

Ethics in Long-term Care: The Ethics of Everyday Living

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Economics And Ethics Of Long Term Care And Disability

**United States. Congress. Pepper
Commission**



Economics And Ethics Of Long Term Care And Disability:

The Economics and Ethics of Long-term Care and Disability Sean Sullivan, Marion Ein Lewin, 1988 **Long-term Care and Access to Health Care** United States. Congress. Pepper Commission, 1990 **Gerontological Nursing** Mary M. Burke, Susan Sherman, 1993 This book makes a compelling argument for an immediate and comprehensive shift in the nursing education curriculum from an acute care to a gerontological nursing care focus It is important reading for deans and faculty in schools of nursing administrators in long term care facilities and community health agencies and for practicing nurses caring for the elderly *Improving the Measurement of Late-Life Disability in Population Surveys* National Research Council, Division of Behavioral and Social Sciences and Education, Committee on Population, Committee on National Statistics, 2009-11-29 *Improving the Measurement of Late Life Disability in Population Surveys* summarizes a workshop organized to draw upon recent advances to improve the measurement of physical and cognitive disability in population surveys of the elderly population The book questions whether or not the measures of activities of daily living and instrumental activities of daily living used in many population surveys are sufficient as the primary survey based indicators of late life disability If not should they be refined or should they be supplemented by other measures of disability in surveys If yes in what ways should disability measures be changed or modified to produce population estimates of late life disability and to monitor trends The book also discusses what further research is needed to advance this effort *The Second Fifty Years* Institute of Medicine, Division of Health Promotion and Disease Prevention, 1992-02-01 Taking its title from the second 50 years of the human life span of about 100 years this book presents wide ranging and practical recommendations for health care providers policymakers and other sectors of society These recommendations range from setting new national policies to changing the way elderly patients are interviewed in the doctor s office and from what exercises older persons should do to how city planners should design our urban environment The bulk of this volume presents the latest research on 13 major health threats to the elderly covering prevalence impact on the older person s life cost and intervention In addition the authors provide a detailed analysis of why older people often do not receive the benefit of prevention programs **Aging and Disabilities** James Callahan, 2019-04-11 This book helps in building a bridge between the networks The topics of aging and disabilities are multifaceted and therefore may be approached in many ways The editors have chosen to focus on four systems and areas of common ground aging developmental disabilities independent living movement and rehabilitation to investigate the emerging intersection of the two fields *The Milbank Quarterly* , 1923 **Current Catalog** National Library of Medicine (U.S.), First multi year cumulation covers six years 1965 70 **Extending Life, Enhancing Life** Institute of Medicine, Division of Health Promotion and Disease Prevention, Committee on a National Research Agenda on Aging, 1991-02-01 Americans are living longer than ever before For many though these extra years have become a bitter gift marred by dementia disability and loss of independence *Extending Life Enhancing Life* sets the course toward practical

solutions to these problems by specifying 15 research priorities in five key areas of investigation Basic biomedicine To understand the fundamental processes of aging Clinical To intervene against common disabilities and maladies of older persons Behavioral and social To build on past successes with behavioral and social interventions Health services delivery To seek answers to the troubling issues of insufficient delivery of health care in the face of increasing health care costs Biomedical ethics To clarify underlying ethical guidelines about life and death decisions Most important the volume firmly establishes the connection between research and its beneficial results for the quality of life for older persons

Clearinghouse Review ,1976 *What Kind of Life?* Daniel Callahan,1995-02-01 A provocative call to rethink America's values in health care Genetic Testing and the Governance of Risk in the Contemporary Economy Lara Khoury,Adelle Blackett,Lukas Vanhonnaeker,2020-10-01 This book addresses emerging questions concerning who should bear responsibility for shouldering risk as well as the viability of existing and experimental governance mechanisms in connection with new technologies Scholars from 14 jurisdictions unite their efforts in this edited collection to provide a comparative analysis of how various legal systems are tackling the challenges produced by the legal aspects of genetic testing in insurance and employment They cover the diverse set of norms that surround this issue and share insights into relevant international regional and national incursions into the field By doing so the authors offer a basis for comparative reflection including on whether transnational standard setting might be useful or necessary for the legal aspects of genetic testing as they relate to the insurance and employment contexts The respective texts cover a broad range of topics including the prevalence of genetic testing in the contexts of insurance and employment and policy factors that might affect this prevalence such as the design of national health or social insurance systems of private insurance schemes or the availability of low cost direct to consumer genetic testing Further the field of genetics is gaining in importance at the international and regional levels Relevant concepts mainly genetic tests and genetic data information have been internationally defined and these definitions have influenced definitions adopted nationally International law also recognizes a special status for human genetic data The authors therefore also consider these definitions and the recognition of the special status of human genetic data within regional and national legal orders They investigate the range of norms that specifically address the use of genetic testing in employment and insurance encompassing international sources including human rights norms that may be binding or non binding as well national statutory regulatory and soft law mechanisms Accordingly some of the texts examine general frameworks relevant to genetic testing in each country including those that stem from general anti discrimination rules and norms protecting rights to autonomy self determination confidentiality and privacy In closing the authors provide an overview of the efficiency of their respective legal regimes approaches specific and generalist to genetic testing or disclosure of genetic information in the employment or insurance contexts including the effect of lack of legal guidance In this regard some of the authors highlight the need for transnational action in the field and make recommendation for future legal

