Volker P Brendel

List of Publications by Year in descending order

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Version: 2024-02-01

64 papers 4,608 citations

34 h-index 62 g-index

68 all docs

68 docs citations

68 times ranked 6694 citing authors

#	Article	IF	Citations
1	TAL Effector-Nucleotide Targeter (TALE-NT) 2.0: tools for TAL effector design and target prediction. Nucleic Acids Research, 2012, 40, W117-W122.	14.5	549
2	Genomewide comparative analysis of alternative splicing in plants. Proceedings of the National Academy of Sciences of the United States of America, 2006, 103, 7175-7180.	7.1	481
3	Engineering a software tool for gene structure prediction in higher organisms. Information and Software Technology, 2005, 47, 965-978.	4.4	251
4	PlantGDB: a resource for comparative plant genomics. Nucleic Acids Research, 2007, 36, D959-D965.	14.5	246
5	Single Nucleotide Polymorphisms of Human STING Can Affect Innate Immune Response to Cyclic Dinucleotides. PLoS ONE, 2013, 8, e77846.	2.5	209
6	Two New Complete Genome Sequences Offer Insight into Host and Tissue Specificity of Plant Pathogenic Xanthomonas spp. Journal of Bacteriology, 2011, 193, 5450-5464.	2.2	189
7	Genome, transcriptome and methylome sequencing of a primitively eusocial wasp reveal a greatly reduced <scp>DNA</scp> methylation system in a social insect. Molecular Ecology, 2016, 25, 1769-1784.	3.9	148
8	Linguistics of Nucleotide Sequences: Morphology and Comparison of Vocabularies. Journal of Biomolecular Structure and Dynamics, 1986, 4, 11-21.	3.5	146
9	Evolutionary Comparisons of RecA-Like Proteins Across All Major Kingdoms of Living Organisms. Journal of Molecular Evolution, 1997, 44, 528-541.	1.8	125
10	Genome-Wide Distribution of Transposed < i > Dissociation < /i > Elements in Maize Â. Plant Cell, 2010, 22, 1667-1685.	6.6	123
11	Terminators of Transcription with RNA Polymerase from (i>Escherichia coli:) What They Look Like and How to Find Them. Journal of Biomolecular Structure and Dynamics, 1986, 3, 705-723.	3.5	118
12	The Maize Genome Contains a Helitron Insertion. Plant Cell, 2003, 15, 381-391.	6.6	115
13	Refined Annotation of the Arabidopsis Genome by Complete Expressed Sequence Tag Mapping. Plant Physiology, 2003, 132, 469-484.	4.8	111
14	Expression and evolution of the phosphoinositide-specific phospholipase C gene family in Arabidopsis thaliana. Plant Physiology and Biochemistry, 2008, 46, 627-637.	5 . 8	106
15	Gene structure prediction from consensus spliced alignment of multiple ESTs matching the same genomic locus. Bioinformatics, 2004, 20, 1157-1169.	4.1	97
16	Comparison of RNA Expression Profiles Based on Maize Expressed Sequence Tag Frequency Analysis and Micro-Array Hybridization. Plant Physiology, 2002, 128, 896-910.	4.8	96
17	Comparative Plant Genomics Resources at PlantGDB. Plant Physiology, 2005, 139, 610-618.	4.8	95
18	Acquisition and Evolution of Plant Pathogenesis–Associated Gene Clusters and Candidate Determinants of Tissue-Specificity in Xanthomonas. PLoS ONE, 2008, 3, e3828.	2.5	89

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19	The Maize Genome Sequencing Project. Plant Physiology, 2002, 130, 1594-1597.	4.8	69
20	Efficient clustering of large EST data sets on parallel computers. Nucleic Acids Research, 2003, 31, 2963-2974.	14.5	69
21	U-richness is a defining feature of plant introns and may function as an intron recognition signal in maize. Plant Molecular Biology, 1998, 36, 573-583.	3.9	65
22	Identification of Biased Amino Acid Substitution Patterns in Human Immunodeficiency Virus Type 1 Isolates from Patients Treated with Protease Inhibitors. Journal of Virology, 1999, 73, 6197-6202.	3.4	65
23	Identification, characterization and molecular phylogeny of U12-dependent introns in the Arabidopsis thaliana genome. Nucleic Acids Research, 2003, 31, 4561-4572.	14.5	61
24	Gene structure prediction by spliced alignment of genomic DNA with protein sequences: increased accuracy by differential splice site scoring11Edited by G. von Heijne. Journal of Molecular Biology, 2000, 297, 1075-1085.	4.2	59
25	GeneSeqer@PlantGDB: gene structure prediction in plant genomes. Nucleic Acids Research, 2003, 31, 3597-3600.	14.5	59
26	The Maize Genetics and Genomics Database. The Community Resource for Access to Diverse Maize Data. Plant Physiology, 2005, 138, 55-58.	4.8	59
27	Cross-species EST alignments reveal novel and conserved alternative splicing events in legumes. BMC Plant Biology, 2008, 8, 17.	3.6	49
28	Quantitative Model of ColE1 Plasmid Copy Number Control. Journal of Molecular Biology, 1993, 229, 860-872.	4.2	46
29	Molecular Characterization and Phylogeny of U2AF35 Homologs in Plants. Plant Physiology, 2006, 140, 624-636.	4.8	46
30	A method to identify distinctive charge configurations in protein sequences, with application to human herpesvirus polypeptides. Journal of Molecular Biology, 1989, 205, 165-177.	4.2	41
31	Discovery and Expression Analysis of Alternative Splicing Events Conserved among Plant SR Proteins. Molecular Biology and Evolution, 2014, 31, 605-613.	8.9	40
32	A combinatorial role for exon, intron and splice site sequences in splicing in maize. Plant Journal, 1997, 11, 1253-1263.	5.7	39
33	PROSET—a fast procedure to create non-redundant sets of protein sequences. Mathematical and Computer Modelling, 1992, 16, 37-43.	2.0	37
34	Too many leucine zippers?. Nature, 1989, 341, 574-575.	27.8	35
35	ZmDB, an integrated database for maize genome research. Nucleic Acids Research, 2003, 31, 244-247.	14.5	33
36	Intervening Sequences Exhibit Distinct Vocabulary. Journal of Biomolecular Structure and Dynamics, 1986, 4, 391-400.	3.5	31

