Kun-Mao Chao

List of Publications by Year in descending order

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100 papers

1,852 citations

304743 22 h-index 38 g-index

106 all docs

106
docs citations

106 times ranked 1764 citing authors

| # | Article | IF | CITATIONS |
|----|---|-----|-----------|
| 1 | A Polynomial-Time Approximation Scheme for Minimum Routing Cost Spanning Trees. SIAM Journal on Computing, 2000, 29, 761-778. | 1.0 | 103 |
| 2 | Aligning two sequences within a specified diagonal band. Bioinformatics, 1992, 8, 481-487. | 4.1 | 94 |
| 3 | Pulmonary Tuberculosis and Delay in Anti-Tuberculous Treatment Are Important Risk Factors for Chronic Obstructive Pulmonary Disease. PLoS ONE, 2012, 7, e37978. | 2.5 | 89 |
| 4 | PhosphoPOINT: a comprehensive human kinase interactome and phospho-protein database. Bioinformatics, 2008, 24, i14-i20. | 4.1 | 88 |
| 5 | Efficient algorithms for locating the length-constrained heaviest segments with applications to biomolecular sequence analysis. Journal of Computer and System Sciences, 2002, 65, 570-586. | 1.2 | 78 |
| 6 | A generalized global alignment algorithm. Bioinformatics, 2003, 19, 228-233. | 4.1 | 77 |
| 7 | Risk factors for pulmonary tuberculosis in patients with chronic obstructive airway disease in Taiwan: a nationwide cohort study. BMC Infectious Diseases, 2013, 13, 194. | 2.9 | 69 |
| 8 | Optimal Duration of Anti-TB Treatment in Patients With Diabetes. Chest, 2015, 147, 520-528. | 0.8 | 60 |
| 9 | Gene Expression-Based Chemical Genomics Identifies Potential Therapeutic Drugs in Hepatocellular Carcinoma. PLoS ONE, 2011, 6, e27186. | 2.5 | 58 |
| 10 | On the generalized constrained longest common subsequence problems. Journal of Combinatorial Optimization, 2011, 21, 383-392. | 1.3 | 56 |
| 11 | Recent Developments in Linear-Space Alignment Methods: A Survey. Journal of Computational Biology, 1994, 1, 271-291. | 1.6 | 51 |
| 12 | Approximation algorithms for some optimum communication spanning tree problems. Discrete Applied Mathematics, 2000, 102, 245-266. | 0.9 | 51 |
| 13 | Constrained Multiple Sequence Alignment Tool Development and Its Application to RNase Family Alignment. Journal of Bioinformatics and Computational Biology, 2003, 01, 267-287. | 0.8 | 44 |
| 14 | A fast algorithm for computing a longest common increasing subsequence. Information Processing Letters, 2005, 93, 249-253. | 0.6 | 38 |
| 15 | MAVG: locating non-overlapping maximum average segments in a given sequence. Bioinformatics, 2003, 19, 151-152. | 4.1 | 36 |
| 16 | Globin Gene Server: A Prototype E-Mail Database Server Featuring Extensive Multiple Alignments and Data Compilation for Electronic Genetic Analysis. Genomics, 1994, 21, 344-353. | 2.9 | 35 |
| 17 | Linear-space algorithms that build local alignments from fragments. Algorithmica, 1995, 13, 106-134. | 1.3 | 33 |
| 18 | An Approximation Algorithm for Haplotype Inference by Maximum Parsimony. Journal of Computational Biology, 2005, 12, 1261-1274. | 1.6 | 32 |

