FACING DEATH



PALLIATIVE CARE

David Clark
Jane Seymour



Reflections On Palliative Care Sociological And Policy Perspectives

David Field

Reflections On Palliative Care Sociological And Policy Perspectives:

Reflections on Palliative Care David Clark, Jane Seymour, 1999 By drawing on a wide range of sources Clark and Seymour offer a set of reflections on the development of palliative care and its place within a wider social context This work considers ethical questions and palliative care s history too Dying: A Social Perspective on the End of Life Alex Broom, 2016-03-03 An inevitable and universal experience dying is experienced by individuals in different ways often related to the character of our relationships family structures gender identities cultural backgrounds and economic means Drawing on extensive qualitative fieldwork with patients carers and health professionals in Australia and the United Kingdom Dying A Social Perspective on the End of Life provides a critical examination of the different spheres of dying in social and cultural context Exploring complex issues such as the politics of assisted dying negotiating medical futility gender and dying the desire for redemption the moralities of the good fight and the lived experience of bodily disintegration this book links novel theoretical ideas within sociology to cutting edge empirical data collected in palliative and end of life care contexts A theoretically engaged understanding of the social mediation of the end of life Dying A Social Perspective on the End of Life also sheds light on the manner in which the end of life can be shaped by major economic cultural and socio cultural shifts including neo liberalism individualisation medicalisation professionalisation and detraditionalisation As such it will appeal to social science health and medical researchers interested in the end of life as well as those working in palliative and end of Palliative care for older people Lieve Van den Block, Gwenda Albers, Sandra Martins Pereira, Bregje life care settings Onwuteaka-Philipsen, Roeline Pasman, Luc Deliens, 2015-05-07 Current projections indicate that by 2050 the number of people aged over 80 years old will rise to 395 million and that by this date 25 30% of people over the age of 85 will show some degree of cognitive decline Palliative care for older people A public health perspective provides a comprehensive account of the current state of palliative care for older people worldwide and illustrates the range of concomitant issues that as the global population ages will ever more acutely shape the decisions of policy makers and care givers The book begins by outlining the range of policies towards palliative care for older people that are found worldwide It follows this by examining an array of socio cultural issues and palliative care initiatives from the care implications of health trajectories of older people to the spiritual requirements of palliative care patients and from the need to encourage compassion towards end of life care within communities to the development of care pathways for older people Palliative care for older people A public health perspective is a valuable resource for professionals and academics in a range of healthcare and public health fields to understand the current state of policy work from around the world The book also highlights the social cultural considerations that influence the difficult decisions that those involved in palliative care face not least patients themselves and offers examples of good practice and recommendations to inspire support and direct healthcare policy and decision making at organisational regional national and international levels Governing Death and Loss Stephen Conway, 2011 Political

economic social cultural and technological changes have led to profound transformations in the ways that death and loss are perceived and managed in contemporary society Over the last few decades the long term shift to chronic illness as a major causal factor has significantly increased the time scale of dying Most people die in institutions and care is typically medical Many communities and ordinary citizens now relinquish control and involvement to experts in the last stages of life However at global and local levels new institutional arrangements are emerging to govern the changing face of death and a new model is being developed to counter claims of the creeping medicalisation of death and dying Emphasising the international context of the issues involved this book illustrates the interlinking nature of society death and loss and gives examples of governance that promotes the empowerment participation and the increasing need for the involvement of ordinary people and communities in differing social and cultural contexts All chapters are written at an accessible level and will appeal to a wide readership Part 1 of the book provides a sociological understanding of the governance of death and loss in international and historical contexts and the implications for practice Part 2 provides examples of good practice drawing upon a sociological To Comfort Always David Clark, 2016-10-06 Palliative medicine was first recognised as a specialist field understanding in 1987 One hundred years earlier London based doctor William Munk published a treatise on easeful death that mapped out the principles of practical spiritual and medical support at the end of life In the intervening years a major process of development took place which led to innovative services new approaches to the study and relief of pain and other symptoms a growing interest in holistic care and a desire to gain more recognition for care at the end of life This book traces the history of palliative medicine from its nineteenth century origins to its modern practice around the world It takes in the changing meaning of euthanasia assesses the role of religious and philanthropic organisations in the creation of homes for the dying and explores how twentieth century doctors created a special focus on end of life care To Comfort Always traces the rise of clinical studies academic programmes and international collaborations to promote palliative care It examines the continuing need to support development with evidence and assesses the dilemmas of unequal access to services and pain relieving drugs as well as the periodic accusations of creeping medicalization within the field This is the first history of its kind and the breadth of information it encompasses makes it an essential resource for those interested in the long term achievements of palliative medicine as well as the challenges that remain Pain and Palliative Care in the Developing World and Marginalized Populations M.r. Rajagapol, 2004-03-31 Essential information for anyone involved in palliative care programs for deprived patients In this comprehensive resource leading healthcare professionals describe pioneering work on the front lines of pain and palliative care service planning and implementation for underserved populations Pain and Palliative Care in the Developing World and Marginalized Populations A Global Challenge explores the challenges and barriers preventing satisfactory pain management for patients who urgently need it This book provides you with true accounts of palliative care programs from around the world to help you meet the needs of disadvantaged clients This essential volume includes a

