

Dataspace4Health In the Luxembourgish Health Ecosystem

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1. EXECUTIVE SUMMARY

- Purpose of the Document: Outline a harmonized view and strategy between the different partners within the Dataspace4Health project to bring Health Data Space and Gaia-X innovations to Luxembourg's healthcare ecosystem and to describe the relation with the upcoming European Health Data Space. Furthermore, this document shall reflect the Luxembourgish view and perception to contribute and innovate within the Gaia-X and Data Space initiatives, including constraints and the health ecosystem itself. Finally, this document shall describe the vision and associated risk assessment on creating, building and running the new Gaia-X compliant Data Space in Production.
- **Key Objectives:** Highlight the focus areas such as regulation, compliance, technical requirements and collaboration between key stakeholders.
- **Strategic Importance:** Explain the potential benefits for Luxembourg's healthcare system, including innovation, data sharing and enhanced patient care.

2. INTRODUCTION

Dataspace4Health (DS4H) is a collaborative project between different partners to build a Health Data Space through interconnecting various stakeholders, to create an open healthcare data exchange ecosystem, focusing on secure data sharing and aiming to contribute to innovation in medical treatments. As part of this project, two concrete use-cases are under development in order to demonstrate its envisioned benefit to Luxembourg.

This document will outline a harmonized view and strategy between the different partners how to bring Health Data Space and Gaia-X innovations to Luxembourg. It takes into account elements of the European Health Data Space (EHDS) as well as Gaia-X (and its potential relation), the Luxembourgish health ecosystem (and its constraints), including the different key stakeholders and their collaboration needs.

Another aspect addressed in this document are the opportunities and potential benefits for the Luxembourgish healthcare system (and beyond) that come with the introduction of an Health Data Space, such as streamlining access to quality data to foster innovations and enhanced patient care due to timely access to accurate data to e.g., prevent insufficient as well as inappropriate care, all through secure and compliant data sharing between different institutions.

Furthermore, this document will address regulatory and compliance considerations, both on nationaland European level, and describe challenges that are identified which may hinder harmonizing compliance.

Finally, this document will propose different approaches to overcome these challenges and to mitigate identified risks, in order to achieve the envisioned open health ecosystem.



3. EHDS & GAIA-X: POTENTIAL FOR LUXEMBOURG

This chapter will describe the EHDS and Gaia-X in more detail. It will also address the relation between both, as well as their relevance to digital health transformation. Additionally, this chapter will describe the Luxembourgish context and the opportunities it may bring to the country. Finally, this chapter will address the objectives of the DS4H project.

3.1. EUROPEAN HEALTH DATA SPACE (EHDS)

The EHDS, Regulation (EU) 2025/327, which entered into force on the 26th of March 2025 and has key milestones towards full implementation defined (see below for more details), is a major European Union (EU) initiative aimed at creating a framework for the secure and efficient sharing of health data within and across EU member states. It is designed to facilitate both primary use (clinical care) and secondary use (research, policymaking and innovation) of health data, while ensuring high levels of privacy and security.

3.1.1. KEY OBJECTIVES

3.1.1.1. EMPOWERMENT OF INDIVIDUALS:

- Access to Personal Health Data: EHDS ensures that individuals across the EU have the right to
 easily access and control their personal health data, such as electronic health records (EHR),
 across borders.
- Portability: Citizens can share their health data with healthcare providers, regardless of the country, improving continuity of care and reducing administrative barriers.

3.1.1.2. IMPROVED HEALTHCARE DELIVERY:

- Cross-border Healthcare: EHDS aims to create a seamless exchange of health data between healthcare providers in different EU member states, allowing for more efficient, timely and personalized care.
- Interoperability Standards: It promotes harmonization of data formats and standards to enable the exchange of data across national health systems.

3.1.1.3. DATA FOR RESEARCH AND INNOVATION:

- Secondary Use of Health Data: EHDS promotes the use of anonymized and pseudonymized health data for research, innovation and policy-making. This enables advancements in medical research, public health and the development of new treatments and technologies.
- Data-Driven Health Solutions: Researchers, companies and policymakers can leverage health data to improve public health outcomes, foster innovation and develop new digital health tools.

3.1.1.4. PRIVACY AND SECURITY:

- GDPR Compliance: EHDS is built around the principles of the General Data Protection Regulation (GDPR), ensuring that health data is handled securely and in line with strict privacy standards.
- Data Sovereignty: EHDS ensures that data remains under the control of individuals and that data protection rights are respected throughout the EU.

3.1.2. COMPONENTS OF THE EHDS



- National Health Data Access Bodies: Each member state is required to establish at least one body to facilitate access to health data for secondary purposes, such as research or policymaking.
- Interoperability Framework: EHDS requires technical standards and infrastructures to ensure that health data can flow smoothly across borders while maintaining high levels of security.
- Cross-border Services: EHDS requires services like ePrescriptions and patient summaries to allow for seamless healthcare across EU countries.

3.1.3. BENEFITS OF THE EHDS

- For Patients: Improved access to personalized healthcare services, even when traveling or living in other EU countries.
- For Healthcare Providers: Enhanced ability to offer more coordinated and efficient care by accessing patient health records in real-time and in the native language of healthcare providers.
- For Researchers and Policy Makers: Easier access to large datasets that can be used for advancing medical research, drug development and public health analysis.
- For Innovators: A supportive environment for developing digital health technologies that rely on secure access to health data.

Defined key milestones towards full implementation of the EHDS, as of the moment of writing this document are:

- March 2025: The EHDS Regulation enters into force, marking the beginning of the transition period.
- March 2027: Deadline for the Commission to adopt several key implementing acts, providing detailed rules for the regulation operationalization
- March 2029: Key parts of the EHDS Regulation will enter into application, including, for primary
 use, the exchange of the first group of priority categories of health data (Patient Summaries,
 ePrescriptions/eDispensations) in all EU Member States. Rules on secondary use will also start to
 apply for most data categories (e.g. data from electronic health records).
- March 2031: For primary use, the exchange of the second group of priority categories of health data (medical images, lab results, and hospital discharge reports) should be operational in all EU Member States. Rules on secondary use will also start to apply for the remaining data categories (e.g. genomic data).
- March 2034: Third countries and international organizations will be able to apply to join HealthData@EU, for the secondary use.

3.1.4. LONG-TERM VISION

The EHDS aims to create a unified European framework that maximizes the value of health data while safeguarding data privacy. It promotes the use of data to drive healthcare innovation, improve patient outcomes and ensures that European citizens benefit from advancements in digital health technologies and treatments.



3.2. GAIA-X

Gaia-X is a European initiative aimed at creating a federated and secure data infrastructure that ensures data sovereignty, interoperability and innovation. The project was launched in 2019 by Germany and France, with the aim of establishing a European cloud ecosystem that can compete with global tech giants while aligning with European values such as privacy, transparency and openness.

3.2.1. KEY OBJECTIVES OF GAIA-X

3.2.1.1. DATA SOVEREIGNTY

- Gaia-X empowers individuals, organizations and countries to maintain control over their own data
 by ensuring that it is processed and stored according to European laws and standards, such as the
 GDPR.
- It provides transparency about where and how data is handled, offering users full control over their information.

3.2.1.2. INTEROPERABILITY AND FEDERATION

- Gaia-X fosters interoperability between different cloud services, ensuring that data can move freely
 across systems, providers and borders without being locked into one proprietary platform.
- It creates a federated ecosystem where multiple cloud service providers, both large and small, can collaborate and share resources while adhering to common standards.

3.2.1.3. SECURITY AND TRUST

- Gaia-X is designed with robust security measures to protect data and ensure trust in the digital infrastructure. Participants must meet stringent security and compliance requirements.
- By offering a transparent and standardized infrastructure, Gaia-X builds trust between data providers and consumers.

