

# Meaningful contextual communication in cancer care

<b>Submission date</b> 07/09/2021	<b>Recruitment status</b> Recruiting	<input checked="" type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
<b>Registration date</b> 15/09/2021	<b>Overall study status</b> Ongoing	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
<b>Last Edited</b> 15/09/2021	<b>Condition category</b> Cancer	<input type="checkbox"/> Individual participant data <input type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims

Contextual care is the continuous, ongoing process of identifying individual patient circumstances (their context) and, if necessary, modifying the plan of care to accommodate those circumstances. It is about essential patient needs that need to be addressed. Being confronted with the reality of having advanced cancer may give rise to life questions, and subsequent struggles such as fear, anger, loss of independence, changing self-image, roles and relationships and failure to find meaning are frequently reported amongst patients. This project will focus on increasing meaningful contextual communication between health care professionals (HCPs) and patients with advanced cancer, considering differences across Europe in both clinical and ethical aspects. The challenge addressed by this project is to develop an intervention founded on a European-wide set of central educational design principles, in order to increase the competence of specialised palliative care HCPs in meaningful contextual communication (what does it mean for this patient at this moment to live with his or her illness?), and as a result to increase QoL of patients with advanced cancer throughout Europe.

### Who can participate?

Adult (over 18 years of age) patients with advanced cancer and their family caregivers under treatment of included multidisciplinary teams in palliative care.

### What does the study involve?

An observational pre-study in which participants will randomly be invited for an interview or focus group.

A communication training for HCPs working in multidisciplinary teams for palliative care (5 steps in 5 months).

HCPs, patients, and their family caregivers will be interviewed before and after the communication training to assess impact of training.

### What are the possible benefits and risks of participating?

Benefits: increased meaningful contextual communication, resulting in increased job satisfaction for HCPs, increased QoL for patients and family caregivers, increased cost-effectiveness.

Risks: None

Where is the study run from?

Radboud University Nijmegen Medical Centre (Netherlands)

When is the study starting and how long is it expected to run for?

September 2021 to June 2028

Who is funding the study?

EU (HORIZON-21)

Who is the main contact?

Dr Anne Wichmann (anne.wichmann@radboudumc.nl)

## Contact information

### Type(s)

Scientific

### Contact name

Dr Anne Wichmann

### ORCID ID

<http://orcid.org/0000-0003-0085-9782>

### Contact details

Geert Grooteplein 10

Nijmegen

Netherlands

6500 HB

+31 628744811

anne.wichmann@radboudumc.nl

## Additional identifiers

### EudraCT/CTIS number

Nil known

### IRAS number

### ClinicalTrials.gov number

Nil known

### Secondary identifying numbers

Nil known

## Study information

### Scientific Title

A mixed-method implementation study of a design-based educational intervention to enhance meaningful contextual communication in care planning with advanced cancer patients and their families across Europe

**Acronym**

onCOntext

**Study objectives**

The introduction of the onCOntext intervention will improve meaningful contextual communication.

**Ethics approval required**

Old ethics approval format

**Ethics approval(s)**

Not provided at time of registration

**Study design**

Interventional uncontrolled pre-post study

**Primary study design**

Interventional

**Secondary study design**

Non randomised study

**Study setting(s)**

Other

**Study type(s)**

Quality of life

**Participant information sheet**

No participant information sheet available

**Health condition(s) or problem(s) studied**

Increasing meaningful contextual communication between multidisciplinary palliative care teams, patients with advanced cancer and their family caregivers.

**Interventions**

The observational cross-sectional study in year 1 comprises of a comprehensive European-wide (The Netherlands, Belgium, Scotland, Switzerland, Poland, Turkey, Spain, Italy) qualitative current practice study amongst HCPs (N=80), patients with advanced cancer (N=80) and their family caregivers (N=80) on i) the perspectives on current practice regarding meaningful contextual communication in palliative care, its integration in care planning and serious late and long-term side effects; ii) how these stakeholders from different European cultures understand meaningful contextual communication in palliative care; iii) how HCPs define the concept 'meaningful contextual communication' and which 'language' they use for it; iv) what makes team-based learning in European palliative care work? For point iv, additionally two focus groups per country (total N=16) will be conducted with 10-15 stakeholders (total N≈200). It will take approximately a year.

The interventional pre- post effect measurement study in year 3 consists of i) prospective interviews conducted amongst HCPs (N≈150), patients (N≈150) and family caregivers (N≈150); ii)

4C coding of audio-recorded consultations (N≈320); iii) a structured retrospective after-death questionnaire study amongst HCP most involved in care (N≈800) and a family caregiver closely involved (N≈800), also about the patient\*.

