# GENETIC INFORMATION

Acquisition, Access, and Control

Edited by Alison K. Thompson and Ruth F. Chadwick

# **Genetic Information Acquisition Access And Control**

**Herman T. Tavani** 

#### **Genetic Information Acquisition Access And Control:**

Genetic Information Alison K. Thompson, Ruth F. Chadwick, 2007-07-23 It is difficult to think of an example of an advancement in the biological sciences that has had an impact on society similar to that of the new genetics Recent developments in biotechnology have occasioned much discussion among academics professionals and lay people alike In particular many questions and concerns have arisen over the acquisi tion access and control of genetic information There are several reasons why the new genetics has commanded such widespread attention and why it is now the subject of con siderable debate Special reference is given in this volume to the implications of genetic information for five different subject areas eugenics the insurance industry the commer cialisation of genetic testing strategies for raising public awareness and the value of theo retical ethical and sociological frameworks in the debate This diverse collection of papers attempts to address and critically discuss issues surrounding the control of and access to genetic information from ethical medical legal and theoretical points of view The first and shortest section of the book attempts to address concerns over the eugenic potential of new biotechnologies It also provides a historical context for the de bate for controversy over the subject of eugenics predates the current debate over genetic information by a considerable length of time Indeed by the time the first patent was is sued for Chakrabarty's strain of oil eating bacteria in the early 1970s the term eugenics had already acquired The Governance of Genetic Information Heather Widdows, Caroline Mullen, 2009-09-03 strong pejorative connotations This volume maps the areas of ethical concern in the debate regarding the governance of genetic information and suggests alternative ethical frameworks and models of regulation in order to inform its restructuring Genetic governance is at the heart of medical and scientific developments and is connected to global exploitation issues of commodification commercialisation and ownership the concepts of property and intellectual property and concerns about individual and communal identity Thus the decisions that are made in the next few years about appropriate models of genetic governance will have knock on effects for other areas of governance In short the final answer to Who Decides in the context of genetic governance will fundamentally shape the ethical constructs of individuals and their networks and relationships in the public sphere Genetic Information Alison K. Thompson, Ruth F. Chadwick, 2013-03-10 It is difficult to think of an example of an advancement in the biological sciences that has had an impact on society similar to that of the new genetics Recent developments in biotechnology have occasioned much discussion among academics professionals and lay people alike In particular many questions and concerns have arisen over the acquisi tion access and control of genetic information There are several reasons why the new genetics has commanded such widespread attention and why it is now the subject of con siderable debate Special reference is given in this volume to the implications of genetic information for five different subject areas eugenics the insurance industry the commer cialisation of genetic testing strategies for raising public awareness and the value of theo retical ethical and sociological frameworks in the debate This diverse collection of papers attempts to

address and critically discuss issues surrounding the control of and access to genetic information from ethical medical legal and theoretical points of view The first and shortest section of the book attempts to address concerns over the eugenic potential of new biotechnologies It also provides a historical context for the de bate for controversy over the subject of eugenics predates the current debate over genetic information by a considerable length of time Indeed by the time the first patent was is sued for Chakrabarty's strain of oil eating bacteria in the early 1970s the term eugenics had already acquired strong pejorative connotations Governing Biobanks Jane Kaye, Susan Gibbons, Catherine Heeney, Andrew Smart, 2012-02-03 Biobanks are proliferating rapidly worldwide because they are powerful tools and organisational structures for undertaking medical research By linking samples to data on the health of individuals it is anticipated that biobanks will be used to explore the relationship between genes environment and lifestyle for many diseases as well as the potential of individually tailored drug treatments based on genetic predisposition However they also raise considerable challenges for existing legal frameworks and research governance structures. This book critically examines the current governance structures in place for biobanks in England and Wales It shows that the technologies techniques and practices involved in biobanking do not always conform neatly to existing legal principles and frameworks that apply to other areas of medical research Using a socio legal approach including interview data gathered from the scientific community this book provides unique insights and makes recommendations about appropriate governance mechanisms for biobanking in the future It also explores the issues around the secondary use of information such as consent and how to protect privacy when biobanks are accessed by a number of different third parties These issues have relevance both within England and Wales and to a wide international audience as well as for other areas where large datasets are used **Encyclopedia of Applied** Ethics, 2012-01-10 The Encyclopedia of Applied Ethics Second Edition Four Volume Set addresses both the physiological and the psychological aspects of human behavior Carefully crafted well written and thoroughly indexed the encyclopedia helps users whether they are students just beginning formal study of the broad field or specialists in a branch of psychology understand the field and how and why humans behave as we do The work is an all encompassing reference providing a comprehensive and definitive review of the field A broad and inclusive table of contents ensures detailed investigation of historical and theoretical material as well as in depth analysis of current issues Several disciplines may be involved in applied ethics one branch of applied ethics for example bioethics is commonly explicated in terms of ethical legal social and philosophical issues Editor in Chief Ruth Chadwick has put together a group of leading contributors ranging from philosophers to practitioners in the particular fields in question to academics from disciplines such as law and economics The 376 chapters are divided into 4 volumes each chapter falling into a subject category including Applied Ethics Bioethics Computers and Information Management Economics Business Environmental Ethics Ethics and Politics Legal Medical Ethics Philosophy Theories Social and Social Media Concise entries ten pages on average provide foundational knowledge of the