3.2.1.4. INNOVATION AND COMPETITIVENESS:

- Gaia-X supports European businesses, governments and institutions in developing and using cloud services that align with European standards.
- It promotes innovation by creating an open and flexible cloud environment, allowing startups, researchers and companies to develop new digital products and services.

3.2.2. GAIA-X FOR THE HEALTH SECTOR

Gaia-X plays a crucial role in transforming the healthcare sector by providing a secure, interoperable and federated cloud infrastructure for managing and sharing health data. The health sector, which deals with highly sensitive information, could benefit greatly from the principles and technologies underpinning Gaia-X.

3.2.3. KEY BENEFITS OF GAIA-X FOR HEALTH

3.2.3.1. SECURE HEALTH DATA SHARING

 Data Sovereignty: Gaia-X ensures that health data remains under the control of the individual or organization that generates it. This is critical in healthcare, where privacy and compliance with regulations like GDPR are essential.



 Encrypted and Federated Data Exchange: Healthcare providers, researchers and institutions can share data securely across borders, enhancing collaboration and innovation without compromising patient privacy or security.

3.2.3.2. INTEROPERABILITY IN HEALTH SYSTEMS

- Gaia-X promotes the technical interoperability of health data systems, enabling seamless
 integration of patient records, clinical data and other health-related information between hospitals,
 doctors and health agencies across Europe.
- This allows trusted healthcare providers to access patient data from different systems in real-time, improving care coordination, especially in cross-border healthcare services.

3.2.3.3. ENABLING INNOVATION IN HEALTHCARE

- Research and Innovation: Through data offerings from healthcare providers, the Gaia-X framework
 is able to facilitate access to large datasets for medical research, public health analysis and the
 development of new treatments. This is crucial for areas like personalized medicine, Artificial
 Intelligence (AI) driven healthcare solutions and the development of digital health applications.
- Faster Medical Advancements: By creating a secure and federated platform, Gaia-X accelerates
 collaboration between pharmaceutical companies, universities and healthcare providers for faster
 drug development and clinical trials.

3.2.3.4. PRIVACY-COMPLIANT HEALTH SERVICES

- GDPR Compliance: Gaia-X ensures that all health data is handled in compliance with European data protection laws, safeguarding patient information while allowing it to be used for legitimate healthcare services and research.
- Digital Health Services: Gaia-X supports the development of secure telemedicine, remote diagnostics and e-health applications by providing a trusted infrastructure for patient data.

3.2.3.5. SUPPORT FOR THE EHDS

- Gaia-X is aligned with the goals of the EHDS, providing the underlying infrastructure for data sharing across healthcare systems in Europe.
- It enables secure access to health data for both primary use (clinical care) and secondary use (research, innovation and policymaking), fostering the vision of a unified European health data ecosystem.

3.2.4. HOW GAIA-X BENEFITS KEY HEALTH STAKEHOLDERS

3.2.4.1. HOSPITALS AND HEALTHCARE PROVIDERS

- Hospitals can securely share patient data across regions and countries, leading to better care coordination, especially in emergencies or when patients move across borders.
- By using a federated infrastructure, hospitals can collaborate with research institutions and other providers without compromising data security.

3.2.4.2. PHARMACEUTICAL COMPANIES AND RESEARCHERS

- Pharmaceutical companies can access anonymized and/or pseudonymized data for research purposes, speeding up clinical trials, drug discovery and personalized medicine.
- Researchers can collaborate with other institutions across Europe through secure data-sharing platforms to advance medical research.

3.2.4.3. GOVERNMENTS AND PUBLIC HEALTH AUTHORITIES

- Governments can use aggregated health data to better understand public health trends, manage pandemics and make data-driven policy decisions.
- Public health authorities can leverage Gaia-X to improve data collection, analysis and sharing between different healthcare institutions and/or different countries.

3.2.4.4. PATIENTS



- The Gaia-X framework allows to enable patients to have control over their own personal health data and can decide when and with whom to share it.
- It enhances patient experience by enabling access to digital health services and better coordination between healthcare providers.

3.2.5. CONCLUSION

Gaia-X offers a revolutionary cloud and data infrastructure that supports data sovereignty, security and innovation, making it especially valuable for the health sector. By providing a federated platform for secure data sharing and collaboration, Gaia-X enables healthcare providers, researchers and governments to deliver improved healthcare services, accelerate medical research and safeguard patient privacy in compliance with European regulations.

3.3. RELATION BETWEEN EHDS AND GAIA-X

The EHDS and Gaia-X are two complementary initiatives that aim to transform the way data is managed, shared and utilized across Europe. They share common goals in terms of ensuring data sovereignty, security and interoperability, but they focus on different aspects of the digital infrastructure. The following section describes how both initiatives relate:

3.3.1. COMMON GOALS

- Data Sovereignty: Both EHDS and Gaia-X emphasize that European citizens and organizations must have full control over their data. EHDS focuses on health data, while Gaia-X addresses a broader range of sectors (including healthcare).
- Security and Privacy: Both initiatives ensure that data is handled in compliance with European regulations, particularly the GDPR. Gaia-X provides the cloud infrastructure to manage data securely, while EHDS establishes frameworks for managing sensitive health data.
- Interoperability: Gaia-X and EHDS aim to create standardized, interoperable systems that allow seamless data exchange across borders, sectors and institutions.

3.3.2. EHDS - HEALTH-FOCUSED DATA ECOSYSTEM

- The EHDS is specifically designed to facilitate the sharing and use of health data across Europe for both primary use (patient care) and secondary use (research, policymaking).
- It focuses on enabling secure access to patient data, creating a digital health ecosystem that improves healthcare delivery and supporting research and innovation in the medical field.
- EHDS needs a robust and secure data infrastructure to manage this sensitive information, which is where Gaia-X could play a crucial role.

3.3.3. GAIA-X - THE FEDERATED CLOUD INFRASTRUCTURE

- Gaia-X could provide the underlying cloud and data infrastructure that enables secure, federated and interoperable data exchange between different systems and organizations across Europe.
- It is a cross-sector initiative designed to ensure that data flows freely, securely and in compliance with European laws. Gaia-X's infrastructure supports various industries, including healthcare, manufacturing, finance and more.
- Gaia-X for Health is one specific application of Gaia-X, providing the framework to manage and exchange health data securely within the EHDS.

3.3.4 HOW EHDS COULD LEVERAGE GAIA-X



- Technical Backbone: EHDS could rely on Gaia-X to provide the technical infrastructure for storing, processing and exchanging health data across borders. Gaia-X's federated architecture ensures that this data remains secure and accessible in compliance with GDPR.
- Interoperability Standards: Gaia-X offers a framework for interoperability across various cloud services and systems, which is crucial for EHDS to achieve its vision of seamless health data exchange across EU member states.
- Security and Compliance: Gaia-X's cloud infrastructure provides enhanced data security, privacy and compliance mechanisms, which are essential for managing sensitive health information under the EHDS framework.

3.3.5. COLLABORATION FOR INNOVATION

- Innovation in Healthcare: EHDS and Gaia-X both foster innovation by enabling researchers, policymakers and healthcare providers to securely access and share health data. This collaboration supports advances in fields such as personalized medicine, medical research and Al-driven healthcare solutions.
- Data-Driven Healthcare: By integrating the capabilities of Gaia-X into EHDS, health data can be analyzed and shared across borders, leading to improved healthcare outcomes and the development of innovative treatments.

