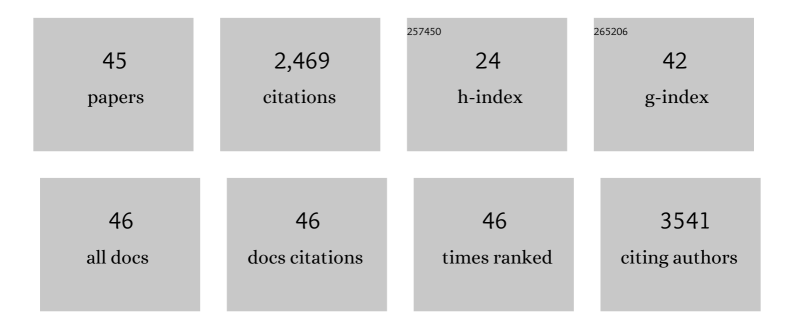
Anna Santure

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	The relevance of pedigrees in the conservation genomics era. Molecular Ecology, 2022, 31, 41-54.	3.9	24
2	Finding the adaptive needles in a populationâ€structured haystack: A case study in a New Zealand mollusc. Journal of Animal Ecology, 2022, 91, 1209-1221.	2.8	3
3	Genetic variance in fitness indicates rapid contemporary adaptive evolution in wild animals. Science, 2022, 376, 1012-1016.	12.6	69
4	Genomic data of different resolutions reveal consistent inbreeding estimates but contrasting homozygosity landscapes for the threatened Aotearoa New Zealand hihi. Molecular Ecology, 2021, 30, 6006-6020.	3.9	25
5	The design and application of a 50 K SNP chip for a threatened Aotearoa New Zealand passerine, the hihi. Molecular Ecology Resources, 2021, , .	4.8	1
6	Expanding the conservation genomics toolbox: Incorporating structural variants to enhance genomic studies for species of conservation concern. Molecular Ecology, 2021, 30, 5949-5965.	3.9	26
7	Who are you? A framework to identify and report genetic sample mixâ€ups. Molecular Ecology Resources, 2021, , .	4.8	0
8	Consequences of space sharing on individual phenotypes in the New Zealand hihi. Evolutionary Ecology, 2020, 34, 821-839.	1.2	5
9	Polygenic basis for adaptive morphological variation in a threatened Aotearoa New Zealand bird, the hihi (<i>Notiomystis cincta</i>). Proceedings of the Royal Society B: Biological Sciences, 2020, 287, 20200948.	2.6	23
10	Hitchhiking consequences for genetic and morphological patterns: the influence of kelp-rafting on a brooding chiton. Biological Journal of the Linnean Society, 2020, 130, 756-770.	1.6	6
11	A comparison of pedigree, genetic and genomic estimates of relatedness for informing pairing decisions in two critically endangered birds: Implications for conservation breeding programmes worldwide. Evolutionary Applications, 2020, 13, 991-1008.	3.1	48
12	Can threatened species adapt in a restored habitat? No expected evolutionary response in lay date for the New Zealand hihi. Evolutionary Applications, 2019, 12, 482-497.	3.1	17
13	Little Adaptive Potential in a Threatened Passerine Bird. Current Biology, 2019, 29, 889-894.e3.	3.9	53
14	Reference Genomes from Distantly Related Species Can Be Used for Discovery of Single Nucleotide Polymorphisms to Inform Conservation Management. Genes, 2019, 10, 9.	2.4	50
15	A highâ€density <scp>SNP</scp> chip for genotyping great tit (<i>Parus major</i>) populations and its application to studying the genetic architecture of exploration behaviour. Molecular Ecology Resources, 2018, 18, 877-891.	4.8	36
16	Wild GWAS—association mapping in natural populations. Molecular Ecology Resources, 2018, 18, 729-738.	4.8	79
17	The Evolutionary Biology, Ecology and Epidemiology of Coccidia of Passerine Birds. Advances in Parasitology, 2018, 99, 35-60.	3.2	8
18	Genetic Kinship Analyses Reveal That Gray's Beaked Whales Strand in Unrelated Groups. Journal of Heredity, 2017, 108, 456-461.	2.4	6

