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Bayesian Methods in Epidemiology



Lyle D. Broemeling



Bayesian Methods In Epidemiology Chapman Hallcrc Biostatistics Series

Robert D. Gibbons, Anup Amatya

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Bayesian Methods in Epidemiology Lyle D. Broemeling, 2013-08-13 Written by a biostatistics expert with over 20 years of experience in the field Bayesian Methods in Epidemiology presents statistical methods used in epidemiology from a Bayesian viewpoint It employs the software package WinBUGS to carry out the analyses and offers the code in the text and for download online The book examines study designs that investigate the association between exposure to risk factors and the occurrence of disease It covers introductory adjustment techniques to compare mortality between states and regression methods to study the association between various risk factors and disease including logistic regression simple and multiple linear regression categorical ordinal regression and nonlinear models. The text also introduces a Bayesian approach for the estimation of survival by life tables and illustrates other approaches to estimate survival including a parametric model based on the Weibull distribution and the Cox proportional hazards nonparametric model Using Bayesian methods to estimate the lead time of the modality the author explains how to screen for a disease among individuals that do not exhibit any symptoms of the disease With many examples and end of chapter exercises this book is the first to introduce epidemiology from a Bayesian perspective It shows epidemiologists how these Bayesian models and techniques are useful in studying the association between disease and exposure to risk factors Bayesian Methods for Measures of Agreement Lyle D. Broemeling, 2009-01-12 Using WinBUGS to implement Bayesian inferences of estimation and testing hypotheses Bayesian Methods for Measures of Agreement presents useful methods for the design and analysis of agreement studies It focuses on agreement among the various players in the diagnostic process The author employs a Bayesian approach to provide statistical inferences Statistical Analysis of Human Growth and Development Yin Bun Cheung, 2013-10-28 Statistical Analysis of Human Growth and Development is an accessible and practical guide to a wide range of basic and advanced statistical methods that are useful for studying human growth and development Designed for nonstatisticians and statisticians new to the analysis of growth and development data the book collects methods scattered throughout the literature and explains how to use them to solve common research problems It also discusses how well a method addresses a specific scientific question and how to interpret and present the analytic results Stata is used to implement the analyses with Stata codes and macros for generating example data sets a detrended Q Q plot and weighted maximum likelihood estimation of binary items available on the book s CRC Press web page After reviewing research designs and basic statistical tools the author discusses the use of existing tools to transform raw data into analyzable variables and back transform them to raw data He covers regression analysis of quantitative binary and censored data as well as the analysis of repeated measurements and clustered data He also describes the development of new growth references and developmental indices the generation of key variables based on longitudinal data and the processes to verify the validity and reliability of measurement tools Looking at the larger picture of research practice the book concludes with coverage of missing values

multiplicity problems and multivariable regression Along with two simulated data sets numerous examples from real experimental and observational studies illustrate the concepts and methods Although the book focuses on examples of anthropometric measurements and changes in cognitive social emotional locomotor and other abilities the ideas are applicable to many other physical and psychosocial phenomena such as lung function and depressive symptoms of Drug Development Eric B. Holmgren, 2013-10-24 Theory of Drug Development presents a formal quantitative framework for understanding drug development that goes beyond simply describing the properties of the statistics in individual studies It examines the drug development process from the perspectives of drug companies and regulatory agencies By quantifying various ideas underlying drug development the book shows how to systematically address problems such as Sizing a phase 2 trial and choosing the range of p values that will trigger a follow up phase 3 trial Deciding whether a drug should receive marketing approval based on its phase 2 3 development program and recent experience with other drugs in the same clinical area Determining the impact of adaptive designs on the quality of drugs that receive marketing approval Designing a phase 3 pivotal study that permits the data driven adjustment of the treatment effect estimate Knowing when enough information has been gathered to show that a drug improves the survival time for the whole patient population Drawing on his extensive work as a statistician in the pharmaceutical industry the author focuses on the efficient development of drugs and the quantification of evidence in drug development He provides a rationale for underpowered phase 2 trials based on the notion of efficiency which leads to the identification of an admissible family of phase 2 designs He also develops a framework for evaluating the strength of evidence generated by clinical trials This approach is based on the ratio of power to type 1 error and transcends typical Bayesian and frequentist statistical analyses Benefit-Risk Assessment in Pharmaceutical Research and Development Andreas Sashegyi, James Felli, Rebecca Noel, 2013-11-27 Many practitioners in the pharmaceutical industry are still largely unfamiliar with benefit risk assessment despite its growing prominence in drug development and commercialization Helping to alleviate this knowledge gap Benefit Risk Assessment in Pharmaceutical Research and Development provides a succinct overview of the key considerations relevant to benefit risk assessment across the pharmaceutical R D spectrum from early clinical development to late stage development to regulatory review to post launch assessment The book first presents interpretations of benefit and risk in the context of a molecule moving from preclinical evaluation into its early testing in humans It next considers benefit and risk characterization and assessment during a molecule s journey from its clinical evaluation in humans through its submission to regulators for marketing approval Throughout these sections the book offers insight into the role of benefit risk assessment in heightening understanding among key stakeholders by shaping questions and guiding discussions among scientists physicians developers and regulatory agencies The book also focuses on a molecule s entry into the marketplace as a drug available for consumption by people It explores the role of benefit risk assessment as the relevance of carefully collected clinical efficacy and safety metrics fades in

