## 7TH PUBLIC HEALTH PALLIATIVE CARE INTERNATIONAL CONFERENCE

# Democratizing caring, dying and grieving participation, action, understanding and evaluation 20-23 SEPTEMBER 2022 - BRUGES, BELGIUM

## **Programme Poster exhibition**

Conference Venue: Bruges Meeting & Convention Centre (BMCC), Beursplein 1, 8000 Bruges

Poster exhibition: Foyer, level 3

#### **Group 1: Poster exhibition on 20-21 September 2022**

#### **ACP & End of Life communication**

140	Tessa	Bergman	Information meetings on end-of-life care organized by GPs and other organizations; do they differ in audience and results?
101	Marcelo	Caballero	Idiolectic Conversation and Symptom Management in Palliative Care Patients
211	Ludovica	De Panfilis	Development and evaluation of an in-hospital service to inform people about end-of-life rights: the SCEGLIERE service
36	Nicholas	Jennings	Medical end-of-life decision-making in a small resource-poor Caribbean country: a mortality follow-back study of home deaths
147	Maria Fidelis	Manalo	Talking about death to advanced cancer patients: not if, but how
171	Fanny	Monnet	Developing an advance care planning website for dementia: Integrating a user-centered approach and patient and public involvement
235	Marta	Perin	Advance Care Planning in Italian palliative care networks: a grounded theory study
173	Galmiche	Perrine	Mirror methodology: using marketing codes for better talking about end-of-life

195	Ramona	Schenell	To make and execute decisions throughout life- a model to facilitate self-determination in residential care.
184	Steffen	Eychmüller	Sense of security in palliative homecare: A qualitative descriptive study on the experiences of patients, family caregivers and outpatient healthcare professionals.
120	Willemijn	Tros	General practitioners' evaluations of optimal timing to initiate ACP for patients with cancer, organ failure or multimorbidity
218	Isabel	Vandenbogaerde	Support from health care professionals in empowering family carers to discuss Advance Care Planning: A population-based survey

# Community

293	Gabriela	Hidalgo	Implementation of community-based palliative care in Primary Health Care in integration with Specialized Outpatient Care
178	Briony	Hudson	Exploring the support needs of the British Muslim Community with palliative care needs – a collaborative qualitative project (TBC)
152	Helen	MacGregor	Supporting compassionate communities in dying, death & bereavement by providing space for orgs to gather, discuss & learn"
259	Steven	Vanderstichelen	Charters are not enough: supporting volunteering at the volunteer level
117	Julia	Verne	A workplace compassionate community approach to support bereaved staff in the two national public health organizations in England

#### COVID

97	Shirley H.	Bush	Experiencing and making sense of illness in an inpatient specialist palliative care unit during a pandemic: an interview study
272	Barbara	Gale	Hospice care - where do we go now?
136	Kathy	Kortes-Miller	Not Just A Visitor; Caregiving experiences in Long Term Care during COVID-19
77	Matteo	Moroni	Elderly Covid -19 patients in nursing home cared by young doctors together with Palliative Care teams. A model beyond pandemic?
208	Stina	Nyblom	Experiences among Swedish patients in specialized palliative home care and their significant others during the Covid-19 pandemic
25	Vilma	Oliveros	Are we now more empathetic, compassionate, and understanding towards difference?

298	Gabriela	Rezende	Impacts of the COVID-19 pandemic on palliative care: a qualitative and multicenter study
90	Ahmed	Sharaf	Opportunities and challenges to family carers end of life care research in the COVID pandemic: A researcher's reflections
82	Katherine	Sleeman	Variation in palliative care service response and association with Covid-19 mortality rates: an ecological study in UK regions
119	Julia	Verne	Learning lessons from COVID-19 experience for palliative care for people with dementia in England
267	Masha	Zee	The relation between distress and feeling appreciated when providing end-of-life care during the COVID-19 pandemic (CO-LIVE).

**Literacy & Education** 

88	Ruth Si Man	Wong	A Systematic Scoping Review on Moral Distress in Undergraduate Medical Students

Various Topics

174	Agathe	CANT-DIOT	How is end-of-life care organized around the world?
93	Marius	Čiurlionis	Home-based palliative care and re-hospitalizations for palliative care patients
49	Ascensión	Doñate-Martínez	Needs for timely palliative care in older patients with non-malignant diseases: a multicenter qualitative study
261	Rodeyns	Julie	Fostering empathy and compassion through art observation
59	Anne	van Driessche	Recruitment and retention in palliative care trials: challenges and strategies
72	Sarah	Yardley	Getting palliative prescription medications right at home, in hospital & hospice: Activity Theory analysis
74	Sarah	Yardley	Reclaiming relationships as a legitimate tool for systems change: A Change Laboratory approach to palliative care
71	Sarah	Yardley	Patient safety in palliative care: improving system design for safe, responsive medication management

46	Tineke	De Keyser	Empowerment of carer in outpatient home care through small chunks of accessible intel"
170	Alfieri	Sara	Multidisciplinary approach implementation in a European comprehensive cancer center

# **Group 2: Poster exhibition on 22-23 September 2022**

#### Cancer

57	Torbjørn	Paulsen	Early referral to a palliative team improves end- of- life care among gynecological cancer patients
192	Tara	Joyce McDonnell	A Controlled Clinical Trial of the effects of Meditation on patient's Perceptions of their Quality of Life in an acute hospital
297	Marysia	M R do Prado De Carlo	Total Pain in The Context of Palliative Care and its Assessment in People with Advanced Lung Cancer
143	Lisa	Shirt	"Patient and Caregiver-reported Acceptability of an "Automatic" Supportive and Palliative Care Referral for Advanced Lung Cancer Patients

