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DELIVERABLE

D6.9 – Data Management Handling Plan – Final version

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Executive Summary

This deliverable is the final version of the Data Management Handling Plan (DMP) for the HosmartAI project, which is funded by the European Union's H2020 Programme under Grant Agreement No. 101016834. The DMP aims to include among other (open) data sources, (open) source code, scientific publications, project deliverables and more. The DMP is used internally by the consortium partners for the effective Data Management Handling in the context of Task 6.5 (M2-M41, Leader: EXYS). Moreover, the DMP is a mandatory requirement for the HosmartAI project to participate in the Horizon 2020 Open Access to scientific peer-reviewed publications and research data. The first version of the DMP (deliverable D6.7) was formulated and delivered in M6 of the project implementation. The second version of the DMP was submitted in M22, while the final version of the DMP (current document) is due in M41.

The HosmartAI project participates in the Pilot on Horizon 2020 Open Research Data. The use of a Data Management Plan is required for all EU H2020 funded projects.

The purpose of the Data Management Plan (DMP) is to provide an analysis of the main elements of the data management policy that have been used by the Consortium regarding the project research data.

It also reflects the current state of the Consortium agreements on data management and must be consistent with the exploitation and IPR requirements. Research data linked to exploitable results will not be put into the open domain if they compromise its commercialisation prospects or have inadequate protection, which is an H2020 obligation. The rest of the research data will be deposited in an open access repository.

The DMP covers the full data management life cycle for the data to be collected, processed and generated by the HosmartAI project. Towards handling research data management, within Publications and Research Data in Horizon 2020, detailing

1. how research data will be handled during & after the project;
2. what data will be collected, processed or generated;
3. what methodology & standards will be applied;
4. whether data will be shared /made open access/ how data will be curated and preserved.

The HosmartAI's DMP is based on the **FAIR principles**¹ (Findable, Accessible, Interoperable, Reusable) [REF-05] and on the Guidelines on Implementation of Open Access to Scientific Publications and Research Data in projects supported by the European Research Council (ERC) under Horizon 2020².

The DMP is finalised and delivered in its definitive form by the present document towards the end of the project. EXYS is leading the activity while all data provision partners are investing effort in safeguarding their proper data management.

¹ The FAIR Guiding Principles for scientific data management and stewardship, March 2016
<https://www.nature.com/articles/sdata201618>

² https://ec.europa.eu/research/participants/data/ref/h2020/other/hi/oa-pilot/h2020-hi-erc-oa-guide_en.pdf

The first version of the DMP included an overview of the datasets to be produced by the project, and the specific conditions that are attached to them. The second version of the HosmartAI DMP delved into the topics related to the research datasets management, continued the work of filling in data in the tables, and collected information about the allocation of resources, which was not included in the previous versions of the DMP. Guidelines on Research Data Alliance (RDA) FAIR Data Maturity Model implementation was also added in the second release, and was refined and enhanced in the final version by providing evidence of compliance with the RDA maturity model FAIR data from HosmartAI's FHIR server. Also in the activities performed to reach this final version, a collaboration with WP2 was carried on to enlighten data security and privacy aspects deployed to pilot datasets. Finally, the DMP was harmonized with the work performed in dissemination activities (Open Calls).

Data Management Strategy: to map all relevant data collections and to establish the data management needs, in the first version of the DMP every Pilot leader was requested to complete a 'Data management survey', which was reported in the Appendix B for that deliverable.

In general, the DMP first evaluated the legal frameworks and the requirements for all pilots, then examined whether there are currently available data to which open access can be granted, always respecting the security and privacy requirements imposed. With regards to the dissemination of the scientific results, the consortium established and promoted open access publications, and partners were encouraged to publish open access articles, to enable researchers to build upon previous research results, foster collaborations, avoid duplication of efforts, and accelerate innovation.

Project members were offered the option of publishing in journals contained/registered in the ROAR³ and/or other repositories (OpenDOAR⁴, OpenAIRE⁵ and Zenodo⁶). Authors' copyright agreements will determine whether scientific publications resulting from the project will adopt the gold or the green model.

In addition to the task leader EXYS, contributors to this deliverable are **TGLV, PhE, UKCM, IRCCS, SERMAS, FIBHULP, CHUL, INTRAS, AHEPA, VUB, UM and PHILIPS.**

The structure of the DMP follows the H2020 Data Management Plan template.

³ ROAR: Research Open Access Repository, <http://roar.eprints.org>

⁴ OpenDOAR: Directory for Open Access Repositories: <https://v2.sherpa.ac.uk/opensoar/>

⁵ <https://www.openaire.eu/>

⁶ <https://zenodo.org>

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0.2	2024-03-11	Carlos Luis Parra Calderon	Evidence of compliance with the RDA maturity model FAIR data from HosmartAI's FHIR server – added to Section 3.4.4.
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0.5	2024-05-27	Luca Gilardi	Final version, before QA and approval
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Definitions, Acronyms and Abbreviations

Acronym/ Abbreviation	Title
ASR	Automatic Speech Recognition
Cath lab	Catheterization laboratory
CCDS	Consensus Coding Sequence
CCTA	Coronary Computed Tomography Angiography
COBIT	Control Objectives for Information and related Technology
CPOE	Computerized Physician Order Entry
CRF	Case Report Form
CSV	Comma-Separated Values
CT	Computed Tomography
DICOM	Digital Imaging and COmmunications in Medicine
DMP	Data Management Plan
DPO	Data Protection Officer
DTA	Data Transfer Agreement
EHR	Electronic Health Record
ERC	European Research Council
FAIR	Findable, Accessible, Interoperable, Reusable
FFRCT	Fractional Flow Reserve derived from CT
FHIR	Fast Healthcare Interoperability Resources ⁷
GDPR	General Data Protection Regulation ⁸
HL7	Health Level Seven
iFR	instantaneous wave-Free ratio Resting index
IMU	Inertial Measurement Unit
IVUS	Intravascular Ultrasound
JSON	JavaScript Object Notation
KPI	Key Performance Indicator
LPD	Swiss Data Protection Law
MIMO	Medical Informatics and digital health Multilingual Ontology
N/A	Data not available by the time of writing this deliverable
OCT	Optical Coherence Tomography
PAM	Privilege Access Management
PGHD	Patient Generated Health Data
PREM	Patient Reported Experience Measure
PROM	Patient Reported Outcome Measure
RDA	Research Data Alliance
RFR	Coronary Physiology Resting Full-Cycle Ratio
SOP	Standard Operating Procedure

⁷ HL7 FHIR (Fast Healthcare Interoperability Resources), <http://hl7.org/fhir/index.html>

⁸ <https://gdpr-info.eu>

Acronym/ Abbreviation	Title
SPSS	Statistical Package for the Social Sciences ⁹
SSH	Secure Shell
SSL / TLS	Secure Sockets Layer / Transport Layer Security
SUS	System Usability Scale
TAM	Technology Acceptance Model
TBD	To be defined/done
TTS	Text-To-Speech
UEQ	User Experience Questionnaire
VPN	Virtual Private Network

Term	Definition
Accessible	Data is Accessible in that it can be always obtained by machines and humans upon appropriate authorization and through a well-defined protocol.
Cohort	In statistics, marketing and demography, a cohort is a group of subjects who share a defining characteristic (typically subjects who experienced a common event in a selected time period, such as birth or graduation).
Findable	Any Data Object should be uniquely and persistently identifiable
Interoperable	The ability of data or tools from non-cooperating resources to integrate or work together with minimal effort. Data Objects can be Interoperable only if: (Meta) data is machine-actionable, (Meta) data formats utilize shared vocabularies and/or ontologies, (Meta) data within the Data Object should thus be both syntactically parseable and semantically machine-accessible.
Pseudoanonymisation	The processing of personal data in such a way that the data can no longer be attributed to a specific data subject without the use of additional information, as long as such additional information is kept separately and subject to technical and organizational measures to ensure non-attribution to an identified or identifiable individual ¹⁰
Re-usable	For Data Objects to be Re-usable they should be sufficiently well-described and rich that it can be automatically (or with minimal human effort) linked or integrated, like-with-like, with other data sources. Published Data Objects should refer to their sources with rich enough metadata and provenance to enable proper citation.

⁹ <https://www.ibm.com/analytics/spss-statistics-software>

¹⁰ GDPR Article 4(3b): <https://www.privacy-regulation.eu/en/article-4-definitions-GDPR.htm>

Term	Definition
Schemaless database	A type of database where each item is saved in its own document with a partial schema, leaving the raw information untouched.

1 Introduction

1.1 Project Information



The HosmartAI vision is a strong, efficient, sustainable and resilient European **Healthcare system** benefiting from the capacities to generate impact of the technology European Stakeholders (SMEs, Research centres, Digital Hubs and Universities).



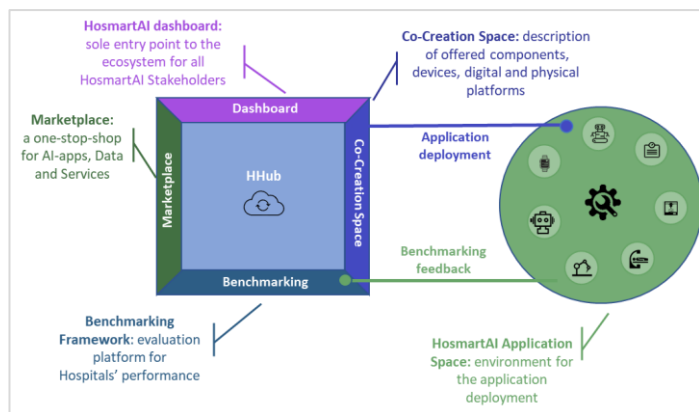
The HosmartAI mission is to guarantee the **integration** of Digital and Robot technologies in new Healthcare environments and the possibility to analyse their benefits by providing an **environment** where digital health care tool providers will be able to design and develop AI solutions as well as a space for the instantiation and deployment of a AI solutions.

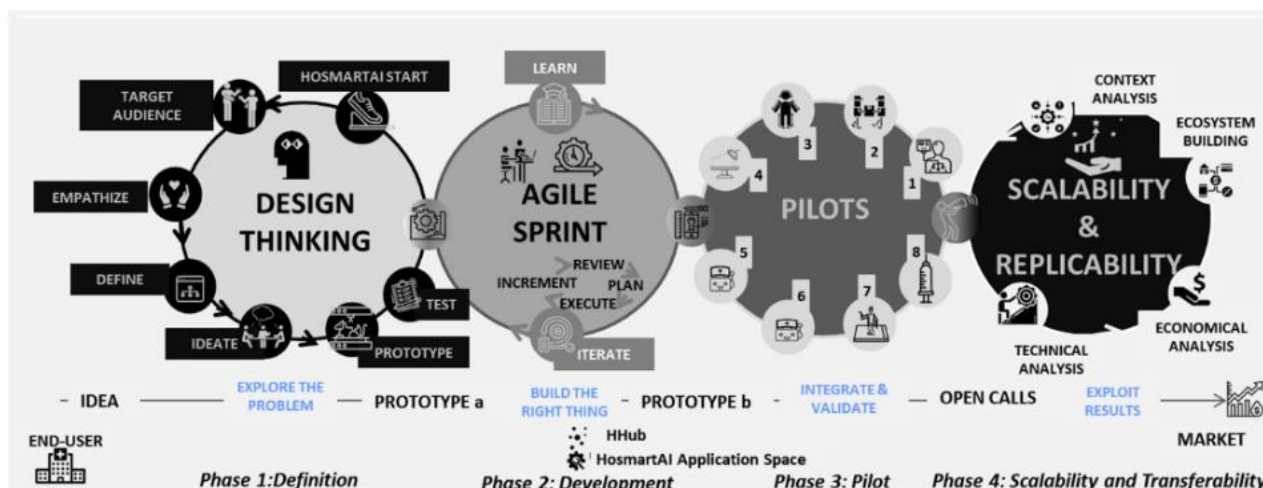
HosmartAI will create a common open Integration **Platform** with the necessary tools to facilitate and measure the benefits of integrating digital technologies (robotics and AI) in the healthcare system.

A central **hub** will offer multifaceted lasting functionalities (Marketplace, Co-creation space, Benchmarking) to healthcare stakeholders, combined

with a collection of methods, tools and solutions to integrate and deploy AI-enabled solutions. The **Benchmarking** tool will promote the adoption in new settings, while enabling a meeting place for technology providers and end-users.

Eight Large-Scale Pilots will implement and evaluate improvements in medical diagnosis, surgical interventions, prevention and treatment of diseases, and support for rehabilitation and long-term care in several Hospital and care settings. The project will target different **medical** aspects or manifestations such as Cancer (Pilot #1, #2 and #8); Gastrointestinal (GI) disorders (Pilot #1); Cardiovascular diseases (Pilot #1, #4, #5 and #7); Thoracic Disorders (Pilot #5); Neurological diseases (Pilot #3); Elderly Care and Neuropsychological Rehabilitation (Pilot #6); Fetal Growth Restriction (FGR) and Prematurity (Pilot #1).





To ensure a user-centred approach, harmonization in the process (e.g. regarding ethical aspects, standardization, and robustness both from a technical and social and healthcare perspective), the **living lab** methodology will be employed. HosmartAI will identify the appropriate instruments (KPI) that measure efficiency without undermining access or quality of care. Liaison and co-operation activities with relevant stakeholders and **open calls** will enable ecosystem building and industrial clustering.

HosmartAI brings together a **consortium** of leading organizations (3 large enterprises, 8 SMEs, 5 hospitals, 4 universities, 2 research centres and 2 associations – see Table 1) along with several more committed organizations (Letters of Support provided).

Table 1: The HosmartAI consortium.

Number ¹¹	Name	Short name
1 (CO)	INTRASOFT INTERNATIONAL SA	INTRA
1.1 (TP)	INTRASOFT INTERNATIONAL SA	INTRA-LU
2	PHILIPS MEDICAL SYSTEMS NEDERLAND BV	PHILIPS
3	VIMAR SPA	VIMAR
4	GREEN COMMUNICATIONS SAS	GC
5	TELEMATIC MEDICAL APPLICATIONS EMPORIA KAI ANAPTIXI PROIONTON TILIATRIKIS MONOPROSOPIKI ETAIRIA PERIORISMENIS EYTHINIS	TMA
6	ECLEXYS SAGL	EXYS
7	F6S NETWORK IRELAND LIMITED	F6S
7.1 (TP)	F6S NETWORK LIMITED	F6S-UK
8	PHARMECONS EASY ACCESS LTD	PhE
9	SMARTSOL SIA	TGLV
10	NINETYONE GMBH	91
11	HEALTH INNOVATION HUB & HOLDING GMBH	EIT
12	UNIVERZITETNI KLINICNI CENTER MARIBOR	UKCM
13	SAN CAMILLO IRCCS SRL	IRCCS

¹¹ CO: Coordinator. TP: linked third party.

Number ¹¹	Name	Short name
14	SERVICIO MADRILENO DE SALUD	SERMAS
14.1 (TP)	FUNDACION PARA LA INVESTIGACION BIOMEDICA DEL HOSPITAL UNIVERSITARIO LA PAZ	FIBHULP
15	CENTRE HOSPITALIER UNIVERSITAIRE DE LIEGE	CHUL
16	PANEPISTIMIAKO GENIKO NOSOKOMEIO THESSALONIKIS AXEPA	AHEPA
17	VRIJE UNIVERSITEIT BRUSSEL	VUB
18	ARISTOTELIO PANEPISTIMIO THESSALONIKIS	AUTH
19	EIDGENOESSISCHE TECHNISCHE HOCHSCHULE ZUERICH	ETHZ
20	UNIVERZA V MARIBORU	UM
21	INSTITUTO TECNOLÓGICO DE CASTILLA Y LEON	ITCL
22	FUNDACION INTRAS	INTRAS
23	ASSOCIATION EUROPEAN FEDERATION FORMEDICAL INFORMATICS	EFMI
24	FEDERATION EUROPEENNE DES HOPITAUX ET DES SOINS DE SANTE	HOPE

1.2 Document Scope

This deliverable aims at deepening the investigation about the research datasets management, continuing the work to collect all data that will be handled by consortium partners in the frame of the HosmartAI project, in particular filling-in the “Allocations of resources” (Chapter 4), which was missing in the first version of the DMP (D6.7), and providing a new section (3.4.4) about Research Data Alliance (RDA) FAIR Data Maturity Model implementation guidelines

The initial phase of data collection was followed by the elaboration of the first release of this document, and of its submission on M6. This deadline was a mandatory requirement by the **Horizon 2020 Open Access Data Management** requirement: once a project has had its funding approved and has started, it must submit a first version of the DMP (as a deliverable) within the first 6 months of the project¹². Other document updates were planned during the project duration, such as the completion of the datasets’ missing information and the allocation of resources for FAIR data management; this allows for accommodating new findings and aligning the DMP to the needs of HosmartAI respecting active regulations.

1.3 Document Structure

This document is comprised of the following chapters:

Chapter 1 is an introduction to the project, the document scope and structure, and gives an overview of the EU-H2020 Open Access programme and of the FAIR principles for data management applied to the HosmartAI project.

Chapter 2 concerns data summary, quotes the Data Management Survey that served as the first version of the DMP, and presents the project’s pilots and purpose of the data collected, the

¹² https://ec.europa.eu/research/participants/docs/h2020-funding-guide/cross-cutting-issues/open-access-data-management/data-management_en.htm

datasets base information, the data types and formats, the physical location of the datasets, their expected sizes, as well as the purposes of the data (data utility) and identification.

Chapter 3 is based on the FAIR principles, and reports about findability of data, provisions for metadata, open accessibility, data interoperability, data reuse through licensing and quality assurance, as well as data cleansing, transforming and analysing.

Chapter 4 illustrates the allocation of resources for data management, i.e. the costs for data FAIR and open access in the HosmartAI project, data management responsibility, costs for long term preservation.

