Sociological
Perspectives

O 2 the
New Genetics

Edited by

Peter Conrad Jonathan Gabe



Sociological Perspectives On The New Genetics

Carole H. Browner, Mabel H. Preloran

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Sociological Perspectives on the New Genetics Peter Conrad, Jonathan Gabe, 2000-01-21 With the advent of the Human Genome Project the new genetics has moved to the cutting edge of science and medicine The development and use of such genetics will have a profound impact on our understanding of disease and behaviour This volume presents new sociological research which explores the structure and production of genetic knowledge its social meaning impact and implication s for The New Genetics and The Public's Health Robin Bunton, Alan Petersen, 2002-09-11 The rapid advancement of society genetic science fuelled by the Human Genome Project and other related initiatives promises a new kind of public health practice based on the pre detection of disease according to calculations of genetic risk This book by two well known sociologists explores the implications of the new genetics for public health as a body of knowledge and a domain of practice assesses the impact of new genetic information and technologies on conceptions of health illness embodiment self and citizenship critically examines the complex discourses surrounding human genetics and public health The New Genetics and The Public's Health addresses the emerging social and political consequences of the new genetics and provides a stimulating critique of current research and practice in public health New Genetics, New Identities Paul Atkinson, Peter Glasner, Helen Greenslade, 2006-12-15 Genetic advocacy groups science and biovalue creating political economies of hope Carlos Novas Patients as public in ethics debates interpreting the role of patient organizations in democracy Annemiek Nelis Gerard de Vries and Rob Hagendijk From scraps and fragments to whole organisms molecular biology clinical research and post genomic bodies Susan E Kelly Fashioning flesh inclusion exclusivity and the potential of genomics Fiona O Neill Mapping origins race and relatedness in population genetics and genetic genealogy Catherine Nash Genetics and the Social Sciences Sahra Gibbon, Carlos Novas, 2007-07-20 This pioneering collection uses Paul Rabinow s concept of biosociality to chart the shifts in social relations and in ideas about nature biology and identity brought about by developments in biomedicine Genetic Governance Robin Bunton, Alan Petersen, 2005-03-18 Ethical and practical issues around genetic research are of major international concern both in academia and in the public domain Questions concerning what interventions are possible and appropriate with the increasing amount of genetic information available challenge our understandings of ourselves our health and wellbeing and the role of medical ethics public health surveillance and risk However there has been little reflection on the socio political effects of this new genetic knowledge and the changes in practice that are currently impacting on our lives Containing contributions from key international researchers this book examines the broader issues of genetic debates and looks at how prediction and risk assessment is being changed in the arenas of health medicine and reproduction bringing new insight on the dangers of surveillance regulation and increased inequality Developed out of the Taylor and Francis journal Critical Public Health the book considers the implications of developments in genetics for contemporary liberal governance as well as for the future of healthcare and public health

Social Theory, Health and Healthcare Orla McDonnell, Maria Lohan, Abbey Hyde, 2009-06-30 Health research education and provision have become increasingly interdisciplinary over the last few years leading health professionals to broaden their knowledge beyond technical aspects of care Practitioners now need a clear understanding of how society can affect health and an appreciation of how societal structures can drive healthcare practices In a clear systematic and accessible style this timely text looks at the social context of health and healthcare by Analysing a wide range of classic and contemporary theories Identifying the relevance of each theory to health Showing how theory has been used in research Outlining the impact of theory on health and health provision Specifically written for health professionals and those engaged in health studies research this book will help students and practitioners alike understand the sociology of health and illness and enable them to critically assess health issues policies and practices Perspectives on Genetic Discrimination Thomas Lemke, 2013-07-18 Over the past 15 years a series of empirical studies in different countries have shown that our increasing genetic knowledge leads to new forms of exclusion disadvantaging and stigmatization The spectrum of this genetic discrimination ranges from disadvantages at work via problems with insurance policies to difficulties with adoption agencies The empirical studies on the problem of genetic discrimination have not gone unnoticed Since the beginning of the 1990s a series of legislative initiatives and statements both on the national level and on the part of international and supranational organizations and commissions have been put forward as ways of protecting people from genetic discrimination This is the first book to critically evaluate the empirical evidence and the theoretical usefulness of the concept of genetic discrimination It discusses the advantages and limitations of adopting the concept and offers a more complex account distinguishing between several dimensions and forms of genetic discrimination Health Travels: Cuban Health(care) On and Off the <u>Island</u> Nancy Burke, 2013 This collection of essays challenges static and binary discourses regarding the Cuban healthcare system bringing together papers that paint a nuanced and dynamic picture of the intricacies of Cuban health care as it is represented and experienced both on the island and around the world Biomedicalization of Alcohol Studies Lorraine Midanik, 2019-01-22 Biomedicalization is seen as the natural outgrowth of continued scientific progress a movement towards improving the quality and quantity of life through scientific inquiries using biomedical perspectives and methods This approach carries with it the assumption that with proper risk assessment detection and treatment our lives can be lengthened improved and indeed more fulfilling Yet critics question biomedicalization s ability to deliver There is concern about how biomedicalization can change our traditional concepts of health as we discover more conditions for which we are at risk and health maintenance is seen as the responsibility of the individual The purpose of the book is to describe assess and critique biomedicalization and its influence as a larger social trend on the health field and specifically in the area of alcohol research policy and programs Chapter 1 gives a broad overview of biomedicalization Chapter 2 lays the groundwork for a historical understanding of how medicalization and biomeidcalization have developed and are expressed in diverse