## **Equity and Access**

160	Clare	Howie	Improving equity of palliative care provision within advanced heart failure via a realist synthesis: Palliative Heart Synthesis
285	Jamilla	Hussain	Equity, diversity and inclusion in palliative and end of life care practice and research: personal reflections
213	Terhi Kukkonen	Kukkonen	Developing the competence of palliative care –project, multidisciplinary cooperation in long distance region

#### **Euthanasia & MAID**

156	Charlotte	Boven	Relatives of cancer patients' experiences regarding the interaction with healthcare providers during a euthanasia process
157	Charlotte	Boven	Healthcare providers' experiences regarding the interaction with family during a euthanasia process: A qualitative study
288	Lavoie	David	Creating narratives to support experiences of medical assistance in dying in Canada: qualitative investigation of nurses' views

193	Gordon	Gubitz	The Canadian Medical Assistance in Dying (MAiD) Curriculum Development Project
132	Kathy	Kortes-Miller	Untold Stories; Accompanying someone who used medical assistance in dying at the end of life
138	Rikmenspoel	Marjolein	Information and support regarding euthanasia: a qualitative study on the needs of nursing home residents in the Netherlands

Family caregiving

83	Kristen	Abbott-Anderson	Conversations and care at end-of-life for persons with dementia: Family care partners share their perspectives
216	Dr Kabindra	Bhagabati	Concerns of the patients' relatives about Breaking Bad News to patients – An experience in a tertiary cancer center in NE India
229	Anneli	Ozanne	Experiences of parenthood when a parent has been diagnosed with ALS
202	F H D SHEHAN	SILVA	Cultural adaptation and validation of Sinhala version of Modified Caregiver Strain Index

#### **Health Promotion**

43	Madeleine	Juhrmann	Exploring a Public Health Palliative Care Approach to Emergency Medical Services
205	Tracey	McConnell	Loneliness at end of life: A public health approach to promoting the health and wellbeing of terminally ill patients and their carers
248	Katarzyna	Patynowska	Promoting the health and wellbeing of family caregivers providing end-of-life care – the role of community Healthcare Assistant
95	Marianne	Tinkler	A framework for developing co-designed integrated service model of palliative care for people living with severe mental illness.
52	Vincent	Van Goethem	Development of a self-management eHealth program to support and empower people with advanced cancer and their family caregivers

## **Literacy & Education**

144	Ashley	Chastain	Development and Testing of the Palliative Care-Related Knowledge, Attitudes & Confidence in Home Health Care
			Questionnaires

37	Zara, Linda	Höök, Frode	Education for palliative representatives increases the quality of palliative care
258	Clément	Meier	Perceptions and knowledge of end-of-life medical situations among older adults in Switzerland
155	Danny	Vereecke	Knowing, being proficient, and being in palliative care starts in education, but does it?

Loss & grief

27	Мо	Blishen	The Wind Telephone - How a disconnected telephone can connect to grief
159	Charlotte	Boven	Walking the walk in connection: Experiences of healthcare providers with bereavement care in oncology

#### **Paediatric Palliative Care**

126	Soham	Bandyopadhyay	A 12-month international, observational study of palliative care in childhood cancer patients
227	Leigh	Donovan	Beyond bricks and mortar: enabling equity of access to pediatric hospice care for Queensland children and families

#### **Poster Lightning Sessions**

## Wednesday 21 September 2022

During Lunch break: level 4, catering hall

49	Ascensión	Doñate-Martínez	Needs for timely palliative care in older patients with non-malignant diseases: a multicenter qualitative study
132	Kathy	Kortes-Miller	Untold Stories; Accompanying someone who used medical assistance in dying at the end of life
297	Marysia	M R do Prado De Carlo	Total Pain in The Context of Palliative Care and its Assessment in People with Advanced Lung Cancer
77	Matteo		Elderly Covid -19 patients in nursing home cared by young doctors together with Palliative Care teams. A model beyond pandemic?

## Thursday 22 September 2022

During Lunch: level 4, catering hall

227	Leigh	Donovan	Beyond bricks and mortar: enabling equity of access to paediatric hospice care for Queensland children and families
193	Gordon	Gubitz	The Canadian Medical Assistance in Dying (MAiD) Curriculum Development Project
25	Vilma	Oliveros	Are we now more empathetic, compassionate, and understanding towards difference?
202	F H D SHEHAN	SILVA	Cultural adaptation and validation of Sinhala version of Modified Caregiver Strain Index
59	Anne	van Driessche	Recruitment and retention in palliative care trials: challenges and strategies

## Friday 23 September 2022

During Lunch: level 4, catering hall

144	Ashley	Chastain	Development and Testing of the Palliative Care-Related Knowledge, Attitudes & Confidence in Home Health Care Questionnaires
285	Jamilla	Hussain	Equity, diversity and inclusion in palliative and end of life care practice and research: personal reflections
43	Madeleine	Juhrmann	Exploring a Public Health Palliative Care Approach to Emergency Medical Services
52	Vincent	Van Goethem	Development of a self-management eHealth program to support and empower people with advanced cancer and their family caregivers
72	Sarah	Yardley	Getting palliative prescription medications right at home, in hospital & hospice: Activity Theory analysis