Chapter 5 concerns the provisions for data security and governance, reporting also the COBIT classification degree for every technical partner.

Chapter 6 presents the ethical and legal aspects linked to data management and data sharing.

Chapter 7 illustrates other issues about data management.

Chapter 8 presents tools and references involved in the management of data.

Chapter 9 lists the references of this document.

Chapter 10 presents the conclusions of this document.

Appendix A reports all data tables related to the datasets.

1.4 Open Access

The European Union (EU) strives to improve access to scientific information and to boost the benefits of public investment in research funded under the EU Framework Programme for Research and Innovation Horizon 2020.

Launched by the European Commission along with the H2020 programme, **Open Access** is the practice of providing online access to scientific information that is free of charge to the reader and is reusable. In the context of research and innovation, scientific information can refer to peer-reviewed scientific research articles or research data.

According to this strategy, in Horizon 2020, a limited pilot action on open access to research data has been implemented so that participating projects are required to develop a Data Management Plan (DMP), in which they specify what data will be open.

According to the H2020 **Article 29.2 of the Model Grant Agreement**¹³ [REF-01][REF-04], each beneficiary must ensure open access to all peer-reviewed scientific publications relating to its results. These open-access requirements are based on a balanced support to both 'Green open access' (immediate or delayed open access that is provided through self-archiving) and 'Gold open access' (immediate open access that is provided by a publisher). Apart from open access to publications, projects must also aim to deposit the research data needed to validate the results presented in the deposited scientific publications, known as "underlying data". In order to

¹³ https://ec.europa.eu/research/participants/data/ref/h2020/grants_manual/amga/h2020-amga_en.pdf#page=242

effectively supply this data, projects need to consider at an early stage how they are going to manage and share the data they create or generate.

Nevertheless, data sharing in the open domain can be restricted if there is a legitimate reason to protect results that can reasonably be expected to be commercially or industrially exploited. Strategies to limit such restrictions will include anonymising or aggregating data, agreeing on a limited embargo period or publishing selected datasets.

From the Data Management point of view, Horizon 2020 strongly suggests that its beneficiaries would make their research data findable, accessible, interoperable and reusable (**FAIR**) to ensure that it is well managed. Good research data management is not a goal in itself, but rather the key conduit leading to knowledge discovery and innovation, and to subsequent data and knowledge integration and reuse. The HosmartAI's DMP follows the FAIR principles [REF-02][REF-03], in particular in the health context [REF-07][REF-08].

1.5 FAIR data concepts

The following two subsections report the FAIR main concepts as illustrated in the FORCE11's Guiding Principles for Findable, Accessible, Interoperable and Re-usable Data Publishing version B1.0 [REF-05].

By adopting all FAIR facets, Data Objects become fully: Findable, Accessible, Interoperable, and Reusable.

The FAIR Data principles aim to ensure that data are shared in a way that enables and enhances reuse by humans and machines. Although FAIR (Findable, Accessible, Interoperable, Reusable) emerged from a workshop for the life-science community, the principles are intended to be applied to data and metadata from all disciplines. Since the formal release via the FORCE11 community, the FAIR data principles have been adopted by several funders and governments worldwide. The European Commission data management guidelines were updated in 2017 to introduce the concept of FAIR [REF-05].

1.5.1 Definitions

- A **Concept** is any defined 'unit of thought' to which we refer in our digital formats.
- A **Data Object** is defined for the purpose of the principles below as: An **Identifiable Data Item with Data elements + Metadata + an Identifier**.
- When we use the term **(Meta) data**, we intend to indicate that the principle is true for **Metadata** as well as for the actual, collected **Data Elements** in the Data Object, and that the principle in question can be independently implemented for both.

1.5.2 FAIR Guiding Principles

1. To be **Findable** any Data Object should be uniquely and persistently identifiable.
 - 1.1. The same Data Object should be re-findable at any point in time; thus, Data Objects should be **persistent**, with emphasis on their metadata.

- 1.2. A Data Object should minimally contain basic machine-actionable metadata that allows it to be distinguished from other Data Objects.
- 1.3. Identifiers for any concept used in Data Objects should therefore be Unique and **Persistent**.
2. Data is **Accessible** in that it can always be obtained by machines and humans.
 - 2.1. Upon appropriate authorization.
 - 2.2. Through a well-defined protocol.
 - 2.3. Thus, machines and humans alike will be able to judge the actual accessibility of each Data Object.
3. Data Objects can be **Interoperable** only if:
 - 3.1. (Meta) data is machine-actionable.
 - 3.2. (Meta) data formats utilize shared vocabularies and/or ontologies.
 - 3.3. (Meta) data within the Data Object should thus be both syntactically parseable and semantically machine-accessible.
4. For Data Objects to be **Re-usable** additional criteria are:
 - 4.1. Data Objects should be compliant with principles 1-3.
 - 4.2. (Meta) data should be sufficiently well-described and rich that it can be automatically (or with minimal human effort) linked or integrated, like-with-like, with other data sources.
 - 4.3. Published Data Objects should refer to their sources with rich enough metadata and provenance to enable proper citation.

To know the maturity level of the application of FAIR principles in this DMP, each pilot, as well as the entire platform, will be checked using the specifications and guidelines of the RDA FAIR Data Maturity Model (<https://www.rd-alliance.org/group/fair-data-maturity-model-wg/outcomes/fair-data-maturity-model-specification-and-guidelines-0>).

See Section 3.4.4.

1.6 Milestones and key achievements in T6.5 Data Management Handling

Table 2: T6.5 achievements history.

Month	Achievements
M2 (February 2021)	Detailed task definition on writing a Data Management Plan (DMP) for the project, based on the FAIR principles
M4 (April 2021)	Identification of the research data to be included into the DMP in the context of the HosmartAI project, and how they will be managed. For this purpose, a survey module was distributed and filled by the contributing partners.

M6 (June 2021)	The first version of the DMP (D6.7) is produced and delivered by M6, Gained access to Open Access publishing (H2020), although not all pilot datasets and information were already included.
M16 (April 2022)	Reached full participation of all pilots, and datasets included.
M22 (October 2022)	First iteration, redaction and delivery of the second version of the Data Management Plan (deliverable D6.8). Also, got missing information about allocation of resources and costs.
M36 (December 2022)	Collaboration with WP2, to enlighten data security and privacy aspects deployed to pilot datasets. Harmonization of DMP with the work performed in dissemination activities (Open Calls).
M39 (March 2024)	EFMI enhanced Section 3.4.4 by providing evidence of compliance with the RDA maturity model FAIR data from HosmartAI's FHIR server.
M41 (May 2024)	Second iteration, redaction and delivery of the final version of the Data Management Plan (deliverable D6.9). [present status: D6.9 closed and submitted to the internal review process]

2 Data summary

As a first step, the following questions inherited from the indications of H2020 Data Management directives were addressed:

- What is the purpose of the data collection/generation and its relation to the objectives of the project?
- What types and formats of data will the project generate/collect?
- Will you re-use any existing data and how?
- What is the origin of the data?
- What is the expected size of the data?
- To whom might it be useful ('data utility')?

2.1 Data Management Survey

For the purpose of collecting information about the research data processed in the frame of the project, a Data Management Survey was developed and proposed to the pilot leaders and technical partners in the first 6 months from project initiation. The involved partners were:

- **AHEPA** for pilot #1
- **CHUL** for pilot #2
- **IRCCS** for pilot #3
- **SERMAS and FIBHULP** for pilot #4
- **UM and UKCM** for pilot #5
- **INTRAS** for pilot #6
- **PHILIPS** for pilot #7
- **VUB** for pilot #8
- **TGLV** for various research data
- **PhE** for various research data

The information collected in the survey was used to complete the first version of the DMP.

Important notice:

This document is the final release of a series of three deliverable documents (D6.7, D6.8 and D6.9) planned during the project. It was deployed to the project partners as a living document, and as such the same was subject to updates during the HosmartAI project. D6.9 is based on the outcomes of D6.7 and D6.8. All 8 pilots have contributed, in particular with information about the allocation of resources and costs in pilot sites, and all the objectives that were still missing in the previous version were fulfilled in this last release of the DMP (D6.9).

2.2 Purpose of the data (HosmartAI pilots)

As explained in the introduction (Section 1.1), eight large-scale pilots have been implemented in the HosmartAI project: improving medical diagnosis, surgical interventions, prevention and treatment of diseases, and support for rehabilitation and long-term care in several hospitals and care settings. These pilots are targeting several **medical** aspects.

Table 3 gives an overview of the eight pilots in relation to the technologies involved.

Table 3: The HosmartAI pilots.

Pilot #	Pilot title	Domain	Main technologies involved	Site	Pilot leader
1	Development of a clinician-friendly, interpretable computer-aided diagnosis system (ICADx) to support and optimise clinical decision making in multi-specialty healthcare environment.	Diagnosis Revolution	Computer-aided diagnosis system.	AHEPA Hospital & Hippokrateio General Hospital of Thessaloniki (Greece)	AHEPA
2	Optimizing the use of radiotherapy	Logistic Improvement	AI algorithm for optimizing patient scheduling.	CHUL Hospital (Belgium)	CHUL
3	Treatment Improvement with the use of innovative technologies and robotics in rehabilitation process	Treatment Improvement	Centralized data collection from wearable devices and environmental sensors.	IRCCS Rehabilitation Centre (Italy)	IRCCS
4	Robotic Systems for minimally Invasive Operation	Surgical support	Robotic system for cardiac catheter navigation, AI and Big Data techniques.	SERMAS Hospital (Spain)	SERMAS
5	Assistive Care in Hospital: Robotic Nurse	Assistive Care	Robotic nurse and Integration of data measured with digital devices.	UKCM Hospital (Slovenia)	UM
6	Assistive Care in Care Centre: Virtual Assistant	Assistive Care	Socially Assistive Robots, eCoach.	INTRAS Care Centre (Spain)	INTRAS
7	Smart Cathlab Assistant	Surgical Support	AI-enabled tools to provide real-time	UZ Brussel (Belgium)	PHILIPS/UZ Brussel

Pilot #	Pilot title	Domain	Main technologies involved	Site	Pilot leader
			clinical decision support and to alleviate the administrative burden in the interventional suite.		
8	Prognosis of cancer patients and their response to treatment combining multi-omics data	Diagnosis and treatment Improvement	General framework to store and analyse raw medical data.	UZ Brussel (Belgium)	VUB/UZ Brussel

The main purpose of HosmartAI’s pilots and the related data outcoming from the HosmartAI project is to improve efficiency in several areas of the medical field (as mentioned before). Moreover, data and datasets are intended to be made open to researchers in the field.

2.3 Datasets, base information

Base information of HosmartAI datasets consists of the type of data collected, the name of the datasets, the related pilots and tasks, as well as the responsible and collaborating partners in charge of managing the different datasets managed by the project.

Reference: Appendix A.1.

2.4 Data types and formats, physical location

This section reports the actual physical location where the original datasets are stored. Moreover, it gives the software tools used for data storage and the data standards used for storing the datasets.

Reference: Appendix A.2.

2.5 Expected sizes and data volumes (nr. of records)

This section gives the approximate size of the datasets used in the HosmartAI project.

Reference: Appendix A.3.

2.6 Data utility and identification

Purpose of data and how they can be re-used (data utility) are important aspects of data management for this project. Reported here is the identifiability of collected information for each dataset and how the data are made accessible to the consortium partners (and whether any restrictions apply).

Reference: Appendix A.4.

3 FAIR

3.1 Making data findable, including provisions for metadata

This section shows if data are discoverable with metadata, identifiable and locatable by means of a standard identification mechanism, reports the identified naming convention, and the search keywords (if any) and the datasets version number to optimize the re-use.

Reference: Appendix A.5.

3.2 Making data openly accessible

All aspects of data open accessibility are covered in this section. This includes which datasets are to be made openly available and in which open repositories those are hosted, as well as the licenses accompanying them, the access identification and the possible restrictions.

Reference: Appendix A.6.

3.3 Making data interoperable

To identify the interoperability of the HosmartAI datasets, metadata vocabularies, ontologies, standards and general methodologies for data interoperability are provided.

Reference: Appendix A.7.

3.4 Increase data re-use (through clarifying licenses)

Re-use of data from the project's datasets are accomplished by answering the following questions:

- How will the data be licensed to permit the widest re-use possible?
- When will the data be made available for re-use? If an embargo is sought to give time to publish or seek patents, specify why and how long this will apply, bearing in mind that research data should be made available as soon as possible.
- Are the data produced and/or used in the project useable by third parties, in particular after the end of the project? If the reuse of some data is restricted, explain why.
- How long is it intended that the data remains re-usable?
- Are data quality assurance processes described?

3.4.1 Data licensing, availability and usability by third parties

This subsection reports the availability, usability and licensing of the project data towards third parties.

Reference: Appendix A.8.

3.4.2 Data Quality Assurance processes

Data Quality Assurance is achieved by specifying for each pilot and task number/dataset the types of analysis that are performed, the person in charge of creating the statistical analysis plan, and how the transformations and analyses on the data are verified.

Reference: Appendix A.9.

3.4.3 Data cleansing, transforming and analysing

Several post-processing activities are envisaged on the project's datasets. Information on data cleansing, transforming and analysing, including the type of data cleaning needed (e.g. correct data types, duplicate removal, add missing info...), the person responsible for data cleaning, the type of data transformation/analysis (e.g. normalization, discretization, ...), the software/tools used for cleaning, transforming, and analysing, where and by whom the analysis will be conducted, and finally, the standards followed for code development/access and re-use (if available).

Reference: Appendix A.10.

3.4.4 Research Data Alliance (RDA) FAIR Data Maturity Model Implementation Guide

Work has been done on the application of the Research Data Alliance (RDA) RDA's FAIR Data Maturity Model [REF-10], following the activities performed Task 6.3 regarding legislation and standardization.

The RDA FAIR Data Maturity Model Working Group has delivered a set of 41 indicators with priorities and guidelines that provide a 'lingua franca' that can be used to make the results of the assessment using those methodologies and tools comparable. The model can function as a tool to be used by various stakeholders to increase the potential for the reuse of research data.

The indicators that are used in the FAIR Data Maturity Model are derived from the FAIR principles and aim to formulate measurable aspects of each principle that can be used by evaluation approaches.

As the scientific community values datasets that underpin research findings, the need to ensure the quality, understanding, and consistency of how these datasets are prepared for others to discover and experience requires a method of measurement. The FAIR Data Maturity Model provides a way for these community-based FAIR assessments to have comparable results and provide consistent feedback as to how well communities are doing in making research data FAIR.

Creating FAIR and AI-ready datasets is transforming the state of AI research practice across disciplines. Considering these activities and given the growth and impact of AI programs for science, it is critical to define at a practical level what FAIR means for AI models. We do this because there is an agreed set of guidelines to FAIRify scientific datasets, from which we will define a compliance model with practical FAIR principles for AI models.

The concept of profiles is being used in the FAIR profile development and adoption community for the practical implementation of FAIR principles in specific domains while ensuring convergence towards a universal model of adoption of the FAIR principles (FAIR convergence).

The approach to the use of the RDA FAIR data maturity model in the HosmartAI project has been made considering that:

To measure the level of compliance with the FAIR principles, we have adapted the model for the project pilot data from a global approach, not analysing the data sets of each pilot one by one.

This analysis will serve as the foundation for an implementation guide for the FAIR data maturity model. The aim of this guide is to facilitate the application of AI and promote the use of data generated by medical devices. The work done in this regard has laid the groundwork for characterizing the mechanisms that will allow progress from the set of indicators of the model to the implementation profiles. This will ultimately lead to improved results of AI models and better utilization of data generated by medical devices in hospitals where AI is used.

In the domain of data collected by IoT devices in healthcare, the aim is to develop a profile that facilitates the sharing and reusability of the data sets, extending discoverability even when combined with the use of AI methods.

A later version of our profiling proposal was considered the FAIR implementation profiling methods being proposed from Go-FAIR, which attempt to maintain the convergence of FAIR methods across different domains [REF-10], taking into account that in addition to FAIR convergence, HosmartAI's priorities with respect to AI and with respect to healthcare IoT devices must be taken into account.

It has been intended to design and implement the two profiles through research on the state of the art of the application of FAIR principles in AI in general and in healthcare in particular, within the timeframe of Task 6.3 development (M5-M41).

In the project, two drivers have been utilized to showcase the achievement of model's indicators.

1. The project's FHIR server guarantees compliance with 26 of the model's indicators. Furthermore, it is possible to cover the remaining 15 indicators through a consensus contribution among the drivers by utilizing semantic standards.
2. On the other hand, the use of the MIMO ontology enables the following indicators to be met:
 - Interoperability I3 RDA-I3-02D: Data includes qualified references to other data.
 - Reusability R1 RDA-R1-01M: A plurality of accurate and relevant attributes is provided to enable reuse.
 - Interoperability I2 RDA-I2-01D: Data uses FAIR-compliant vocabularies. In this case, the FAIR compliance level of the MIMO ontology in its integration into the HosmartAI Platform is the evidence for this.

Evidence of compliance with the RDA maturity model FAIR data from HosmartAI's FHIR server.

(RDA-F1-01M): Metadata is identified by a persistent identifier. The assessment detail indicates that the persistence of an identifier is determined by the policies of the naming authority, suggesting the need for a durable link to the metadata. HosmartAI's FHIR server provision meets this by assigning a globally unique and persistent identifier to both metadata and data, which aligns with the FAIR principle of making data findable with persistent identifiers.

(RDA-F1-01D): Data is identified by a persistent identifier. There's no specific assessment detail provided, but based on the context, this indicator emphasizes the importance of persistent identifiers for data, like the metadata requirement. HosmartAI's approach to using globally unique and persistent identifiers for data supports this need, ensuring data is easily findable.