Foreword written by a world leader in palliative care Jan Stjernsward Former Chief of the Cancer and Palliative Care Program of the World Health Organization and currently International Director of the Oxford International Centre for Palliative Care in the United Kingdom Pain and Palliative Care in the Developing World and Marginalized Populations A Global Challenge addresses issues of vital importance for the global health care community such as Why do so many people in the developing world suffer excruciating pain for months and years when simple inexpensive medication could make them comfortable They get MRI scans why don't they have access to palliative care Why do some palliative care programs fail to reach the needy How could a palliative care delivery system be adapted to local needs Why are medical and nursing students not taught the fundamentals of pain management What direction should palliative care education take Could health care resources be channeled to deliver care in a more just and equitable manner This book chronicles the efforts of ambitious pain management care professionals to confront these questions working toward an end to needless preventable pain and suffering It examines their programs and acknowledges their successes and failures to date with commentaries by international experts This indispensable manual discusses palliative care programs in developing countries such as India Chile Argentina Saudi Arabia Thailand Hong Kong Malaysia and others Pain and Palliative Care in the Developing World and Marginalized Populations also offers an important look at pain management programs geared toward several specific underserved populations in both developing and developed countries including Native Americans and inmates in a New Zealand prison Illustrated with figures graphs and tables this book is essential for practitioners and officials in both palliative and public health care All proceeds from sales of this book will be used to support the growth of palliative care programs in Developing Holistic Care for Long-term Conditions Carl Margereson, Steve Trenoweth, 2009-10-16 Comprising four India sections this text introduces the policy and background to caring for people with chronic illness as well as the psychosocial impact of long term conditions Essential skills for practice are explored including holistic assessment symptom control and the promotion of effective partnership between client and carer in supporting coping recovery and end of life care There is an emphasis on maximising individual health potential and resilience with the role of nutrition exercise complementary therapy and spirituality considered The focus is on client centred care which addresses the whole person mind and body The extensive final section presents examples of key health issues where UK national guidelines have been published including Long term neurological conditions Diabetes Mental health Cancer Coronary heart disease Older people Abjectly Boundless Trudy Rudge, 2016-03-16 Within a variety of practice environments health professionals often experience feelings of disgust and repulsion towards the presence of an abject object Cadaverous sick disabled bodies troubled minds wounds vomit and so forth are all part of health and care work and threaten the clean and proper bodies of those who undertake it yet this unclean side of health work is rarely accounted for in academic literature This volume employs the work of Julia Kristeva through a range of case studies drawn from care and nursing settings around the world It brings together work from researchers and practitioners within the social and health sciences the caring professions and psychotherapy to expose and highlight the important impact of the concept of abjection which historically has been silenced in the health sciences The New Politics of Disablement Michael Oliver, Colin Barnes, 2012-05-08 Disability luminary Mike Oliver is joined by Colin Barnes in this agenda setting response to a capitalist society faced with globalisation financial instability and lower public expenditure A timely new edition which reignites the debate on the nature of disability and reasserts the political power of Nursing Older Adults Reed, Jan, Clarke, Charlotte, MacFarlane, Ann, 2011-06-01 the academic field of disability studies This timely textbook aims to provide adult nurses with the principles and practice insights needed to deliver exceptional care in partnership with older adults The Ethics of Palliative Care H. ten Have, David Clark, 2002 This volume provides a picture of palliative care ethics in the European context It should interest those involved in the delivery and management of palliative care services as well as students and researchers Living, Dying, Death, and Bereavement (Volume Two) David E. Balk, 2020-10-21 This two volume book offers extensive interviews with persons who have made significant contributions to thanatology the study of dying death loss and grief The book s in depth conversations provide compelling life stories of interest to clinicians researchers and educated lay persons and to specialists interested in oral history as a means of gaining rich understandings of persons lives Several disciplines that contribute to thanatology are represented in this book such as psychology religious studies art literature history social work nursing theology education psychiatry sociology philosophy and anthropology The book is unique no other text offers such a comprehensive insightful and personal review of work in the thanatology field The salience of thanatology is obvious when we consider several topics including the aging demographics of most countries the leading causes of death the devastation of COVID 19 the realities of how most persons die the growth both of hospice and of efforts within medicine to ensure that a good death becomes the norm of medical practice and increases in the number of countries and states permitting physician assisted suicide This second volume includes conversations with 16 thanatologists a rich extensive bibliography an index of names and subjects and a biographical sketch of the author The experts interviewed in this volume include Danai Papadatou Holly Prigerson Jack Jordan Illene Cupit Heather Servaty Seib Irwin Sandler Simon Shimshon Rubin Carla Sofka Harold Ivan Smith and Phyllis Kosminsky Exploring Disability Colin Barnes, Geof Mercer, 2018-05-21 The second edition of this widely used text has been carefully rewritten to ensure that it is up to date with cutting edge debates evidence and policy changes Since the book s initial publication there has been an expansion of interest in disability in the social sciences and disability has come to play an increasingly prominent role in political debates The new edition takes account of all these developments and also gives greater emphasis to global issues in order to reflect the increasing and intensifying interdependence of nation states in the twenty first century The authors examine amongst other issues the changing nature of the concept of disability key debates in the sociology of health and illness the politicisation of disability social policy and the cultural and media representation of