3.3.6. SUPPORT FOR THE EUROPEAN HEALTH DATA ECOSYSTEM

- EHDS and Gaia-X both contribute to the creation of a unified European health data ecosystem that
 facilitates secure, interoperable and privacy-preserving data flows. This ecosystem enhances
 healthcare, research and policymaking while respecting European values such as transparency
 and data sovereignty.
- Gaia-X's infrastructure aligns with the EHDS vision by providing the secure, scalable and interoperable cloud services necessary for health data to be shared and used efficiently across the EU.

3.3.7. PRACTICAL IMPLEMENTATION

- In practical terms, Gaia-X could provide the cloud services and data-sharing platforms that allow healthcare providers, researchers and governments to access and analyze health data securely and efficiently within the framework set by EHDS.
- Healthcare providers would use the EHDS to share patient records across borders, while
 researchers could access anonymized and/or pseudonymized data for medical research, all
 facilitated by the Gaia-X infrastructure.

3.3.8. CONCLUSION

The EHDS and Gaia-X are closely related, with EHDS focusing on the secure and compliant sharing of health data across Europe and Gaia-X providing the underlying federated cloud infrastructure. Together, they form a powerful ecosystem that ensures European data sovereignty, privacy and security, while fostering innovation and collaboration in healthcare and beyond. The DS4H project serves as some sort of a pilot for the EHDS implementation, to create and achieve a Health Data Space on a smaller scale – namely in the Luxembourgish health ecosystem.



3.4. RELEVANCE OF EHDS AND GAIA-X TO DIGITAL HEALTH TRANSFORMATION

The EHDS and Gaia-X are pivotal to the ongoing digital health transformation in Europe. They enable a more integrated, efficient and secure approach to managing health data and delivering healthcare services. The following section will describe the relevance of both initiatives to digital health transformation:

3.4.1. DATA-DRIVEN HEALTHCARE

- EHDS enables the secure and seamless exchange of health data across borders, fostering a datadriven healthcare system where health records, lab results and diagnostic data can follow the patient across the EU. This supports continuity of care, improving patient outcomes and making healthcare more personalized.
- Gaia-X provides the infrastructure for securely managing, storing and analyzing these large datasets, making data easily accessible to healthcare providers, researchers and institutions. The ability to access comprehensive, real-time health data is crucial for improving diagnosis, treatment and care coordination.

3.4.2. PERSONALIZED MEDICINE AND INNOVATION

- Personalized Medicine: With EHDS, healthcare providers can access a patient's full medical history, genomic data and other health-related information, enabling more tailored treatments. This personalized approach to care can result in more accurate diagnoses and better outcomes.
- Gaia-X enables the secure exchange of data across different healthcare providers and research
 institutions, allowing for innovation in Al-driven healthcare solutions, drug development and
 personalized therapies. These innovations are made possible by the availability of large, secure
 datasets provided by EHDS, managed by Gaia-X.

3.4.3. INTEROPERABILITY ACROSS HEALTH SYSTEMS

- EHDS aims to break down silos between and within national health systems, promoting interoperability and the standardization of health data across Europe. This ensures that health information can flow freely and securely between healthcare providers, patients and researchers.
- Gaia-X plays a key role by creating a federated infrastructure that supports the technical
 interoperability of health systems, enabling data from various cloud providers and platforms to be
 accessed and shared in a standardized way. This is essential for creating a unified digital health
 ecosystem.

3.4.4. ENHANCED HEALTHCARE EFFICIENCY

- EHDS contributes to greater healthcare efficiency by digitizing health records, enabling telemedicine and streamlining administrative processes like ePrescriptions and cross-border patient summaries. This reduces inefficiencies in healthcare delivery and facilitates quicker, more coordinated care.
- Gaia-X ensures that healthcare data can be securely processed, stored and shared between
 different healthcare providers, improving collaboration, reducing redundancies and optimizing the
 healthcare supply chain. This is especially relevant in managing complex healthcare systems,
 where multiple stakeholders need access to the same data in real time.

3.4.5. SUPPORTING HEALTHCARE RESEARCH AND INNOVATION



- Secondary Use of Health Data: EHDS enables secondary use of health data for research, innovation and policymaking. Researchers can securely access anonymized or pseudonymized data to conduct studies, develop new treatments or address public health challenges.
- Gaia-X provides the infrastructure needed for this type of large-scale data collection, exchange and
 analysis while ensuring that data privacy and security are maintained. Through its federated cloud,
 Gaia-X allows researchers from different countries to collaborate without compromising sensitive
 health information. This promotes innovation and the development of cutting-edge healthcare
 technologies.

3.4.6. PRIVACY AND DATA SECURITY IN HEALTHCARE

- Both EHDS and Gaia-X emphasize the importance of data security and privacy, critical for healthcare, where sensitive personal data is involved. GDPR compliance and data sovereignty are central tenets of both initiatives, ensuring that health data is protected while being shared and used responsibly.
- Gaia-X provides secure, GDPR-compliant cloud infrastructure that ensures health data is stored
 and processed in line with European regulations. This fosters trust in digital health systems,
 encouraging wider adoption of digital health services among healthcare providers and patients.

3.4.7. TELEMEDICINE AND REMOTE HEALTHCARE

- EHDS supports the use of telemedicine, enabling remote consultations, diagnostics and treatment. With patient data available across borders, telemedicine services can be offered with greater confidence, improving access to healthcare in rural or underserved areas.
- Gaia-X ensures that the technical infrastructure supporting telemedicine is secure, interoperable
 and scalable, facilitating the growth of remote healthcare solutions. This is particularly relevant postCOVID-19, as telehealth has become a vital tool for healthcare delivery.

3.4.8. CROSS-BORDER HEALTHCARE AND MOBILITY

- EHDS allows for the secure exchange of patient health data across EU borders, ensuring that
 healthcare providers have access to patient information, regardless of where the patient is located.
 This is essential for improving healthcare access and continuity for cross-border mobility within the
 EU.
- Gaia-X ensures that the infrastructure needed to support this cross-border data exchange is in
 place, providing seamless integration between different healthcare systems while maintaining the
 highest standards of data security.

3.4.9. BUILDING TRUST IN DIGITAL HEALTH ECOSYSTEMS

- Trust is critical in digital health transformation. EHDS and Gaia-X together create a trusted environment for data sharing and collaboration. By adhering to strict European standards for security, privacy and compliance, they provide assurance to healthcare providers, patients and regulators that health data is handled responsibly.
- This trust is essential for encouraging broader adoption of digital health technologies, from EHRs to Al-powered diagnostics and digital therapeutics.

3.4.10. SUPPORTING THE EUROPEAN DIGITAL HEALTH VISION

• The combination of EHDS and Gaia-X aligns with the broader vision of a European Health Union, where health data is shared securely and used effectively across member states to improve patient care and foster innovation.



 By providing a unified infrastructure (Gaia-X) and clear regulatory framework (EHDS), Europe can lead the world in digital health transformation, leveraging data to improve healthcare services, public health outcomes and economic growth.

3.4.11. CONCLUSION

The EHDS and Gaia-X initiatives are central to Europe's digital health transformation. By enabling secure, interoperable and innovative use of health data, they drive improvements in patient care, healthcare efficiency, research and innovation, all while ensuring data sovereignty and privacy. Together, they create a robust framework for building a future-focused, data-driven healthcare system that benefits both patients and healthcare providers across Europe.

3.5. CONTEXT IN LUXEMBOURG

In Luxembourg, the context for implementing the EHDS and Gaia-X initiatives is shaped by the country's commitment to developing a highly innovative, secure and patient-centered healthcare system. Luxembourg is positioned as a digital and financial hub in Europe, with a strong emphasis on digital transformation, data privacy and cybersecurity. This creates an ideal environment for integrating these European initiatives into its healthcare sector.

The key elements of Luxembourg's Context for DS4H, EHDS and Gaia-X in health are presented in the following chapters.

