

<<生物信息学>>

图书基本信息

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内容概要

“精要速览系列(Instant Notes Series)”是国外教材“Best Seller”榜的上榜教材。该系列结构新颖，视角独特；重点明确，脉络分明；图表简明清晰；英文自然易懂，被国内多所重点院校选用作为双语教材。

先锋版是继“现代生物学精要速览”之后推出的跨学科的升级版本。

本书是该系列中的《生物信息学》分册，全书共15章，全面系统地概括了该学科的核心内容和前沿动态，涉及生物学数据的获得与处理、系统发生学、芯片数据分析、蛋白质组数据分析、生物信息学的应用等主要内容。

本书是指导大学生快速掌握生物信息学基础知识的优秀教材，也是辅助教师授课的极佳教学参考书，同时可供生命科学相关专业的研究生参考。

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书籍目录

AbbreviationsPrefaceSection A-Overview of bioinformatics A1 The scope of bioinformatics A2 Bioinformatics and the Internet A3 Useful bioinformatics sites on the WWWSection B- Data acquisition B1 Sequencing DNA,RNA and proteins B2 Determination of protein structure B3 Gene and protein expression data B4 Protein interaction dataSection C-Databases-contents,structure and annotation C1 File formats C2 Annotated sequence databases C3 Genome and organism-specific databases C4 Miscellaneous databasesSection D-Retrieval of biological data D1 Data retrieval with Entrez and DBGET/LinkDB D2 Data retrieval with SRS(sequence retrieval system)Section E-Searching sequence databases by sequence E1 Sequence similarity searches E2 Amino acid substitution matrices E3 Databases searches:FASTA and BLAST E4 Sequence filters E5 Iterative database searches and PSI-BLASTSection F-Multiple sequence alignment and family relationships F1 Multiple sequence alignment and family relationships F2 Protein families and pattern databases F3 Protein domain familiesSection G-Phylogenetics G1 Phylogenetics,cladistics and ontology G2 Building phylogenetic trees G3 Evolution of macromolecular sequencesSection H-Sequence annotation H1 Principles of genome annotation H2 Annotation tools and resourcesSection I-Structural bioinformatics I1 Conceptual models of protein structure I2 The relationship of protein three-dimensional structure to protein function I3 The evolution of protein structure and function I4 Obtaining,viewing and analyzing structural data I5 Structural alignment I6 Classification of proteins of known three-dimensional structure:CATH and SCOP I7 Introduction to protein structure prediction I8 Structure prediction by comparative modeling I9 Secondary structure prediction I10 Advanced protein structure prediction and prediction strategiesSection J-Microarray data analysis J1 Microarray data:analysis methods J2 Microarray data:tools and resources J3 Sequence sampling and SAGESection K-Proteomic data analysis K1 Analyzing data from 2D-PAGE gels K2 Analyzing protein mass spectrometry dataSection L-Higher-order systems L1 Modeling and reconstructing molecular pathways L2 Protein interaction informatics L3 Higher-order modelsSection M-Cheminformatics in biology M1 Conventions for representing molecular pathways M2 Cheminformatics resourcesSection N-Bioinformatics in the pharmaceutical industry N1 Bioinformatics and drug discovery N2 Pharmainformatics resourcesSection O-Basic principles of computing in bioinformatics O1 Running computer software O2 Computer operating systems O3 Software downloading and installation O4 Database managementFurther readingGlossaryIndex

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