disability As well as providing an excellent overview of the literature in the area the book develops an understanding of disability that has implications for both sociology and society The second edition of Exploring Disability will be indispensable for students across the social sciences and in health and social care who really want to understand the issues facing disabled **Textbook of Palliative Medicine** Eduardo Bruera, Irene Higginson, Charles F von people and disabling societies Gunten, 2009-01-30 Textbook of Palliative Medicine provides an alternative truly international approach to this rapidly growing specialty This textbook fills a niche with its evidence based multi professional approach and global perspective ensured by the international team of editors and contributing authors In the absence of an international curriculum for the study of palliative medicine this textbook provides essential guidance for those both embarking upon a career in palliative medicine or already established in the field and the structure and content have been constructed very much with this in mind With an emphasis on providing a service anywhere in the world including the important issue of palliative care in the developing nations Textbook of Palliative Medicine offers a genuine alternative to the narrative approach of its competitors and is an ideal complement to them It is essential reading for all palliative care physicians in training and in practice as well as palliative care nurses and other health professionals in the palliative care team Researching Later Life and Ageing Miranda Leontowitsch, 2016-11-09 This collection on researching later life and ageing critically reflects upon the qualitative methods used in gaining knowledge of under researched groups of older people and sets out future research agendas

Palliative Care and Communication Anne-Mei The, 2002 Palliative Care and Communication seeks explanations for the sense of optimism found among patients with small cell bronchial carcinoma or lung cancer Over a period of five years the author used ethnographic research methods to monitor the illness processes of patients from the time of receiving bad news until their death from the perspectives of the various people involved the patient family and relatives the doctors and nursing staff Based on narrative descriptions interspersed with observations the author demonstrates why things happen the way they do in practice The book explores informal codes ambiguous messages the dilemma between professional detachment and personal involvement patterns of information and communication during the phases of the illness the latent realisation of approaching death and the ambivalence of patients with regard to knowing and not knowing Anne Mei The s engaging writing style and extensive use of narrative make this book accessible to a wide range of readers doctors nurses policy makers patients relatives researchers in the field of health care and social scientists It is suitable for use on various branches of medical and nursing training to illustrate the process of communication with patients during the terminal phase of a patient s illness The Oxford Handbook of Ethics at the End of Life Stuart J. Youngner, Robert M. Arnold, 2016-09-09 This handbook explores the topic of death and dying from the late twentieth to the early twenty first centuries with particular emphasis on the United States In this period technology has radically changed medical practices and the way we die as structures of power have been reshaped by the rights claims of African Americans women gays students and most relevant

