1. Record Nr. UNINA9910877867003321 Autore Hamdan Mahmoud <1947-> Titolo Proteomics today: protein assessment and biomarkers using mass spectrometry, 2D electrophoresis, and microarray technology // Mahmoud Hamdan, Pier Giorgio Righetti Hoboken, N.J., : John Wiley & Sons, 2005 Pubbl/distr/stampa **ISBN** 1-280-27548-0 9786610275489 0-470-24684-7 0-471-70910-7 0-471-70915-8 Descrizione fisica 1 online resource (446 p.) Collana Wiley-Interscience series in mass spectrometry Altri autori (Persone) RighettiP. G Disciplina 572/.6 Soggetti Proteins - Spectra Mass spectrometry **Proteomics** 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. PROTEOMICS TODAY; CONTENTS: PREFACE TO PART I: Nota di contenuto ACKNOWLEDGMENT: I INTRODUCTION TO PART I: 1 INSTRUMENTATION AND DEVELOPMENTS; 1.1 Introduction; 1.2 Ionization Techniques for Macromolecules: 1.2.1 (252)Cf Plasma Desorption Ionization: 1.2.2 Fast Atom/Ion Bombardment; 1.2.3 Type of Fragment Ions and Nomenclature; 1.3 Examples on Analytical Solutions Based on FAB-MS; 1.3.1 Detection of Abnormalities in Hemoglobin; 1.3.2 Glycoprotein Structure Determination: 1.3.3 Early Scanning Functions on Sector Machines; 1.4 Electrospray Ionization; 1.5 Matrix-Assisted Laser **Desorption Ionization** 1.5.1 MALDI at High Pressure 1.5.2 Desorption Ionization on Silicon (DIOS); 1.5.3 Delayed Extraction; 1.6 Ion Detection; 1.6.1 Microchannel Plates (MCPs); 1.6.2 Cryogenic Detectors; 1.6.2.1 Superconducting Tunnel Junction; 1.6.2.2 Thermal Detectors; 1.7 Types of Analyzers; 1.7.1 Quadrupole Mass Filter; 1.7.2 Three-Dimensional Quadrupole Ion

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Sommario/riassunto

The last few years have seen an unprecedented drive toward the application of proteomics to resolving challenging biomedical and biochemical tasks. Separation techniques combined with modern mass spectrometry are playing a central role in this drive. This book discusses the increasingly important role of mass spectrometry in proteomic research, and emphasizes recent advances in the existing technology and describes the advantages and pitfalls as well.* Provides a scientifically valid method for analyzing the approximatey 500,000 proteins that are encoded in the human genome* Explains t