(RDA-F1-02M): Metadata is identified by a globally unique identifier. The detailed assessment highlights the importance of global uniqueness for identifiers to avoid conflicts and ensure

worldwide findability. HosmartAI's FHIR server provision addresses this by assigning identifiers that are both globally unique and persistent, which enhances the metadata's findability on a global scale.

(RDA-F1-02D): Data is identified by a globally unique identifier. Like the previous row, this emphasizes the need for data identifiers to be globally unique. HosmartAI's use of such identifiers for data as well ensures compliance with this principle, facilitating the global findability and accessibility of data.

(RDA-F2-01M): Rich metadata is provided to allow discovery. The assessment details suggest evaluating this indicator by checking the comprehensiveness and richness of the metadata in facilitating discovery. HosmartAI's provision underlines the use of rich metadata to describe data, which aligns with the principle of enhancing data discovery through detailed metadata.

(RDA-F3-01M): Metadata includes the identifier for the data. This is evaluated by verifying that metadata explicitly includes the data's identifier, ensuring a direct link between them. HosmartAI meets this requirement by ensuring metadata clearly and explicitly includes the identifier for the associated data, thereby enhancing the metadata's utility in identifying and accessing the data.

(RDA-F4-01M): Metadata is offered in a way that it can be indexed. The evaluation involves verifying the accessibility of metadata for indexing purposes. HosmartAI's approach, where metadata is registered or indexed in a searchable resource, fulfils this criterion by making metadata readily available for search and indexing, thereby supporting the data's findability.

(RDA-A1-01M): Metadata contains information to enable the user to access the data. This indicator is assessed by examining metadata for information that facilitates data access. HosmartAI's metadata assignment with globally unique and persistent identifiers likely supports this by providing essential information to access the corresponding data.

(RDA-A1-02M & RDA-A1-02D): These indicators focus on the manual accessibility of metadata and data, respectively, suggesting that both should be accessible through human intervention, such as browsing or searching. HosmartAI's provision for assigning globally unique and persistent identifiers to metadata and data supports this accessibility by ensuring they can be located and accessed manually.

(RDA-A1-03M): Metadata identifier resolves to a metadata record. The assessment involves checking the resolution of the metadata identifier to an actual metadata record. HosmartAI's strategy of using globally unique and persistent identifiers for metadata ensures that each identifier directly leads to the corresponding metadata record, fulfilling this requirement.

(RDA-A1-03D): Data identifier resolves to a digital object. This is evaluated by invoking the identifier, which should lead directly to the digital object it represents. HosmartAI's practice of assigning globally unique and persistent identifiers ensures that each data identifier can be resolved to its respective digital object, facilitating direct access to data.

(RDA-A1-04M & RDA-A1-04D): These indicators emphasize the importance of accessing metadata and data through standardized protocols. The assessment involves checking the protocols used for access to ensure they are standardized and widely adopted. HosmartAI's provision does not specify

the protocols but given the emphasis on unique and persistent identifiers, it suggests an underlying infrastructure that supports standardized access protocols.

(RDA-A1-05D): Data can be accessed automatically (i.e., by a computer program). This indicator is evaluated by the ability to resolve the data identifier through an automated process, ensuring that data can be accessed without human intervention. HosmartAI's approach of using unique and persistent identifiers likely supports automated access, though specific mechanisms or protocols for this are not detailed.

(RDA-A1.1-01M & RDA-A1.1-01D): Both indicators highlight that metadata and data are accessible through a free access protocol. The assessment checks for the openness and cost-free nature of the access protocol. HosmartAI's provision indicates the use of an open, free, and universally implementable protocol for accessing both metadata and data, which aligns with the principles of accessible and interoperable data sharing.

(RDA-A1.2-01D): Data is accessible through an access protocol that allows for an authenticated and authorized request. This indicator focuses on the security aspect of data access, ensuring that data can be accessed securely through authentication and authorization mechanisms. HosmartAI addresses this by implementing a protocol that supports authenticated and authorized requests, which ensures controlled access to sensitive or restricted data.

(RDA-A2-01M): Metadata is guaranteed to remain available even when the data is no longer available. This indicator focuses on the longevity and availability of metadata independent of the data. HosmartAI's provision that metadata should be accessible even when the data is not, supports this requirement, ensuring that metadata remains a valuable resource for understanding, discovery, and context even in the absence of the data.

(RDA-I1-01M & RDA-I1-01D): These indicators stress the use of knowledge representation expressed in a formal, accessible, shared, and broadly applicable language for both metadata and data. This is essential for interoperability and the effective sharing of data. HosmartAI addresses this through its commitment to using a formal, accessible, shared, and broadly applicable language for both metadata and data, which facilitates understanding and reuse across diverse systems and by different users.

(RDA-I1-02M & RDA-I1-02D): Metadata and data use machine-understandable knowledge representation. The evaluation of these indicators would look at whether the knowledge representation formats employed are compatible with machine processing, enhancing automated integration and use. HosmartAI's provision suggests a uniform approach to using formal, accessible, shared, and broadly applicable languages that are likely machine-understandable, promoting interoperability and the potential for automated processing.

(RDA-I2-01M & RDA-I2-01D): Both indicators focus on the use of FAIR-compliant vocabularies for metadata and data. This involves verifying that the vocabularies used are aligned with FAIR principles, ensuring they are findable, accessible, interoperable, and reusable. HosmartAI's commitment to using vocabularies that follow the FAIR principles reinforces the interoperability and reusability of the data and metadata, facilitating their integration and use in various contexts.

(RDA-I3-01M/D to RDA-I3-04M): These indicators focus on the inclusion of references within metadata and data. The indicators emphasize the importance of metadata and data referencing each other and other relevant resources, both directly and with qualifications, to enhance interoperability and context understanding. HosmartAI's approach, where metadata and data include qualified references to each other and to other resources, aligns with these indicators, facilitating richer connections and understanding across datasets.

(RDA-R1-01M): This indicator highlights the richness of the description with a plurality of accurate and relevant attributes for metadata. HosmartAI's provision that metadata and data are described with a rich set of attributes supports the creation of detailed and informative descriptions that enhance findability and reusability.

(RDA-R1.1-01M to RDA-R1.1-03M): These rows address the inclusion of licensing and reuse information within metadata. HosmartAI ensures that metadata and data are released with clear and accessible licensing information, which may include standard reuse licenses and machine-understandable formats, facilitating understanding and compliance with reuse conditions.

(RDA-R1.2-01M & RDA-R1.2-02M): These indicators focus on the inclusion of detailed provenance information within metadata, according to a standard that supports automatic understanding. HosmartAI's commitment to associating metadata and data with detailed provenance information enhances transparency and trust in the data's origins and processing history.

(RDA-R1.3-01M/D to RDA-R1.3-02M/D): These rows emphasize compliance with community standards and machine-understandable standards for metadata and data. HosmartAI addresses these needs by ensuring that metadata and data not only meet domain-relevant community standards but also are expressed in formats that support machine understanding, promoting interoperability and reusability within and across communities.

4 Allocation of resources

This chapter reports the expected costs for making data FAIR in the HosmartAI project and for long term preservation, and how these costs will be covered (taking into consideration that the costs related to open access to research data are eligible as part of the Horizon 2020 grant). The responsible partner for data management is also indicated, when applicable.

4.1 Costs of FAIR and non-FAIR data in HosmartAI

Table 4: Costs for making data FAIR.

Pilot #	Expected costs for making data FAIR
1	Institute dedicated resources.
2	MosaiQ Machine. The relevant data are free by DICOM port for ELEKTRA. Electronic Health Record. History of patient (OMNIPRO). TMA: Data is extracted from CHUL data source through REST queries. Data is transformed to FHIR model with combination of open source / proprietary software. Data is stored to FHIR server.
3	N/A ¹⁴
4	N/A
5	Retrospective data will not be shared. The cost of transforming data to FAIR would include: <ol style="list-style-type: none"> 1. Extraction of data from the specific clinical cohorts (manual extraction via queries) 2. Transformation to FHIR model (automatic, proprietary software) 3. Iterative process (half-automated): Verification of level of anonymity and anonymization (i.e. removal of risks related to re-identification). Open-source tools can be used to decrease the cost. Prospectively collected data can be shared under FAIR conditions. The cost of transforming data to FAIR will include: <ol style="list-style-type: none"> 1. Extraction of data from the specific clinical cohorts (automatic, since prospective Patient generated health data – PGHD - will be stored in FHIR) 2. Creation of meta description for the dataset for the given cohort. Open tools can be used to partially automate the process. 3. Iterative process (half-automated): Verification of level of anonymity (half-automated) and anonymization (i.e., removal of risks related to re-identification). Open-source tools can be used to decrease the cost. 4. Storage of datasets. The use of predefined cohorts published on an open repository (e.g. EOSC) can further mitigate the cost.
6	INTRAS: <ul style="list-style-type: none"> - Extraction of Grador data, both patient and outcome data will be done semi-automatically following a convertible scheme to FHIR.

¹⁴ N/A = Data not available by the time of writing this deliverable.

Pilot #	Expected costs for making data FAIR
	<ul style="list-style-type: none"> - Transformation to FHIR model (automatic, https://hapifhir.hhub.hosmartai.eu/auth). <p>AUTH: Data collected prospectively using iPrognosis tools, namely iMAT, could be shared under FAIR conditions. The cost of transforming data to FAIR would include:</p> <ol style="list-style-type: none"> 1. Extraction of data from the storage databases (hosted either on the cloud or on the pilot’s infrastructure). 2. Transformation to the FHIR model. 3. Creation of meta description of the dataset(s). 4. Storage of the dataset(s). <p>TMA:</p> <ul style="list-style-type: none"> – Extraction data from E-pokratis sensors – Mapping data Conversion to FHIR model <p>ITCL:</p> <ul style="list-style-type: none"> – Data transformation from iPrognosis to FHIR server. – Creation of new anonymization id patient data on FHIR server. – Save configuration profiles for the apps for each patient. <p>Storage to FHIR server</p>
7	N/A
8	<p>Sufficient meta data labelling needs to be done. For genetic and imaging data, it is FAIR as the data is mostly pulled from public databases. We have to make the data via search engine. Patient data from UZ Brussel hospital, cancer variant Database that is similar to rare diseases at the moment. We need to incorporate this data in EU <u>health data space</u>¹⁵ for that inter operatable ontology needs to be defined GONE. Reproducible is not possible as it is a patient data. It should not cost anything as it is done at hospital level. We are ready with all the information we can provide to go into the EU health data space.</p>

4.2 Responsible partner for data management

Table 5: Data management responsible.

Pilot #	Data management responsible partner
1	Data Manager and/or PI of the partner responsible for this dataset.
2	<p>TMA: Collect and store data from the machine. Patient is the data owner, ITCL is the data controller while CHUL is the data processor. ITCL performs data pre-processing and validation.</p> <p>CHUL: Generate anonymized datasheet.</p>
3	N/A
4	N/A
5	N/A

¹⁵ https://health.ec.europa.eu/ehealth-digital-health-and-care/european-health-data-space_en

Pilot #	Data management responsible partner
6	<p>INTRAS: responsible for the data collected from Gradior sessions on its servers as well as for generating the anonymized data for sending to the HosmartAI platform.</p> <p>AUTH: responsible for managing the data collected from iPrognosis tools (performing any processing required by the iPrognosis tools and/or sending it to storage infrastructure).</p> <p>TMA: responsible for data collection from E-pokratis sensors, validation check and pre-processing (convert to FHIR model and store).</p> <p>ITCL: responsible of web integration (frontend and backend) between different platforms and convert iPrognosis data to the FHIR model and storage on HosmartAI platform. Deployment of mobile and pepper robot applications for care center.</p> <p>GC: save data of APK sessions on the blockchain with anonymized data.</p>
7	<p>Data controllers: Jean-francois Argacha (UZ Brussel), Bert Vandeloo (UZ Brussel) and senior Cathlab staff (UZ Brussel). Data processor: Jean-francois Argacha (UZ Brussel), Bert Vandeloo (UZ Brussel), UZ Brussel IT, Philips IT specialist in charge of the study.</p>
8	<p>All relevant personal and clinical data of the participating patients will be processed by the PI of this study or anyone working directly under his supervision. Only those investigators with a medical background (physicians) and physicians caring for the patient, will have access to personal and clinical data. All other involved parties and investigators will only be able to view anonymized data.</p>

4.3 Costs for long term preservations

Table 6: Long term preservation costs.

Pilot #	Long term preservation costs
1	No costs will exist since the data are and will remain stored in AUTH's dedicated resources.
2	CHUL hospital must analyse if the management of the anonymized datasheet will have long term preservation costs.
3	N/A
4	N/A
5	The process of exports of anonymized and full de-identifiable data is done per request and is not stored separately by the UKCM. In HosmartAI, the PGHD (pseudo anonymized and fully identifiable) will be stored on a dedicated on-site FHIR server in a private area network (PAN). The access to the server will be IP and MAC restricted. HosmartAI KPIs will be stored in the HosmartAI FHIR server (this is fully anonymized data). The long term-preservation costs include the allocation of a new FHIR server infrastructure (6-10k euro) and operational costs, and costs related to the maintenance of the server (2-4k euro per year).

Pilot #	Long term preservation costs
6	Current legislation requires that personal information included in this study be kept for 5 to 10 years.
7	Data sent outside UZB to Philips will be pseudonymized. Patient administrative data will be deleted.
8	Current legislation requires that personal information included in this study be kept for 20 to 30 years if this data is also part of their medical record at UZB hospital. The data will be stored at the hospital level.

5 Data security

This chapter aims to answer the following questions:

- What provisions are in place for data security (including data recovery as well as secure storage and transfer of sensitive data)?
- Is the data safely stored in certified repositories for long term preservation and curation?

5.1 Provisions for data security and governance

Classification

For the assessment of governance and security, the maturity of the process is classified in maturity levels using values that are inherited from the COBIT standardization guidelines¹⁶. Those values are:

Table 7: Report on the COBIT classification degree for data security provision for every technical partner.

Acronym	Name	Description
I	Initial	Process not specified, based on spontaneous initiative that is poorly controlled and reactive.
M	Managed	Process is planned, documented and monitored at the project level but not integrated in a broader scope at organization level.
D	Defined	Proactive process active at organization level.
Q	Quantitatively Managed	The process is measured and controlled/ verified.
O	Optimizing	Focus is on continuous process and improvement.

Table 8: Data security.

Partner short name	Classification	Code of conduct used for each task	Levels of data security in place.	Levels of security that primary data used in the project will undergo	Name and email of the data privacy officer(s)
EXYS	Q	Data handling procedures at EXYS are defined according to GDPR and LPD (Swiss data protection law)	Technical measures: <ul style="list-style-type: none"> • Data storage in dedicated servers segmented on network environment • Authorized access with data access audit logs • VPN regulated access to the data processing data centre • Data communication 	Whenever required by the project, data will be stored in firewall-protected computers with strong authentication.	Angelo Consoli; angelo.consoli@ecl-exys.com

¹⁶ ISACA®, [COBIT® 2019 Framework: Governance and Management Objectives](#), USA 2018

Partner short name	Classification	Code of conduct used for each task	Levels of data security in place.	Levels of security that primary data used in the project will undergo	Name and email of the data privacy officer(s)
			and transfer protected by Secure Socket Layer (SSL) and Transport Layer Security (TLS) protocols.		
AUTH	N/A	Data are handled according to GDPR and international ethical guidelines (inter alia, the Word Medical Association Declaration of Helsinki), mandated by the AUTH Research Ethics Committee. Ethical approval must be obtained before data collection involving human beings.	Technical measures include: <ul style="list-style-type: none"> • Data storage in firewall-protected computers • Authorized access with data access audit logs • Data transfer via Secure Shell (SSH) or Secure Socket Layer (SSL) protocols. 	<ul style="list-style-type: none"> • Data will be stored in firewall-protected computers with authorized access. • Access will be limited to members of the AUTH research team. • In case of data transfer, this will take place via SSH or SSL protocols. • Pseudo-anonymised data will be stored separately from records including subjects' personally identifiable information (e.g. signed consent forms) 	Ms. Kornilia Skarpeta data.protection@auth.gr
VIMAR	M	Data are handled according to GDPR.	Technical measures include: <ul style="list-style-type: none"> • Data storage in firewall-protected computers. • Authorized access with data access audit logs. • Data transfer via Secure Shell (SSH) or Secure Socket Layer (SSL) protocols. 	<ul style="list-style-type: none"> • Data will be stored in firewall-protected computers with authorized access. • Access will be limited to authorized members of the team. • In case of data transfer, this will take place via SSL protocols. • Pseudo-anonymised data will be stored separately from records including subjects' personally identifiable information. 	Beni Luigi Giancesin beni.giancesin@vimar.com

Partner short name	Classification	Code of conduct used for each task	Levels of data security in place.	Levels of security that primary data used in the project will undergo	Name and email of the data privacy officer(s)
PhE	N/A	Data analysis will be handled according to GDPR following the PhE Standard Operating Procedure of Process & Handling of Personal Data.	Cloud – One Drive	Data will be password protected and accessed only by the PhE HOSMARTAI dedicated team.	George Tsonis Tsonislaw@yahoo.com
CHUL	Q	Data are handled according to GDPR mandated by both DPO and the CHU Liège Ethics Committee			Ghislaine.Dumont@chuliege.be
ITCL	Q	Data are handled according to GDPR and international ethical guidelines. Ethical approval must be obtained before data collection involving human beings.	Technical measures include: <ul style="list-style-type: none"> • Data storage in firewall-protected computers • Authorized access with data access audit logs • Data transfer via Secure Shell (SSH) or Secure Socket Layer (SSL) protocols. • Encryption of data. 	Data will be stored in firewall-protected computers with authorized access. Access will be limited to members of the ITCL research team. In case of data transfer, this will take place via SSH or SSL protocols. Pseudo-anonymised data will be stored separately from records including subjects' personally identifiable information (e.g. signed consent forms) Critical data will be encrypted for better patient safety.	Angel Lopez, angel.lopez@itcl.es
VUB	I/M	Data are handled according to GDPR and international ethical guidelines, as well as internal UZ Brussel patient data security rules.	The data will remain in the tightly controlled and firewalled UZ Brussel ICT system, with only local or VPN access to limited authorised subsystems.	Limited access by authorized personnel only, no access from outside nor to outside from virtual machine harbouring the data	Data protection office dpo@vub.be

