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Autore	Carvalho Rogerio Atem de
Titolo	Nanosatellites : space and ground technologies, operations and economics // edited by Rogerio Atem de Carvalho, Reference Center for Embedded and Aerospace Systems (CRSEA), Polo de Inovacao Campos dos Goytacazes (PICG), Instituto Federal Fluminense (IFF), Brazil, Jaime Estela, Spectrum Aerospace Group, Germering, Germany, Martin Langer, Institute of Astronautics, Technical University of Munich, Garching, Germany and Orbital Oracle Technologies GmbH, Munich, Germany
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ISBN	1-119-04206-2 1-119-04205-4 1-119-04204-6
Descrizione fisica	1 online resource (xxxvi, 670 pages), 16 unnumbered pages of plates : illustrations
Disciplina	629.46
Soggetti	Microspacecraft
Lingua di pubblicazione	Inglese
Formato	Materiale a stampa
Livello bibliografico	Monografia
Nota di bibliografia	Includes bibliographical references and index.
Sommario/riassunto	"Nano-Satellites : Space and Ground Technologies, Operations and Economics comprehensively presents the latest research on the fast developing area of nano-satellites. It begins with a brief history of nano-satellites and introduces nano-satellite technologies, payloads for nano-satellite and explains how these are deployed into space. Part two provides an overview on ground segment and operations and part three focuses on the regulations, policies, economics and future trends. With contributions from global experts, Nano-Satellites: Space and Ground Technologies, Operations and Economics, is a comprehensive reference for researchers and practitioners working with nano-satellites in the aerospace industry"--

2. Record Nr.	UNINA9910825591603321
Titolo	Algorithms in computational molecular biology : techniques, approaches and applications / / edited by Mourad Elloumi, Albert Y. Zomaya
Pubbl/distr/stampa	Hoboken, N.J., : Wiley, 2011
ISBN	9786612939631 9781282939639 1282939637 9780470892091 0470892099 9780470892107 0470892102
Descrizione fisica	1 online resource (1085 p.)
Collana	Wiley series on bioinformatics
Altri autori (Persone)	ElloumiMourad ZomayaAlbert Y
Disciplina	572.80285
Soggetti	Biophysics - Mathematical models
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 and index.
Nota di contenuto	ALGORITHMS INCOMPUTATIONALMOLECULAR BIOLOGY; CONTENTS; PREFACE; CONTRIBUTORS; I STRINGS PROCESSING AND APPLICATION TO BIOLOGICAL SEQUENCES; 1 STRING DATA STRUCTURES FOR COMPUTATIONAL MOLECULAR BIOLOGY; 1.1 Introduction; 1.2 Main String Indexing Data Structures; 1.2.1 Suffix Trees; 1.2.2 Suffix Arrays; 1.3 Index Structures for Weighted Strings; 1.4 Index Structures for Indeterminate Strings; 1.5 String Data Structures in Memory Hierarchies; 1.6 Conclusions; References; 2 EFFICIENT RESTRICTED-CASE ALGORITHMS FOR PROBLEMS IN COMPUTATIONAL BIOLOGY; 2.1 The Need for Special Cases 2.2 Assessing Efficient Solvability Options for General Problems and Special Cases2.3 String and Sequence Problems; 2.4 Shortest Common Superstring; 2.4.1 Solving the General Problem; 2.4.2 Special Case: SCSt for Short Strings Over Small Alphabets; 2.4.3 Discussion; 2.5 Longest Common Subsequence; 2.5.1 Solving the General Problem; 2.5.2

Special Case: LCS of Similar Sequences; 2.5.3 Special Case: LCS Under Symbol-Occurrence Restrictions; 2.5.4 Discussion; 2.6 Common Approximate Substring; 2.6.1 Solving the General Problem; 2.6.2 Special Case: Common Approximate String; 2.6.3 Discussion 2.7 ConclusionReferences; 3 FINITE AUTOMATA IN PATTERN MATCHING; 3.1 Introduction; 3.1.1 Preliminaries; 3.2 Direct Use of DFA in Stringology; 3.2.1 Forward Automata; 3.2.2 Degenerate Strings; 3.2.3 Indexing Automata; 3.2.4 Filtering Automata; 3.2.5 Backward Automata; 3.2.6 Automata with Fail Function; 3.3 NFA Simulation; 3.3.1 Basic Simulation Method; 3.3.2 Bit Parallelism; 3.3.3 Dynamic Programming; 3.3.4 Basic Simulation Method with Deterministic State Cache; 3.4 Finite Automaton as Model of Computation; 3.5 Finite Automata Composition; 3.6 Summary; References 4 NEW DEVELOPMENTS IN PROCESSING OF DEGENERATE SEQUENCES4.1 Introduction; 4.1.1 Degenerate Primer Design Problem; 4.2 Background; 4.3 Basic Definitions; 4.4 Repetitive Structures in Degenerate Strings; 4.4.1 Using the Masking Technique; 4.4.2 Computing the Smallest Cover of the Degenerate String  $x$ ; 4.4.3 Computing Maximal Local Covers of  $x$ ; 4.4.4 Computing All Covers of  $x$ ; 4.4.5 Computing the Seeds of  $x$ ; 4.5 Conservative String Covering in Degenerate Strings; 4.5.1 Finding Constrained Pattern  $p$  in Degenerate String  $T$ ; 4.5.2 Computing  $\gamma$ -Conservative Covers of Degenerate Strings 4.5.3 Computing  $\gamma$ -Conservative Seeds of Degenerate Strings4.6 Conclusion; References; 5 EXACT SEARCH ALGORITHMS FOR BIOLOGICAL SEQUENCES; 5.1 Introduction; 5.2 Single Pattern Matching Algorithms; 5.2.1 Algorithms for DNA Sequences; 5.2.2 Algorithms for Amino Acids; 5.3 Algorithms for Multiple Patterns; 5.3.1 Trie-Based Algorithms; 5.3.2 Filtering Algorithms; 5.3.3 Other Algorithms; 5.4 Application of Exact Set Pattern Matching for Read Mapping; 5.4.1 MPSCAN: An Efficient Exact Set Pattern Matching Tool for DNA/RNA Sequences; 5.4.2 Other Solutions for Mapping Reads 5.4.3 Comparison of Mapping Solutions

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## Sommario/riassunto

This book represents the most comprehensive and up-to-date collection of information on the topic of computational molecular biology. Bringing the most recent research into the forefront of discussion, Algorithms in Computational Molecular Biology studies the most important and useful algorithms currently being used in the field, and provides related problems. It also succeeds where other titles have failed, in offering a wide range of information from the introductory fundamentals right up to the latest, most advanced levels of study.

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