Record Nr. UNINA9910828479303321 **Titolo** Bioinformatics: the impact of accurate quantification on proteomic and genetic analysis and research / / edited by Yu Liu, PhD Pubbl/distr/stampa Waretown, NJ:,: Apple Academic Press, Inc. Boca Raton, FL:,: CRC Press,, [2014] ©2014 **ISBN** 0-429-15779-7 1-4822-4662-7 Edizione [First edition.] 1 online resource (406 p.) Descrizione fisica Disciplina 570.285 Soggetti Bioinformatics Computational biology Lingua di pubblicazione Inglese **Formato** Materiale a stampa Livello bibliografico Monografia Description based upon print version of record. Note generali Includes bibliographical references at the end of each chapters. Nota di bibliografia Nota di contenuto Front Cover: About the Editor: Contents: Acknowledgments and How to Cite; List of Contributors; Introduction; Part I: RNA-Seq; Chapter 1: The Bench Scientist's Guide to Statistical Analysis of RNA-Seq Data; Chapter 2: Assembly of Non-Unique Insertion Content Using Next-Generation Sequencing; Chapter 3: RSEM: Accurate Transcript Quantification from RNA-Seg Data With or Without a Reference Genome: Part II: Microarray: Chapter 4: A Regression System for Estimation of Errors Introduced by Confocal Imaging into Gene Expression Data in Situ Chapter 5: SPACE: An Algorithm to Predict and Quantify Alternatively Spliced Isoforms Using MicroarraysChapter 6: Link-Based Quantitative Methods to Identify Differentially Coexpressed Genes and Gene Pairs; Chapter 7: Dimension Reduction with Gene Expression Data Using Targeted Variable Importance Measurement; Part III: GWAS; Chapter 8: Genome-Wide Association Study of Stevens-Johnson Syndrome and

Toxic Epidermal Necrolysis in Europe; Chapter 9: Genotyping Common

Chapter 10: Learning Genetic Epistasis Using Bayesian Network Scoring

Association Studies on vWF and FVIII Plasma Levels; Part IV: Proteomics;

and Rare Variation Using Overlapping Pool Sequencing

CriteriaChapter 11: Combined Analysis of Three Genome-Wide

	Chapter 12: Statistical Methods for Quantitative Mass Spectrometry Proteomic Experiments with Labeling; Chapter 13: MRCQuant: An Accurate LC-MS Relative Isotopic Quantification Algorithm on TOF Instruments; Author Notes
Sommario/riassunto	The book introduces bioinformatic and statistical methodology and shows approaches to bias correction and error estimation. It also presents quantitative methods for genome and proteome analysis.