Partner short name	Classification	Code of conduct used for each task	Levels of data security in place.	Levels of security that primary data used in the project will undergo	Name and email of the data privacy officer(s)
UZB, VUB	?	Data will be handled according to EU 2016/679 GDPR.	Data storage in firewall-protected computers Authorized access with data access audit logs Data transfer via Secure Shell (SSH) or Secure Socket Layer (SSL) protocols.	Data will be stored in firewall-protected computers with authorized access. Access will be limited to authorized members of the team. In case of data transfer, this will take place via SSL protocols.	Luc Maes lucm@uzbrussel.be
INTRAS	Q	Data handling procedures at INTRAS are defined according to GDPR	Technical measures: - Data storage in dedicated servers segmented on network environment - Authorized access with data access audit logs - VPN regulated access to the data processing data centre Data communication and transfer protected by Secure Socket Layer (SSL) and Transport Layer Security (TLS) protocols.	- Data will be stored in firewall-protected computers with authorized access. - Access will be limited to members of the INTRAS research team. - In case of data transfer, this will take place via SSH or SSL protocols. Pseudo-anonymised data will be stored separately from records including subjects' personally identifiable information (e.g. signed consent forms)	Francisco Mallo Vázquez <fmv@intras.es>
UM	M	Data are handled according to UM's privacy policy (https://feri.um.si/en/about-us/privacy-policy/)	Technical measures include: • Data storage in firewall-protected computers • Authorized access with data access audit logs • Data transfer via Secure Shell (SSH) or Secure Socket Layer (SSL) protocols	• UM will not store sensitive data	Doc. Dr. Miha Dvojmoč (dpo@um.si), if contacted please always refer project and the main contact person of the project (izidor.mlakar@um.si)

Partner short name	Classification	Code of conduct used for each task	Levels of data security in place.	Levels of security that primary data used in the project will undergo	Name and email of the data privacy officer(s)
UKCM	M	Data are handled according to GDPR and international ethical guidelines (inter alia, the World Medical Association Declaration of Helsinki), mandated by the UKCM's and National Ethics Committee. Ethical approval must be obtained before data collection involving human beings.	Technical measures include: <ul style="list-style-type: none"> • Data storage in firewall-protected computers. • IP and MAC restricted access • Authorized access with data access audit logs • Data transfer via Secure Shell (SSH) or Secure Socket Layer (SSL) protocols. 	<ul style="list-style-type: none"> • Data will be stored in firewall-protected computers with authorized access. • Access will be limited to members of the UKCM and UM research team. A specific DTA will be established. • In case of data transfer, this will take place via SSH or SSL protocols. • Pseudo-anonymised data will be stored separately from records including subjects' personally identifiable information (e.g. signed consent forms) 	mag. Klara Mihaldinec, (dpo@ukc-mb.si)
91	Q	Data handling procedures are defined according to HIPAA and GDPR	Data storage in HIPAA compliant cloud, high-end encryption with endpoint threat detection and multi factor authentication	<ul style="list-style-type: none"> • Data will be stored and managed according to the highest levels of protection and encryption as required by HIPAA and GDPR, with firewall, restricted access, and endpoint cybersecurity threat detection 	Durim Krasniqi Durim@91.life
SERMAS	O	Best Practices set out by data privacy officers	Data storage in servers that require authorization for access.	<ul style="list-style-type: none"> • Access by a password. • Anonymized data retrieval and transfer. 	FUNDACIÓN PARA LA INVESTIGACIÓN BIOMÉDICA DEL HOSPITAL UNIVERSITARIO LA PAZ (FIBHULP). protecciondedatos@idipaz.es
GC	Q	Data handling procedures are defined according to GDPR	Technical measures include: <ul style="list-style-type: none"> • Wireless communications are WPA2/WPA3 encrypted in 	<ul style="list-style-type: none"> • Data will be stored in authorized access local edge server. • The server will be confined and 	Green Communications contact@green-communications.fr

Partner short name	Classification	Code of conduct used for each task	Levels of data security in place.	Levels of security that primary data used in the project will undergo	Name and email of the data privacy officer(s)
			<p>access and backhaul.</p> <ul style="list-style-type: none"> • Remote access is provided through secure IP tunnels (VPN) or secure shell (SSH). • Blockchain application uses use TLS (Transport Layer Security) and HTTPS cryptographic protocols. • The blockchain node can be physically isolated from the Internet 	<p>isolated from the internet or connected.</p> <ul style="list-style-type: none"> • when connected, the node is protected by a firewall and remote access is ensured via VPN and SSH by authenticated users. 	

6 Ethical and legal aspects

Ethical and legal aspects which are part of the HosmartAI project are treated in this chapter. The following questions will be addressed:

- Are there any ethical or legal issues that can have an impact on data sharing? These can also be discussed in the context of the ethics review. If relevant, include references to ethics deliverables and the ethics chapter in the Description of the Action (DoA).
- Is an informed consent for data sharing and long-term preservation included in questionnaires dealing with personal data?

6.1 Ethical issues for data sharing

Data Protection is a fundamental human right, as enshrined in the EU Charter of Fundamental Rights, aimed at providing any individual¹⁷ with control over the way their personal information is collected and used. Article 8(1) of the Charter of Fundamental Rights of the European Union (the ‘Charter’) and Article 16(1) of the Treaty on the Functioning of the European Union (TFEU) grant everyone the right to the protection of their personal data. Data protection is a central issue for research ethics.

Whenever personal data is collected, there are both ethical and legal obligations to ensure that participants’ information is properly protected. This is fundamental to safeguarding their rights and freedoms, and minimising the ethics risks related to the data processing. In HosmartAI data security is provided on all levels. Only authorized users will have access to digital information. The project will adopt recommendations and standards provided by ENISA (European Union Agency for Cybersecurity) [REF-06]. It is the goal of all project partners to mitigate the risk for all participating patients

Publication of Results HosmartAI complies with the highest ethical standards. Researchers, authors, sponsors, editors and publishers all have ethical obligations regarding the publication and dissemination of the research results.

6.1.1 Ethical review

This subsection deals with the type of ethical review needed for each pilot.

Reference: Appendix A.11.

6.2 Legal issues for data sharing

The GDPR provides the basic legal framework for personal data processing and therefore data sharing. Particular attention must be paid to research involving sensitive data such as health data, which according to GDPR must not be processed unless the data subject has given explicit consent.

¹⁷ An individual is an identifiable natural person is one who can be identified, directly or indirectly, in particular by reference to an identifier such as a name, an identification number, location data, an online identifier or to one or more factors specific to the physical, physiological, genetic, mental, economic, cultural or social identity of that natural person (Art. 2(a) EU General Data Protection Regulation (GDPR)).

This imposes obligations on researchers to provide research subjects with detailed information about what will happen to the personal data that they collect.

In HosmartAI, all data processing complies with EU law as well as national data laws. It is ensured that any partners, contractors or service providers that process research data at HosmartAI partners' request and on their behalf comply with the GDPR and the H2020 ethics standards. Special attention is given to a good balance between research objectives and the means used to achieve them.

Types of Personal Data Processed During the lifetime of HosmartAI two categories of personal and sensitive data are/may be generated or collected: (1) "Data related to stakeholders" (individuals working for the consortium partners or in any way professionally involved with the project, etc.): Information on these data subjects, such as contact details (e.g. e-mails and names), their signatures, authorship of deliverables, etc. is collected and processed by all partners of the Project. (2) "Patient's health data in pilots" (sensitive data or "special personal data" pursuant to Article 9 GDPR):

Informed Consent When personal data is used, informed consent is the cornerstone of research ethics. The lawful basis for the processing of personal data related to stakeholders, under the GDPR, is that each data subject working for a project partner has given consent to the processing of their personal data (GDPR Article 6 (1)(a)) and that the processing is necessary for the performance of a contract - namely, the data subjects' employment agreements with each project partner (GDPR Article 6(1)(b)). At the end of the project, files containing personal information of data subjects working for Project partners will be maintained by each project partner. Any partner will have the right to continue to maintain its copy of the contact data of employees working for HosmartAI partners unless employees have requested a deletion of the contact data. Mailing lists of the project will be deleted only after the very final payment and assessment from the European Commission. Data subjects' contact details will be shared only with project members and only for the time needed to execute the Grant Agreement and/or complete the project. Authorship information may be made publicly available with the consent of the data subjects once the application becomes publicly or commercially available.

Whenever personal data is collected from patients, the patients' informed consent must be sought by means of a procedure that meets the minimum standards of the GDPR. This requires consent to be given by a clear affirmative act. For consent to data processing to be 'informed', the data subject must be provided with detailed information about the envisaged data processing in an intelligible and easily accessible form, using clear and plain language. The researchers at the pilots will explain to patients (i) what the research is about; (ii) what their participation in the project will entail and what risks may be involved. The partner will give information as to whether data will be shared with, or transferred to, third parties and for what purposes and for how long the data will be retained before being destroyed. The patients will also be informed about the right to withdraw consent or access their data. They will also be told the procedures to follow should they wish to do so. They will also receive information on their right to lodge a complaint with a supervisory authority. The data subjects must also be made aware if data are to be used for any other purposes, and if it is to be shared with research partners or transferred to organisations outside

the EU. Records documenting the informed consent procedure will be kept, including the information sheets and consent forms provided to research participants. The consent process(es) and the information provided to the data subjects will cover all the data processing activities related to their participation in HosmartAI. If in the course of the HosmartAI research project, any significant changes to methodology or processing arrangements that have a bearing on the data subjects' rights or the use of their data should occur, the data subjects will be made aware of the intended changes, and their express consent for further use of the data will be sought.

Privacy by Design To innovate ethically and responsibly, researchers and developers apply the concept of 'privacy by design', which provides a framework for focusing the design of systems, databases and processes with respect to data subjects' fundamental rights. A wider concept of 'data protection by design', now included in the GDPR, requires the implementation of appropriate technical and organisational measures to give effect to the GDPR's core data-protection principles. Data protection by design is one of the best ways to address the ethics concerns that arise within a research project. Minimisation of data is essential in this respect. Data processing must be lawful, fair and transparent. It should involve only data that are necessary and proportionate to achieve the specific task or purpose for which they are collected.

Deletion and Archiving of Data Personal data will only be kept as long as is necessary for the purposes for which they are collected, or in accordance with the established auditing, archiving or retention provisions of HosmartAI. As soon as the research data is no longer needed, or subject to an established retention period, the data will be deleted. Data retained for auditing processes will be stored securely and further processed for those purposes only. Research data held in the cloud or by a third-party service provider, will also be held together with any back-ups.

Reuse of Data A potential later use of the HosmartAI platform may permit medical researchers to use data sets for the purpose of conducting medical research. The procedure for this eventuality has not yet been addressed by the project partners. As a result of this effort, ethical and legal considerations may arise with respect to large scale or big data processing, and will be discussed in the last version of the DMP.

7 Other issues about data management

This chapter reports on other possible issues related to data management in the HosmartAI project. For instance, the use of other national/funder/sectorial/departmental procedures for data management.

Pilot #	Other issues related to data management
1	No other issue identified.
2	CHUL hospital has to establish and manage the anonymized datasheet.
3	No other issue identified.
4	No other issue identified.
5	UKCM prefers the decentralized approach in which data is stored within the pilot site. If sharing of prospective data is established, UKCM must create and anonymize data. Individual DTAs with clearly identified intent for the use of 'raw' data and how data will be handled must be negotiated. Fully anonymized cohorts can be shared openly for research.
6	No other issue identified.
7	No other issue identified.
8	No other issue identified.

8 Tools and references

- The Research Data Alliance provides a [Metadata Standards Directory](#) that can be searched for discipline-specific standards and associated tools.
- Research Data Alliance FAIR Data Maturity Model (<https://www.rd-alliance.org/group/fair-data-maturity-model-wg/outcomes/fair-data-maturity-model-specification-and-guidelines-0>).
- The [EUDAT B2SHARE](#) tool includes a built-in license wizard that facilitates the selection of an adequate license for research data.
- Useful listings of repositories include:
 - [Registry of Research Data Repositories](#)
 - Some repositories like [Zenodo](#), an OpenAIRE and CERN collaboration), allow researchers to deposit both publications and data, while providing tools to link them.
- Other useful tools include DMP online and platforms for making individual scientific observations available such as [ScienceMatters](#).
- Mosaiq Machine data for radiotherapy by FHIR.
- Omnipro¹⁸. IT CHUL hospital transfer the anonymized therapy data by FHIR.
- FHIR4FAIR Implementation Guide (HL7 project underway, ballot scheduled for September 2021) (<http://build.fhir.org/ig/HL7/fhir-forfair/>)
- ROAR: Research Open Access Repository, <http://roar.eprints.org>
- OpenDOAR: Directory for Open Access Repositories: <https://v2.sherpa.ac.uk/opensoar/>
- OpenAIRE: <https://www.openaire.eu/>
- Zenodo: <https://zenodo.org/>
- Elekta MosaiQ Radiation Oncology: <https://www.elekta.com/software-solutions/care-management/mosaiq-radiation-oncology/>
- GRADIOR: Computer-based cognitive rehabilitation program.
- [EVA Corpus](#): A Corpus for Analysing Linguistic and Paralinguistic Features in Multi-Speaker Spontaneous Conversations
- HBASE, MONGO
- K-Anonymity: <https://github.com/Nuclearstar/K-Anonymity>
- ARX - Open Source Data Anonymization Software: <https://github.com/arx-deidentifier/arx>
- European Open Science Cloud (EOSC): <https://eosc-portal.eu/>

¹⁸ <https://cabinetprive.xperthis.com/omnipro/>

9 References

[REF-01]	AGA - Annotated Model Grant Agreement, H2020 Programme, v. 5.2., June 2019
[REF-02]	Template for the Data Management Plan , H2020 Programme
[REF-03]	FAIR Data Management template Summary table , H2020 programme
[REF-04]	H2020 Model Grant Agreement - Article 29.2
[REF-05]	FORCE11's Guiding Principles for Findable, Accessible, Interoperable and Re-usable Data Publishing (https://www.force11.org/fairprinciples)
[REF-06]	ENISA: European Union Agency for Cybersecurity (https://www.enisa.europa.eu/)
[REF-07]	Jaime Delgado, FAIR4Health - Report on Security and Privacy in FAIR processes, Horizon 2020 project, grant agreement No 824666
[REF-08]	Jaime Delgado and al., Approaches to the integration of TRUST and FAIR principles, Universitat Politècnica de Catalunya (UPC BarcelonaTECH)
[REF-09]	Research Data Alliance (RDA) FAIR Data Maturity Model, https://www.rd-alliance.org/groups/fair-data-maturity-model-wg
[REF-10]	Data Intelligence 2(2020), 158-170. doi: 10.1162/dint_a_00038

10 Conclusions

The last (final) version of the HosmartAI DMP completes the investigations related to the research datasets management, initiated in the first version of the DMP (D6.7) and continued in the second version (D6.8). This document finalizes the work of filling in data in the tables and sections, some of which was still missing in the second version of the DMP.

Section 3.4.4 on the RDA FAIR Data Maturity Model implementation guidelines, was enhanced by providing evidence of compliance with the RDA maturity model FAIR data from HosmartAI's FHIR server. The work described in 3.4.4 was carried on in Task 6.3, through the implementation of two profiles, and provide a quantitative self-assessment model for measuring the maturity level of the datasets.

During the last stage of the DMP implementation, a collaboration was undertaken with WP2 to elucidate some data security and privacy aspects deployed to pilot datasets. In addition, the DMP was harmonized considering the work performed in the same WP about dissemination activities (Open Calls).

The maturity level of the pilot datasets, measured quantitatively using the RDA FAIR Data Maturity Model guidelines and implemented in Task T6.3, concludes the work done in T6.5 and leads to this final version of the DMP.

Appendix A Datasets

Preliminary remark:

It was a common decision within Pilot 6, during the last stage of the DMP implementation, that the datasets DS3.2, DS3.3 and DS3.4 do not fit the purpose of pilot 3. Hence, they were removed from the list of the managed research datasets, but they were kept in the following tables for convenience, appearing crossed out (in strikethrough characters).

A.1 Datasets base information

Table 9: Datasets base information.

