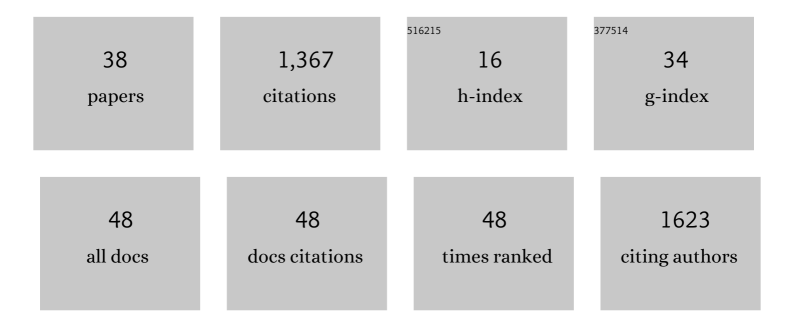
## Gabriel Rodrigues Alves Margarido

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/3509547/publications.pdf

Version: 2024-02-01



## GABRIEL RODRIGUES ALVES

#	Article	IF	CITATIONS
1	Means, motive and opportunity for biological invasions: Genetic introgression in a fungal pathogen. Molecular Ecology, 2023, 32, 2428-2442.	2.0	10
2	Genomic prediction with allele dosage information in highly polyploid species. Theoretical and Applied Genetics, 2022, 135, 723-739.	1.8	13
3	Transcriptome changes in the developing sugarcane culm associated with high yield and early-season high sugar content. Theoretical and Applied Genetics, 2022, 135, 1619-1636.	1.8	1
4	Limited allele-specific gene expression in highly polyploid sugarcane. Genome Research, 2022, 32, 297-308.	2.4	8
5	Allele expression biases in mixed-ploid sugarcane accessions. Scientific Reports, 2022, 12, .	1.6	1
6	Long-term comparison between index selection and optimal independent culling in plant breeding programs with genomic prediction. PLoS ONE, 2021, 16, e0235554.	1.1	11
7	The Wild Sugarcane and Sorghum Kinomes: Insights Into Expansion, Diversification, and Expression Patterns. Frontiers in Plant Science, 2021, 12, 668623.	1.7	18
8	A Missense Mutation in the MYBPH Gene Is Associated With Abdominal Fat Traits in Meat-Type Chickens. Frontiers in Genetics, 2021, 12, 698163.	1.1	4
9	Modern Approaches for Transcriptome Analyses in Plants. Advances in Experimental Medicine and Biology, 2021, 1346, 11-50.	0.8	Ο
10	Temporal Gene Expression in Apical Culms Shows Early Changes in Cell Wall Biosynthesis Genes in Sugarcane. Frontiers in Plant Science, 2021, 12, 736797.	1.7	1
11	Differential expression in leaves of Saccharum genotypes contrasting in biomass production provides evidence of genes involved in carbon partitioning. BMC Genomics, 2020, 21, 673.	1.2	10
12	Genomic prediction applied to multiple traits and environments in second season maize hybrids. Heredity, 2020, 125, 60-72.	1.2	12
13	Boosting predictive ability of tropical maize hybrids via genotypeâ€byâ€environment interaction under multivariate GBLUP models. Crop Science, 2020, 60, 3049-3065.	0.8	15
14	Single nucleotide polymorphism calling and imputation strategies for costâ€effective genotyping in a tropical maize breeding program. Crop Science, 2020, 60, 3066-3082.	0.8	4
15	Time-series expression profiling of sugarcane leaves infected with Puccinia kuehnii reveals an ineffective defense system leading to susceptibility. Plant Cell Reports, 2020, 39, 873-889.	2.8	25
16	Genomic resources for energy cane breeding in the post genomics era. Computational and Structural Biotechnology Journal, 2019, 17, 1404-1414.	1.9	38
17	Evidence for Strong Kinship Influence on the Extent of Linkage Disequilibrium in Cultivated Common Beans. Genes, 2019, 10, 5.	1.0	32
18	Assembly of the 373k gene space of the polyploid sugarcane genome reveals reservoirs of functional diversity in the world's leading biomass crop. GigaScience, 2019, 8, .	3.3	106

