12
40	Co-occurrence of ecologically similar species of Hawaiian spiders reveals critical early phase of adaptive radiation. BMC Evolutionary Biology, 2018, 18, 100.	3.2	20
41	Ancient mitogenomes of Phoenicians from Sardinia and Lebanon: A story of settlement, integration, and female mobility. PLoS ONE, 2018, 13, e0190169.	1.1	40
42	The Proprotein Convertases in Hypercholesterolemia and Cardiovascular Diseases: Emphasis on Proprotein Convertase Subtilisin/Kexin 9. Pharmacological Reviews, 2017, 69, 33-52.	7.1	90
43	Synchronous genetic turnovers across Western Eurasia in Late Pleistocene collared lemmings. Global Change Biology, 2016, 22, 1710-1721.	4.2	45
44	First complete mitochondrial genome data from ancient South American camelids - The mystery of the chilihueques from Isla Mocha (Chile). Scientific Reports, 2016, 6, 38708.	1.6	9
45	Geographically contrasting biodiversity reductions in a widespread New Zealand seabird. Molecular Ecology, 2015, 24, 4605-4616.	2.0	22
46	Complete Mitochondrial Genomes of New Zealand's First Dogs. PLoS ONE, 2015, 10, e0138536.	1.1	26
47	From cheek swabs to consensus sequences: an A to Z protocol for high-throughput DNA sequencing of complete human mitochondrial genomes. BMC Genomics, 2014, 15, 68.	1.2	27
48	Integrating multiple lines of evidence into historical biogeography hypothesis testing: a <i>Bison bison</i> case study. Proceedings of the Royal Society B: Biological Sciences, 2014, 281, 20132782.	1.2	41
49	Extinction and recolonization of coastal megafauna following human arrival in New Zealand. Proceedings of the Royal Society B: Biological Sciences, 2014, 281, 20140097.	1.2	53
50	Ancient mitochondrial <scp>DNA</scp> and the genetic history of <scp>E</scp> urasian beaver ( <i><scp>C</scp>astor fiber</i> ) in <scp>E</scp> urope. Molecular Ecology, 2014, 23, 1717-1729.	2.0	24
51	Effects of late quaternary climate change on <scp>P</scp> alearctic shrews. Global Change Biology, 2013, 19, 1865-1874.	4.2	24
52	Losing ground: past history and future fate of <scp>A</scp> rctic small mammals in a changing climate. Global Change Biology, 2013, 19, 1854-1864.	4.2	46
53	The Genetic Diversity of the Nguni Breed of African Cattle (Bos spp.): Complete Mitochondrial Genomes of Haplogroup T1. PLoS ONE, 2013, 8, e71956.	1.1	36
54	Complete mitochondrial DNA genome sequences from the first New Zealanders. Proceedings of the National Academy of Sciences of the United States of America, 2012, 109, 18350-18354.	3.3	51

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55	TempNet: a method to display statistical parsimony networks for heterochronous DNA sequence data. Methods in Ecology and Evolution, 2011, 2, 663-667.	2.2	98
56	Influence of Climate Warming on Arctic Mammals? New Insights from Ancient DNA Studies of the Collared Lemming Dicrostonyx torquatus. PLoS ONE, 2010, 5, e10447.	1.1	48
57	A population history of Tokelau – genetic variation and change in atoll populations. Journal of Island and Coastal Archaeology, 0, , 1-18.	0.6	0
58	Chromosome-level genome assembly of a benthic associated Syngnathiformes species: the common dragonet, Callionymus lyra. GigaByte, 0, 2020, 1-10.	0.0	7