\* all participating teams retrospectively report all deaths of patients over a three month period, about whom the HCP and family caregiver will also answer questions. Using a three-month period limits recall bias and has been tested previously.

## **Intervention Type**

Behavioural

## **Primary outcome measure**

1. Observational pre-study: current practice, perspectives, cultures measured amongst HCPs, patients, family caregivers and relevant experts using interviews (N=80 per stakeholder group)

Main interventional study:

2. Experiences of HCPs, patients and their family caregivers with current care measured using interviews at baseline (T0) and 3-6 months post-intervention (T1)

3. Contextual communication measured in audio-taped consultations using 4C coding, at baseline (T0) and 3-6 months post-intervention (T1)

## **Secondary outcome measures**

1. HCP competence measured using questionnaire (SCCS), at baseline (T0) and 3-6 months post-intervention (T1)

2. HCP job satisfaction measured using questionnaire (NEXT study), at baseline (T0) and 3-6 months post-intervention (T1)

3. Patient QoL measured using questionnaires (EQ5D-5L, FACIT-Sp, ICECAP-SCM) at baseline (T0) and 3-6 months post-intervention (T1)

4. Family caregiver burden measured using questionnaire (CRA) at baseline (T0) and 3-6 months post-intervention (T1)

5. Costs measured using consultation length and questionnaire (RUD) at baseline (T0) and 3-6 months post-intervention (T1)

## **Overall study start date**

01/09/2021

## **Completion date**

01/06/2028

# **Eligibility**

## **Key inclusion criteria**

Adult (>18 years of age) patients with advanced cancer and their family caregivers under treatment of included multidisciplinary teams in palliative care

## **Participant type(s)**

Patient

## **Age group**

Adult

**Lower age limit**

18 Years

**Sex**

Both

**Target number of participants**

800

**Key exclusion criteria**

1. Patients with intellectual disabilities
2. Children

**Date of first enrolment**

01/06/2025

**Date of final enrolment**

31/12/2026

**Locations****Countries of recruitment**

Belgium

Italy

Netherlands

Poland

Scotland

Spain

Switzerland

Türkiye

United Kingdom

**Study participating centre**

Radboud university medical center

Netherlands

6500 HB

**Study participating centre**

**UEDIN**  
Edinburgh  
United Kingdom  
EH8 9AG

**Study participating centre**  
**UANTWERPEN**  
Antwerp  
Belgium  
2610

**Study participating centre**  
**UZ UNIVERSITY OF ZIELONA GORA**  
Poland  
65-046

**Study participating centre**  
**PALLIATIVE HEALTH SERVICES ASSOCIATION**  
Türkiye  
06520

**Study participating centre**  
**UNIBO**  
Italy  
40126

**Study participating centre**  
**CENTRAL UNIVERSITY OF CATALONIA**  
Spain  
08500 Vic

**Study participating centre**  
**UNIVERSITAET BERN**  
Switzerland  
3012 Bern

**Sponsor information**

**Organisation**

Radboud University Nijmegen Medical Centre

**Sponsor details**

Geert Grooteplein 10

Nijmegen

Netherlands

6500 HB

+31 (0)24 361 1111

digitalezorg@radboudumc.nl

**Sponsor type**

Hospital/treatment centre

**Website**

<https://www.radboudumc.nl/EN/Pages/default.aspx>

**ROR**

<https://ror.org/05wg1m734>

**Funder(s)****Funder type**

Government

**Funder Name**

European Commission

**Alternative Name(s)**

European Union, Comisión Europea, Europäische Kommission, EU-Kommissionen, Euroopa Komisjoni, Ευρωπαϊκή Επιτροπή, Европейската комисия, Evropské komise, Commission européenne, Choimisiúin Eorpaigh, Europskoj komisiji, Commissione europea, La Commissione europea, Eiropas Komisiju, Europos Komisijos, Európai Bizottságról, Europese Commissie, Komisja Europejska, Comissão Europeia, Comisia Europeană, Európskej komisii, Evropski komisiji, Euroopan komission, Europeiska kommissionen, EC, EU

**Funding Body Type**

Government organisation

**Funding Body Subtype**

National government

**Location**

# Results and Publications

## Publication and dissemination plan

Planned publication in a high-impact peer-reviewed journal.

## Intention to publish date

01/06/2029

## Individual participant data (IPD) sharing plan

The current data sharing plans for this study are unknown and will be available at a later date

## IPD sharing plan summary

Data sharing statement to be made available at a later date