Code	Type of data to be collected / name of the dataset	Pilot Nr.	Task Nr.	Responsible partner	Collaborating partners
DS1.1	Cardiac ultrasound video recordings	1	3.1, 5.2	AHEPA	AUTH
DS1.2	Capsule endoscopy video recordings	1	3.1, 5.2	AHEPA	AUTH
DS1.3	Cardiotocography variables and results, biometric data, medical history data	1	3.1, 5.2	AUTH	N/A
DS1.4	Coronary computed tomography angiography (CCTA) variables, biometric data, medical history data	1	3.1, 5.2	AHEPA	AUTH
DS2.1	Data related to hours spent by specialists.	2	3.2	CHUL	ITCL
DS2.2	Retrospective patient schedule data and precondition of the treatment	2	3.2	CHUL	ITCL
DS2.3	Data linked to radiotherapy machines (tumours treatment indication, maintenance, building location)	2	1.2	CHUL	N/A
DS2.4	PROMs/PREMs	2	5.2	CHUL	UM
DS2.5	Prospective patient clinical data	2	5.2	CHUL	ITCL
DS2.6	Patient personal data (address, preferences, ...)	2	5.2	CHUL	ITCL
DS2.7	Retrospective EHR	2	3.2	CHUL	ITCL

Code	Type of data to be collected / name of the dataset	Pilot Nr.	Task Nr.	Responsible partner	Collaborating partners
DS2.8	Data from radiotherapy services, infrastructure. Patient's satisfaction	2	3.2	ITCL	TMA
DS3.1	Smart home data: Consumption/production of instantaneous electricity and consumption logs, Human presence/access control, Devices' activation and connected loads	3	3.3	IRCCS	VIMAR
DS3.2	IMU data captured by iPrognosis smartphone application	3,6	5.2	AUTH	INTRAS
DS3.3	Voice related time and spectral features captured by iPrognosis smartphone application	3,6	5.2	AUTH	INTRAS
DS3.4	Key taps press and release timestamps captured by iPrognosis smartphone virtual keyboard	3,6	5.2	AUTH	INTRAS
DS3.5	Angles of elbows and shoulders and scores achieved at the tests included in the iPrognosis iMAT tool	3,6	5.2	AUTH	INTRAS
DS3.6	Results of rehabilitation sessions with Gradior and care plans (editor and patient data)	6	3.5	INTRAS	ITCL/AUTH
DS4.1	Demographic patient data (age, gender, atrial size, etc.)	4	5.2	SERMAS	91
DS4.2	3D navigation system data (intracardiac signals, geometry)	4	5.2	SERMAS	91
DS5.1	Patient recordings and features extracted from interaction with patients	5	5.2	UM	UKCM, ITCL
DS5.2	Biometric data (e.g., blood pressure and heart rate)	5	5.2	UKCM	UM

Code	Type of data to be collected / name of the dataset	Pilot Nr.	Task Nr.	Responsible partner	Collaborating partners
DS5.3	Retrospective electronic health records	5	5.2	UKCM	UM
DS5.4	PREMs related to clinical staff (depends on T1.4)	5	5.2	UKCM	UM
DS5.5	PREMs related patients (PAM, SUS-SI/TAM, UEQ)	5	5.2	UM	UKCM
DS5.6	Datasets for facial expression and emotion recognition	5	3.5	UM	N/A
DS5.7	Datasets for ASR and TTS in Slovenian	5	3.5	UM	N/A
DS5.8	Datasets for ASR and TTS in French	5	3.5	UM	N/A
DS5.9	EVA Corpus ¹⁹ , data set of conversational expression	5	3.5	UM	N/A
DS5.10	Video recordings of third persons	5	3.5	ITCL	UM
DS5.11	Recording of a skeleton representation of the patient doing breathing exercises	5	3.5	ITCL	UM
DS5.12	Patient's facial information	5	3.5	UM	N/A
DS6.1	Patient's physical, medical and cognitive status. Vital signs with smartwatches sensor data.	6	3.5	ITCL	INTRAS / TMA
DS7.1	Administrative data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	7	3.6	Cardiology department of University Hospital Brussels, VUB	Philips Image Guided Therapy Systems (Philips)
DS7.2	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	7	3.6	Cardiology department of University Hospital Brussels, VUB	Philips Image Guided Therapy Systems (Philips)
DS7.3	Coronary angiogram imaging data of patients who undergone a coronary	7	3.6	Cardiology department of University	Philips Image Guided Therapy

¹⁹ <https://www.iasr.org/iasr/home/cijc/a-corpus-for-analyzing-linguistic-and-paralinguistic-features-in-multi-speaker-spontaneous-conversations-eva-corpus>

Code	Type of data to be collected / name of the dataset	Pilot Nr.	Task Nr.	Responsible partner	Collaborating partners
	angiogram/coronary intervention @ UZB			Hospital Brussels, VUB	Systems (Philips)
DS7.4	Coronary physiology data of patients evaluated by a resting index measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary intervention @ UZB	7	3.6	Cardiology department of University Hospital Brussels, VUB	Philips Image Guided Therapy Systems (Philips)
DS7.5	Intravascular imaging data of patient evaluated by either OCT or IVUS technique before and after a coronary intervention	7	3.6	Cardiology department of University Hospital Brussels, VUB	Philips Image Guided Therapy Systems (Philips)
DS7.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention	7	3.6	Cardiology department of University Hospital Brussels, VUB	Philips Image Guided Therapy Systems (Philips)
DS7.7	Full prospective UZB data set of administrative, clinical, coronary angiogram, coronary physiology, intravascular imaging and coronary CT data.	7	3.6	Cardiology department of University Hospital Brussels, VUB	Philips Image Guided Therapy Systems (Philips)
DS8.1	Image, gene, phenotype and pathology data for glioma patients	8	N/A	VUB	UZ Brussel
DS.O.1	All pilot data with Common KPIs (Economic & PROMs/PREMs)	PhE	5.3	All pilots	PhE

A.2 Data types and formats, physical location

Table 10: Data types, formats and physical location.

Code	Type of data to be collected / name of the dataset	Physical location where primary (original) data is stored	Software/ tools used for storage of data?	Format and type of data standards used to store the data
DS1.1	Cardiac ultrasound video recordings	At the edge, dedicated infrastructure for the pilot	N/A	DICOM, HL7-compatible resource
DS1.2	Capsule endoscopy video recordings	At the edge, dedicated infrastructure for the pilot	N/A	MP4, HL7-compatible resource
DS1.3	Cardiotocography variables and results, biometric data, medical history data	At the edge, dedicated infrastructure for the pilot	N/A	N/A
DS1.4	Coronary computed tomography angiography (CCTA) variables, biometric data, medical history data	At the edge, dedicated infrastructure for the pilot	N/A	DICOM partially (others TB)
DS2.1	Data related to hours spent by specialists.	Private cloud of the University of Liège. This infrastructure is based on ESX virtualized environment. This infrastructure is certified ISO27001 and ISO9001 annually since 2016.	UltrAgenda	HL7-compatible resource
DS2.2	Retrospective patient schedule data and precondition of the treatment	As DS2.1	Mosaiq OIS	Json

Code	Type of data to be collected / name of the dataset	Physical location where primary (original) data is stored	Software/ tools used for storage of data?	Format and type of data standards used to store the data
DS2.3	Data linked to radiotherapy machines (tumours treatment indication, maintenance, building location)	As DS2.1	Mosaiq OIS	Json
DS2.4	PROMs/PREMs	As DS2.1	Questionnaires stored in Excel file	Excel format
DS2.5	Prospective patient clinical data	As DS2.1	EHR	Json
DS2.6	Retrospective EHR	As DS2.1	EHR	Json
DS2.7	Data from radiotherapy services, infrastructure. Patient's satisfaction, preferences....	At the edge, dedicated infrastructure for the pilot	Data Store (, RDBMS)	Media Resource and formats of RDBMS observations
DS3.1	Smart home data: Consumption/product ion of instantaneous electricity and consumption logs, Human presence/access control, Devices activation and connected loads	On VIMAR cloud infrastructure	InfluxDB	JSON
DS3.2	IMU data captured by iPrognosis smartphone application	Cloud infrastructure for the pilots	Schemaless database	JSON, HL7-compatible resource
DS3.3	Voice-related time and spectral features captured by iPrognosis smartphone application	Cloud infrastructure for the pilots	Schemaless database	JSON, HL7-compatible resource

Code	Type of data to be collected / name of the dataset	Physical location where primary (original) data is stored	Software/ tools used for storage of data?	Format and type of data standards used to store the data
DS3.4	Key taps press and release timestamps captured by iPrognosis smartphone virtual keyboard	Cloud infrastructure for the pilots	Schemaless database	JSON, HL7-compatible resource
DS3.5	Angles of elbows and shoulders and scores achieved at the tests included in the iPrognosis iMAT tool	Cloud infrastructure for the pilots	Schemaless database	JSON, HL7-compatible resource
DS3.6	Results of rehabilitation sessions with Grador and care plans (editor and patient data)	Cloud infrastructure for the pilots	Relational databases	JSON
DS4.1	Demographic patient data (age, gender, atrial size, etc.)	SERMAS	Server (provided by 91)	N/A
DS4.2	3D navigation system data (intracardiac signals, geometry)	SERMAS Cloud, Navigation system internal memory	Server (provided by 91)	*.xls or *.xlsx files
DS5.1	Patient recordings and features extracted from interaction with patients	At the edge, dedicated infrastructure for the pilot.	Open FHIR Server	FHIR Media Resource FHIR Observation and FHIR Compositions
DS5.2	Biometric data (e.g., blood pressure and heart rate)	At the edge, dedicated infrastructure for the pilot	Open FHIR Server	FHIR Diagnostic Report, and Observation resource
DS5.3	Retrospective electronic health records	At the edge, interface with existing IT system of the hospital	Proprietary HI7 Compliant Medis Server	N/A
DS5.4	PREMs related to clinical staff (depends on T1.4)	A GDPR compliant online survey infrastructure (https://1ka.arne)	SPSS or similar, Open FHIR Server	FHIR Questionnaire, FHIR QuestionnaireResp

Code	Type of data to be collected / name of the dataset	Physical location where primary (original) data is stored	Software/ tools used for storage of data?	Format and type of data standards used to store the data
		s.si/index.php?lang_id=2) or UM's CHATBOT for collecting PROMs/PREMs		onse, FHIR Composition
DS5.5	PREMs related patients (PAM, SUS-SI/TAM, UEQ)	A GDPR compliant online survey infrastructure (https://1ka.arnes.si/index.php?lang_id=2), Open FHIR Server	SPSS or similar, Open FHIR Server	FHIR Questionnaire, FHIR QuestionnaireResponse, FHIR Composition
DS5.6	Datasets for facial expression and emotion recognition	UM's internal infrastructure	N/A	N/A
DS5.7	Datasets for ASR and TTS in Slovenian	UM's internal infrastructure	N/A	N/A
DS5.8	Datasets for ASR and TTS in French	UM's internal infrastructure	N/A	N/A
DS5.9	EVA Corpus, data set of conversational expression	UM's internal infrastructure And CLARIN.SI Repository	N/A	N/A
DS5.10	Video recordings of third persons	At the edge in the hospital, dedicated infrastructure for the pilot	N/A	MP4
DS5.11	Recording of the skeleton of the patient doing breathing exercises	At the edge in the hospital, dedicated infrastructure for the pilot	N/A	MP4
DS5.12	Patient's facial information	At the edge, dedicated	Data Store (HBASE,	FAPS, AUs

Code	Type of data to be collected / name of the dataset	Physical location where primary (original) data is stored	Software/ tools used for storage of data?	Format and type of data standards used to store the data
		infrastructure for the pilot	MONGO, RDBMS)	
DS6.1	Patient's physical, medical and cognitive status. Vital signs with sensors data	At the edge, dedicated infrastructure for the pilot	Data Storage FHIR and RDBMS	Media Resource and formats RDBMS and json for FHIR server.
DS7.1	Administrative data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	UZB, VUB	Electronic health records system branded PRIMUZ	xlsx files
DS7.2	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	UZB, VUB	Electronic health records system branded PRIMUZ	xlsx files
DS7.3	Coronary angiogram imaging data of patients who undergone a coronary angiogram/coronary intervention @ UZB	UZB, VUB	Philips cathlabs over the Philips IntelliSpace CardioVascular (ISCV) Portal	DICOM
DS7.4	Coronary physiology data of patients evaluated by a resting index measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary intervention @ UZB	UZB, VUB	Coroventis platform (RFR/FFR) and Volcano platform (iFR/FFR)	Data extracted in .dat or .xls files
DS7.5	Intravascular imaging data of patient evaluated by either	UZB, VUB	Abbott OPTIS platform (OCT)	N/A

Code	Type of data to be collected / name of the dataset	Physical location where primary (original) data is stored	Software/ tools used for storage of data?	Format and type of data standards used to store the data
	OCT or IVUS technique before and after a coronary intervention		(Integrated platform) Volcano platform (IVUS) (non-integrated platform)	
DS7.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention	UZB, VUB	Philips (CT) Heartflow (FFRCT)	N/A
DS7.7	Full prospective UZB data set of administrative, clinical, coronary angiogram, coronary physiology, intravascular imaging and coronary CT data.	Philips AI smart-cathlab prototype working at UZB, VUB	N/A	N/A
DS8.1	Image, gene, phenotype and pathology data for glioma patients	Separate in-hospital databases, with final central integration on site	PRIMUZ, XNAT	RDMS (various), JSON
DS.O.1	All pilot data with Common KPIs (Economic & PROMs/PREMs)	The original data at each pilot's repository. The analysis of PhE will be stored in cloud space of PhE (One Drive)	R, Stata	Econometric data, *dta, *xlsx,

A.3 Expected data sizes and volumes

Table 11: Datasets expected size.

Code	Type of data to be collected / name of the dataset	Record size and expected data volume (n. of records)
DS1.1	Cardiac ultrasound video recordings	N/A
DS1.2	Capsule endoscopy video recordings	Recordings from 60 patients
DS1.3	Cardiotocography variables and results, biometric data, medical history data	N/A
DS1.4	Coronary computed tomography angiography (CCTA) variables, biometric data, medical history data	N/A
DS2.1	Data related to hours spent by specialists.	N/A
DS2.2	Retrospective patient schedule data and precondition of the treatment	3 Patients anonymized
DS2.3	Data linked to radiotherapy machines (tumours treatment indication, maintenance, building location)	Data from 5 machines
DS2.4	PROMs/PREMs	N/A
DS2.5	Prospective patient clinical data	N/A
DS2.6	Patient personal data (address, preferences, ...)	20 patients
DS2.7	Retrospective EHR	3 Patients anonymized
DS2.8	Data from radiotherapy services, infrastructure. Patient's satisfaction	N/A
DS3.1	Smart home data: Consumption/production of instantaneous electricity and consumption logs, Human presence/access control, Devices' activation and connected loads	N/A
DS3.2	IMU data captured by iPrognosis smartphone application	N/A
DS3.3	Voice related time and spectral features captured by iPrognosis smartphone application	N/A
DS3.4	Key taps press and release timestamps captured by iPrognosis smartphone virtual keyboard	N/A
DS3.5	Angles of elbows and shoulders and scores achieved at the tests included in the iPrognosis iMAT tool	10 KB per user

Code	Type of data to be collected / name of the dataset	Record size and expected data volume (n. of records)
DS3.6	Results of rehabilitation sessions with Grador and care plans (editor and patient data)	Up to 10 MB per patient
DS4.1	Demographic patient data (age, gender, atrial size, etc.)	50
DS4.2	3D navigation system data (intracardiac signals, geometry)	50
DS5.1	Patient recordings and features extracted from interaction with patients	1 recording per interaction, assuming 5 minutes of interaction of up to 300 MB per recording. Feature files per recording are 15kb per 5 minutes for the handcrafted features and roughly 2.5GB MB per 5-10 minutes for fully low-level feature extraction
DS5.2	Biometric data (e.g., blood pressure and heart rate)	> 10 kb per FHIR resource
DS5.3	Retrospective electronic health records	Up to 100MB per patient
DS5.4	PREMs related to clinical staff (depends on T1.4)	Up to 2 MB per patient
DS5.5	PREMs related patients (e.g., PAM, SUS-SI/TAM, UEQ) and PROs	Up to 2 MB per patient
DS5.6	Datasets for facial expression and emotion recognition	> 10GB
DS5.7	Datasets for ASR and TTS in Slovenian	We estimate at least 20h of speech
DS5.8	Datasets for ASR and TTS in French	We estimate at least 20h of speech
DS5.9	EVA Corpus, data set of conversational expression	2GB per annotated 1h recording
DS5.10	Video recordings of third persons	0. No size occupied as the video frames are deleted as processed
DS5.11	Recording of a skeleton representation of the patient doing breathing exercises	Up to 200MB per patient per exercise set
DS5.12	Patient's facial information	N/A
DS6.1	Patient's physical, medical and cognitive status. Vital signs with sensors data.	At least 1 recording per day per patient for mental and physical status.

Code	Type of data to be collected / name of the dataset	Record size and expected data volume (n. of records)
		At least 1 recording per hour for each vital signal and patient with smartwatch.
DS7.1	Administrative data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	Around 1100 coronary angiograms including 350 CA with PCI performed in cathlab 3 and 4 between 01/11/2020 and 01/05/2021
DS7.2	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	Around 1100 coronary angiograms including 350 CA with PCI performed in cathlab 3 and 4 between 01/11/2020 and 01/05/2021
DS7.3	Coronary angiogram imaging data of patients who undergone a coronary angiogram/coronary intervention @ UZB	Around 1100 coronary angiograms including 350 CA with PCI performed in cathlab 3 and 4 between 01/11/2020 and 01/05/2021
DS7.4	Coronary physiology data of patients evaluated by a resting index measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary intervention @ UZB	Around 150 single point and 20 motorized pullbacks of physiological evaluations by FFR/iFR/RFR performed in cathlab 3 and 4 between 01/11/2020 and 01/05/2021
DS7.5	Intravascular imaging data of patient evaluated by either OCT or IVUS technique before and after a coronary intervention	Around 20 OCT and 10 IVUS performed in cathlab 3 and 4 between 01/11/2020 and 01/05/2021
DS7.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention	Around 300 CTA performed in patient referred for an invasive angiogram performed in cathlab 3 and 4 between 01/11/2020 and 01/05/2021
DS7.7	Full prospective UZB data set of administrative, clinical, coronary angiogram, coronary physiology, intravascular imaging and coronary CT data.	Validation study of the AI prototype effects on key performance indicators (hospital and health care productivity). Prospective cohort of 100 patients.
DS8.1	Image, gene, phenotype and pathology data for glioma patients	50 patients/year

Code	Type of data to be collected / name of the dataset	Record size and expected data volume (n. of records)
DS.O.1	All pilot data with Common KPIs (Economic & PROMs/PREMs)	Based on input provided by pilots

A.4 Data utility and identification

Table 12: Purpose and re-use of data (data utility) and identification.