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| 19 | Title is missing!. Journal of Combinatorial Optimization, 1999, 3, 199-211. | 1.3 | 30 |
| 20 | Positive and negative regulatory elements of the rabbit embryonic l̃µ-globin gene revealed by an improved multiple alignment program and functional analysis. DNA Sequence, 1993, 4, 163-176. | 0.7 | 26 |
| 21 | Strontium ranelate-laden near-infrared photothermal-inspired methylcellulose hydrogel for arthritis treatment. Materials Science and Engineering C, 2021, 123, 111980. | 7.3 | 25 |
| 22 | SMART: Statistical Metabolomics Analysis—An R Tool. Analytical Chemistry, 2016, 88, 6334-6341. | 6.5 | 24 |
| 23 | Light graphs with small routing cost. Networks, 2002, 39, 130-138. | 2.7 | 23 |
| 24 | A Polynomial Time Approximation Scheme for Optimal Product-Requirement Communication Spanning Trees. Journal of Algorithms, 2000, 36, 182-204. | 0.9 | 21 |
| 25 | A greedier approach for finding tag SNPs. Bioinformatics, 2006, 22, 685-691. | 4.1 | 21 |
| 26 | Constrained sequence alignment. Bulletin of Mathematical Biology, 1993, 55, 503-524. | 1.9 | 19 |
| 27 | Optimal algorithms for locating the longest and shortest segments satisfying a sum or an average constraint. Information Processing Letters, 2005, 96, 197-201. | 0.6 | 19 |
| 28 | A fault-tolerant method for HLA typing with PacBio data. BMC Bioinformatics, 2014, 15, 296. | 2.6 | 18 |
| 29 | Locating well-conserved regions within a pairwise alignment. Bioinformatics, 1993, 9, 387-396. | 4.1 | 17 |
| 30 | Selecting additional tag SNPs for tolerating missing data in genotyping. BMC Bioinformatics, 2005, 6, 263. | 2.6 | 17 |
| 31 | Distant HNF1 Site as a Master Control for the Human Class I Alcohol Dehydrogenase Gene Expression*. Journal of Biological Chemistry, 2006, 281, 19809-19821. | 3.4 | 17 |
| 32 | The Swap Edges of a Multiple-Sources Routing Tree. Algorithmica, 2008, 50, 299-311. Improved algorithms for the Ammilmath altimg="sill-gif" overflow="scroll" | 1.3 | 16 |
| 33 | xmins:xocs= http://www.eisevier.com/xmi/xocs/dtd xmins:xs= http://www.w3.org/2001/XMLSchema xmlns:xsi="http://www.w3.org/2001/XMLSchema-instance" xmlns="http://www.elsevier.com/xml/ja/dtd" xmlns:ja="http://www.elsevier.com/xml/ja/dtd" xmlns:mml="http://www.w3.org/1998/Math/MathML" xmlns:tb="http://www.elsevier.com/xml/common/table/dtd" | 0.9 | 15 |
| 34 | xmlns:sb="http://www.elsevier.com/xml/common/struct-bib/dtd" xmlns:ce="http://www.elsevier.com/x On the range maximum-sum segment query problem. Discrete Applied Mathematics, 2007, 155, 2043-2052. | 0.9 | 15 |
| 35 | Linear-Time Algorithms for the Multiple Gene Duplication Problems. IEEE/ACM Transactions on Computational Biology and Bioinformatics, $2011,8,260\text{-}265$. | 3.0 | 15 |
| 36 | Finding all sorting tandem duplication random loss operations. Journal of Discrete Algorithms, 2011, 9, 32-48. | 0.7 | 14 |

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| 37 | Finding All Approximate Gapped Palindromes. Lecture Notes in Computer Science, 2009, , 1084-1093. | 1.3 | 14 |
| 38 | Efficient methods for generating optimal single and multiple spaced seeds. , 0, , . | | 13 |
| 39 | Algorithms for finding the weight-constrained k longest paths in a tree and the length-constrained k maximum-sum segments of a sequence. Theoretical Computer Science, 2008, 407, 349-358. | 0.9 | 13 |
| 40 | Urinary tuberculosis is associated with the development of urothelial carcinoma but not renal cell carcinoma: a nationwide cohort study in Taiwan. British Journal of Cancer, 2013, 109, 2933-2940. | 6.4 | 13 |
| 41 | On the Range Maximum-Sum Segment Query Problem. Lecture Notes in Computer Science, 2004, , 294-305. | 1.3 | 13 |
| 42 | Efficient combination of multiple word models for improved sequence comparison. Bioinformatics, 2004, 20, 2529-2533. | 4.1 | 12 |
| 43 | Finding a Length-Constrained Maximum-Density Path in a Tree. Journal of Combinatorial Optimization, 2005, 9, 147-156. | 1.3 | 11 |
| 44 | The backup 2â€center and backup 2â€median problems on trees. Networks, 2009, 53, 39-49. | 2.7 | 11 |
| 45 | Facilitated and Controlled Strontium Ranelate Delivery Using GCS-HA Nanocarriers Embedded into PEGDA Coupled with Decortication Driven Spinal Regeneration. International Journal of Nanomedicine, 2021, Volume 16, 4209-4224. | 6.7 | 11 |
| 46 | CNVDetector: locating copy number variations using array CGH data. Bioinformatics, 2008, 24, 2773-2775. | 4.1 | 10 |
| 47 | Hardness of comparing two run-length encoded strings. Journal of Complexity, 2010, 26, 364-374. | 1.3 | 10 |
| 48 | FINDING ALL APPROXIMATE GAPPED PALINDROMES. International Journal of Foundations of Computer Science, 2010, 21, 925-939. | 1.1 | 10 |
| 49 | In-silico drug screening and potential target identification for hepatocellular carcinoma using Support Vector Machines based on drug screening result. Gene, 2013, 518, 201-208. | 2.2 | 9 |
| 50 | <scp><i>M</i></scp> <i>ycobacterium tuberculosis</i> nucleic acid amplification tests reduce nosocomial tuberculosis exposure in intensive care units: A nationwide cohort study. Respirology, 2015, 20, 1233-1240. | 2.3 | 9 |
| 51 | A tight analysis of the Katriel–Bodlaender algorithm for online topological ordering. Theoretical Computer Science, 2007, 389, 182-189. | 0.9 | 8 |
| 52 | Improving the performance of identifying contributors for XML keyword search. SIGMOD Record, 2011, 40, 5-10. | 1.2 | 8 |
| 53 | On computing all suboptimal alignments. Information Sciences, 1998, 105, 189-207. | 6.9 | 7 |
| 54 | On the uniform edge-partition of a tree. Discrete Applied Mathematics, 2007, 155, 1213-1223. | 0.9 | 6 |