the wake of real world use and evidence of effectiveness and safety Bringing together the expertise of 15 contributors from academia and the industry this book offers an easy to read guide to the various facets of benefit risk assessment in the major stages of pharmaceutical R D Suitable for those in both technical and managerial roles it enables readers to communicate more effectively across their development chain as well as rationally and thoughtfully embed benefit risk assessment into Data Analysis with Competing Risks and Intermediate States Ronald B. Geskus, 2015-07-14 This practical and thorough book explains when and how to use models and techniques for the analysis of competing risks and intermediate states It covers the most recent insights on estimation techniques and discusses in detail how to interpret the obtained results Each chapter includes standard exercises a software section on SAS Stata and R and computer practicals that allow readers to practice with the techniques The book s website provides the R code for the computer practicals along with other material Inference Principles for Biostatisticians Ian C. Marschner, 2014-12-11 Designed for students training to become biostatisticians as well as practicing biostatisticians Inference Principles for Biostatisticians presents the theoretical and conceptual foundations of biostatistics It covers the theoretical underpinnings essential to understanding subsequent core methodologies in the field Drawing on his extensive experience teaching graduate level biostatistics courses and working in the pharmaceutical industry the author explains the main principles of statistical inference with many examples and exercises Extended examples illustrate key concepts in depth using a specific biostatistical context In addition the author uses simulation to reinforce the repeated sampling interpretation of numerous statistical concepts Reducing the computational complexities he provides simple R functions for conducting simulation studies This text gives graduate students with diverse backgrounds across the health medical social and mathematical sciences a solid unified foundation in the principles of statistical inference This groundwork will lead students to develop a thorough understanding of biostatistical methodology Statistical Methods for Healthcare Performance Monitoring Alex Bottle, Paul Aylin, 2016-08-05 Healthcare is important to everyone yet large variations in its quality have been well documented both between and within many countries With demand and expenditure rising it s more crucial than ever to know how well the healthcare system and all its components from staff member to regional network are performing This requires data which inevitably differ in form and quality It also requires statistical methods the output of which needs to be presented so that it can be understood by whoever needs it to make decisions Statistical Methods for Healthcare Performance Monitoring covers measuring quality types of data risk adjustment defining good and bad performance statistical monitoring presenting the results to different audiences and evaluating the monitoring system itself Using examples from around the world it brings all the issues and perspectives together in a largely non technical way for clinicians managers and methodologists Statistical Methods for Healthcare Performance Monitoring is aimed at statisticians and researchers who need to know how to measure and compare performance health service regulators health service managers

with responsibilities for monitoring performance and quality improvement scientists including those involved in clinical audits Cancer Clinical Trials Stephen L. George, Xiaofei Wang, Herbert Pang, 2016-08-19 Cancer Clinical Trials Current and Controversial Issues in Design and Analysis provides statisticians with an understanding of the critical challenges currently encountered in oncology trials Well known statisticians from academic institutions regulatory and government agencies such as the U S FDA and National Cancer Institute and the pharmaceutical industry share their extensive experiences in cancer clinical trials and present examples taken from actual trials The book covers topics that are often perplexing and sometimes controversial in cancer clinical trials Most of the issues addressed are also important for clinical trials in other settings After discussing general topics the book focuses on aspects of early and late phase clinical trials It also explores personalized medicine including biomarker based clinical trials adaptive clinical trial designs and dynamic treatment regimes Statistical Methods for Drug Safety Robert D. Gibbons, Anup Amatya, 2015-07-28 Explore Important Tools for High Quality Work in Pharmaceutical Safety Statistical Methods for Drug Safety presents a wide variety of statistical approaches for analyzing pharmacoepidemiologic data It covers both commonly used techniques such as proportional reporting ratios for the analysis of spontaneous adverse event reports and newer approaches such as the use of marginal structural models for controlling dynamic selection bias in the analysis of large scale longitudinal observational data Choose the Right Statistical Approach for Analyzing Your Drug Safety Data The book describes linear and non linear mixed effects models discrete time survival models and new approaches to the meta analysis of rare binary adverse events It explores research involving the re analysis of complete longitudinal patient records from randomized clinical trials The book discusses causal inference models including propensity score matching marginal structural models and differential effects as well as mixed effects Poisson regression models for analyzing ecological data such as county level adverse event rates The authors also cover numerous other methods useful for the analysis of within subject and between subject variation in adverse events abstracted from large scale medical claims databases electronic health records and additional observational data streams Advance Statistical Practice in Pharmacoepidemiology Authored by two professors at the forefront of developing new statistical methodologies to address pharmacoepidemiologic problems this book provides a cohesive compendium of statistical methods that pharmacoepidemiologists can readily use in their work It also encourages statistical scientists to develop new methods that go beyond the foundation covered in the text

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