3.5.1. DIGITAL HEALTH INFRASTRUCTURE IN LUXEMBOURG

3.5.1.1. AGENCE ESANTÉ:

The central eHealth agency in Luxembourg, Agence eSanté, plays a crucial role in managing the country's health information systems, including the National eHealth Platform (eSanté), which facilitates the secure exchange of patient data between healthcare providers. Other digital health services offered by Agence eSanté include:

- Electronic Health Record (Dossier de Soins Partagé DSP): a platform for sharing and exchanging
 data in the health sector including the electronic health record. Agence eSanté is in the progress of
 updating the DSP into a new generation (NG) in line with the EHDS regulation.
- Electronic Vaccination Record (Carnet de Vaccination Electronique CVE)
- HealthNet: the framework for the secure interconnection network between the different actors in the
 health sector. It is a secure high-speed infrastructure network that provides various security
 services, such as Reverse Proxy, Web Proxy, Email Gateway
- Secure messaging

Additionally, Agence eSanté created a Master Plan for Health Information Systems (SDSI, currently Version 3) defining a national strategy for the interoperability of health information systems. In regards to the EHDS, Agence eSanté has a leading role for the country's implementation in terms of primary use of data.

3.5.1.2. LUXEMBOURG'S EHEALTH STRATEGY:

In 2020 the ministry of Health and Social Security, the Agence eSanté and the Caisse Nationale de Santé (CNS) formalized a proposal for a national eHealth strategy 2021-2028 to help accelerate digitalization in the healthcare sector. The proposed national eHealth strategy 2021-2028 states: "Let's mobilize the potential of digitalization to serve healthcare professionals and patients within the framework of clear governance, to modernize our healthcare system by developing secure systems that ensure the sharing of healthcare data." It is articulated around six key strategic priorities (1 Facilitate



patient follow-up for professionals, 2) Engage / involve the patient, 3) Simplify administrative procedures for all, 4) Support players as they upgrade their skills, 5) Adopt methodical, integrated governance, 6) Facilitate secure data sharing and access). Luxembourg has been actively investing in digital health technologies, with initiatives to promote telemedicine, digital health records and personalized medicine. EHDS and Gaia-X align with Luxembourg's goal to create a connected health ecosystem where health data is shared securely across systems, which the DS4H project pilots for.

3.5.1.3. LUXEMBOURG'S NATIONAL INTEROPERABILITY FRAMEWORK:

The country is working to ensure that health data can be exchanged across different healthcare providers. This framework supports the objectives of both EHDS (seamless data exchange) and Gaia-X (interoperability and secure infrastructure).

3.5.1.4. THE STATE INFORMATION TECHNOLOGY CENTER (LE CENTRE DES TECHNOLOGIES DE L'INFORMATION DE L'ÉTAT - CTIE):

The national administration responsible for IT services for the Luxembourgish government, ministries and administrations. CTIE plays a role on the Data Governance Act (DGA) and – as of writing of this document – is likely to have the EHDS technical infrastructure under them, in additional to Agence eSanté and Luxembourg National Data Service (LNDS).

3.5.1.5. NATIONAL INSTITUTE FOR HEALTH AND SOCIAL SECURITY (INSPECTION GÉNÉRALE DE LA SÉCURITÉ SOCIALE - IGSS):

The Luxembourg Microdata Platform on Labour and Social Protection (LMDP) allows for sharing of pseudonymized health data, for example with research organizations.

3.5.1.6. LUXEMBOURG IT FOR HEALTHCARE (LUXITH):

Purpose to implement and operate the shared IT services, software and infrastructures of its members. In additional, they are responsible for implementing the hospital sector's IT strategic plan, which provides for a gradual pooling of IT skills from hospital establishments towards a single system for all hospitals and, if possible, interoperable with the systems of other players in the health sector. LUXITH aims to set up a Health Content Management (HCM) system in 2026 in Luxembourg. The HCM aims to facilitate the archiving and sharing of medical and healthcare documentation, connecting existing Hospital Information Systems (HIS), offering electronic signature and protecting data against cyberattacks. Moreover, through the HCM, hospitals should benefit from a more centralized infrastructure, connection to the DSP and the integration of medical imaging management.

3.5.1.7. DATA PROTECTION COMMISSION:

Luxembourg's Commission Nationale pour la Protection des Données (CNPD) launched a regulatory sandbox on artificial intelligence (AI), a technology whose rapid progress is raising concerns about privacy and the protection of personal data. The isolated digital environment allows innovators in the Luxembourg ecosystem to test AI systems for a limited time before they are put on the market. The aim is to help them develop AI applications that comply with the GDPR and therefore respect the privacy of the people concerned.

3.5.2. REGULATORY AND COMPLIANCE FRAMEWORK

- GDPR Leadership: Luxembourg, as a part of the EU, complies with the GDPR. The protection of
 personal data, especially sensitive health data, is a priority and both EHDS and Gaia-X align with
 these GDPR standards.
- National Health Regulations: The Ministry of Health and Social Security (Ministère de la Santé et de la Sécurité sociale) oversees the compliance of healthcare providers with both national and



- European regulations. The Ministry plays a key role in ensuring that the implementation of EHDS meets the necessary regulatory and security standards.
- CNPD is responsible for overseeing data privacy regulations at the national level. The CNPD will
 be critical in ensuring that health data sharing under EHDS and Gaia-X is compliant with data
 privacy laws.
- Health studies (such as clinical trials) conducted in Luxembourg are subject to ethical evaluation
 from the National Research Ethics Committee (Comité National d'Ethique de Recherche CNER).
 The CNER operates according to the rules established by the ICH, the Grand-ducal Regulation of
 30 May 2005, the law on hospitals of 18 March 2018 and in compliance with the Declaration of
 Helsinki.

3.5.3. COLLABORATION BETWEEN KEY STAKEHOLDERS

- **Ministry of Health and Social Security:** The Ministry drives the digital transformation of Luxembourg's healthcare system, aligning with European initiatives such as EHDS and Gaia-X to promote innovation and enhance public health services.
- CNS: Luxembourg's national health insurance system, CNS, manages large amounts of health data and would benefit from more streamlined data access and sharing. By integrating EHDS and Gaia-X, CNS could enhance healthcare reimbursements, patient care and policy development based on data insights.
- Hospitals and Healthcare Providers: Hospitals and doctors in Luxembourg need access to
 interoperable health data systems to improve the quality of care and patient outcomes. Gaia-X and
 EHDS provide a platform for secure and standardized data exchange across providers and borders.
- Luxembourg Institute of Health (LIH) and University of Luxembourg: Both play crucial roles in
 medical research and innovation. Access to shared health data through EHDS and secure
 infrastructure via Gaia-X enables advanced research in areas such as personalized medicine,
 public health and epidemiology.

3.5.4. HEALTHCARE INNOVATION AND RESEARCH

- Personalized Medicine: Luxembourg is a leader in personalized medicine, with institutions like the
 Luxembourg Centre for Systems Biomedicine (LCSB), the Integrated Biobank of Luxembourg (IBBL)
 and the National Center of Translational Cancer Research (NCTCR). Additionally, Luxembourg is
 a front-runner in innovative health projects, such as Clinnova, CoLive Voice and Personalized
 Functional Profiling (PFP). EHDS can provide more robust access to health data for personalized
 treatment, while Gaia-X can provide the infrastructure to handle complex, sensitive datasets.
- Clinical Research and Data Sharing: The LIH and other research institutions rely on access to large datasets for public health studies and clinical trials. EHDS will enable them to access health data across borders, while Gaia-X ensures data security and compliance with EU standards.

