

1. Record Nr.	UNISA996200356903316
Titolo	Algorithms in Bioinformatics [[electronic resource]] : 15th International Workshop, WABI 2015, Atlanta, GA, USA, September 10-12, 2015, Proceedings / / edited by Mihai Pop, Hélène Touzet
Pubbl/distr/stampa	Berlin, Heidelberg : , : Springer Berlin Heidelberg : , : Imprint : Springer, , 2015
ISBN	3-662-48221-5
Edizione	[1st ed. 2015.]
Descrizione fisica	1 online resource (XX, 328 p. 85 illus.)
Collana	Lecture Notes in Bioinformatics ; ; 9289
Disciplina	572.80285
Soggetti	Bioinformatics Algorithms Computer science—Mathematics Data mining Proteins Computational Biology/Bioinformatics Algorithm Analysis and Problem Complexity Discrete Mathematics in Computer Science Data Mining and Knowledge Discovery Protein Structure
Lingua di pubblicazione	Inglese
Formato	Materiale a stampa
Livello bibliografico	Monografia
Note generali	Includes index.
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Sommario/riassunto

This book constitutes the refereed proceedings of the 15th International Workshop on Algorithms in Bioinformatics, WABI 2015, held in Atlanta, GA, USA, in September 2015. The 23 full papers presented were carefully reviewed and selected from 56 submissions. The selected papers cover a wide range of topics from networks to phylogenetic studies, sequence and genome analysis, comparative genomics, and RNA structure.
