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| Descrizione fisica | 1 online resource (XVI, 225 p. 66 illus., 35 illus. in color.) |
| Collana | Lecture Notes in Bioinformatics ; ; 11488 |
| Disciplina | 570.285 |
| Soggetti | Bioinformatics Arithmetic and logic units, Computer Algorithms Artificial intelligence Numerical analysis Computer science—Mathematics Computational Biology/Bioinformatics Arithmetic and Logic Structures Algorithm Analysis and Problem Complexity Artificial Intelligence Numeric Computing Discrete Mathematics in Computer Science |
| Lingua di pubblicazione | Inglese |
| Formato | Materiale a stampa |
| Livello bibliografico | Monografia |
| Nota di bibliografia | Includes bibliographical references and index. |
| Nota di contenuto | Invited Talk -- New Divide-and-Conquer Techniques for Large-Scale Phylogenetic Estimation -- Biological Networks and Graph Algorithms -- New Polynomial-Time Algorithm around the Scaffolding Problem -- Enumerating Dominant Pathways in Biological Networks by Information Flow Analysis -- Comparing Different Graphlet Measures for Evaluating Network Model Fits to BioGRID PPI Networks -- Graph-Theoretic Partitioning of RNAs and Classification of Pseudoknots -- PathRacer: Racing Profile HMM Paths on Assembly Graph -- Genome |

Rearrangement, Assembly and Classification -- A Uniform Theory of Adequate Subgraphs for the Genome Median, Halving, and Aliquoting Problems -- Lightweight Metagenomic Classification via eBWT -- MULKSG: MULTiple K Simultaneous Graph Assembly -- Counting Sorting Scenarios and Intermediate Genomes for the Rank Distance -- Generalizations of the Genomic Rank Distance to Indels -- Sequence Analysis, Phylogenetics and Other Biological Processes -- Using INC within Divide-and-Conquer Phylogeny Estimation -- Predicting Methylation from Sequence and Gene Expression Using Deep Learning with Attention -- A Mathematical Model for Enhancer Activation Kinetics During Cell Differentiation -- Transcript Abundance Estimation and the Lamina Packing Problem -- Efficient Algorithms for Finding Edit-Distance Based Motifs.

Sommario/riassunto

This book constitutes the proceedings of the 6th International Conference on Algorithms for Computational Biology, ACoB 2019, held in Berkeley, CA, USA, in May 2019. The 15 full papers presented together with 1 invited paper were carefully reviewed and selected from 30 submissions. They are organized in the following topical sections: Biological networks and graph algorithms; genome rearrangement, assembly and classification; sequence analysis, phylogenetics and other biological processes.