3.5.5. CROSS-BORDER HEALTHCARE

- Luxembourg's Geopolitical Position: As a small country surrounded by larger EU countries (France, Germany and Belgium), Luxembourg's healthcare system interacts frequently with cross-border patients and care institutions. EHDS can facilitate cross-border healthcare by ensuring that patient data is accessible when citizens seek medical treatment in other EU countries.
- Medical Tourism and Workforce Mobility: Luxembourg experiences a high level of workforce
 mobility due to its proximity to neighboring countries. EHDS can support mobility of patients and
 healthcare professionals by making patient data accessible across EU borders, improving care
 coordination for cross-border workers and patients.



3.5.6. TELEMEDICINE AND DIGITAL HEALTH SERVICES

- Adoption of Telemedicine: The COVID-19 pandemic accelerated the adoption of and (temporarily) allowed for telemedicine in Luxembourg. EHDS and Gaia-X can further enhance telemedicine by providing secure and standardized access to health data, ensuring that healthcare professionals have the information needed to offer remote consultations in a secure environment.
- **ePrescriptions:** Luxembourg has made progress in enabling ePrescriptions, which are aligned with the EHDS' goal of creating standardized digital health services. Gaia-X may provide the technical infrastructure for securely managing and transmitting ePrescriptions across borders.

3.5.7. CYBERSECURITY AND DATA SOVEREIGNTY

- Focus on Cybersecurity: Luxembourg has a strong focus on cybersecurity, particularly in the financial and governmental sectors. The Cybersecurity Competence Center and national strategies support the secure handling of sensitive health data. Gaia-X's emphasis on secure cloud infrastructure aligns well with Luxembourg's cybersecurity objectives for the health sector.
- Data Sovereignty: Gaia-X ensures that health data remains under European control, preventing
 dependency on non-EU cloud providers. This is critical for Luxembourg, given its focus on data
 sovereignty and alignment with European privacy standards.

3.5.8. ECONOMIC OPPORTUNITIES AND DIGITAL TRANSFORMATION

- **HealthTech Industry Growth**: Luxembourg is positioning itself as a hub for HealthTech startups and companies. By adopting EHDS and Gaia-X, Luxembourg could attract new businesses and innovators focused on digital health, biotechnology and Al-driven healthcare solutions.
- **Digital Transformation in Healthcare:** With initiatives like Digital Luxembourg, the government aims to lead in digital transformation. EHDS and Gaia-X will be key enablers in digitizing the healthcare sector, making Luxembourg a model for eHealth solutions in Europe.

3.5.9. PROOF OF CONCEPT FOR EHDS AND GAIA-X

- Luxembourg as a Testbed: Luxembourg's size and agile digital infrastructure make it an ideal location for pilot projects and proof of concept (PoC) trials for EHDS and Gaia-X implementations, such as the DS4H project with its two concrete use-cases. The country could serve as a testing ground for large-scale health data exchange and cloud infrastructure deployment.
- Multi-Stakeholder Collaboration: The coordinated involvement of public institutions (Ministry of Health and Social Security, CNS), private healthcare providers and research institutions (LIH, University of Luxembourg) is key to making a PoC for EHDS and Gaia-X successful in Luxembourg.

3.5.10. CONCLUSION

Luxembourg is uniquely positioned to be at the forefront of digital health transformation by integrating EHDS and Gaia-X innovations into its healthcare system. With strong regulatory frameworks, a commitment to data security and sovereignty and a collaborative approach among key stakeholders, Luxembourg can leverage these initiatives to improve healthcare services, foster medical research and become a leader in digital health and personalized medicine within Europe.



3.6 OPPORTUNITIES FOR LUXEMBOURG

Luxembourg has a significant opportunity to become a pioneer in the implementation of EHDS and Gaia-X innovations due to its advanced digital infrastructure, supportive regulatory environment and collaborative healthcare ecosystem (and limited size thereof). By leveraging these advantages, Luxembourg can position itself as a leader in digital health transformation and reap both healthcare and economic benefits.

The key opportunities for Luxembourg to pioneer EHDS and Gaia-X innovations are presented in the following chapters.

3.6.1. ADVANCED DIGITAL INFRASTRUCTURE AND AGILITY

- Luxembourg has a highly developed digital infrastructure with strong internet connectivity, cuttingedge data centers and a commitment to secure cloud services. This makes the country well-suited for the deployment of Gaia-X infrastructure to securely manage and share health data.
- Luxembourg's small size and centralized healthcare system allow for quick implementation and integration of new technologies. This agility can enable rapid adoption of EHDS initiatives and serve as a model for larger EU countries.

3.6.2. SUPPORTIVE REGULATORY ENVIRONMENT

- Luxembourg's robust adherence to European privacy regulations, including GDPR, ensures that
 any health data initiative is fully compliant with data protection standards. This makes Luxembourg
 an ideal location for testing EHDS innovations, which require strict compliance with data privacy
 laws.
- With Luxembourg's government and its regulatory bodies, such as the CNPD, prioritizing data sovereignty and cybersecurity, Luxembourg can lead by example in ensuring secure and ethical management of health data under EHDS.

3.6.3. HEALTHCARE INNOVATION HUB

- Luxembourg is already home to innovative health institutions like the LIH (including the NCTCR
 and IBBL) and the LCSB. These institutions focus on personalized medicine, biomedical research
 and health data analytics. By incorporating EHDS and Gaia-X, Luxembourg can boost its research
 capabilities through secure and interoperable health data sharing.
- Luxembourg can attract HealthTech startups and companies by offering a secure, federated
 platform to test and develop new digital health solutions. This could stimulate growth in the
 HealthTech sector, making Luxembourg a leader in Al-driven healthcare innovations.

3.6.4. CROSS-BORDER HEALTHCARE AND MOBILITY

- Due to Luxembourg's central location in Europe and its close interaction with neighboring countries, there is a high demand for cross-border healthcare services. EHDS can facilitate seamless patient data exchange across borders, ensuring continuity of care for patients traveling or working in other EU countries.
- Luxembourg can lead the way in implementing cross-border health solutions, making it a key player
 in the development of a more connected and integrated European healthcare system.

3.6.5. LEADERSHIP IN PERSONALIZED MEDICINE AND RESEARCH

• Luxembourg is already pioneering personalized medicine initiatives, which rely heavily on access to big health data and secure, interoperable systems. By embracing EHDS and Gaia-X,



- Luxembourg can become a hub for personalized treatment, leveraging secure data sharing to enhance research on genomics, public health and chronic disease management.
- Gaia-X will enable Luxembourg to provide cloud infrastructure that allows researchers to collaborate on health data across borders, facilitating large-scale clinical trials and the development of precision medicine.

3.6.6. TELEMEDICINE AND DIGITAL HEALTH LEADERSHIP

- The rise of telemedicine in Luxembourg, accelerated by the COVID-19 pandemic, creates an
 opportunity for Luxembourg to lead in the development of secure and interoperable digital health
 services. EHDS and Gaia-X can enhance the quality and security of telemedicine, making
 Luxembourg a model for other EU nations.
- Luxembourg can pioneer the deployment of ePrescriptions, remote patient monitoring and AI-driven diagnostics, supported by secure data sharing between healthcare providers and patients.

3.6.7. ECONOMIC GROWTH AND ATTRACTING INVESTMENT

- By positioning itself as a leader in digital health transformation, Luxembourg can attract European funding and investment from the private sector, particularly in HealthTech, biotechnology and digital therapeutics.
- Luxembourg can become a European hub for HealthTech innovation, attracting startups, researchers and investors who want to test new technologies in a compliant and secure environment.