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
DS1.1	Cardiac ultrasound video recordings	Development and evaluation of AI-assisted cardiology diagnosis tool	Partially identifiable since it will be interlinked with a specific patient. We could interlink extracted features and medical images/videos.	Access to pseudo-anonymised data is allowed with no restrictions, unless otherwise decided (e.g., in case of pending publication or patent). Request must be issued by each individual partner to AHEPA.
DS1.2	Capsule endoscopy video recordings	Development and evaluation of AI-assisted gastroenterology diagnosis tool	Partially identifiable since it will be interlinked with a specific patient. We could interlink extracted features and medical images/videos.	Access to pseudo-anonymised data is allowed with no restrictions, unless otherwise decided (e.g., in case of pending publication or patent). Request must be issued by each individual partner to AHEPA.
DS1.3	Cardiotocography variables and results, biometric data, medical history data	Development and evaluation of AI-assisted diagnosis tool	Possibly anonymous data, but interconnected with hospital data in case of needed patient identification	It is not available/accessible outside pilot #1 AUTH.
DS1.4	Coronary computed	Development and evaluation of AI-	Possibly anonymous data,	It is not available/accessible

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
	tomography angiography (CCTA) variables, biometric data, medical history data	assisted diagnosis tool	but interconnected with hospital data in case of needed patient identification	outside pilot #1 AHEPA/AUTH.
DS2.1	Data related to hours spent by specialists.	Create patient's schedule and assess patient's satisfaction	Anonymized	None
DS2.2	Retrospective patient schedule data and precondition of the treatment	Develop AI model and digital twin.	Anonymized	Under the conditions of the CA as assessed in D8.3 SELP Impact Assessment.
DS2.3	Data linked to radiotherapy machines (tumours treatment indication, maintenance, building location)	Develop AI model and digital twin Validation.	Fully identifiable	None
DS2.4	PROMs/PREMs	Validation (chatbot)	Anonymized	PROM and PREM questionnaire no restriction, however how chatbot data will be shared with partners
DS2.5	Prospective patient clinical data	Validation	Fully identifiable, should not be shared.	Private patient information is not available/accessible outside pilot #2.

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
DS2.6	Patient personal data (address, preferences, ...)	Validation	Fully identifiable, should not be shared.	Private patient information is not available/accessible outside pilot #2.
DS2.7	Retrospective EHR	Develop AI model and digital twin	Anonymized	Under the conditions of the CA as assessed in D8.3 SELP Impact Assessment.
DS2.8	Data from radiotherapy services, infrastructure. Patient's satisfaction	Development algorithm for Optimization Scheduler.	Fully identifiable, should not be shared.	Private patient information is not available/accessible outside pilot #2. What we can share in extracted is features + pattern identification outcome since this is anonymized and will be also exploited in publications
DS3.1	Smart home data: Consumption/production of instantaneous electricity and consumption logs, Human presence/access control, Devices' activation and connected loads	Monitor of the environment	Fully identifiable since it will be interlinked with a specific location	Data belongs to the owner of the location where the home automation devices are installed. The owner defines access to the data

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
DS3.2	IMU data captured by iPrognosis smartphone application	Remote monitoring of people with Parkinson's disease (PwP) via the iPrognosis application	Pseudo-anonymised	Access to pseudo-anonymised data is allowed with no restrictions, unless otherwise decided (e.g., in case of pending publication or patent). Data access roles will be granted.
DS3.3	Voice-related time and spectral features captured by iPrognosis smartphone application	Remote monitoring of PwP via the iPrognosis application	Pseudo-anonymised	Access to pseudo-anonymised data is allowed with no restrictions, unless otherwise decided (e.g., in case of pending publication or patent). Data access roles will be granted.
DS3.4	Key taps press and release timestamps captured by iPrognosis smartphone virtual keyboard	Remote monitoring of PwP via the iPrognosis application	Pseudo-anonymised	Access to pseudo-anonymised data is allowed with no restrictions, unless otherwise decided (e.g., in case of pending publication or patent). Data access roles will be granted.
DS3.5	Angles of elbows and shoulders and scores achieved at the tests included in the iPrognosis iMAT tool	Remote assessment of the motor status of PwP via the iPrognosis iMAT application	Pseudo-anonymised	Access to pseudo-anonymised data is allowed with no restrictions, unless otherwise decided (e.g., in case of pending publication or patent). Data access roles will be granted.

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
DS3.6	Results of rehabilitation sessions with Grador and care plans (editor and patient data)	Develop AI intervention model and Validation	Pseudo-anonymised	Access to pseudo-anonymised data is allowed with no restrictions, unless otherwise decided (e.g., in case of pending publication or patent). Data access roles will be granted.
DS4.1	Demographic patient data (age, gender, atrial size, etc.)	Population characterization	Pseudoanonymized	Data will not be available for no other person outside pilot #4.
DS4.2	3D navigation system data (intracardiac signals, geometry)	Data source for AI analysis	Pseudoanonymized	Data will not be available for no other person outside pilot #4.
DS5.1	Patient recordings and features extracted from interaction with patients	Support for Spoken Language Interaction, Classification of psychological distress (symptoms of depression) and action recognition	Fully identifiable, should not be shared. We can think of sharing extracted features	It is not available/accessible outside pilot #5 UM/UKCM. Special DTA is signed with UKCM as data owner and UM as data processor prior to the trial. <i>What we can share is extracted features + classification outcome since this is anonymized and will be also exploited in publications</i>

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
DS5.2	Biometric data (e.g., blood pressure and heart rate)	Impact of PGHD on CCDS Efficiency and improved management of clinical parameters	Partially identifiable since it will be interlinked with a specific patient. We could interlink extracted features and biometric data as a data set.	It is not available/accessible outside pilot #5 UM/UKCM. Special DTA with UKCM as data owner and UM as data processor will be prepared and signed prior to the trial. We will consider, however, creating datasets, such as, correlating interaction features and monitored biomarkers with moods/emotions/p psychological distress, to be shared openly for research
DS5.3	Retrospective electronic health records	Impact of CPOE on clinical routine, CCDS Efficiency and improved management of clinical parameters	Fully identifiable since it will be interlinked with a specific patient. Cannot be shared.	It is not available/accessible outside pilot #5 UM/UKCM. DTA with UKCM as data owner and UM as data processor will be prepared and signed prior to the trial.
DS5.4	PREMs related to clinical staff (depends on T1.4)	Impact of Social robotics system on various aspects of clinical workflow, including staff satisfaction and workload	Anonymized, however, a code will be provided for comparison prior to, during and after the intervention	Statistical data and cohorts are available to be shared with the consortium and wider, for research

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
DS5.5	PREMs related patients (e.g., PAM, SUS-SI/TAM, UEQ) and PROs	Impact of Social robotics system on quality of care	Anonymized, however, a code will be provided for comparison prior to, during and after the intervention	Statistical data and cohorts available to be shared with the consortium, and wider, for research
DS5.6	Datasets for facial expression and emotion recognition	Development of sensing AI to collect and classify symptoms of depression from facial expressions, speech and text.	Public dataset, please consider the individual licenses and restrictions	The data is openly available, request must be issued by each individual partner to the specific owner of the data set
DS5.7	Datasets for ASR and TTS in Slovenian	Support for Spoken Language Interaction	Proprietary dataset owned by UM. It is Background we bring into the project.	The data is not publicly open. Access can be granted on an individual basis (bilateral agreements which may include charges)
DS5.8	Datasets for ASR and TTS in French	Support for Spoken Language Interaction	Proprietary dataset owned by UM and public datasets. It is background we bring into the project.	Access to public datasets must be managed by individual partners, access to UM's closed datasets can be granted on an individual basis (bilateral agreements which may include charges)
DS5.9	EVA Corpus, data set of conversational expression	Support for Spoken Language Interaction	Proprietary dataset owned by UM. It is background we bring into the project.	Access via CLARIN.SI repository

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
			It is already publicly available.	
DS5.10	Video recordings of third persons	The recorded data is sent to a model to determine if a patient is making a gesture to call a doctor or if somebody falls	Yes, and the video frames are deleted as they are processed	The data is not available as it would affect the patient's privacy
DS5.11	Recording of a skeleton representation of the patient doing breathing exercises	Testing the monitoring of the breathing exercise	Only the skeleton is saved, no information of the patient is available	UM has access to the edge server where the data is stored
DS5.12	Patient's facial information	N/A	N/A	N/A
DS6.1	Patient's physical, medical and cognitive status. Vital signs with sensors data.	Support for identification of abnormal pattern recognition.	Fully identifiable, should not be shared.	Private patient information is not available/accessible outside pilot #6. What we can share in extracted is features + pattern identification outcome since this is anonymized and will be also exploited in publications
DS7.1	Administrative data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	Retrospective dataset starting from 11/2020 used to develop an AI prototype to alleviate the administrative burden in the	Pseudo anonymization. Data will be anonymized before sending to third party (Philips) but the encrypting key will be stored by	Data will be not available/accessible outside pilot #7

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
		interventional suite by an automatic procedure tracking.	the owner of the data (UZB, VUB)	
DS7.2	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	Retrospective dataset starting from 11/2020 used to develop an AI prototype to alleviate the administrative burden in the interventional suite by an automatic procedure tracking.	Pseudo anonymization. Data will be anonymized before sending to third party (Philips) but the encrypting key will be stored by the owner of the data (UZB, VUB)	Data will not be available/accessible outside pilot #7
DS7.3	Coronary angiogram imaging data of patients who undergone a coronary angiogram/coronary intervention @ UZB	Retrospective dataset starting from 11/2020 used to develop an AI prototype to ensure an automatic logging a smart reporting of both imaging and patient X-ray dosimetry and a facilitated coronary angiogram interpretation by calculation of severity scores	Pseudo anonymization. Data will be anonymized before sending to third party (Philips) but the encrypting key will be stored by the owner of the data (UZB, VUB)	Data will not be available/accessible outside pilot #7
DS7.4	Coronary physiology data of patients evaluated by a resting index	Retrospective dataset starting from 11/2020 used to develop an AI prototype	Pseudo anonymization. Data will be anonymized before sending to third	Data will not be available/accessible outside pilot #7

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
	measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary intervention @ UZB	to ensure a clinical decision support.	party (Philips) but the encrypting key will be stored by the owner of the data (UZB, VUB)	
DS7.5	Intravascular imaging data of patient evaluated by either OCT or IVUS technique before and after a coronary intervention	Retrospective dataset starting from 11/2020 used to develop an AI prototype to ensure a clinical decision support.	Pseudo anonymization. Data will be anonymized before sending to third party (Philips) but the encrypting key will be stored by the owner of the data (UZB, VUB)	Data will not be available/accessible outside pilot #7
DS7.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention	Retrospective dataset starting from 11/2020 used to develop an AI prototype to ensure a clinical decision support.	Pseudo anonymization. Data will be anonymized before sending to third party (Philips) but the encrypting key will be stored by the owner of the data (UZB, VUB)	Data will not be available/accessible outside pilot #7
DS7.7	Full prospective UZB data set of administrative, clinical, coronary angiogram,	Prospective validation of AI prototype to ensure automatic procedure tracking, an	Non anonymized data will be treated on site (UZB, VUB) by AI smart cathlab Philips prototype	Data will not be available/accessible outside pilot #7

Code	Type of data to be collected / name of the dataset	Purpose and re-use of the data (data utility)	Collected information will be identifiable?	How will the data be made accessible to other partners in consortium? Are there any restrictions?
	coronary physiology, intravascular imaging and coronary CT data.	automatic logging, a smart reporting and help to clinical decision support.	Data required to be sent for an external analysis by a third party will be anonymized but the encrypting key will be stored by the owner of the data (UZB, VUB)	
DS8.1	Image, gene, phenotype and pathology data for glioma patients	Dataset for highlighting relationships between different data types and identifying tumour type	The data will be identifiable and will not be shared as is, but procedures to make them (partially) available will be pursued	The data is restricted, we will pursue ways to make the data (partially) available to the consortium via pseudo anonymization; work is ongoing with UZ Brussel.
DS.O.1	All pilot data with Common KPIs (Economic & PROMs/PREMs)	To identify the efficiency in terms of cost-effectiveness of the new AI technologies vs. the previous state of the art of all pilot data	Patient level data will be anonymized, only the intervention will be known to the statistician/health economist.	The primary data is the property of each pilot, and the results of the economic/PROM/PREM analysis will be available to all partners. Coding of the economic part is the property of PhE.

A.5 Data findability, metadata

Table 13: Making data findable, metadata.

Code	Type of data to be collected / name of the dataset	Data identification and versioning	Naming conventions	Search keywords for re-use	Metadata
DS1.1	Cardiac ultrasound video recordings	N/A	N/A	N/A	N/A
DS1.2	Capsule endoscopy video recordings	N/A	N/A	N/A	N/A
DS1.3	Cardiotocography variables and results, biometric data, medical history data	N/A	N/A	N/A	N/A
DS1.4	Coronary computed tomography angiography (CCTA) variables, biometric data, medical history data	N/A	N/A	N/A	N/A
DS2.1	Data related to hours spent by specialists.	By date, by system (legacy vs AI). TBC whether this can be published or not.	N/A	Radiotherapy, throughput, scheduling, satisfaction	N/A
DS2.2	Retrospective patient schedule data and precondition of the treatment	Sample dataset.	N/A	N/A	N/A
DS2.3	Data linked to radiotherapy machines (tumours treatment indication,	By site	N/A	N/A	N/A

Code	Type of data to be collected / name of the dataset	Data identification and versioning	Naming conventions	Search keywords for re-use	Metadata
	maintenance, building location)				
DS2.4	PROMs/PREMs	Those data should not be persisted/shared beyond the project's scope.	N/A	N/A	N/A
DS2.5	Prospective patient clinical data	Internal use only	N/A	N/A	N/A
DS2.6	Patient personal data (address, preferences, ...)	Internal use only	N/A	N/A	N/A
DS2.7	Retrospective EHR	Sample dataset.	N/A	N/A	N/A
DS2.8	Data from radiotherapy services, infrastructure. Patient's satisfaction	N/A	N/A	N/A	N/A
DS3.1	Smart home data: Consumption/production of instantaneous electricity and consumption logs, Human presence/access control, Devices' activation and connected loads	N/A	N/A	N/A	N/A
DS3.2	IMU data captured by iPrognosis	Still TBD	Still TBD	Still TBD	Still TBD

Code	Type of data to be collected / name of the dataset	Data identification and versioning	Naming conventions	Search keywords for re-use	Metadata
	smartphone application				
DS3.3	Voice-related time and spectral features captured by iPrognosis smartphone application	Still TBD	Still TBD	Still TBD	Still TBD
DS3.4	Key taps press and release timestamps captured by iPrognosis smartphone virtual keyboard	Still TBD	Still TBD	Still TBD	Still TBD
DS3.5	Angles of elbows and shoulders and scores achieved at the tests included in the iPrognosis iMAT tool	N/A	N/A	N/A	N/A
DS3.6	Results of rehabilitation sessions with Grador and care plans (editor and patient data)	by date, by cognitive area, by patient's id	N/A	N/A	N/A
DS4.1	Demographic patient data (age, gender, atrial size, etc.)	N/A	N/A	N/A	N/A
DS4.2	3D navigation system data (intracardiac signals, geometry)	N/A	N/A	N/A	N/A
DS5.1	Patient recordings and features extracted from	Patient, and Department, ID (internal hospital ID),	Action Units, Acoustic units and concepts, Linguistic units	Depressive or anxiety disorder	Depending on the open data repository

Code	Type of data to be collected / name of the dataset	Data identification and versioning	Naming conventions	Search keywords for re-use	Metadata
	interaction with patients	Date-Time, Unit ID, Room ID	and concerts, XPOS for morphology, ICD-10 for disease classification		
DS5.2	Biometric data (e.g. blood pressure and heart rate)	Patient, and Department, ID (internal hospital ID), Date-Time, Monitor ID	LOINC, ICD-10	LOINC Features, ICD-10 disease, procedure	Depending on the open data repository
DS5.3	Retrospective electronic health records	N/A	N/A	N/A	N/A
DS5.4	PREMs related to clinical staff (depends on T1.4)	Staff-ID, Department ID, Date-Time	LOINC, ICD-10 or custom if the tool is not supported	N/A	N/A
DS5.5	PREMs related patients (e.g. PAM, SUS-SI/TAM, UEQ) and PROs	Patient, and Department, ID (internal hospital ID), Date-Time	LOINC, ICD-10 or custom if the tool is not supported	N/A	N/A
DS5.6	Datasets for facial expression and emotion recognition	N/A	N/A	N/A	N/A
DS5.7	Datasets for ASR and TTS in Slovenian	N/A	N/A	N/A	N/A
DS5.8	Datasets for ASR and TTS in French	N/A	N/A	N/A	N/A
DS5.9	EVA Corpus, data set of conversational expression	N/A	N/A	N/A	N/A

Code	Type of data to be collected / name of the dataset	Data identification and versioning	Naming conventions	Search keywords for re-use	Metadata
DS5.10	Video recording of third persons	N/A	N/A	N/A	N/A
DS5.11	Recording of a skeleton representation of the patient doing breathing exercises	N/A	N/A	N/A	N/A
DS5.12	Patient's facial information	By patient name or ID	N/A	N/A	N/A
DS6.1	Patient's physical, medical and cognitive status. Vital signs with sensors data.	Age, Body Height, General status. Heartbeat. Oxygen, Blood pressure, Body temperature, Urine, Glucose, Body weight, Stool.	LOINC	LOINC	N/A
DS7.1	Administrative data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	N/A	N/A	N/A	N/A
DS7.2	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	N/A	N/A	N/A	N/A
DS7.3	Coronary angiogram imaging data of patients who undergone a coronary angiogram/coron	N/A	N/A	N/A	N/A

Code	Type of data to be collected / name of the dataset	Data identification and versioning	Naming conventions	Search keywords for re-use	Metadata
	ary intervention @ UZB				
DS7.4	Coronary physiology data of patients evaluated by a resting index measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary intervention @ UZB	N/A	N/A	N/A	N/A
DS7.5	Intravascular imaging data of patient evaluated by either OCT or IVUS technique before and after a coronary intervention	N/A	N/A	N/A	N/A
DS7.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention	N/A	N/A	N/A	N/A
DS7.7	Full prospective UZB data set of administrative, clinical, coronary angiogram, coronary physiology, intravascular	N/A	N/A	N/A	N/A

Code	Type of data to be collected / name of the dataset	Data identification and versioning	Naming conventions	Search keywords for re-use	Metadata
	imaging and coronary CT data.				
DS8.1	Image, gene, phenotype and pathology data for glioma patients	Unique IDs via patient numbers, versioning for local systems exists.	N/A	N/A	N/A

A.6 Data open accessibility

Table 14: Making data open accessible.