fields such as aging psychiatry mental health and women s health Chapter 3 focuses in depth on alcoholism and assesses the development and assumptions underlying the two movements that have greatly influenced the substance abuse field the medicalization of deviance and the growth of the disease model of alcoholism Chapter 4 discusses the origins and development of the National Institute on Alcohol Abuse and Alcoholism NIAAA from its inception in 1970 Chapter 5 illustrates the growing biomedicalization that has occurred in the alcohol field prior to NIAAA s movement to the National Institute of Health NIH Chapter 6 assesses how Sweden has handled alcohol problems and currently funds alcohol research Chapter 7 concludes with a rationale for an expanded discourse between social scientists and biomedical researchers working on social problems particularly alcohol issues This volume will stimulate discussion of the processes by which social problems and specifically alcohol issues are framed managed and studied It will hold particular interest for researchers and students in the areas of alcohol studies social science and social welfare Lorraine Midanik is a professor in the School of Social Welfare University of California Berkeley **Rethinking Social Policy** Gail Lewis, Sharon Gewirtz, John Clarke, 2000-03-28 Rethinking Social Policy is a comprehensive introduction to and analysis of the complex mixture of problems and possibilities within the study of social policy Contributors at the cutting edge of social policy analysis reflect upon the implications of new social and theoretical movements for welfare and the study of social policy Topics covered include criminology and crime control race class and gender poverty and sexuality the body and the emotions violence work and welfare in Europe Examples are drawn from a variety of welfare sectors such as social services and community care health education employment and criminal justice This is a course reader for The Open University course D860 Rethinking The Sociology of Healthcare Safety and Quality Davina Allen, Jeffrey Braithwaite, Jane Sandall, Justin Social Practice Waring, 2016-10-17 The Sociology of Healthcare Safety and Quality presents a series of research informed readings on the sociological contributions of technologies practices experiences and organizational quality and safety across a range of healthcare contexts Represents the first collection of peer reviewed research articles showcasing ways that sociology can contribute to the ongoing policy concern of healthcare safety and quality Features original contributions from leading experts in healthcare related fields from three continents Reveals the state of the art in sociological analyses of contemporary healthcare safety and quality along with future directions in the field Offers sociological insights from the perspectives of managers clinicians and patients Routledge Handbook of Genomics, Health and Society Sahra Gibbon, Barbara Prainsack, Stephen Hilgartner, Janelle Lamoreaux, 2018-04-17 The Handbook provides an essential resource at the interface of Genomics Health and Society and forms a crucial research tool for both new students and established scholars across biomedicine and social sciences Building from and extending the first Routledge Handbook of Genetics and Society the book offers a comprehensive introduction to pivotal themes within the field an overview of the current state of the art knowledge on genomics science and society and an outline of emerging areas of research Key themes addressed include the way

