| Record Nr.<br>Autore<br>Titolo<br>Pubbl/distr/stampa | UNINA9910830637803321<br>McLachlan Geoffrey J. <1946-><br>Analyzing microarray gene expression data [[electronic resource] /] /<br>Geoffrey J. McLachlan, Kim-Anh Do, Christopher Ambroise<br>Hoboken, N.J., : Wiley-Interscience, c2004   |
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| ISBN   | 1-280-25332-0<br>9786610253326<br>0-470-35030-X<br>0-471-72612-5<br>0-471-72842-X  |
| Descrizione fisica                                   | 1 online resource (366 p.)   |
| Collana  | Wiley series in probability and statistics   |
| Altri autori (Persone)                               | DoKim-Anh <1960-><br>AmbroiseChristophe <1969->  |
| Disciplina   | 572.8636<br>572.865  |
| Soggetti   | DNA microarrays - Statistical methods<br>Gene expression - Statistical methods   |
| 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                                    | Analyzing Microarray Gene Expression Data; Contents; Preface; 1<br>Microarrays in Gene Expression Studies; 1.1 Introduction; 1.2<br>Background Biology; 1.2.1 Genome, Genotype, and Gene Expression;<br>1.2.2 Of Wild-Types and Other Alleles; 1.2.3 Aspects of Underlying<br>Biology and Physiochemistry; 1.3 Polymerase Chain Reaction; 1.4 cDNA;<br>1.4.1 Expressed Sequence Tag; 1.5 Microarray Technology and<br>Application; 1.5.1 History of Microarray Development; 1.5.2 Tools of<br>Microarray Technology; 1.5.3 Limitations of Microarray Technology;<br>1.5.4 Oligonucleotides versus cDNA Arrays<br>1.5.5 SAGE: Another Method for Detecting and Measuring Gene<br>Expression Levels1.5.6 Emerging Technologies; 1.6 Sampling of<br>Relevant Research Entities and Public Resources; 2 Cleaning and<br>Normalization; 2.1 Introduction; 2.2 Cleaning Procedures; 2.2.1 Image<br>Processing to Extract Information; 2.2.2 Missing Value Estimation; 2.2.3<br>Sources of Nonlinearity; 2.3 Normalization and Plotting Procedures for<br>Oligonucleotide Arrays; 2.3.1 Global Approaches for Oligonucleotide |

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|                    | Array Data; 2.3.2 Spiked Standard Approaches; 2.3.3 Geometric Mean<br>and Linear Regression Normalization for Multiple Arrays<br>2.3.4 Nonlinear Normalization for Multiple Arrays Using Smooth<br>Curves2.4 Normalization Methods for cDNA Microarray Data; 2.4.1<br>Single-Array Normalization; 2.4.2 Multiple Slides Normalization; 2.4.3<br>ANOVA and Related Methods for Normalization; 2.4.4 Mixed-Model<br>Method for Normalization; 2.4.5 SNOMAD; 2.5 Transformations and<br>Replication; 2.5.1 Importance of Replication; 2.5.2 Transformations;<br>2.6 Analysis of the Alon Data Set; 2.7 Comparison of Normalization<br>Strategies and Discussion; 3 Some Cluster Analysis Methods; 3.1<br>Introduction; 3.2 Reduction in the Dimension of the Feature Space<br>3.3 Cluster Analysis3.4 Some Hierarchical Agglomerative Techniques;<br>3.5 k-Means Clustering; 3.6 Cluster Analysis with No A Priori Metric;<br>3.7 Clustering via Finite Mixture Models; 3.7.1 Definition; 3.7.2<br>Advantages of Model-Based Clustering; 3.8 Fitting Mixture Models Via<br>the EM Algorithm; 3.9.1 E-Step; 3.8.2 M-Step; 3.8.3 Choice of Starting<br>Values for the EM Algorithm; 3.9 Clustering Via Normal Mixtures; 3.9.1<br>Heteroscedastic Components; 3.9.2 Homoscedastic Components; 3.9.3<br>Spherical Components; 3.9.4 Choice of Root; 3.9.5 Available Software;<br>3.10 Mixtures of Factor Analyzers3.12 Choice of Clustering Solution;<br>3.13 Classification ML Approach; 3.14 Mixture Models for Clinical and<br>Microarray Data; 3.14.1 Unconditional Approach; 3.14.2 Conditional<br>Approach; 3.15 Choice of the Number of Components in a Mixture<br>Model; 3.15.1 Order of a Mixture Model; 3.15.2 Approaches for<br>Assessing Mixture Order; 3.15.3 Bayesian Information Criterion; 3.15.4<br>Integrated Classification Likelihood Criterion; 3.16 Resampling<br>Approach; 3.17 Other Resampling Approaches for Number of Clusters;<br>3.17.1 The Gap Statistic<br>3.17.2 The Clest Method for the Number of Clusters |
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| Sommario/riassunto | A multi-discipline, hands-on guide to microarray analysis of biological<br>processes Analyzing Microarray Gene Expression Data provides a<br>comprehensive review of available methodologies for the analysis of<br>data derived from the latest DNA microarray technologies. Designed for<br>biostatisticians entering the field of microarray analysis as well as<br>biologists seeking to more effectively analyze their own experimental<br>data, the text features a unique interdisciplinary approach and a<br>combined academic and practical perspective that offers readers the<br>most complete and applied coverage of the subject   |