Record Nr. UNINA9910454291203321 Selected topics in post-genome knowledge discovery [[electronic Titolo resource] /] / editor[s], Limsoon Wong, Louxin Zhang Pubbl/distr/stampa Singapore, : Singapore University Press Singapore;; River Edge, NJ,: World Scientific Pub. Co., c2004 **ISBN** 1-281-93473-9 9786611934736 981-279-484-0 Descrizione fisica 1 online resource (176 p.) Collana Lecture notes series;; vol. 3 Altri autori (Persone) WongLimsoon <1965-> ZhangLouxin Disciplina 572.8 572.80285 Genetics - Mathematics Soggetti Genetics Electronic books. Lingua di pubblicazione Inglese **Formato** Materiale a stampa Livello bibliografico Monografia Note generali Description based upon print version of record. Nota di bibliografia Includes bibliographical references. Nota di contenuto Contents : Foreword : Preface ; Dynamic Programming Strategies for Analyzing Biomolecular Sequences ; 2. Elementary Dynamic-Programming ; 1. Introduction Algorithms ; 2.1. Fibonacci numbers ; 2.2. The maximum-sum substring problem 2.3. Longest increasing subsequence 2.4. Longest common subsequence 3. Sequence Alignment ; 3.1. Global alignment ; 3.2. Local alignment ; 3.3. Affine gap penalties ; 3.4. Space-saving strategies ; 3.5. Multiple sequence alignment ; The Representation Comparison and Prediction of Protein Pathways ; 1. Introduction 2. Online Pathway Resources 3. Pathway Representation ; 3.1. Pathway Space ; 3.2. SLIPR Format ; 4. Pathway Comparison

; 4.1. Comparing Individual Components 4.2. Aligning Two Pathways Using Dynamic Programming ; 4.3. Pathway Database Comparison 4.4. One Implementation: PM\_search Documentation 5. Orthologous Pathway Prediction ; 6. Discussion : 6.1. Theoretical Issues on Evolutionary Study of Pathways ; 6.2. Establishing a Relational Pathway Database and its Web Interfaces ; 6.3. Pathway Prediction and Beyond Gene Network Inference and Biopathway Modeling 1. Introduction ; 2. Gene Network Inference from Microarray Data ; 2.1. Boolean **Network Model** : 2.2. Bayesian Network Model ; 3. Modeling and Simulation ; 3.1. Architecture for Biopathway Modeling ; 3.2. How to Model Biopathways 3.3. Genomic Object Net and Biopathway Databases Towards Simulation

## Sommario/riassunto

The Institute for Mathematical Sciences at the National University of Singapore organized a program on "Post-Genome Knowledge Discovery" from January to June 2002. The program focused on the computational and statistical analysis of sequences and genetics, and the mathematical modeling of complex biological interactions, which are critical to the accurate annotation of genomic sequences, the study of the interplay between genes and proteins, and the study of the genetic variability of species. As part of the program, tutorials for graduate students and newcomers to this transdisciplinary area