here patients Respecting patients values has been recognized as the essential moral component of clinical decision making Technology s promise has been seen to have a dark side it prolongs the dying process For the first time in history human beings have the ability control the timing of death With this ability comes a responsibility that is awesome and inescapable How we understand and manage this responsibility is the theme of this volume The book comprises six sections Section I examines how the law has helped shape clinical practice emphasizing the roles of rights and patient autonomy Section II focuses on specific clinical issues including death and dying in children continuous sedation as a way to relieve suffering at the end of life and the problem of prognostication in patients who are thought to be dying Section III considers psychosocial and cultural issues Section IV discusses death and dying among various vulnerable populations such as the elderly and persons with disabilities Section V deals with physician assisted suicide and active euthanasia lethal injection Finally Section VI looks at hospice and palliative care as a way to address the psychosocial and ethical problems of death and dying

EBOOK: Death's Dominion: Ethics at the End of Life Simon Woods, 2006-12-16 I enjoyed reading this book very much It is very readable and well argued using real life cases and thought experiments as well The book provides the reader with a short history of and an overview of the most important issues in modern palliative care Various theoretical discussions are clearly set out such as the relationship between the hospice movement and modern palliative care between palliative care and health care in general between palliative sedation and euthanasia and the guestion whether euthanasia can be part of palliative care The author starts with exploring the existing debates and then develops his own arguments in a balanced and well structured way Medicine Health Care and Philosophy The text of this book is accessible the philosophical and ethical arguments are clearly articulated and relevant ethical principles are integrated into the critique of the issues making this a very useful book for nurses working in palliative as well as in general care Nursing Ethics It is crucially important for any student or researcher who is seriously consideringethical and policy matters at the end of life to embrace and tackle intellectually the issues that Woods raises in this book I would happily recommend it Journal of Medical Ethics What constitutes a good death Is it possible to arrange a good death Is killing compatible with caring This book looks at death and the issues and ethical dilemmas faced at the end of life It addresses the central issues in the field such as Withholding and withdrawing treatment Euthanasia and assisted suicide Terminal sedation The role of autonomy Palliative care Drawing on a philosophical framework the author explores end of life issues in order to reflect on the nature of the good death and how this may be achieved The book considers whether it is permissible or desirable to influence the quality of dying offering palliative sedation as a possible alternative to terminal sedation the argument is extended to examine why some forms of assisted dying can be shown to be compatible with the ideas of palliative care Consideration is also given to future developments such as life extension techniques and the ethical questions that that these techniques might raise As such the book follows in the ongoing philosophical tradition to critique and analyse current thought on the topic of death encouraging

self reflection in the reader and offering suggestions for practice in end of life care Death's Dominion is key reading for students and professionals involved in care of the dying as well as those with an interest in the philosophical issues surrounding end of life care EBOOK: Loss, Change and Bereavement in Palliative Care Pam Firth, Gill Luff, David Oliviere, 2004-12-16 For anyone seeking to develop their understanding of loss and change whether in a palliative care of general or social care setting this book contains much useful material which can be taken selectively or in its entirety Hospise Information Bulletin How do professionals meet the needs of bereaved people How do professionals undertake best practice with individuals groups families and communities What are the implications for employing research to influence practice This book provides a resource for working with a complex range of loss situations and includes chapters on childhood bereavement and individual and family responses to loss and change It contains the most up to date work in the field presented by experienced practitioners and researchers and is relevant not only for those working in specialist palliative care settings but for professionals in general health and social care sectors Strong links are maintained between research and good practice throughout the book These are reinforced by the coherent integration of international research material and the latest thinking about loss and bereavement Experts and clinicians draw upon their knowledge and practice whilst the essential perspective of the service user is central to this book Loss Change and Bereavement in Palliative Care provides essential reading for a range of professional health and social care disciplines practising at postgraduate or post registration qualification level It challenges readers at an advanced level on issues of loss change and bereavement Contributors Lesley Adshead Jenny Altschuler Peter Beresford Grace Christ Suzy Croft Pam Firth Shirley Firth Richard Harding Felicity Hearn Jennie Lester Gill Luff Linda Machin Jan McLaren David Oliviere Ann Quinn Phyllis Silverman Jean Walker Karen Wilman