GABRIEL RODRIGUES ALVES

#	Article	IF	CITATIONS
19	Differential Gene Expression Among Genotypes of the Genus Saccharum Contrasting in Biomass Production. Proceedings (mdpi), 2019, 36, .	0.2	0
20	Genomic prediction applied to high-biomass sorghum for bioenergy production. Molecular Breeding, 2018, 38, 49.	1.0	28
21	A fully automated pipeline for quantitative genotype calling from next generation sequencing data in autopolyploids. BMC Bioinformatics, 2018, 19, 398.	1.2	62
22	Linkage Disequilibrium and Population Structure in Wild and Cultivated Populations of Rubber Tree (Hevea brasiliensis). Frontiers in Plant Science, 2018, 9, 815.	1.7	20
23	GBS-based single dosage markers for linkage and QTL mapping allow gene mining for yield-related traits in sugarcane. BMC Genomics, 2017, 18, 72.	1.2	91
24	Restriction site associated <scp>DNA</scp> ( <scp>RAD</scp> ) for de novo sequencing and marker discovery in sugarcane borer, <i>Diatraea saccharalis</i> Fab. (Lepidoptera: Crambidae). Molecular Ecology Resources, 2017, 17, 454-465.	2.2	5
25	Molecular markers for conservation genetic resources of four Passiflora species. Scientia Horticulturae, 2016, 212, 251-261.	1.7	5
26	ConPADE: Genome Assembly Ploidy Estimation from Next-Generation Sequencing Data. PLoS Computational Biology, 2015, 11, e1004229.	1.5	41
27	Multi-trait multi-environment quantitative trait loci mapping for a sugarcane commercial cross provides insights on the inheritance of important traits. Molecular Breeding, 2015, 35, 175.	1.0	19
28	A model for quantitative trait loci mapping, linkage phase, and segregation pattern estimation for a full-sib progeny. Tree Genetics and Genomes, 2014, 10, 791-801.	0.6	34
29	Studying the genetic basis of drought tolerance in sorghum by managed stress trials and adjustments for phenological and plant height differences. Theoretical and Applied Genetics, 2012, 124, 1389-1402.	1.8	98
30	A mixed model QTL analysis for sugarcane multiple-harvest-location trial data. Theoretical and Applied Genetics, 2012, 124, 835-849.	1.8	73
31	The Biotechnology Roadmap for Sugarcane Improvement. Tropical Plant Biology, 2010, 3, 75-87.	1.0	62
32	Precision of distances and ordering of microsatellite markers in consensus linkage maps of chromosomes 1, 3 and 4 from two reciprocal chicken populations using bootstrap sampling. Genetics and Molecular Research, 2010, 9, 1357-1376.	0.3	4
33	Evaluation of algorithms used to order markers on genetic maps. Heredity, 2009, 103, 494-502.	1.2	41
34	An Integrated Molecular Map of Yellow Passion Fruit Based on Simultaneous Maximum-likelihood Estimation of Linkage and Linkage Phases. Journal of the American Society for Horticultural Science, 2008, 133, 35-41.	0.5	44
35	Comparação dos algoritmos delineação rápida em cadeia e seriação, para a construção de mapas genéticos. Pesquisa Agropecuaria Brasileira, 2008, 43, 505-512.	0.9	1
36	OneMap: software for genetic mapping in outcrossing species. Hereditas, 2007, 144, 78-79.	0.5	326

#	Article	IF	CITATIONS
37	Functional integrated genetic linkage map based on EST-markers for a sugarcane (Saccharum spp.) commercial cross. Molecular Breeding, 2007, 20, 189-208.	1.0	79
38	Improvements in the sequencing and assembly of plant genomes. GigaByte, 0, 2021, 1-10.	0.0	15