3.6.8. PROOF OF CONCEPT AND SCALABLE SOLUTIONS

- Luxembourg is well-suited to serve as a testbed for EHDS and Gaia-X innovations, given its small size, efficient regulatory framework and agile healthcare system. By successfully piloting projects such as cross-border data sharing, personalized medicine trials and telemedicine solutions, Luxembourg can demonstrate scalable solutions that can be expanded to larger EU countries.
- A successful PoC in Luxembourg would reinforce the country's reputation as a leader in digital health and position it as a model for the European Health Union.

3.6.9. COLLABORATION WITH KEY STAKEHOLDERS

- Luxembourg has a strong network of collaborative stakeholders in healthcare, research and government, including the Ministry of Health and Social Security, CNS, LIH, Hospitals and Agence eSanté. This makes the country ideal for fostering collaboration between public and private sectors to achieve a harmonized digital health strategy.
- The collaboration with Gaia-X's federated infrastructure will allow Luxembourg's stakeholders to lead in the secure sharing of health data, ensuring that data sovereignty and compliance are maintained while enhancing patient care and research capabilities.

3.6.10. CONCLUSION

Luxembourg's advanced digital infrastructure, commitment to data privacy, leadership in personalized medicine and its central position in Europe offer a unique opportunity for the country to pioneer the implementation of EHDS and Gaia-X innovations. By leading in digital health transformation, Luxembourg can improve healthcare services, attract investment and become a model for other EU nations, driving the future of European healthcare.



3.7 OBJECTIVE OF THE PROJECTS

The objective of establishing a harmonized strategy is to develop a PoC for the implementation of Health Data Space and Gaia-X innovations within the healthcare sector in Luxembourg, as part of the DS4H project. This will also serve as preparational work for early identification of potential barriers for future EHDS implementation and will allow to propose actions to overcome these. This strategy aims to ensure alignment across key stakeholders—including healthcare providers, government bodies, research institutions and technical experts—in terms of regulatory compliance, data security, interoperability and privacy standards. The PoC will serve as a blueprint for demonstrating how Luxembourg can leverage these innovations to enhance healthcare delivery, research capabilities and cross-border data exchange, while adhering to European data protection regulations.



4. STAKEHOLDER ANALYSIS

The health system in Luxembourg involves around a large variety of key stakeholders, both public and private, with different roles and responsibilities, collaborative needs and balancing aspects. The different stakeholders each playing a critical role in the delivery, management, innovation and regulation of healthcare services.

This chapter will briefly highlight the identified stakeholders, without going into details about their different roles and responsibilities, collaborative needs and balancing aspects.

These details can be found in Annex I, in a table format.

The stakeholders identified are:

- Ministry of Health and Social Security
- CNS
- Agence eSanté
- Key Healthcare Providers
- Union of Sickness Funds (Union des Caisses de Maladie UCM)
- IGSS
- LNDS
- CNER
- Health Information Security Body (Organe de Sécurité Informatique en Santé OSIS)
- Professional Associations and Trade Unions
- Patient (Associations)
- Health Professions Council (Conseil Supérieur des Professions de Santé)
- Health Scientific Council (Conseil scientifique du domaine de la santé)
- Health Inspection (Division de l'Inspection Sanitaire)
- National Health Observatory (L'Observatoire national de la santé)
- The Pharmaceutical Industry
- Academic and Research Institutions
- Public Health Institute (Direction de la Santé Publique)

4.1 CONCLUSION

The healthcare system in Luxembourg operates with a multi-stakeholder approach, where governmental bodies regulate and fund healthcare, providers deliver services and patients have a say in their care. Research institutions introduce health innovations. The system is built on the principles of universal health coverage, with significant emphasis on data privacy, patient rights and quality healthcare delivery. These stakeholders work collaboratively to ensure that healthcare services in Luxembourg are efficient, equitable and accessible to all.

The success of a unified eHealth strategy in Luxembourg relies on this collaboration across all stakeholders, each contributing their expertise to balance regulatory compliance, technical requirements and operational needs. By working together, these stakeholders ensure that eHealth solutions are secure and lead to overall better health outcomes.



5. REGULATORY AND COMPLIANCE CONSIDERATIONS

Different European- and national regulations and directives need to be adhered – and are of relevance – to the (e)Health system in Luxembourg.

This chapter will briefly highlight the different regulations and directives, without going into details about their different key requirements and aspects, and how they are of relevance to (e)Health.

These details can be found in Annex II, in a table format.

The different regulations and directives are:

- EHDS European Health Data Space, Regulation (EU) 2025/327
- GDPR General Data Protection Regulation, (EU) 2016/679
- NIS2 Network and Information Security 2 Directive, (EU) 2022/2555
- ePrivacy Directive, (EU) 2002/58/EC
- Patients' Rights in Cross-Border Healthcare, Directive 2011/24/EU
- DGA Data Governance Act, Regulation (EU) 2022/868
- Al Act, Regulation (EU) 2024/1689
- MDR Medical Device Regulation, (EU) 2017/745
- IVDR In Vitro Diagnostic Regulation, (EU) 2017/746
- Luxembourg Data Protection Law (Law of 1 August 2018)
- Law on Electronic Communications and Data Security (Law of 30 May 2005, as amended)
- Law on Hospitals and Medical Establishments (Law of 8 March 2018)
- Law on Patient Rights and Obligations (Law of 24 July 2014)
- Law on Health Professions (Law of 10 August 1991, as amended)

5.1 CONCLUSION

The GDPR, EHDS and other EU regulations like the NIS2, DGA, AI Act, IVDR and MDR create a strong legal and operational framework for data protection, sovereignty and secure (usage of) health data (exchange) in the EU. Together, they balance the need for efficient and accessible healthcare across the EU with the protection of sensitive health data, ensuring that patients' rights to privacy and security are upheld while enabling innovation and improved healthcare outcomes.

Luxembourg's legal framework for data protection and healthcare regulations is well-developed, integrating both national laws and EU directives. The framework ensures that health data is processed securely, patients' rights are protected and healthcare providers comply with stringent data security and privacy standards. National authorities like the CNPD, Ministry of Health and Social Security and IGSS play critical roles in overseeing compliance, while healthcare providers, insurance bodies and eHealth platforms like Agence eSanté are required to maintain high levels of data protection and operational efficiency.



6. CHALLENGES IN HARMONIZING COMPLIANCE

Harmonizing compliance in Luxembourg's healthcare system, especially with regard to data protection and eHealth regulations, presents several challenges. These arise primarily from the need to align national regulations with EU regulations and directives, ensure smooth cross-border healthcare and data exchange and manage the complexities of implementing interoperable digital health systems.

Additionally, healthcare providers, regulators, insurers and eHealth platforms must coordinate closely to ensure the lawful and efficient handling of sensitive health data.

A detailed look at the (regulatory) barriers and challenges in harmonizing compliance will be outlined in the next subchapters.

6.1. BARRIERS TO DATA PROTECTION AND PRIVACY

Challenge: Complexity of GDPR implementation and obtaining ethical approval for (usage of) health data.

The GDPR sets high standards for data protection, particularly for sensitive health data, but applying these standards consistently across various healthcare providers, insurers and eHealth platforms can be difficult. GDPR requires:

- Explicit consent from patients for health data processing.
- Strict rules around data minimization and purpose limitation.
- Right to be forgotten and right to portability, which are harder to manage with complex health records.

6.1.1. POTENTIAL BARRIERS

- Interpretation of GDPR Across Sectors: Different healthcare entities (e.g., hospitals, insurance providers) may interpret the application of GDPR requirements differently, leading to inconsistent compliance practices.
- Data Access and Control: Patients may demand control over their health data, but managing access control across multiple healthcare providers can lead to technical and operational complexities.
- Anonymization and Pseudonymization: Ensuring that health data is properly anonymized or pseudonymized for research or public health purposes without compromising patient privacy is difficult.