Code	Type of data to be collected / name of the dataset	Dataset made available openly?	Which open repository (or other available medium)?	Licenses, access identification, restrictions
DS1.1	Cardiac ultrasound video recordings	YES (unless otherwise decided due to IPRs)	Zenodo	N/A
DS1.2	Capsule endoscopy video recordings	YES (unless otherwise decided due to IPRs)	Zenodo	N/A
DS1.3	Cardiotocography variables and results, biometric data, medical history data	N/A	N/A	N/A
DS1.4	Coronary computed tomography angiography (CCTA) variables, biometric data, medical history data	N/A	N/A	N/A
DS2.1	Data related to hours spent by specialists.	Yes	N/A	N/A

Code	Type of data to be collected / name of the dataset	Dataset made available openly?	Which open repository (or other available medium)?	Licenses, access identification, restrictions
DS2.2	Retrospective patient schedule data and precondition of the treatment	No	N/A	N/A
DS2.3	Data linked to radiotherapy machines (tumours treatment indication, maintenance, building location)	No	N/A	N/A
DS2.4	PROMs/PREMs	No	N/A	N/A
DS2.5	Prospective patient clinical data	No	N/A	N/A
DS2.6	Patient personal data (address, preferences, ...)	No	N/A	N/A
DS2.7	Retrospective EHR	No	N/A	N/A
DS2.8	Data from radiotherapy services, infrastructure. Patient's satisfaction	No	N/A	N/A
DS3.1	Smart home data: Consumption/production of instantaneous electricity and consumption logs, Human presence/access control, Devices	Data belongs to the owner of the location where the home automation devices are installed. The owner defines access to the data.	None	The owner defines access to the data.

Code	Type of data to be collected / name of the dataset	Dataset made available openly?	Which open repository (or other available medium)?	Licenses, access identification, restrictions
	activation and connected loads			
DS3.2	IMU data captured by iPrognosis smartphone application	YES (unless otherwise decided due to IPRs)	Zenode	Still TBD
DS3.3	Voice-related time and spectral features captured by iPrognosis smartphone application	YES (unless otherwise decided due to IPRs)	Zenode	Still TBD
DS3.4	Key taps press and release timestamps captured by iPrognosis smartphone virtual keyboard	YES (unless otherwise decided due to IPRs)	Zenode	Still TBD
DS3.5	Angles of elbows and shoulders and scores achieved at the tests included in the iPrognosis iMAT tool	N/A	N/A	N/A
DS3.6	Results of rehabilitation sessions with Grador and care plans (editor and patient data)	YES (but considering the limitations related to the IPR/Consortium Agreement)	N/A	N/A
DS4.1	Demographic patient data (age, gender, atrial size, etc.)	No	N/A	N/A
DS4.2	3D navigation system data (intracardiac	No	N/A	N/A

Code	Type of data to be collected / name of the dataset	Dataset made available openly?	Which open repository (or other available medium)?	Licenses, access identification, restrictions
	signals, geometry)			
DS5.1	Patient recordings and features extracted from interaction with patients	NO for recording, YES for features and patient is replaced by a random number during anonymization	EOSC, Zenodo	Preferred open for research (e.g. CC-BY-NC)
DS5.2	Biometric data (e.g., blood pressure and heart rate)	YES for cohorts	EOSC, Zenodo	Preferred open for research
DS5.3	Retrospective electronic health records	NO	N/A	N/A
DS5.4	PREMs related to clinical staff (depends on T1.4)	YES (but considering anonymization)	Zenodo	Preferred open for research (e.g. CC-BY-NC)
DS5.5	PREMs related patients (e.g., PAM, SUS-SI/TAM, UEQ) and PROs	YES (but considering anonymization)	Zenodo	Preferred open for research (e.g. CC-BY-NC)
DS5.6	Datasets for facial expression and emotion recognition	YES	e.g., jaffe, FER-2013, MMI, Cohn-Kanade, RaFD, FERG, EMOTIC, Affect data, SemEval-2017 Task 4, DailyDialog, The MPLab GENKI Database, FFECTIVA-MIT Facial Expression	Licenses are granted on individual requests, not handled by UM but the owners of data sets

Code	Type of data to be collected / name of the dataset	Dataset made available openly?	Which open repository (or other available medium)?	Licenses, access identification, restrictions
			Dataset (AM-FED), Grounded Emotions, Reldi, etc.	
DS5.7	Datasets for ASR and TTS in Slovenian	NO		Access can be granted on individual basis (bilateral agreements which may include charges)
DS5.8	Datasets for ASR and TTS in French	NO		Access can be granted on individual basis (bilateral agreements which may include charges)
DS5.9	EVA Corpus, data set of conversational expression	YES	Clarin.SI	CC-BY 4.0 License
DS5.10	Video recordings of third persons	NO	N/A	N/A
DS5.11	Recording of a skeleton representation of the patient doing breathing exercises	NO	N/A	N/A
DS5.12	Patient's facial information	NO	N/A	N/A
DS6.1	Patient's physical, medical and cognitive status. Vital signs with sensors data.	NO	N/A	N/A
DS7.1	Administrative data of patients scheduled for a	N/A	N/A	N/A

Code	Type of data to be collected / name of the dataset	Dataset made available openly?	Which open repository (or other available medium)?	Licenses, access identification, restrictions
	coronary angiogram/coronary intervention @ UZB			
DS7.2	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	N/A	N/A	N/A
DS7.3	Coronary angiogram imaging data of patients who undergone a coronary angiogram/coronary intervention @ UZB	N/A	N/A	N/A
DS7.4	Coronary physiology data of patients evaluated by a resting index measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary intervention @ UZB	N/A	N/A	N/A
DS7.5	Intravascular imaging data of patient evaluated by either OCT or IVUS technique	N/A	N/A	N/A

Code	Type of data to be collected / name of the dataset	Dataset made available openly?	Which open repository (or other available medium)?	Licenses, access identification, restrictions
	before and after a coronary intervention			
DS7.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention	N/A	N/A	N/A
DS7.7	Full prospective UZB data set of administrative, clinical, coronary angiogram, coronary physiology, intravascular imaging and coronary CT data.	N/A	N/A	N/A
DS8.1	Image, gene, phenotype and pathology data for glioma patients	No	Not relevant	N/A
DS.O.1	All pilot data with Common KPIs (Economic & PROMs/PREMs)	Applicable for pilot specifics	Applicable for pilot specifics	Applicable for pilot specifics

A.7 Data interoperability

Table 15: Data interoperability.

Code	Type of data to be collected / name of the dataset	Metadata vocabularies, ontologies, standards and methodologies for data interoperability
DS1.1	Cardiac ultrasound video recordings	HL7
DS1.2	Capsule endoscopy video recordings	HL7
DS1.3	Cardiotocography variables and results, biometric data, medical history data	N/A
DS1.4	Coronary computed tomography angiography (CCTA) variables, biometric data, medical history data	N/A
DS2.1	Data related to hours spent by specialists.	N/A
DS2.2	Retrospective patient schedule data and precondition of the treatment	JSON
DS2.3	Data linked to radiotherapy machines (tumours treatment indication, maintenance, building location)	JSON
DS2.4	PROMs/PREMs	CSV, HL7 FHIR
DS2.5	Prospective patient clinical data	HL7 FHIR
DS2.6	Patient personal data (address, preferences, ...)	N/A
DS2.7	Retrospective EHR	JSON
DS2.8	Data from radiotherapy services, infrastructure. Patient's satisfaction	RDBMS, HL7 FHIR
DS3.1	Smart home data: Consumption/production of instantaneous electricity and consumption logs, Human presence/access control, Devices' activation and connected loads	N/A
DS3.2	IMU data captured by iPrognosis smartphone application	HL7
DS3.3	Voice related time and spectral features captured by iPrognosis smartphone application	HL7

Code	Type of data to be collected / name of the dataset	Metadata vocabularies, ontologies, standards and methodologies for data interoperability
DS3.4	Key taps press and release timestamps captured by iPrognosis smartphone virtual keyboard	HL7
DS3.5	Angles of elbows and shoulders and scores achieved at the tests included in the iPrognosis iMAT tool	FHIR
DS3.6	Results of rehabilitation sessions with Grador and care plans (editor and patient data)	FHIR v4
DS4.1	Demographic patient data (age, gender, atrial size, etc.)	N/A
DS4.2	3D navigation system data (intracardiac signals, geometry)	N/A
DS5.1	Patient recordings and features extracted from interaction with patients	FHIR v4
DS5.2	Biometric data (e.g., blood pressure and heart rate)	FHIR v4
DS5.3	Retrospective electronic health records	N/A
DS5.4	PREMs related to clinical staff (depends on T1.4)	FHIR v4
DS5.5	PREMs related patients (e.g., PAM, SUS-SI/TAM, UEQ) and PROs	FHIR v4
DS5.6	Datasets for facial expression and emotion recognition	N/A
DS5.7	Datasets for ASR and TTS in Slovenian	
DS5.8	Datasets for ASR and TTS in French	N/A
DS5.9	EVA Corpus, data set of conversational expression	N/A
DS5.10	Video recordings of third persons	N/A
DS5.11	Recording of a skeleton representation of the patient doing breathing exercises	N/A
DS5.12	Patient's facial information	FHIR v4
DS6.1	Patient's physical, medical and cognitive status.	RDBMS, HL7 FHIR

Code	Type of data to be collected / name of the dataset	Metadata vocabularies, ontologies, standards and methodologies for data interoperability
	Vital signs with sensors data.	
DS7.1	Administrative data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	N/A
DS7.2	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	N/A
DS7.3	Coronary angiogram imaging data of patients who undergone a coronary angiogram/coronary intervention @ UZB	N/A
DS7.4	Coronary physiology data of patients evaluated by a resting index measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary intervention @ UZB	N/A
DS7.5	Intravascular imaging data of patient evaluated by either OCT or IVUS technique before and after a coronary intervention	N/A
DS7.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention	N/A
DS7.7	Full prospective UZB data set of administrative, clinical, coronary angiogram, coronary physiology, intravascular imaging and coronary CT data.	N/A
DS8.1	Image, gene, phenotype and pathology data for glioma patients	N/A
DS.O.1	All pilot data with Common KPIs (Economic & PROMs/PREMs)	Applicable for pilot specifics

A.8 Data licensing, availability and usability

Table 16: Data licensing, availability and usability by third parties.

Code	Type of data to be collected / name of the dataset	Availability and usability of data by third parties. Licensing
DS1.1	Cardiac ultrasound video recordings	N/A
DS1.2	Capsule endoscopy video recordings	N/A
DS1.3	Cardiotocography variables and results, biometric data, medical history data	N/A
DS1.4	Coronary computed tomography angiography (CCTA) variables, biometric data, medical history data	N/A
DS2.1	Data related to hours spent by specialists.	Consortium agreement
DS2.2	Retrospective patient schedule data and precondition of the treatment	Consortium agreement
DS2.3	Data linked to radiotherapy machines (tumours treatment indication, maintenance, building location)	Consortium agreement
DS2.4	PROMs/PREMs	Internal use only
DS2.5	Prospective patient clinical data	Internal use only
DS2.6	Patient personal data (address, preferences, ...)	Internal use only
DS2.7	Retrospective EHR	Consortium agreement
DS2.8	Data from radiotherapy services, infrastructure. Patient's satisfaction.	N/A
DS3.1	Smart home data: Consumption/production of instantaneous electricity and consumption logs, Human presence/access control, Devices' activation and connected loads	N/A
DS3.2	IMU data captured by iPrognosis smartphone application	Still TBD
DS3.3	Voice related time and spectral features captured by iPrognosis smartphone application	Still TBD

Code	Type of data to be collected / name of the dataset	Availability and usability of data by third parties. Licensing
DS3.4	Key taps press and release timestamps captured by iPrognosis smartphone virtual keyboard	Still TBD
DS3.5	Angles of elbows and shoulders and scores achieved at the tests included in the iPrognosis iMAT tool	N/A
DS3.6	Results of rehabilitation sessions with Grador and care plans (editor and patient data)	Consortium agreement
DS4.1	Demographic patient data (age, gender, atrial size, etc.)	N/A
DS4.2	3D navigation system data (intracardiac signals, geometry)	N/A
DS5.1	Patient recordings and features extracted from interaction with patients	N/A for videos, for features and cohorts CC BY-NC 4.0 is preferred
DS5.2	Biometric data (e.g., blood pressure and heart rate)	CC BY-NC 4.0 is preferred
DS5.3	Retrospective electronic health records	N/A
DS5.4	PREMs related to clinical staff (depends on T1.4)	CC BY-NC 4.0 is preferred
DS5.5	PREMs related patients (e.g., PAM, SUS-SI/TAM, UEQ) and PROs	CC BY-NC 4.0 is preferred
DS5.6	Datasets for facial expression and emotion recognition	Licenses are granted on individual requests, not handled by UM but the owners of data sets
DS5.7	Datasets for ASR and TTS in Slovenian	Access can be granted on individual basis (bilateral agreements which may include charges)
DS5.8	Datasets for ASR and TTS in French	Access can be granted on individual basis (bilateral agreements which may include charges)
DS5.9	EVA Corpus, data set of conversational expression	CC-BY 4.0 License
DS5.10	Video recordings of third persons	N/A
DS5.11	Recording of a skeleton representation of the patient doing breathing exercises	N/A

Code	Type of data to be collected / name of the dataset	Availability and usability of data by third parties. Licensing
DS5.12	Patient's facial information	N/A
DS6.1	Patient's physical, medical and cognitive status. Vital signs with sensors data.	N/A
DS7.1	Administrative data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	N/A
DS7.2	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	N/A
DS7.3	Coronary angiogram imaging data of patients who undergone a coronary angiogram/coronary intervention @ UZB	N/A
DS7.4	Coronary physiology data of patients evaluated by a resting index measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary intervention @ UZB	N/A
DS7.5	Intravascular imaging data of patient evaluated by either OCT or IVUS technique before and after a coronary intervention	N/A
DS7.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention	N/A
DS7.7	Full prospective UZB data set of administrative, clinical, coronary angiogram, coronary physiology, intravascular imaging and coronary CT data.	N/A
DS8.1	Image, gene, phenotype and pathology data for glioma patients	N/A
DS.O.1	All pilot data with Common KPIs (Economic & PROMs/PREMs)	Applicable for pilot specifics

A.9 Data quality assurance

Table 17: Data quality assurance.

Pilot nr	Task nr	Type of analysis	Will you work according to specific protocol(s)? If yes, which one(s)?	Who will create the statistical analysis plan? (partner short name; person name; email)	How will data transformation & analysis be verified?
1	5.2	N/A	N/A	AUTH; person(s) N/A	Peer-review
2	3.2	Protocols published for the pilot 2 Clinical studies	Verification and validation plan	CHU de Liège, Patrick Duflot	System test, Unit test, Manual test
2	5.2	N/A	Prospective study protocol is under preparation	CHU de Liège, Patrick Duflot	As defined in clinical study protocol
2	5.3	As defined by PhE	As defined by PhE	As defined by PhE	As defined by PhE
3	3.3	N/A	N/A	N/A	N/A
4	5.2	N/A	N/A	N/A	N/A
5	3.5	Protocols published for the pilot 5 Clinical studies	Verification and validation plan, Data Monitoring Plan	UM, GC	Peer-review, Manual test
6	3.5	Protocols shared by PhE	Protocols shared by PhE	ITCL	INTRAS as define in the study protocol approved.
7	3.6	Administrative data of patients scheduled for a coronary angiogram/coronary	Retrospective study protocol (registry) is under preparation	Philips Image Guided Therapy Systems, Netherlands	N/A

Pilot nr	Task nr	Type of analysis	Will you work according to specific protocol(s)? If yes, which one(s)?	Who will create the statistical analysis plan? (partner short name; person name; email)	How will data transformation & analysis be verified?
		intervention @ UZB			
7	3.6	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	Retrospective study protocol (registry) is under preparation	Philips Image Guided Therapy Systems, Netherlands	N/A
7	3.6	Coronary angiogram imaging data of patients who undergone a coronary angiogram/coronary intervention @ UZB	Retrospective study protocol (registry) is under preparation	Philips Image Guided Therapy Systems, Netherlands	N/A
7	3.6	Coronary physiology data of patients evaluated by a resting index measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary	Retrospective study protocol (registry) is under preparation	Philips Image Guided Therapy Systems, Netherlands	N/A

Pilot nr	Task nr	Type of analysis	Will you work according to specific protocol(s)? If yes, which one(s)?	Who will create the statistical analysis plan? (partner short name; person name; email)	How will data transformation & analysis be verified?
		intervention @ UZB			
7	3.6	Intravascular imaging data of patient evaluated by either OCT or IVUS technique before and after a coronary intervention@ UZB	Retrospective study protocol (registry) is under preparation	Philips Image Guided Therapy Systems, Netherlands	N/A
7	3.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention@ UZB	Retrospective study protocol (registry) is under preparation	Philips Image Guided Therapy Systems, Netherlands	N/A
7	3.6	Full prospective UZB data set of administrative, clinical, coronary angiogram, coronary physiology, intravascular imaging and	Prospective study protocol (comparative observational study) is under preparation	Primary investigator UZB, VUB	Comparative analysis of different key performance indicators and patient reported outcome and experience measures (PREMS and PROMS) evolution

Pilot nr	Task nr	Type of analysis	Will you work according to specific protocol(s)? If yes, which one(s)?	Who will create the statistical analysis plan? (partner short name; person name; email)	How will data transformation & analysis be verified?
		coronary CT of data.			before (control period) and after implantation of AI smart cathlab prototype (study period)
8	N/A	Image, gene, phenotype and pathology data for glioma patients	To be determined in collaboration with UZ Brussel ICT	VUB	Local testing, feedback from specialists, eventual peer review
PhE	5.3	Based on Quality control Standard Operating Procedure (SOP) followed by PhE for all projects/deliverables	Protocol & CRF has been prepared by PhE and will be shared with all pilots in order to follow the same	PhE project team / Eugena Stamuli, Declan O'Byrne, Magda Chatzikou	N/A

A.10 Data cleansing, transforming and analysing

Table 18: Data cleansing, transforming and analysing.

