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| Edizione                | [1st ed. 2014.]  |
| Descrizione fisica      | 1 online resource (XVIII, 409 p. 115 illus.)   |
| Collana                 | Lecture Notes in Bioinformatics ; ; 8492   |
| Disciplina              | 570.285  |
| Soggetti                | Bioinformatics<br>Data mining<br>Pattern recognition<br>Biomathematics<br>Computational Biology/Bioinformatics<br>Data Mining and Knowledge Discovery<br>Pattern Recognition<br>Mathematical and Computational Biology<br>Conference papers and proceedings.   |
| 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 author index.  |
| Nota di contenuto       | Predicting Disease Risks Using Feature Selection Based on Random<br>Forest and Support Vector Machine Phylogenetic Bias in the<br>Likelihood Method Caused by Missing Data Coupled with Among-Site<br>Rate Variation: An Analytical Approach An Eigen decomposition<br>Method for Protein Structure Alignment Functional Interplay between<br>Hemagglutinin and Neuraminidase of Pandemic 2009 H1N1 from the<br>Perspective of Virus Evolution Predicting Protein Submitochondrial<br>Locations Using a K-Nearest Neighbors Method Based on the Bit-Score<br>Weighted Euclidean Distance Algorithms Implemented for Cancer<br>Gene Searching and Classifications Dysregulated microRNA Profile in<br>HeLa Cell Lines Induced by Lupeol A Simulation for Proportional<br>Biological Operational Mu-Circuit Computational Prediction of<br>Human Saliva-Secreted Proteins A Parallel Scheme for Three- |

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Dimensional Reconstruction in Large-Field Electron Tomography -- An Improved Correlation Method Based on Rotation Invariant Feature for Automatic Particle Selection -- An Effective Algorithm for Peptide de novo Sequencing from Mixture MS/MS Spectra -- Identifying Spurious Interactions in the Protein-Protein Interaction Networks Using Local Similarity Preserving Embedding -- Multiple RNA Interaction with Suboptimal Solutions -- Application of Consensus String Matching in the Diagnosis of Allelic Heterogeneity (Extended Abstract) -- Continuous Time Bavesian Networks for Gene Network Reconstruction: A Comparative Study on Time Course Data -- Drug Target Identification Based on Structural Output Controllability of Complex Networks --NovoGMET: De Novo Peptide Sequencing Using Graphs with Multiple Edge Types (GMET) for ETD/ECD Spectra -- Duplication Cost Diameters -- Computational Identification of De-Centric Genetic Regulatory Relationships from Functional Genomic Data -- Classification of Mutations by Functional Impact Type: Gain of Function, Loss of Function and Switch of Function -- Network Analysis of Human Disease Comorbidity Patterns Based on Large-Scale Data Mining --Identification of Essential Proteins by Using Complexes and Interaction Network -- GenoScan: Genomic Scanner for Putative miRNA Precursors -- Searching SNP Combinations Related to Evolutionary Information of Human Populations on HapMap Data -- 2D Pharmacophore Query Generation -- Structure-Based Analysis of Protein Binding Pockets Using Von Neumann Entropy -- A New Mathematical Model for Inbreeding Depression in Large Populations -- dSpliceType: A Multivariate Model for Detecting Various Types of Differential Splicing Events Using RNA-Seq -- Conformational Transitions and Principal Geodesic Analysis on the Positive Semidefinite Matrix Manifold -- Joint Analysis of Functional and Phylogenetic Composition for Human Microbiome Data -- schematikon: Detailed Sequence-Structure Relationships from Mining a Non-redundant Protein Structure Database (Extended Abstract) -- PNImodeler: Web Server for Inferring Protein Binding Nucleotides from Sequence Data -- A MCI Decision Support System Based on Ontology -- Context Similarity Based Feature Selection Methods for Protein Interaction Article Classification --Network Propagation Reveals Novel Genetic Features Predicting Drug Response of Cancer Cell Lines -- Splice Site Prediction Using Support Vector Machine with Markov Model and Codon Information --Exploiting Topic Modeling to Boost Metagenomic Sequences Binning --Network-Based Method for Identifying Overlapping Mutated Driver Pathways in Cancer -- Completing a Bacterial Genome with in silico and Wet Lab Approaches -- Protein Function Prediction: A Global Prediction Method with Multiple Data Sources -- A microRNA-Gene Network in Ovarian Cancer from Genome-Wide QTL Analysis -- K-Profiles Nonlinear Clustering -- Estrogen Induced RNA Polymerase II Stalling in Breast Cancer Cell Line MCF7 -- A Knowledge-Driven Approach in Constructing a Large-Scale Drug-Side Effect Relationship Knowledge Base for Computational Drug Discovery -- Systems Biology Approach to Understand Seed Composition -- Prediction of the Cooperative cisregulatory Elements for Broadly Expressed Neuronal Genes in Caenorhabditis Elegans -- Improving the Mapping of the Smith-Waterman Sequence Database Search Algorithm onto CUDA GPUs --VFP: A Visual Tool for Predicting Gene-Fusion Base on Analyzing Single-end RNA-Sequence -- A Novel Method for Identifying Essential Proteins from Active PPI Networks -- RAUR: Re-alignment of Unmapped Reads with Base Quality Score -- PIGS: Improved Estimates of Identityby-Descent Probabilities by Probabilistic IBD Graph Sampling --Clustering PPI Data through Improved Synchronization-Based

|                    | Hierarchical Clustering Method Order Decay in Transcription<br>Regulation in Type 1 Diabetes Simulated Regression Algorithm for<br>Transcriptome Quantification.   |
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| Sommario/riassunto | This book constitutes the refereed proceedings of the 10th<br>International Symposium on Bioinformatics Research and Applications,<br>ISBRA 2014, held in Zhangjiajie, China, in June 2014. The 33 revised<br>full papers and 31 one-page abstracts included in this volume were<br>carefully reviewed and selected from 119 submissions. The papers<br>cover a wide range of topics in bioinformatics and computational<br>biology and their applications including the development of<br>experimental or commercial systems. |