In addition to the abovementioned challenges in regards to complying with and implementation/interpretation of GDPR requirements, for the secondary usage of health data, ethical approval (or exemption) is required. The process to request an opinion for a new study, or for an amendment of an existing study is a manual effort and can be rather time-consuming, complex and requires large amounts of paperwork.

While ethical considerations to protect an individual involved in a clinical study is of great importance, timely access to health data for research purposes is equally important and crucial to enhance medical innovations. Both aspects need to be well balanced.

6.1.2. COLLABORATIVE SOLUTIONS

- Joint Guidelines and Standards: Collaborative development of standardized guidelines by CNPD, Agence eSanté, CNER and healthcare providers will ensure uniform GDPR implementation. These guidelines should clarify key aspects like consent management, data retention and anonymization and pseudonymization processes.
- Shared Training Initiatives: Healthcare providers, insurers and tech companies could create joint training programs for staff to ensure a unified understanding of GDPR compliance when handling health data.



- Centralized Consent Management Platforms: Developing a centralized consent management tool
 within the DSP system would allow patients to easily manage their consent, ensuring GDPRcompliant data access across the system.
- Automate the process to request, review and grant, reject or exempt ethical assessments to new
 clinical research studies to make it less cumbersome. Create a link to the Centralized Consent
 Management Platform within the DSP, where individual citizens can consent to sharing their data,
 also for secondary research purposes, either in anonymized or pseudonymized form. Allow for a
 broad consent, that the individual consents that their data can be used for any future medical
 research project, without having to consent for each specific research project.

6.2. INTEROPERABILITY AND DATA EXCHANGE

Challenge: Achieving interoperability across healthcare systems

One of the most significant challenges in eHealth is achieving interoperability between disparate systems. Luxembourg must ensure that health data can be shared securely and efficiently between healthcare providers, insurers and across borders, particularly through platforms like MyHealth@EU.

6.2.1. POTENTIAL BARRIERS

- Different Data Standards: Variations in the data formats and standards used by healthcare providers make it difficult to ensure interoperability between hospitals, clinics and national health databases.
- Cross-Border Healthcare Data Exchange: Differences in regulations and infrastructure between Luxembourg and neighboring countries may slow down cross-border healthcare data sharing.
- Technical Barriers to Integrating Legacy Systems: Older systems used by some healthcare providers may not be easily integrated with new eHealth platforms, requiring significant investments in infrastructure upgrades.

6.2.2. COLLABORATIVE SOLUTIONS

- National Interoperability Framework: Agence eSanté should lead the development of a national interoperability framework in collaboration with Ministry of Health and Social Security, CNS and IT providers. This would standardize data formats, exchange protocols and ensure that all systems can communicate with each other.
- Cross-Border Healthcare Agreements: Luxembourg should actively collaborate with neighboring countries to create specific bilateral agreements that facilitate data exchange under GDPR and cross-border healthcare directives.
- Funding and Support for Legacy System Upgrades: The Ministry of Health and Social Security and IGSS could provide incentives or funding programs for smaller healthcare providers to upgrade outdated systems, ensuring they are compatible with national and EU-level eHealth infrastructures.

6.3 BALANCING DATA SECURITY AND ACCESSIBILITY

Challenge: Ensuring data security while maintaining accessibility for patients and providers

Healthcare data is highly sensitive and maintaining robust security measures (such as encryption and access control), while ensuring that healthcare providers have timely access to data, can be difficult. Cybersecurity risks are increasing, particularly as the reliance on digital health platforms grows.

6.3.1. POTENTIAL BARRIERS

- Striking a Balance Between Security and Usability: Implementing stringent security protocols (e.g., multi-factor authentication, encryption) can slow down access to health records in emergency situations or for cross-border care.
- Cybersecurity Threats: Healthcare organizations are increasingly targeted by cyberattacks.
 Complying with the NIS2 Directive to ensure cybersecurity is a complex and resource-intensive process.



6.3.2. COLLABORATIVE SOLUTIONS

- Cybersecurity Best Practices Across the Sector: Ministry of Health and Social Security, in collaboration with CNPD and Agence eSanté, should establish national cybersecurity guidelines for healthcare providers, ensuring they adopt secure systems while maintaining usability. This could include minimum encryption standards, threat detection systems and secure data-sharing practices.
- Emergency Access Protocols: Develop specific protocols that allow healthcare providers to quickly access patient records during emergencies, while maintaining GDPR and security compliance. This could include specific exemptions for data access during critical care.
- Sector-Wide Cybersecurity Response Teams: Establish cybersecurity response teams within Luxembourg's healthcare sector, consisting of representatives from CNS, hospitals and Agence eSanté. These teams would coordinate efforts to prevent, detect and respond to cybersecurity incidents.

6.4. CROSS-BORDER DATA EXCHANGE AND SOVEREIGNTY

Challenge: Aligning national data protection rules with EU-wide frameworks

While the GDPR provides a unified framework for data protection across the EU, individual countries like Luxembourg may implement certain derogations (exceptions) for national data sovereignty. Coordinating health data exchange for cross-border healthcare (under Directive 2011/24/EU) with varying national interpretations of GDPR poses a challenge.

6.4.1. POTENTIAL BARRIERS

- Differing National Data Laws: While GDPR applies EU-wide, individual member states have different interpretations or national laws that can affect cross-border data sharing, particularly for healthcare.
- Fragmentation in Cross-Border Healthcare Reimbursement Systems: Differences in healthcare reimbursement rules between countries can make it difficult for patients to access care abroad and have their data and treatment reimbursed smoothly.

6.4.2. COLLABORATIVE SOLUTIONS

- Bilateral and Multilateral Data-Sharing Agreements: Luxembourg should work closely with neighboring countries to establish bilateral agreements that outline clear rules for cross-border data exchange. These agreements should also include provisions for aligning healthcare reimbursement systems.
- Standardizing Cross-Border eHealth Systems: Adherence to the EHDS, including MyHealth@EU will ensure that Luxembourg's eHealth platforms are interoperable with other EU member states.
 This will streamline data exchange for cross-border care, particularly in managing ePrescriptions and patient summaries.
- Regular Dialogue on Sovereignty Issues: Establish regular dialogues between Luxembourg's CNPD and the Ministry of Health and Social Security with their counterparts in neighboring countries to address data sovereignty concerns and ensure that GDPR is interpreted consistently, especially in cross-border care scenarios.

6.5. HEALTHCARE PROFESSIONALS AND DIGITAL TRANSFORMATION

Challenge: Ensuring healthcare providers are adequately trained and supported

The adoption of eHealth platforms, telemedicine and other digital health solutions requires that healthcare providers are well-versed in using these technologies, while ensuring compliance with complex legal and regulatory frameworks.



6.5.1. POTENTIAL BARRIERS

- Resistance to Adoption: Healthcare professionals may be hesitant to adopt new digital tools due to a lack of training or concerns about the time and effort required to comply with digital data entry and security procedures.
- Varying Levels of Digital Literacy: Different levels of digital literacy among healthcare workers can lead to inconsistent implementation of eHealth solutions and may create potential compliance gaps, particularly in terms of data security.

6.5.2. COLLABORATIVE SOLUTIONS

- Comprehensive Training Programs: Agence eSanté and professional bodies such as the Health Professions Council should develop comprehensive training and certification programs on the use of eHealth platforms, telemedicine and data protection compliance. These programs should be mandatory for healthcare workers to ensure consistent digital literacy across the sector.
- Integration of Digital Tools in Clinical Practice: Healthcare providers should collaborate with digital health providers and CNS to ensure that eHealth tools are seamlessly integrated into daily workflows. This reduces the burden on healthcare professionals and increases adoption.

