Victor V Solovyev

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Community structure and metabolism through reconstruction of microbial genomes from the environment. Nature, 2004, 428, 37-43.	13.7	2,045
2	Insights into social insects from the genome of the honeybee Apis mellifera. Nature, 2006, 443, 931-949.	13.7	1,648
3	The genome of the model beetle and pest Tribolium castaneum. Nature, 2008, 452, 949-955.	13.7	1,255
4	The Genome Sequence of Taurine Cattle: A Window to Ruminant Biology and Evolution. Science, 2009, 324, 522-528.	6.0	1,038
5	The Genome of the Sea Urchin Strongylocentrotus purpuratus. Science, 2006, 314, 941-952.	6.0	1,018
6	Ab initio Gene Finding in Drosophila Genomic DNA. Genome Research, 2000, 10, 516-522.	2.4	987
7	A High Quality Draft Consensus Sequence of the Genome of a Heterozygous Grapevine Variety. PLoS ONE, 2007, 2, e1326.	1.1	945
8	Genome Sequence of the Pea Aphid Acyrthosiphon pisum. PLoS Biology, 2010, 8, e1000313.	2.6	913
9	Functional and Evolutionary Insights from the Genomes of Three Parasitoid <i>Nasonia</i> Species. Science, 2010, 327, 343-348.	6.0	808
10	Automatic annotation of eukaryotic genes, pseudogenes and promoters. Genome Biology, 2006, 7, S10.	13.9	749
11	The ctenophore genome and the evolutionary origins of neural systems. Nature, 2014, 510, 109-114.	13.7	606
12	Analysis of canonical and non-canonical splice sites in mammalian genomes. Nucleic Acids Research, 2000, 28, 4364-4375.	6.5	520
13	Assemblathon 1: A competitive assessment of de novo short read assembly methods. Genome Research, 2011, 21, 2224-2241.	2.4	443
14	Finding the missing honey bee genes: lessons learned from a genome upgrade. BMC Genomics, 2014, 15, 86.	1.2	375
15	Predicting internal exons by oligonucleotide composition and discriminant analysis of spliceable open reading frames. Nucleic Acids Research, 1994, 22, 5156-5163.	6.5	333
16	Expression of Msl-2 causes assembly of dosage compensation regulators on the X chromosomes and female lethality in Drosophila. Cell, 1995, 81, 867-877.	13.5	310
17	The DNA sequence and biology of human chromosome 19. Nature, 2004, 428, 529-535.	13.7	298
18	Rice SNP-seek database update: new SNPs, indels, and queries. Nucleic Acids Research, 2017, 45, D1075-D1081.	6.5	290

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19	Prediction of Protein Secondary Structure by Combining Nearest-neighbor Algorithms and Multiple Sequence Alignments. Journal of Molecular Biology, 1995, 247, 11-15.	2.0	280
20	Community-wide assessment of GPCR structure modelling and ligand docking: GPCR Dock 2008. Nature Reviews Drug Discovery, 2009, 8, 455-463.	21.5	260
21	PlantProm: a database of plant promoter sequences. Nucleic Acids Research, 2003, 31, 114-117.	6.5	240
22	SpliceDB: database of canonical and non-canonical mammalian splice sites. Nucleic Acids Research, 2001, 29, 255-259.	6.5	210
23	Recognition of prokaryotic and eukaryotic promoters using convolutional deep learning neural networks. PLoS ONE, 2017, 12, e0171410.	1.1	183
24	Identification of Promoter Regions and Regulatory Sites. Methods in Molecular Biology, 2010, 674, 57-83.	0.4	137
25	A novel type of RNase III family proteins in eukaryotes. Gene, 2000, 245, 213-221.	1.0	120
26	Sequence alignment kernel for recognition of promoter regions. Bioinformatics, 2003, 19, 1964-1971.	1.8	116
27	Protein secondary structure prediction using local alignments. Journal of Molecular Biology, 1997, 268, 31-36.	2.0	115
28	PromH: promoters identification using orthologous genomic sequences. Nucleic Acids Research, 2003, 31, 3540-3545.	6.5	113
29	Plant promoter prediction with confidence estimation. Nucleic Acids Research, 2005, 33, 1069-1076.	6.5	106
30	Alignathon: a competitive assessment of whole-genome alignment methods. Genome Research, 2014, 24, 2077-2089.	2.4	102
31	Promoter analysis and prediction in the human genome using sequence-based deep learning models. Bioinformatics, 2019, 35, 2730-2737.	1.8	89
32	Automated Whole-Genome Multiple Alignment of Rat, Mouse, and Human. Genome Research, 2004, 14, 685-692.	2.4	79
33	Karect: accurate correction of substitution, insertion and deletion errors for next-generation sequencing data. Bioinformatics, 2015, 31, 3421-3428.	1.8	75
34	Performance assessment of promoter predictions on ENCODE regions in the EGASP experiment. Genome Biology, 2006, 7, S3.	13.9	61
35	nGASP – the nematode genome annotation assessment project. BMC Bioinformatics, 2008, 9, 549.	1.2	61
36	TSSPlant: a new tool for prediction of plant Pol II promoters. Nucleic Acids Research, 2017, 45, gkw1353.	6.5	59