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37	Changes in gene DNA methylation and expression networks accompany caste specialization and ageâ€related physiological changes in a social insect. Molecular Ecology, 2019, 28, 1975-1993.	3.9	30
38	The Xa7 resistance gene guards the rice susceptibility gene SWEET14 against exploitation by the bacterial blight pathogen. Plant Communications, 2021, 2, 100164.	7.7	30
39	Common introns within orthologous genes: software and application to plants. Briefings in Bioinformatics, 2009, 10, 631-644.	6.5	29
40	Prediction of splice sites in plant pre-mRNA from sequence properties. Journal of Molecular Biology, 1998, 276, 85-104.	4.2	28
41	Community-based gene structure annotation. Trends in Plant Science, 2005, 10, 9-14.	8.8	24
42	ParsEval: parallel comparison and analysis of gene structure annotations. BMC Bioinformatics, 2012, 13, 187.	2.6	24
43	Simple and efficient profiling of transcription initiation and transcript levels with STRIPE-seq. Genome Research, 2020, 30, 910-923.	5.5	23
44	Comparative EST Analyses in Plant Systems. Methods in Enzymology, 2005, 395, 400-418.	1.0	22
45	Kinetics of complementary RNA-RNA interaction involved in plasmid ColE1 copy number control. Journal of Molecular Biology, 1989, 208, 245-255.	4.2	21
46	yrGATE: a web-based gene-structure annotation tool for the identification and dissemination of eukaryotic genes. Genome Biology, 2006, 7, R58.	9.6	21
47	Promoter Architecture and Sex-Specific Gene Expression in <i>Daphnia pulex</i> . Genetics, 2016, 204, 593-612.	2.9	20
48	xGDB: open-source computational infrastructure for the integrated evaluation and analysis of genome features. Genome Biology, 2006, 7, R111.	9.6	17
49	Pairwise statistical significance and empirical determination of effective gap opening penalties for protein local sequence alignment. International Journal of Computational Biology and Drug Design, 2008, 1, 347.	0.3	16
50	Test of the combinatorial model of intron recognition in a native maize gene. Plant Molecular Biology, 1999, 41, 637-644.	3.9	15
51	Tracembler – software for in-silico chromosome walking in unassembled genomes. BMC Bioinformatics, 2007, 8, 151.	2.6	12
52	Kinetic Analysis of Adsorption Processes. SIAM Journal on Applied Mathematics, 1987, 47, 1306-1319.	1.8	11
53	The Next Generation of Training for Arabidopsis Researchers: Bioinformatics and Quantitative Biology. Plant Physiology, 2017, 175, 1499-1509.	4.8	11
54	Flexible analysis of TSS mapping data and detection of TSS shifts with TSRexploreR. NAR Genomics and Bioinformatics, 2021, 3, lqab051.	3.2	11

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55	A comparative analysis of distinctive features of yeast protein sequences. Yeast, 1993, 9, 1287-1298.	1.7	10
56	SRAssembler: Selective Recursive local Assembly of homologous genomic regions. BMC Bioinformatics, 2019, 20, 371.	2.6	8
57	Computational modeling of gene structure in Arabidopsis thaliana. Plant Molecular Biology, 2002, 48, 49-58.	3.9	8
58	On modes of recombination, replication, and segregation of the higher plant mitochondrial genome. Journal of Theoretical Biology, 1987, 125, 163-176.	1.7	4
59	Gene Structure Annotation at PlantGDB. , 2007, 406, 521-533.		4
60	Tools and applications for integrative analysis of DNA methylation in social insects. Molecular Ecology Resources, 2022, 22, 1656-1674.	4.8	4
61	From small RNA discoveries to a new paradigm in computational genomics?. New Phytologist, 2018, 220, 659-660.	7.3	3
62	Using RAMPAGE to Identify and Annotate Promoters in Insect Genomes. Methods in Molecular Biology, 2019, 1858, 99-116.	0.9	3
63	xGDBvm: A Web GUI-Driven Workflow for Annotating Eukaryotic Genomes in the Cloud. Plant Cell, 2016, 28, 840-854.	6.6	2
64	iLoci: robust evaluation of genome content and organization for provisional and mature genome assemblies. NAR Genomics and Bioinformatics, 2022, 4, lqac013.	3.2	O