developments Stakeholders and Ethics in Healthcare Lisa A. Martinelli, 2022-03-20 This ground breaking book uses organizational ethics and stakeholder theory to explore the ethical accountability of leadership in healthcare organizations to their distinct vulnerable stakeholder communities The book begins with a discussion of the moral agency of healthcare organizations and introduces stakeholder theory It then looks at key ethical challenges in relation to the confidentiality and privacy of healthcare data before turning to child health and interventions around issues such as obesity maltreatment and parenting The book ends by focusing on ethics of care in relation to older people and people with disabilities An insightful contribution to thinking about ethics for contemporary healthcare management and leadership this interdisciplinary book is of interest to readers with a background in healthcare business and management law bioethics and theology **The Long**

Term Care Crisis Carroll L. Estes, James H. Swan, 1993 This book examines the implementation of the prospective payment system PPS in the US for Medicare hospital reimbursement which started in 1983 The authors discuss the impact of the PPS on health care provision and conclude that rather than improving conditions for the elderly in their transition from hospital to community and decreasing escalating health costs the PPS has restructured the system with the result that the greater financial burden is placed on informal caregivers community and home health care agencies and the elderly themselves

Last Rights Barbara Logue, 1993-01-01 Several recent trends demographic social and economic are increasing the incidence and public support of deliberate death among the old and sick The number of elderly people is at an all time high Medical technology can prolong the lives of these men and women but it cannot keep them independent free of pain or competent to manage their own affairs Never before have so many been so sick for so long Changes in the size and structure of the American family make long term home care almost impossible while exorbitant health costs and scarce resources limit the provision of highquality care by paid workers Researcher Barbara J Logue refers to any deliberate behavior that causes death as death control She points out that death control like birth control involves very serious and deeply personal decisions choices that ought to be made on the basis of reliable information with care and compassion without coercion She goes on to assert that like birth control death control should be performed legally and with the help of the medical community Without access to assisted deaths many frail old people are left to endure wretched deaths Some die slowly of neglect or malnutrition in nursing homes Others take their own lives sometimes by violent means while they can still act independently because they are afraid that no one will help them if they wait until they are very sick Some die alone fearful of implicating friends or relatives in their back alley euthanasia While some die too soon because they are poor uneducated or hard to manage others are overtreated and their deaths delayed because doctors and hospitals want to increase their income or fear prosecution if they help a patient die Barbara Logue describes these and many other hazards inherent in our present long term care system noting that all too often the system causes suffering instead of alleviating it After assessing the alternatives she urges that we must make compassionate death control as available as birth control We must regulate and monitor it like any other medical

procedure taking steps to minimize the risks while maximizing the benefits National Library of Medicine Current Catalog
National Library of Medicine (U.S.), **Perspectives on Disability** Mark Nagler,1993 Contains 55 articles by noted
authorities focusing on opportunities challenges of the disabled in employment education society Includes legal ethical
medical psychological sexual issues **In re Martin (Martin v. Martin), 450 MICH 204 (1995)** ,1995 99699 99700

Handbook on Ethical Issues in Aging Tanya F. Johnson,1999-06-30 Johnson addresses ethical issues in aging in a variety
of contexts the social cultural environment physical health care mental health care social health care legal care and spiritual
care Because long term aging has created a new generation of older adults some new issues are emerging which need to be
addressed from an ethical perspective elder abuse physician assisted suicide dementia intergenerational equity guardianship
and living wills A wide range of experts including physicians philosophers lawyers social workers nurses sociologists public
health persons theologians historians and ethicists share their insights on the ethical issues and dilemmas older adults in
American society are facing or are likely to face over the life course Of interest to undergraduate and graduate faculty and
students in sociology social work and social services practitioners policymakers and academic and professional libraries

Disability in America Committee on a National Agenda for the Prevention of Disabilities,1991-01-15 This report focuses
on preventing potentially disabling conditions from developing into disabilities and on minimizing the effects of such
conditions on a person s productivity and quality of life It describes disability as a social and public health issue and not just a
physical condition The report begins with an executive summary an introduction which discusses prevention issues in general
and defines concepts and a list of 27 recommendations Subsequent chapters discuss 1 the magnitude and dimensions of
disability in the United States 2 a conceptual approach to disability prevention and use of the tools and principles of
epidemiology 3 major areas of disability developmental disabilities injury related disabilities chronic diseases and aging and
secondary conditions associated with primary disabling conditions 4 government and private sector programs concerned with
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contain a paper by Saad Z Nagi titled Disability Concepts Revisited Implications for Prevention a statement of one committee
member dissenting from this majority report of the Committee on a National Agenda for the Prevention of Disabilities a
response to the dissenting statement by committee members and committee biographies Approximately 375 references JDD

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