field Each article will features suggested readings pointing readers to additional sources for more information a list of related websites a 5 10 word glossary and a definition paragraph and cross references to related articles in the encyclopedia Newly expanded editorial board and a host of international contributors from the US Australia Belgium Canada France Germany Ireland Israel Japan Sweden and the United Kingdom The 376 chapters are divided into 4 volumes each chapter falling into a subject category including Applied Ethics Bioethics Computers and Information Management Economics Business Environmental Ethics Ethics and Politics Legal Medical Ethics Philosophy Theories Social and Social Media

Genetic Data and the Law Mark Taylor, 2012-03-08 Research using genetic data raises various concerns relating to privacy protection Many of these concerns can also apply to research that uses other personal data but not with the same implications for failure The norms of exclusivity associated with a private life go beyond the current legal concept of personal data to include genetic data that relates to multiple identifiable individuals simultaneously and anonymous data that could be associated with any number of individuals in different but reasonably foreseeable contexts It is the possibilities and implications of association that are significant and these possibilities can only be assessed if one considers the interpretive potential of data They are missed if one fixates upon its interpretive pedigree or misunderstands the meaning and significance of identification This book demonstrates how the public interest in research using genetic data might be reconciled with the public interest in proper privacy protection Human Genes and Neoliberal Governance Antoinette Rouvroy, 2007-12-13 Original and interdisciplinary this is the first book to explore the relationship between a neoliberal mode of governance and the so called genetic revolution Looking at the knowledge power relations in the post genomic era and addressing the pressing issues of genetic privacy and discrimination in the context of neoliberal governance this book demonstrates and explains the mechanisms of mutual production between biotechnology and cultural political economic and legal frameworks In the first part Antoinette Rouvroy explores the social political and economic conditions and consequences of this new perceptual regime In the second she pursues her analysis through a consideration of the impact of geneticization on political support of the welfare state and on the operation of private health and life insurances Genetics and neoliberalism she argues are complicit in fostering the belief that social and economic patterns have a fixed nature beyond the reach of democratic deliberation whilst the characteristics of individuals are unusually plastic and within the scope of individual choice and responsibility This book will be of interest to all students of law sociology and politics Genetic Democracy Veikko Launis, Juha Räikkä, 2007-12-20 Genetic Democracy involves an in depth analysis of the ethical social and philosophical issues related to modern genetic research and gene technology The aim of the book is to introduce systematic research on the social and ethical impacts of the use and development of genetically modified organisms GMOs as well as the acquisition use and storage of human genetic information HGI The book contributes to enhancing public discussion and reaching fair and democratic decision making practices in GMO and HGI use and development both on local and global level

