Marta Kasprzak

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Labeled Graphs in Life Sciences—Two Important Applications. Mechanisms and Machine Science, 2022, , 201-217.	0.3	0
2	Genome-scale <i>de novo</i> assembly using ALGA. Bioinformatics, 2021, 37, 1644-1651.	1.8	7
3	Graph algorithms for DNA sequencing – origins, current models and the future. European Journal of Operational Research, 2018, 264, 799-812.	3.5	16
4	Classification of deÂBruijn-based labeled digraphs. Discrete Applied Mathematics, 2018, 234, 86-92.	0.5	6
5	GRASShopPER—An algorithm for de novo assembly based on GPU alignments. PLoS ONE, 2018, 13, e0202355.	1.1	3
6	Tabu Search for the RNA Partial Degradation Problem. International Journal of Applied Mathematics and Computer Science, 2017, 27, 401-415.	1.5	1
7	High-order statistical compressor for long-term storage of DNA sequencing data. RAIRO - Operations Research, 2016, 50, 351-361.	1.0	2
8	Structural alignment of protein descriptors – a combinatorial model. BMC Bioinformatics, 2016, 17, 383.	1.2	4
9	An Integrated Approach (CLuster Analysis Integration Method) to Combine Expression Data and Protein–Protein Interaction Networks in Agrigenomics: Application on <i>Arabidopsis thaliana</i> . OMICS A Journal of Integrative Biology, 2014, 18, 155-165.	1.0	16
10	Dna Sequence Assembly Involving an Acyclic Graph Model. Foundations of Computing and Decision Sciences, 2013, 38, 25-34.	0.5	5
11	Ties between Graph Theory and Biology. Discrete Mathematics and Its Applications, 2013, , 1559-1579.	0.1	0
12	Reduced-by-matching Graphs: Toward Simplifying Hamiltonian Circuit Problem. Fundamenta Informaticae, 2012, 118, 225-244.	0.3	5
13	Complexity Issues in Computational Biology. Fundamenta Informaticae, 2012, 118, 385-401.	0.3	5
14	Highly Efficient Parallel Approach to the Next-Generation DNA Sequencing. Lecture Notes in Computer Science, 2012, , 262-271.	1.0	0
15	RNA Partial Degradation Problem: Motivation, Complexity, Algorithm. Journal of Computational Biology, 2011, 18, 821-834.	0.8	8
16	The simplified partial digest problem: Approximation and a graph-theoretic model. European Journal of Operational Research, 2011, 208, 142-152.	3.5	4
17	Whole genome assembly from 454 sequencing output via modified DNA graph concept. Computational Biology and Chemistry, 2009, 33, 224-230.	1.1	18
18	On the approximability of the Simplified Partial Digest Problem. Discrete Applied Mathematics, 2009, 157, 3586-3592.	0.5	2

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19	Finding Hamiltonian circuits in quasi-adjoint graphs. Discrete Applied Mathematics, 2008, 156, 2573-2580.	0.5	10
20	A new algorithm for genome assembly from short reads. , 2008, , .		1
21	Parallel Implementation of the Novel Approach to Genome Assembly. , 2008, , .		0
22	Simplified Partial Digest Problem: Enumerative and Dynamic Programming Algorithms. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2007, 4, 668-680.	1.9	10
23	A polynomial time equivalence between DNA sequencing and the exact perfect matching problem. Discrete Optimization, 2007, 4, 154-162.	0.6	4
24	Computational complexity of isothermic DNA sequencing by hybridization. Discrete Applied Mathematics, 2006, 154, 718-729.	0.5	13
25	Dealing with repetitions in sequencing by hybridization. Computational Biology and Chemistry, 2006, 30, 313-320.	1.1	11
26	Selected combinatorial problems of computational biology. European Journal of Operational Research, 2005, 161, 585-597.	3.5	22
27	Evolutionary Approaches to DNA Sequencing with Errors. Annals of Operations Research, 2005, 138, 67-78.	2.6	13
28	Combinatorial optimization in DNA mapping — a computational thread of the Simplified Partial Digest Problem. RAIRO - Operations Research, 2005, 39, 227-241.	1.0	9
29	Sequencing by hybridization with isothermic oligonucleotide libraries. Discrete Applied Mathematics, 2004, 145, 40-51.	0.5	16
30	Tabu search algorithm for DNA sequencing by hybridization with isothermic libraries. Computational Biology and Chemistry, 2004, 28, 11-19.	1.1	14
31	DNA Sequencing—Tabu and Scatter Search Combined. INFORMS Journal on Computing, 2004, 16, 232-240.	1.0	24
32	Complexity of DNA sequencing by hybridization. Theoretical Computer Science, 2003, 290, 1459-1473.	0.5	49
33	A heuristic managing errors for DNA sequencing. Bioinformatics, 2002, 18, 652-660.	1.8	36
34	On the recognition of de Bruijn graphs and their induced subgraphs. Discrete Mathematics, 2002, 245, 81-92.	0.4	13
35	Hybrid Genetic Algorithm for DNA Sequencing with Errors. Journal of Heuristics, 2002, 8, 495-502.	1.1	38