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19	Social and spatial effects on genetic variation between foraging flocks in a wild bird population. Molecular Ecology, 2017, 26, 5807-5819.	3.9	8
20	Recombination: the good, the bad and the variable. Philosophical Transactions of the Royal Society B: Biological Sciences, 2017, 372, 20170279.	4.0	39
21	Variation in recombination frequency and distribution across eukaryotes: patterns and processes. Philosophical Transactions of the Royal Society B: Biological Sciences, 2017, 372, 20160455.	4.0	306
22	An ecological model organism flies into the genomics era. Molecular Ecology Resources, 2016, 16, 379-381.	4.8	0
23	Building strong relationships between conservation genetics and primary industry leads to mutually beneficial genomic advances. Molecular Ecology, 2016, 25, 5267-5281.	3.9	16
24	Replicated analysis of the genetic architecture of quantitative traits in two wild great tit populations. Molecular Ecology, 2015, 24, 6148-6162.	3.9	61
25	Determinants of male floating behaviour and floater reproduction in a threatened population of the hihi (<i>Notiomystis cincta</i>). Evolutionary Applications, 2015, 8, 796-806.	3.1	19
26	No evidence for <scp>MHC</scp> class lâ€based disassortative mating in a wild population of great tits. Journal of Evolutionary Biology, 2015, 28, 642-654.	1.7	19
27	Transposable elements as agents of rapid adaptation may explain the genetic paradox of invasive species. Molecular Ecology, 2015, 24, 2241-2252.	3.9	178
28	The influence of nonrandom extra-pair paternity on heritability estimates derived from wild pedigrees. Evolution; International Journal of Organic Evolution, 2015, 69, 1336-1344.	2.3	42
29	Highly Variable Recombinational Landscape Modulates Efficacy of Natural Selection in Birds. Genome Biology and Evolution, 2014, 6, 2061-2075.	2.5	36
30	Replicated high-density genetic maps of two great tit populations reveal fine-scale genomic departures from sex-equal recombination rates. Heredity, 2014, 112, 307-316.	2.6	53
31	Genomic dissection of variation in clutch size and egg mass in a wild great tit (<i>Parus major</i>) population. Molecular Ecology, 2013, 22, 3949-3962.	3.9	93
32	Partitioning of genetic variation across the genome using multimarker methods in a wild bird population. Molecular Ecology, 2013, 22, 3963-3980.	3.9	78
33	Fewer invited talks by women in evolutionary biology symposia. Journal of Evolutionary Biology, 2013, 26, 2063-2069.	1.7	120
34	FINE-SCALE GENETIC STRUCTURE IN A WILD BIRD POPULATION: THE ROLE OF LIMITED DISPERSAL AND ENVIRONMENTALLY BASED SELECTION AS CAUSAL FACTORS. Evolution; International Journal of Organic Evolution, 2013, 67, 3488-3500.	2.3	44
35	Genomic Imprinting Leads to Less Selectively Maintained Polymorphism on X Chromosomes. Genetics, 2012, 192, 1455-1464.	2.9	2
36	The design and crossâ€population application of a genomeâ€wide SNP chip for the great tit <i>Parus major</i> . Molecular Ecology Resources, 2012, 12, 753-770.	4.8	56

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37	High genetic diversity in the remnant island population of hihi and the genetic consequences of re-introduction. Molecular Ecology, 2011, 20, 29-45.	3.9	63
38	Characterisation of the transcriptome of a wild great tit Parus major population by next generation sequencing. BMC Genomics, 2011, 12, 283.	2.8	67
39	Quantitative Genetics of Genomic Imprinting: A Comparison of Simple Variance Derivations, the Effects of Inbreeding, and Response to Selection. G3: Genes, Genomes, Genetics, 2011, 1, 131-142.	1.8	12
40	Genome mapping in intensively studied wild vertebrate populations. Trends in Genetics, 2010, 26, 275-284.	6.7	85
41	On the use of large marker panels to estimate inbreeding and relatedness: empirical and simulation studies of a pedigreed zebra finch population typed at 771 SNPs. Molecular Ecology, 2010, 19, 1439-1451.	3.9	130
42	The Joint Effects of Selection and Dominance on the <i>Q</i> ST â^' <i>F</i> ST Contrast. Genetics, 2009, 181, 259-276.	2.9	18
43	Parentage and Sibship Inference From Multilocus Genotype Data Under Polygamy. Genetics, 2009, 181, 1579-1594.	2.9	400
44	Influence of Mom and Dad: Quantitative Genetic Models for Maternal Effects and Genomic Imprinting. Genetics, 2006, 173, 2297-2316.	2.9	26
45	Population structure in the barn swallow, Hirundo rustica: a comparison between neutral DNA markers and quantitative traits. Biological Journal of the Linnean Society, 0, 99, 306-314.	1.6	19