Code	Type of data to be collected / name of the dataset	Type of data cleaning needed (e.g., correct data types, remove duplicates, add missing info...)	Person responsible for data cleaning (partner short name; person name; email)	Type of data transformation/analysis (e.g., normalization, discretization, ...)	Software/tools used for cleaning, transform, and analyse	Where/by whom will the analysis be conducted?	Standards followed for code development / access and re-use
DS1.1	Cardiac ultrasound video recordings	N/A	N/A	N/A	N/A	N/A	N/A
DS1.2	Capsule endoscopy video recordings	N/A	N/A	N/A	N/A	N/A	N/A
DS1.3	Cardiotocography variables and results, biometric data, medical history data	N/A	N/A	N/A	N/A	N/A	N/A
DS1.4	Coronary computed tomography angiography (CCTA) variables, biometric data, medical history data	N/A	N/A	N/A	N/A	N/A	N/A
DS2.1	Data related to hours spent by specialists.	N/A	N/A	N/A	N/A	N/A	N/A
DS2.2	Retrospective patient schedule data and precondition of the treatment	Extraction and alignment from subsystems.	CHUL	Mapping to FHIR (responsible TMA)	None	In HosmartAI platform	
DS2.3	Data linked to radiotherapy machines (tumours treatment indication,	Extraction and alignment from subsystems.	CHUL	Mapping to FHIR (responsible TMA)	None	In HosmartAI platform	

Code	Type of data to be collected / name of the dataset	Type of data cleaning needed (e.g., correct data types, remove duplicates, add missing info...)	Person responsible for data cleaning (partner short name; person name; email)	Type of data transformation/analysis (e.g., normalization, discretization, ...)	Software/tools used for cleaning, transform, and analyse	Where/by whom will the analysis be conducted?	Standards followed for code development / access and re-use
	maintenance, building location)						
DS2.4	PROMs/PREMs	N/A	ITCL	N/A	N/A	N/A	N/A
DS2.5	Prospective patient clinical data	N/A	N/A	N/A	N/A	N/A	N/A
DS2.6	Patient personal data (address, preferences, ...)	N/A	N/A	N/A	N/A	N/A	N/A
DS2.7	Retrospective EHR	Extraction and alignment from subsystems.	CHUL	Mapping to FHIR (responsible TMA)	N/A	N/A	
DS2.8	Data from radiotherapy services, infrastructure. Patient's satisfaction	N/A	N/A	N/A	N/A	N/A	N/A
DS3.1	Smart home data: Consumption/production of instantaneous electricity and consumption logs, Human presence/access control, Devices' activation and connected loads	Outlier detection, inference on missing data	VIMAR	Normalization	N/A	N/A	N/A
DS3.2	IMU data captured by iPrognosis smartphone application	Still TBD	Still TBD	Still TBD	Still TBD	Still TBD	Still TBD

Code	Type of data to be collected / name of the dataset	Type of data cleaning needed (e.g., correct data types, remove duplicates, add missing info...)	Person responsible for data cleaning (partner short name; person name; email)	Type of data transformation/analysis (e.g., normalization, discretization, ...)	Software/tools used for cleaning, transform, and analyse	Where/by whom will the analysis be conducted?	Standards followed for code development / access and re-use
DS3.3	Voice-related time and spectral features captured by iPrognosis smartphone application	Still TBD	Still TBD	Still TBD	Still TBD	Still TBD	Still TBD
DS3.4	Key taps press and release timestamps captured by iPrognosis smartphone virtual keyboard	Still TBD	Still TBD	Still TBD	Still TBD	Still TBD	Still TBD
DS3.5	Angles of elbows and shoulders and scores achieved at the tests included in the iPrognosis iMAT tool	N/A	N/A	N/A	N/A	N/A	N/A
DS3.6	Results of rehabilitation sessions with Gradior and care plans (editor and patient data)	Extraction and alignment from subsystems	INTRAS	Mapping to FHIR (responsible TMA)	N/A	In HosmartAI platform	N/A
DS4.1	Demographic patient data (age, gender, atrial size, etc.)	N/A	N/A	N/A	N/A	N/A	N/A
DS4.2	3D navigation system data (intracardiac signals, geometry)	N/A	N/A	N/A	N/A	N/A	N/A

Code	Type of data to be collected / name of the dataset	Type of data cleaning needed (e.g., correct data types, remove duplicates, add missing info...)	Person responsible for data cleaning (partner short name; person name; email)	Type of data transformation/analysis (e.g., normalization, discretization, ...)	Software/tools used for cleaning, transform, and analyse	Where/by whom will the analysis be conducted?	Standards followed for code development / access and re-use
DS5.1	Patient recordings and features extracted from interaction with patients	Presence of face and audio. Data cleaning will be carried out by UM	UM	N/A	OpenCV and other tools developed by UM	At the edge, semi-automatic	N/A
DS5.2	Biometric data (e.g., blood pressure and heart rate)	Data cleaning will be performed by the data controller (UKCM)	UKCM	N/A	N/A	N/A	N/A
DS5.3	Retrospective electronic health records	N/A	N/A	N/A	N/A	N/A	N/A
DS5.4	PREMs related to clinical staff (depends on T1.4)	Data cleaning, such as removal of partial answers, duplication, etc. will be performed by the data controller (UKCM)	UM, UKCM	N/A	N/A	At the edge, semi-automatic	N/A
DS5.5	PREMs related patients (PAM, SUS-SI/TAM, UEQ)	Data cleaning will be performed by the data controller (UKCM)	UM, UKCM	N/A	N/A	At the edge, semi-automatic	N/A
DS5.6	Datasets for facial expression and	N/A	N/A	N/A	N/A	N/A	N/A

Code	Type of data to be collected / name of the dataset	Type of data cleaning needed (e.g., correct data types, remove duplicates, add missing info...)	Person responsible for data cleaning (partner short name; person name; email)	Type of data transformation/analysis (e.g., normalization, discretization, ...)	Software/tools used for cleaning, transform, and analyse	Where/by whom will the analysis be conducted?	Standards followed for code development / access and re-use
	emotion recognition						
DS5.7	Datasets for ASR and TTS in Slovenian	N/A	N/A	N/A	N/A	N/A	N/A
DS5.8	Datasets for ASR and TTS in French	N/A	N/A	N/A	N/A	N/A	N/A
DS5.9	EVA Corpus, data set of conversational expression	N/A	N/A	N/A	N/A	N/A	N/A
DS5.10	Video recordings of third persons	N/A	N/A	N/A	N/A	N/A	N/A
DS5.11	Recording of a skeleton representation of the patient doing breathing exercises	N/A	N/A	N/A	N/A	N/A	N/A
DS5.12	Patient's facial information	N/A	N/A	N/A	N/A	N/A	N/A
DS6.1	Patient's physical, medical and cognitive status. Vital signs with sensors data.	Gradior data cleaning will be performed by data owner (INTRAS) TMA, AUTH and ITCL.	INTRAS, TMA, AUTH and ITCL will be responsible for data cleaning and preparation.	N/A	Excel/SQLSERVER	N/A	N/A
DS7.1	Administrative data of patients scheduled for a coronary angiogram/coronary	Data cleaning will be performed by the data	UZB primary investigator will be responsible for data cleaning and preparation	Manual removal of outlier patients (low image quality)	Excel	Philips Image Guided Therapy Systems, Netherlands	N/A

Code	Type of data to be collected / name of the dataset	Type of data cleaning needed (e.g., correct data types, remove duplicates, add missing info...)	Person responsible for data cleaning (partner short name; person name; email)	Type of data transformation/analysis (e.g., normalization, discretization, ...)	Software/tools used for cleaning, transform, and analyse	Where/by whom will the analysis be conducted?	Standards followed for code development / access and re-use
	intervention @ UZB	owner (UZB)					
DS7.2	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	Data cleaning will be performed by the data owner (UZB)	UZB primary investigator will be responsible for data cleaning and preparation	Manual removal of outlier patients (low image quality)	Excel	Philips Image Guided Therapy Systems, Netherlands	N/A
DS7.3	Coronary angiogram imaging data of patients who undergone a coronary angiogram/coronary intervention @ UZB	Data cleaning will be performed by the data owner (UZB)	UZB primary investigator will be responsible for data cleaning and preparation	Manual removal of outlier patients (low image quality)	Excel	Philips Image Guided Therapy Systems, Netherlands	N/A
DS7.4	Coronary physiology data of patients evaluated by a resting index measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary intervention @ UZB	Data cleaning will be performed by the data owner (UZB)	UZB primary investigator will be responsible for data cleaning and preparation	Manual removal of outlier patients (low physiology curves quality)	Coroventis software and Virtual stenting algorithm (VSA) for the interoperation of RFR/FFR pullback created by JF Argacha and Jean Decamp (I-depot number 123060 Date 16-04-2020)	Philips Image Guided Therapy Systems, Netherlands And JF Argacha, cardiology department, UZB, VUB, Brussel	N/A
DS7.5	Intravascular imaging data of patient	Data cleaning will be	UZB primary investigator will be	Manual removal of outlier	Optis and volcano software	Philips Image Guided Therapy	N/A

Code	Type of data to be collected / name of the dataset	Type of data cleaning needed (e.g., correct data types, remove duplicates, add missing info...)	Person responsible for data cleaning (partner short name; person name; email)	Type of data transformation/analysis (e.g., normalization, discretization, ...)	Software/tools used for cleaning, transform, and analyse	Where/by whom will the analysis be conducted?	Standards followed for code development / access and re-use
	evaluated by either OCT or IVUS technique before and after a coronary intervention	performed by the data owner (UZB)	responsible for data cleaning and preparation	patients (low image quality)		Systems, Netherlands	
DS7.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention	Data cleaning will be performed by the data owner (UZB)	UZB primary investigator will be responsible for data cleaning and preparation	Manual removal of outlier patients (low image quality)	Philips and Heartflow software	Philips Image Guided Therapy Systems, Netherlands	N/A
DS7.7	Full prospective UZB data set of administrative, clinical, coronary angiogram, coronary physiology, intravascular imaging and coronary CT data.	Not applicable (prospective inclusion)	Not applicable (prospective inclusion)	Not applicable (prospective inclusion)	Not applicable (prospective inclusion)	Philips Image Guided Therapy Systems, Netherlands And Cardiology department UZB, VUB	N/A
DS8.1	Image, gene, phenotype and pathology data for glioma patients	Connecting sample information per patient across databases, machine learning on data	To be hired	Integration , neural networks	Python, pytorchporch, sklearn, XNAT framework	VUB	N/A

Code	Type of data to be collected / name of the dataset	Type of data cleaning needed (e.g., correct data types, remove duplicates, add missing info...)	Person responsible for data cleaning (partner short name; person name; email)	Type of data transformation/analysis (e.g., normalization, discretization, ...)	Software/tools used for cleaning, transform, and analyse	Where/by whom will the analysis be conducted?	Standards followed for code development / access and re-use
DS.O.1	All pilot data with Common KPIs (Economic & PROMs/PREMs)	Data cleaning will be performed by each data owner (pilot). PhE will perform the analysis on clean datasets	N/A	Economic & PRO/PREM analysis (economic evaluation, cost consequence analysis, cost-utility analysis, patients' quality of life measurement, bootstrapping, regression, etc	Stata, Excel, maybe SAS if necessary	PhE	Following ISPOR guidelines for economic evaluation and Dolan's publication on EQ-5D analysis

A.11 Ethical review

Table 19: Ethical review.

Code	Type of data to be collected / name of the dataset	Type of ethical review needed
DS1.1	Cardiac ultrasound video recordings	Clinical protocol, Ethical approval by Institutional Research Ethics Board
DS1.2	Capsule endoscopy video recordings	Clinical protocol, Ethical approval by Institutional Research Ethics Board
DS1.3	Cardiotocography variables and results, biometric data, medical history data	Clinical protocol and ethical approval from the hospital administration
DS1.4	Coronary computed tomography angiography (CCTA) variables, biometric data, medical history data	Clinical protocol and ethical approval from the hospital administration
DS2.1	Data related to hours spent by specialists.	N/A

Code	Type of data to be collected / name of the dataset	Type of ethical review needed
DS2.2	Retrospective patient schedule data and precondition of the treatment	DPO approval
DS2.3	Data linked to radiotherapy machines (tumours treatment indication, maintenance, building location)	N/A
DS2.4	PROMs/PREMs	Ethical approval
DS2.5	Prospective patient clinical data	Ethical approval
DS2.6	Patient personal data (address, preferences, ...)	Ethical approval
DS2.7	Retrospective EHR	DPO approval
DS2.8	Data from radiotherapy services, infrastructure. Patient's satisfaction	N/A
DS3.1	Smart home data: Consumption/production of instantaneous electricity and consumption logs, Human presence/access control, Devices' activation and connected loads	Clinical protocol, Ethical approval by the national review board
DS3.2	IMU data captured by iPrognosis smartphone application	Clinical protocol, Ethical approval by Research Ethics Committee
DS3.3	Voice related time and spectral features captured by iPrognosis smartphone application	Clinical protocol, Ethical approval by Research Ethics Committee
DS3.4	Key taps press and release timestamps captured by iPrognosis smartphone virtual keyboard	Clinical protocol, Ethical approval by Research Ethics Committee
DS3.5	Angles of elbows and shoulders and scores achieved at the tests included in the iPrognosis iMAT tool	Clinical protocol, Ethical approval by Research Ethics Committee
DS3.6	Results of rehabilitation sessions with Grador and care plans (editor and patient data)	Clinical protocol, Ethical approval by Research Ethics Committee
DS4.1	Demographic patient data (age, gender, atrial size, etc.)	N/A
DS4.2	3D navigation system data (intracardiac signals, geometry)	N/A
DS5.1	Audio-Visual recordings of patients	Clinical protocol, Ethical approval by the hospital's Ethics Committee
DS5.2	Biometric data (e.g., blood pressure and heart rate)	Clinical protocol, Ethical approval by the hospital's Ethics Committee

Code	Type of data to be collected / name of the dataset	Type of ethical review needed
DS5.3	Retrospective electronic health records	Clinical protocol, Ethical approval by the hospital's Ethics Committee
DS5.4	PREMs related to clinical staff (depends on T1.4)	Clinical protocol, Ethical approval by the hospital's Ethics Committee
DS5.5	PREMs related patients (e.g., PAM, SUS-SI/TAM, UEQ) and PROs	Clinical protocol, Ethical approval by the hospital's Ethics Committee
DS5.6	Datasets for facial expression and emotion recognition	N/A
DS5.7	Datasets for ASR and TTS in Slovenian	N/A
DS5.8	Datasets for ASR and TTS in French	N/A
DS5.9	EVA Corpus, data set of conversational expression	N/A
DS5.10	Video recordings of third persons	Clinical protocol, Ethical approval by the hospital's Ethics Committee
DS5.11	Recording of a skeleton representation of the patient doing breathing exercises	N/A
DS5.12	Patient's facial information	N/A
DS6.1	Patient's physical, medical and cognitive status. Vital signs with sensors data	Ethical approved by the ethics committed region.
DS7.1	Administrative data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	Clinical protocol (patient registry). Ethical approval by the hospital review board will be asked.
DS7.2	Clinical data of patients scheduled for a coronary angiogram/coronary intervention @ UZB	Clinical protocol (patient registry). Ethical approval by the hospital review board will be asked.
DS7.3	Coronary angiogram imaging data of patients who undergone a coronary angiogram/coronary intervention @ UZB	Clinical protocol (patient registry). Ethical approval by the hospital review board will be asked.

Code	Type of data to be collected / name of the dataset	Type of ethical review needed
DS7.4	Coronary physiology data of patients evaluated by a resting index measure (iFR/RFR) or a hyperemic index (FFR) either during a manual or a motorized wire pullback and performed before and after a coronary intervention @ UZB	Clinical protocol (patient registry). Ethical approval by the hospital review board will be asked.
DS7.5	Intravascular imaging data of patient evaluated by either OCT or IVUS technique before and after a coronary intervention	Clinical protocol (patient registry). Ethical approval by the hospital review board will be asked.
DS7.6	Coronary CT data including FFRCT computation of patient referred for an invasive coronary angiogram and/or a coronary intervention	N/A
DS7.7	Full prospective UZB data set of administrative, clinical, coronary angiogram, coronary physiology, intravascular imaging and coronary CT data.	Clinical protocol (prospective comparative study). Ethical approval by the hospital review board will be asked.
DS8.1	Image, gene, phenotype and pathology data for glioma patients	Already performed and approved.
DS.O.1	All pilot data with Common KPIs (Economic & PROMs/PREMs)	The economic & PROMs/PREMs data will be included in the protocol.

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