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Collana	Lecture Notes in Bioinformatics ; ; 3692
Disciplina	572.0285
Soggetti	Life sciences Computer programming Algorithms Data structures (Computer science) Computers Computer science—Mathematics Life Sciences, general Programming Techniques Algorithm Analysis and Problem Complexity Data Structures Computation by Abstract Devices Discrete Mathematics in Computer Science
Lingua di pubblicazione	Inglese
Formato	Materiale a stampa
Livello bibliografico	Monografia
Note generali	Bibliographic Level Mode of Issuance: Monograph
Nota di bibliografia	Includes bibliographical references and index.
Nota di contenuto	Expression -- Spectral Clustering Gene Ontology Terms to Group Genes by Function -- Dynamic De-Novo Prediction of microRNAs Associated with Cell Conditions: A Search Pruned by Expression -- Clustering Gene Expression Series with Prior Knowledge -- A Linear Time Biclustering Algorithm for Time Series Gene Expression Data -- Time-Window Analysis of Developmental Gene Expression Data with Multiple Genetic Backgrounds -- Phylogeny -- A Lookahead Branch-and-Bound Algorithm for the Maximum Quartet Consistency Problem -- Computing the Quartet Distance Between Trees of Arbitrary Degree

-- Using Semi-definite Programming to Enhance Supertree Resolvability  
-- An Efficient Reduction from Constrained to Unconstrained Maximum Agreement Subtree -- Pattern Identification in Biogeography -- On the Complexity of Several Haplotyping Problems -- A Hidden Markov Technique for Haplotype Reconstruction -- Algorithms for Imperfect Phylogeny Haplotyping (IPPH) with a Single Homoplasy or Recombination Event -- Networks -- A Faster Algorithm for Detecting Network Motifs -- Reaction Motifs in Metabolic Networks -- Reconstructing Metabolic Networks Using Interval Analysis -- Genome Rearrangements -- A 1.375-Approximation Algorithm for Sorting by Transpositions -- A New Tight Upper Bound on the Transposition Distance -- Perfect Sorting by Reversals Is Not Always Difficult -- Minimum Recombination Histories by Branch and Bound -- Sequences -- A Unifying Framework for Seed Sensitivity and Its Application to Subset Seeds -- Generalized Planted (l,d)-Motif Problem with Negative Set -- Alignment of Tandem Repeats with Excision, Duplication, Substitution and Indels (EDSI) -- The Peres-Shields Order Estimator for Fixed and Variable Length Markov Models with Applications to DNA Sequence Similarity -- Multiple Structural RNA Alignment with Lagrangian Relaxation -- Faster Algorithms for Optimal Multiple Sequence Alignment Based on Pairwise Comparisons -- Ortholog Clustering on a Multipartite Graph -- Linear Time Algorithm for Parsing RNA Secondary Structure -- A Compressed Format for Collections of Phylogenetic Trees and Improved Consensus Performance -- Structure -- Optimal Protein Threading by Cost-Splitting -- Efficient Parameterized Algorithm for Biopolymer Structure-Sequence Alignment -- Rotamer-Pair Energy Calculations Using a Trie Data Structure -- Improved Maintenance of Molecular Surfaces Using Dynamic Graph Connectivity -- The Main Structural Regularities of the Sandwich Proteins -- Discovery of Protein Substructures in EM Maps.

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