genomic based DNA technologies have become incorporated into diverse arenas of clinical practice and research whilst also extending beyond the clinic the role of genomics in contemporary bioeconomies how challenges in the governance of medical genomics can both reconfigure and stabilise regulatory processes and jurisdictional boundaries how questions of diversity and justice are situated across different national and transnational terrains of genomic research and how genomics informs and is shaped by developments in fields such as epigenetics synthetic biology stem cell microbial and animal model research Chapters 13 and 28 of this book are freely available as downloadable Open Access PDFs at http www taylorfrancis com under Genetic Databases Oonagh a Creative Commons Attribution Non Commercial No Derivatives CC BY NC ND 4 0 license Corrigan, Richard Tutton, 2004-07-31 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 Trames ,2007 Genetics and Primary Care Imran Rafi, John Spicer, 2018-05-08 Increasingly primary care professionals are faced with challenges in dealing with patients who have been affected by a genetic disorder or whose family history is of concern A basic understanding of clinical genetics and the role of the genetics centres leads to greater confidence in the management of these patients. This book is an ideal introduction to the principles of genetics It outlines the key influences that will affect primary care including screening programmes the role of genetics education such as the RCGP Genetics Curriculum and national guidelines It provides information on basic clinical genetics and includes some of the more common clinical genetic conditions seen in primary care such as cystic fibrosis breast cancer and the haemoglobinopathies It addresses some of the key ethical issues that may be faced including patient confidentiality the ethics of reproductive genetic medicine and relevant medico legal cases The wider societal impact of genetics is also discussed An introduction to the increasing impact of genetics into primary care this book is invaluable for every primary healthcare professional Ageing, Dementia and the Social Mind Paul Higgs, Chris Gilleard, 2017-06-28 A groundbreaking exploration of the sociology of dementia with contributions from distinguished international scholars and practitioners Organised around the four themes of personhood care social representations and social differentiation Provides a critical look at dementia and demonstrates how sociology and other disciplines can help us understand its social context as well as the challenges it poses Contributing authors explore the social terrain responding in part to Paul Higgs and Chris

Gilleard's highly influential work on ageing Breaks new ground in giving specific attention to the social and cultural dimensions of responses to dementia **Crossing Over** Frank Timmermans, 2005 Crossing Over promotes reflection on the socio ethical and policy and regulatory aspects of genomics and biotechnology This collection of essays illuminates the common themes surrounding this hotly contested issue and considers the politics public perception ethics media representations and future possibilities of genomics Crime, Genes, Neuroscience and Cyberspace Tim Owen, 2017-09-18 This book applies Owen's unique genetic social framework to the study of crime and criminal behaviour with an emphasis on cybercrime Moving beyond challenges which confront contemporary criminological theorizing such as the stagnation of critical criminology the relativistic nihilism of the cultural turn posthumanism and virtual criminology the author codifies and applies the latest version of the framework to the study of crime both in and out of cyberspace Drawing upon evolutionary psychology behavioural genetics and the philosophy of Heidegger he introduces new terms such as Neuro Agency and notions of Embodied Cognition into criminological theorizing Adopting a soft compatibilist approach to free will and Realist ontology Owen s meta theoretical focus provides a new direction for criminological theorizing in particular in the direction of the conceptualization and prediction of cyber violence Exciting and timely this book will appeal to scholars and advanced students of criminology law sociology social policy psychology philosophy policing and forensic investigation

Neurogenetic Diagnoses, the Power of Hope, and the Limits of Today's Medicine Carole H. Browner, Mabel H. Preloran, 2009-12-18 Amid intense debate over the consequences of decoding the human genome and the impact of such technology on our lives these lucid richly textured jargon free case studies explore the diverse meanings and impacts of genetic diagnoses for patients enduring currently incurable ultimately fatal neurodegenerative diseases and for their family caregivers and clinicians Science in Society Matthew David, 2017-03-16 Science Technoscience has moved to centre stage in debates over change power and justice in twenty first century societies This text provides a general framework for understanding combining and applying the rich range of approaches that exist within sociology about science in particular the role and limitations of science in generating knowledge and the relationship between scientific knowledge and social progress Drawing on case studies from the past up until today s new genetics this is a clear even handed and comprehensive introduction to the field

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