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37	Analysis of mutation rates in the SMCY/SMCX genes shows that mammalian evolution is male driven. Mammalian Genome, 1997, 8, 134-138.	1.0	52
38	Abundance of plastid DNA insertions in nuclear genomes of rice and Arabidopsis. Plant Molecular Biology, 2003, 52, 923-934.	2.0	51
39	SNP-Seek II: A resource for allele mining and analysis of big genomic data in Oryza sativa. Current Plant Biology, 2016, 7-8, 16-25.	2.3	48
40	INFOGENE: a database of known gene structures and predicted genes and proteins in sequences of genome sequencing projects. Nucleic Acids Research, 1999, 27, 248-250.	6.5	45
41	Nsite, NsiteH and NsiteM computer tools for studying transcription regulatory elements. Bioinformatics, 2015, 31, 3544-3545.	1.8	41
42	Predicting α-helix and Î ² -strand segments of globular proteins. Bioinformatics, 1994, 10, 661-669.	1.8	37
43	Pecularities of immunoglobulin gene structures as a basis for somatic mutation emergence. FEBS Letters, 1987, 214, 87-91.	1.3	34
44	Assignment of position-specific error probability to primary DNA sequence data. Nucleic Acids Research, 1994, 22, 1272-1280.	6.5	34
45	New insights from Opisthorchis felineus genome: update on genomics of the epidemiologically important liver flukes. BMC Genomics, 2019, 20, 399.	1.2	29
46	Evidence-based gene models for structural and functional annotations of the oil palm genome. Biology Direct, 2017, 12, 21.	1.9	24
47	Intercalary heterochromatin in Drosophila. Chromosoma, 1988, 97, 247-253.	1.0	23
48	A novel method of protein sequence classification based on oligopeptide frequency analysis and its application to search for functional sites and to domain localization. Bioinformatics, 1993, 9, 17-24.	1.8	19
49	Nucleotide patterns aiding in prediction of eukaryotic promoters. PLoS ONE, 2017, 12, e0187243.	1.1	19
50	Fractal graphical representation and analysis of DNA and protein sequences. BioSystems, 1993, 30, 137-160.	0.9	17
51	Towards the integration of genomics, epidemiological and clinical data. Genome Medicine, 2011, 3, 48.	3.6	15
52	Prediction of Metabolic Pathway Involvement in Prokaryotic UniProtKB Data by Association Rule Mining. PLoS ONE, 2016, 11, e0158896.	1.1	13
53	Epigeneticâ€genetic chromatin footprinting identifies novel and subjectâ€specific genes active in prefrontal cortex neurons. FASEB Journal, 2019, 33, 8161-8173.	0.2	12
54	Conformational changes in the globin family during evolution. Journal of Molecular Evolution, 1984, 21, 42-53.	0.8	10

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55	New elements of glucocorticoid-receptor binding sites of hormone-regulated genes. Biochimica Et Biophysica Acta Gene Regulatory Mechanisms, 1991, 1089, 367-376.	2.4	9
56	A simple method for the calculation of low energy packings of α-helices—A threshold approximation. I. The use of the method to estimate the effects of amino acid substitutions, deletions and insertions in globins. Journal of Theoretical Biology, 1984, 110, 67-91.	0.8	4
57	The template RNAs of RNA polymerases can have compact secondary structure, formed by long double helices with partial violations of the complementarity. FEBS Letters, 1984, 165, 72-78.	1.3	4
58	The PlantProm DB: Recent Updates. , 2012, , .		2
59	Genome Sequence of a Multidrug-Resistant Strain of Stenotrophomonas maltophilia with Carbapenem Resistance, Isolated from King Abdullah Medical City, Makkah, Saudi Arabia. Genome Announcements, 2015, 3, .	0.8	2
60	Rule Mining Techniques to Predict Prokaryotic Metabolic Pathways. Methods in Molecular Biology, 2017, 1613, 311-331.	0.4	2
61	Evolution and performance of electron and photon triggers in ATLAS in the year 2011. , 2012, , .		1
62	INTENSIFICATION OF NANOPOROUS GLASS PRODUCING BY POWERFUL ULTRASONIC FIELDS. Integrated Ferroelectrics, 2008, 103, 60-65.	0.3	0
63	Genome-Wide Prokaryotic Promoter Recognition Based on Sequence Alignment Kernel. Lecture Notes in Computer Science, 2003, , 386-396.	1.0	Ο