There are currently few European texts which address the issues involved in a theoretical and systematical manner Genetic Democracy has been written from the viewpoint of social and political philosophy rather than that of traditional bioethics There is a clear need for a throughout and authoritative philosophical and ethical analysis of the issues involved in genetic research and gene technology The book will appeal to philosophers social scientists genetics professionals policy makers academics industrial organisations and human rights organisations as well as university students and legal scholars The book will have a broad appeal across Europe Asia and America since many states are currently considering policy responses to many of the practices discussed in the books e g human biobanks Human Genome News ,1989 Genetic Databases Oonagh Corrigan, Richard Tutton, 2004 Genetic Databases offers a timely analysis of the underlying tensions contradictions and limitations of the current regulatory frameworks for and policy debates about genetic databases Drawing on original empirical research and theoretical debates in the fields of sociology anthropology and legal studies the contributors to this book challenge the prevailing orthodoxy of informed consent and explore the relationship between personal privacy and the public good They also consider the multiple meanings attached to human tissue and the role of public consultations and commercial involvement in the creation and use of genetic databases. The authors argue that policy and regulatory frameworks produce a representation of participation that is often at odds with the experiences and understandings of those taking part The findings present a serious challenge for public policy to provide mechanisms to safeguard the welfare of individuals participating in genetic databases **Informed Insurance Choice?** Leander D. Loacker, 2015-04-30 The direction and clarity of the author's argument is commendably clear. Thus it is clear at the outset that he is mainly concerned with pre contractual information duties as they affect consumers and thus standard form contracts although he argu

Ethics, Law and Society Jennifer Gunning,2017-07-28 This key collection brings together a selection of papers commissioned and published by the Cardiff Centre for Ethics Law Society It incorporates contributions from a group of international experts along with a selection of short opinion pieces written in response to specific ethical issues The collection addresses issues arising in biomedical and medical ethics ranging from assisted reproductive technologies to the role of clinical ethics committees It examines broader societal issues with particular emphasis on sustainability and the environment and also focuses on issues of human rights in current global contexts The contributors collect responses to issues arising from high profile cases such as the legitimacy of war in Iraq to physician related suicide The volume will provide a valuable resource for practitioners and academics with an interest in ethics across a range of disciplines

Analysing Citizenship Talk Heiko Hausendorf, Alfons Bora, 2006-02-15 Citizenship talk refers to various types of discourse initiated to make citizens take part in politically and socially contested decision making processes citizen participation Citizenship has accordingly become one of the dazzling key words whenever the democratic deficit of modern societies is moaned about Asking for citizenship to be conceived of as a communicative achievement the present book shows that

sociolinguistics and pragmatics can essentially contribute to this interdisciplinary up to date issue of research the volume offers a theoretically innovative concept of communicated citizenship and it presents a set of methodological approaches suited to deal with this concept at an empirical level including contributions from Conversation Analysis Critical Discourse Analysis Social Positioning Theory Speech Act Theory and Ethnography Furthermore concrete data and empirical analyses are provided which take up the case of decision making processes around the application of modern green biotechnology GMO field trials The volume thus illustrates the kind of findings and results that can be expected from this new and promising approach towards citizenship talk Ethics, Computing, and Genomics Herman T. Tavani, 2006 Comprised of eighteen chapters contributed by experts in the fields of biology computer science information technology law and philosophy Ethics Computing and Genomics provides instructors with a flexible resource for undergraduate and graduate courses in an exciting new field of applied ethics computational genomics. The chapters are organized in a way that takes the reader from a discussion of conceptual frameworks and methodological perspectives including ethical theory to an in depth analysis of controversial issues involving privacy and confidentiality information consent and intellectual property The volume concludes with some predictions about the future of computational genomics including the role that nanotechnology will likely play as biotechnologies and information technologies continue to converge Implementation of the Data Protection Directive in Relation to Medical Research in Europe D. Townend, S. Rouille-Mirza, J. Wright, D. Beyleveld, 2017-11-28 The Data Protection and Medical Research in Europe PRIVIREAL series focuses on the Privacy in Research Ethics and Law EC funded project examining the implementation of Directive 95 46 EC on data protection in relation to medical research and the role of ethics committees in European countries The series consists of five separate volumes following the complete development of the PRIVIREAL project This volume relates to the first stage of this project concerning the implementation of the Data Protection Directive in particular in the area of medical research It contains reports from 26 European countries on the implementation of the Directive or the data protection regime all with a specific focus on issues and questions relating to medical research Presenting a unique resource for all those involved in data protection medical research and their implications for each other this title provides a valuable insight into the actual workings across Europe including both the New Member States and the Newly Associated Member States Brave New World? Celia Deane-Drummond, 2003-11-01 One of the key issues facing us in the next millennium is the ability to manipulate the genetics of living organisms. The possibility of manipulating human genetics raises many theological ethical and socio political issues. These include specific decisions about whether the technology will be developed how it will be applied and more general questions about the technical manipulation of natural processes From a theological perspective the human genome project not only challenges particular doctrines such as that of creation eschatology and anthropology but also raises particular issues of social justice and medical ethics The purpose of this book is to bring together the collective expertise of theologians scientists and social scientists in order to provide a