<u>Researching Palliative Care</u> David Field,2001 There are many texts available on research methods but few that are related directly to palliative treatment This book fills the gap in the literature and provides a useful resource for students engaged in such activity

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Table of Contents Reflections On Palliative Care Sociological And Policy Perspectives

- 1. Understanding the eBook Reflections On Palliative Care Sociological And Policy Perspectives
 - The Rise of Digital Reading Reflections On Palliative Care Sociological And Policy Perspectives
 - Advantages of eBooks Over Traditional Books
- 2. Identifying Reflections On Palliative Care Sociological And Policy Perspectives
 - Exploring Different Genres
 - Considering Fiction vs. Non-Fiction
 - Determining Your Reading Goals
- 3. Choosing the Right eBook Platform
 - Popular eBook Platforms
 - Features to Look for in an Reflections On Palliative Care Sociological And Policy Perspectives
 - User-Friendly Interface
- 4. Exploring eBook Recommendations from Reflections On Palliative Care Sociological And Policy Perspectives
 - Personalized Recommendations
 - Reflections On Palliative Care Sociological And Policy Perspectives User Reviews and Ratings
 - Reflections On Palliative Care Sociological And Policy Perspectives and Bestseller Lists
- 5. Accessing Reflections On Palliative Care Sociological And Policy Perspectives Free and Paid eBooks
 - Reflections On Palliative Care Sociological And Policy Perspectives Public Domain eBooks
 - Reflections On Palliative Care Sociological And Policy Perspectives eBook Subscription Services
 - Reflections On Palliative Care Sociological And Policy Perspectives Budget-Friendly Options

Reflections On Palliative Care Sociological And Policy Perspectives

- 6. Navigating Reflections On Palliative Care Sociological And Policy Perspectives eBook Formats
 - o ePub, PDF, MOBI, and More
 - Reflections On Palliative Care Sociological And Policy Perspectives Compatibility with Devices
 - Reflections On Palliative Care Sociological And Policy Perspectives Enhanced eBook Features
- 7. Enhancing Your Reading Experience
 - Adjustable Fonts and Text Sizes of Reflections On Palliative Care Sociological And Policy Perspectives
 - Highlighting and Note-Taking Reflections On Palliative Care Sociological And Policy Perspectives
 - Interactive Elements Reflections On Palliative Care Sociological And Policy Perspectives
- 8. Staying Engaged with Reflections On Palliative Care Sociological And Policy Perspectives
 - Joining Online Reading Communities
 - o Participating in Virtual Book Clubs
 - Following Authors and Publishers Reflections On Palliative Care Sociological And Policy Perspectives
- 9. Balancing eBooks and Physical Books Reflections On Palliative Care Sociological And Policy Perspectives
 - Benefits of a Digital Library
 - o Creating a Diverse Reading Collection Reflections On Palliative Care Sociological And Policy Perspectives
- 10. Overcoming Reading Challenges
 - Dealing with Digital Eye Strain
 - Minimizing Distractions
 - Managing Screen Time
- 11. Cultivating a Reading Routine Reflections On Palliative Care Sociological And Policy Perspectives
 - Setting Reading Goals Reflections On Palliative Care Sociological And Policy Perspectives
 - Carving Out Dedicated Reading Time
- 12. Sourcing Reliable Information of Reflections On Palliative Care Sociological And Policy Perspectives
 - Fact-Checking eBook Content of Reflections On Palliative Care Sociological And Policy Perspectives
 - Distinguishing Credible Sources
- 13. Promoting Lifelong Learning
 - Utilizing eBooks for Skill Development
 - Exploring Educational eBooks
- 14. Embracing eBook Trends
 - Integration of Multimedia Elements

• Interactive and Gamified eBooks

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