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| 55 | A Compact and Efficient Labeling Scheme for XML Documents. Lecture Notes in Computer Science, 2013, , 269-283. | 1.3 | 6 |
| 56 | The Extent of Linkage Disequilibrium and Computational Challenges of Single Nucleotide Polymorphisms in Genome-Wide Association Studies. Current Drug Metabolism, 2011, 12, 498-506. | 1.2 | 5 |
| 57 | The impact of diabetes mellitus and its control on the development of tuberculosis: a nationwide longitudinal study in Taiwan. Pharmacoepidemiology and Drug Safety, 2013, 22, 995-1003. | 1.9 | 5 |
| 58 | Approximate Matching for Run-Length Encoded Strings Is 3sum-Hard. Lecture Notes in Computer Science, 2009, , 168-179. | 1.3 | 5 |
| 59 | A new framework for the selection of tag SNPs by multimarker haplotypes. Journal of Biomedical Informatics, 2008, 41, 953-961. | 4.3 | 4 |
| 60 | A Fully Compressed Algorithm for Computing the Edit Distance of Run-Length Encoded Strings. Algorithmica, 2013, 65, 354-370. | 1.3 | 4 |
| 61 | A Takagi-Sugeno fuzzy neural network-based algorithm with single-channel EEG signal for the discrimination between light and deep sleep stages. , 2016, , . | | 4 |
| 62 | On the influenza vaccination policy through mathematical modeling. International Journal of Infectious Diseases, 2020, 98, 71-79. | 3.3 | 4 |
| 63 | The Generalized Popular Condensation Problem. Lecture Notes in Computer Science, 2014, , 606-617. | 1.3 | 4 |
| 64 | Optimal algorithms for the average-constrained maximum-sum segment problem. Information Processing Letters, 2009, 109, 171-174. | 0.6 | 3 |
| 65 | On Locating Disjoint Segments with Maximum Sum of Densities. Lecture Notes in Computer Science, 2006, , 300-307. | 1.3 | 3 |
| 66 | Proof of a Conjecture About Minimum Spanning Tree Cycle Intersection. Discrete Applied Mathematics, 2022, 321, 19-23. | 0.9 | 3 |
| 67 | A tight bound on the min-ratio edge-partitioning problem of a tree. Discrete Applied Mathematics, 2010, 158, 1471-1478. | 0.9 | 2 |
| 68 | Influence of Pressure Control Levels on the Pulse Pressure Variations. Shock, 2011, 36, 628-632. | 2.1 | 2 |
| 69 | Efficient retrieval of approximate palindromes in a run-length encoded string. Theoretical Computer Science, 2012, 432, 28-37. | 0.9 | 2 |
| 70 | Data-driven interdisciplinary mathematical modelling quantitatively unveils competition dynamics of co-circulating influenza strains. Journal of Translational Medicine, 2017, 15, 163. | 4.4 | 2 |
| 71 | Asymptotic Limits of a New Type of Maximization Recurrence with an Application to Bioinformatics. Lecture Notes in Computer Science, 2012, , 177-188. | 1.3 | 2 |
| 72 | Preserving Inversion Phylogeny Reconstruction. Lecture Notes in Computer Science, 2012, , 1-13. | 1.3 | 2 |