forum for critique and public debate focused on the human genome project It is hoped that the results presented in this book offer a sophisticated theological and ethical response The Ethics and Governance of Human Genetic Databases Matti Häyry, Ruth Chadwick, Vilhjálmur Árnason, Gardar Árnason, 2007-04-19 The Medical Biobank of Ume in Sweden deCODE's Health Sector Database in Iceland the Estonian Genome Project and the UK Biobank contain health data and genetic data from large populations Some include genealogical or lifestyle information They are resources for research in human genetics and medicine exploring interaction between genes lifestyle environmental factors and health and diseases The collection storage and use of this data raise ethical legal and social issues In this book first published in 2007 bioethics scholars examine whether existing ethical frameworks and social policies reflect people's concerns and how they may need to change in light of new scientific and technological developments The ethical issues of social justice genetic discrimination informational privacy trust in science and consent to participation in database research are analyzed whilst an empirical survey conducted in the four countries demonstrates public views of privacy and related moral values in the context of human genetic databases The Ethics of Screening in Health Care and Medicine Niklas Juth, Christian Munthe, 2011-09-20 Medical or health oriented screening programs are amongst the most debated aspects of health care and public health practices in health care and public health ethics as well as health policy discussions In spite of this most treatments of screening in the research literature restrict themselves to isolated scientific aspects sometimes complemented by economic analyses or loose speculations regarding policy aspects At the same time recent advances in medical genetics and technology as well as a rapidly growing societal focus on public health concerns inspires an increase in suggested or recently started screening programs This book involves an in depth analysis of the ethical political and philosophical issues related to health oriented screening programs It explores the considerations that arise when heath care interacts with other societal institutions on a large scale as is the case with screening What values may be promoted or compromised by screening programs What conflicts of values do typically arise both internally and in relation to the goals of health care on the one hand and the goals of public health and the general society on the other What aspects of screening are relevant for determining whether it should be undertaken or not and how it should be organised in order to remain defensible What implications does the ethics of screening have for health care ethics as a whole These questions are addressed by applying philosophical methods of conceptual analysis as well as models and theories from moral and political philosophy medical ethics and public health ethics to a large number of ongoing and proposed screening programs which makes this book the first comprehensive work on the ethics of screening Analyses and suggestions are made that are of potential interest to health care staff medical researchers policy makers and the general public Genes and Insurance Marcus Radetzki, Marian Radetzki, Niklas Juth, 2003-11-06 The result of two key social developments in recent years are examined here the partial dismantling of the welfare state and the progress of genetics Genetic insights are increasingly valuable for risk assessment

and insurers would like to use these insights to help determine premiums Combined with the fact that social welfare is being curtailed this could potentially create an uninsured high risk population Along with considerations of autonomy and privacy this is the basis for an ethical critique of insurer s access to information The result has often been regulation of such information but the authors argues that due to adverse selection regulation will not solve these problems and this may jeopardize the survival of private personal insurance Instead we should look towards the resurrection of social insurance a key component of the welfare state This will interest academic researchers as well as professionals involved with genetics and insurance Genetic Information Values and Rights Niklas Juth, 2005

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