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| 73 | Identifying Approximate Palindromes in Run-Length Encoded Strings. Lecture Notes in Computer Science, 2010, , 339-350. | 1.3 | 2 |
| 74 | A Fully Compressed Algorithm for Computing the Edit Distance of Run-Length Encoded Strings. Lecture Notes in Computer Science, 2010, , 415-426. | 1.3 | 2 |
| 75 | Forming Plurality at Minimum Cost. Lecture Notes in Computer Science, 2015, , 77-88. | 1.3 | 2 |
| 76 | An optimal algorithm for querying tree structures and its applications in bioinformatics. SIGMOD Record, 2004, 33, 21-26. | 1.2 | 1 |
| 77 | MINKOWSKI SUM SELECTION AND FINDING. International Journal of Computational Geometry and Applications, 2011, 21, 283-311. | 0.5 | 1 |
| 78 | Efficient algorithms for local ranking. Information Processing Letters, 2012, 112, 517-522. | 0.6 | 1 |
| 79 | A linear-time algorithm for finding an edge-partition with max-min ratio at most two. Discrete Applied Mathematics, 2013, 161, 932-943. | 0.9 | 1 |
| 80 | Minkowski Sum Selection and Finding. Lecture Notes in Computer Science, 2008, , 460-471. | 1.3 | 1 |
| 81 | Faster Algorithms for Searching Relevant Matches in XML Databases. Lecture Notes in Computer Science, 2010, , 290-297. | 1.3 | 1 |
| 82 | Locating Valid SLCAs for XML Keyword Search with NOT Semantics. SIGMOD Record, 2014, 43, 29-34. | 1.2 | 1 |
| 83 | A Fixed-Parameter Algorithm for Detecting a Singleton Attractor in an AND/OR Boolean Network with Bounded Treewidth. IEICE Transactions on Fundamentals of Electronics, Communications and Computer Sciences, 2015, E98.A, 384-390. | 0.3 | 1 |
| 84 | Fast algorithms for aligning sequences with restricted affine gap penalties. Lecture Notes in Computer Science, 1997, , 264-273. | 1.3 | 0 |
| 85 | Approximation algorithms for the optimization problems of SNPs and haplotypes. , 0, , . | | 0 |
| 86 | The 2-radius and 2-radiian problems on trees. Theoretical Computer Science, 2008, 407, 524-531. | 0.9 | 0 |
| 87 | On Locating Disjoint Segments with Maximum Sum of Densities. Algorithmica, 2009, 54, 107-117. | 1.3 | 0 |
| 88 | Pattern identification in a haplotype block. , 0, , 23-35. | | 0 |
| 89 | GUEST EDITORS' FOREWORD. International Journal of Computational Geometry and Applications, 2013, 23, 425-426. | 0.5 | 0 |
| 90 | Preface Algorithms and Computation (ISAAC 2012). Algorithmica, 2014, 70, 694-695. | 1.3 | 0 |

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| 91 | Finding a Length-Constrained Maximum-Density Path in a Tree. Lecture Notes in Computer Science, 2003, , 78-87. | 1.3 | 0 |
| 92 | DYNAMIC PROGRAMMING STRATEGIES FOR ANALYZING BIOMOLECULAR SEQUENCES. Lecture Notes Series, Institute for Mathematical Sciences, 2004, , 1-23. | 0.2 | 0 |
| 93 | Maximum-scoring Segment with Length Restrictions. , 2008, , 506-507. | | 0 |
| 94 | Maximum-Density Segment. , 2008, , 502-504. | | 0 |
| 95 | Finding All Sorting Tandem Duplication Random Loss Operations. Lecture Notes in Computer Science, 2009, , 301-313. | 1.3 | 0 |
| 96 | Maximum-Average Segments. , 2015, , 1-3. | | 0 |
| 97 | Maximum-Sum Segments. , 2015, , 1-3. | | 0 |
| 98 | Maximum-Sum Segments. , 2016, , 1242-1244. | | 0 |
| 99 | Maximum-Average Segments. , 2016, , 1241-1242. | | 0 |
| 100 | Algorithms for Computing the Length-Constrained Max-Score Segments with Applications to DNA Copy Number Data Analysis., 2007,, 834-845. | | 0 |