6.6. CONCLUSION

Harmonizing compliance in Luxembourg's healthcare system, particularly around data protection, interoperability and eHealth adoption, presents several challenges. These challenges are rooted in balancing GDPR requirements, national sovereignty and the operational needs of healthcare providers.

Through collaboration among stakeholders — including the Ministry of Health and Social Security, CNPD, CNS, Agence eSanté, healthcare providers and patients — Luxembourg can overcome these barriers. Coordinated efforts in training, data-sharing agreements, cybersecurity and interoperability standards will ensure a unified, compliant and secure digital healthcare system that aligns with both national and European regulations.



7. PROPOSED APPROACH

Harmonizing compliance across all stakeholders in Luxembourg's healthcare system, particularly concerning data privacy, consent management and cross-border health data sharing, requires a structured pathway that integrates regulatory frameworks, technological solutions and collaborative governance. The goal is to align the operations of healthcare providers, insurers, government bodies and patients with both Luxembourg's national laws and EU regulations, ensuring that health data is securely handled, accessible when needed and properly managed across borders.

The proposed approach for harmonizing compliance is descript in the next sections.

7.1. ESTABLISH A SELF-LEARNING HEALTH SYSTEM

Objective: Create a self-leaning healthcare system, in which high-quality health data is (securely and in adherence with all relevant regulations) transferred from care- to research institutions, accelerating medical innovation and returning new insights back to the healthcare providers to enhance patient wellbeing.

The two concrete use-cases as part of the DS4H PoC, for diabetes and oncology respectively, intend to demonstrate how exchange of health data from care- to research institutions allows for data-driven (AI) innovations, from which the actual patient seen in the care institute can ultimately benefit (in the future).

7.1.1. KEY ACTIONS

- 1. Promote a Culture of Continuous Learning:
 - Foster a culture of continuous learning and improvement among healthcare providers.
 - Encourage the use of data and analytics to drive decision-making and improve patient care.
- 2. Engage Stakeholders in the Self-Learning Process:
 - Involve healthcare providers, researchers and patients in the self-learning process.
 - Create feedback loops to continuously gather and apply evidence in real-time to guide care.
- 3. Enhance Data Governance and Security:
 - Strengthen data governance policies to ensure the secure and ethical use of health data.
 - Implement robust security measures to protect patient data and maintain compliance with regulatory standards.
- 4. Foster Collaboration and Innovation:
 - Encourage collaboration among healthcare providers, researchers, and technology partners.
 - Support innovative projects that leverage data-driven insights to improve healthcare delivery and patient outcomes.

7.1.2. BENEFITS

This will close the loop between primary use of health data on one hand, and secondary use of health data on the other hand – and thus goes beyond EHDS.

7.2. ESTABLISH A MULTI-STAKEHOLDER GOVERNANCE FRAMEWORK

Objective: Create a centralized governance structure to oversee the alignment of data privacy, consent management and cross-border data sharing across all stakeholders, ensuring transparency, accountability and consistency.



7.2.1. KEY ACTIONS

7.2.1.1. FORM A NATIONAL EHEALTH STEERING COMMITTEE

Members: Ministry of Health and Social Security, CNS, Agence eSanté, CNPD, Healthcare Providers, Patients' Associations and Professional Councils.

Role: Oversee the harmonization of regulatory compliance, propose unified standards and ensure continuous alignment between national and EU regulations (e.g., GDPR, EHDS).

Responsibilities:

- Develop a national roadmap for eHealth compliance, including data privacy, consent management and cross-border sharing.
- Monitor the integration of digital health tools and ensure ongoing collaboration between stakeholders.

7.2.1.2. APPOINT A COMPLIANCE TASK FORCE

Role: Ensure GDPR and EHDS compliance across all sectors.

Responsibilities: Provide recommendations on how to integrate best practices on data protection into daily healthcare operations, oversee audits and address potential regulatory barriers.

7.2.2. BENEFITS:

This centralized governance framework will streamline decision-making and ensure that all stakeholders work together under a unified approach, preventing fragmented compliance efforts.

It is expected that over time, in line with the EHDS implementation, this will be addressed (to a certain extend).

7.3. DEVELOP A NATIONAL STANDARD FOR CONSENT MANAGEMENT

Objective: Implement a centralized consent management system that complies with GDPR requirements and allows patients to control access to their health data across different healthcare providers and systems, both nationally and cross-border.

7.3.1. KEY ACTIONS

7.3.1.1. IMPLEMENT A CENTRALIZED CONSENT MANAGEMENT SYSTEM (CMS)

Lead: Agence eSanté in collaboration with CNPD and Ministry of Health and Social Security.

Role: Enable patients to manage their consent for data access in real-time, through an online platform (integrated with the DSP).

Functionality:

- Patients can view, grant or withdraw consent for healthcare providers to access specific health records. Also for secondary usage of their health data, patients should have the possibility within the CMS to content (or not) to sharing their data for research purposes, either in anonymized or pseudonymized form, as well as being able to choose to give consent for a specific research project, or rather a broad consent.
- Patients can give consent for cross-border sharing of health data in compliance with Directive 2011/24/EU.
- Consent status is updated across all healthcare providers in real-time to ensure that only authorized personnel has access to patient data.



7.3.1.2. STANDARDIZE CONSENT FORMS AND PROCEDURES

- Develop standardized, multilingual consent forms (aligned with GDPR) for both national and crossborder healthcare.
- Ensure that patients understand their rights and the scope of data sharing, with simple opt-in/optout mechanisms.

7.3.1.3. CREATE AWARENESS CAMPAIGNS FOR PATIENTS AND PROVIDERS

- Run public awareness campaigns to educate patients on how to manage their health data and rights under GDPR.
- Train healthcare providers on how to integrate the CMS into their workflows.

7.3.2. BENEFITS

A centralized consent management system provides transparency for patients, enhances their control over personal data and ensures legal compliance across national and cross-border healthcare environments.

Although patient consent will be a central aspect of the EHDS, this proposal will go beyond the expected minimal requirements from the regulation.

7.4. ENSURE INTEROPERABILITY AND SECURE CROSS-BORDER HEALTH DATA SHARING

7.4.1. OBJECTIVE

Enable interoperable and secure health data exchange within Luxembourg and across EU borders, ensuring that healthcare providers can access and share patient data securely when needed, particularly for cross-border healthcare services.

7.4.2. KEY ACTIONS

7.4.2.1. ADOPT COMMON DATA STANDARDS FOR INTEROPERABILITY

Lead: Agence eSanté and Ministry of Health and Social Security, in collaboration with EU bodies (e.g., MyHealth@EU)

- Implement EU-wide health data standards (HL7 FHIR, ICD, SNOMED CT) for EHRs to ensure interoperability and compatibility with cross-border health systems.
- Ensure that all healthcare providers in Luxembourg adopt these standards for easy data sharing across systems.

7.4.2.2. STRENGTHEN CROSS-BORDER DATA EXCHANGE INFRASTRUCTURE

- Integrate Luxembourg's national eHealth systems (DSP, ePrescriptions) with the MyHealth@EU platform to facilitate secure cross-border data sharing.
- Ensure real-time exchange of patient summaries, ePrescriptions and diagnostic data with healthcare providers in other EU member states, ensuring compliance with GDPR and EHDS.
- Adopt EU Digital COVID Certificate infrastructure as a model for future cross-border health data sharing.

7.4.2.3. ESTABLISH CROSS-BORDER DATA-SHARING AGREEMENTS

- Work with neighboring countries to establish bilateral or multilateral agreements that define clear protocols for secure data sharing in line with GDPR and national data sovereignty laws.
- These agreements should include specific provisions on data handling, consent requirements and secure data transfer protocols.



7.4.2.4. ENHANCE DATA SECURITY MEASURES

- Implement end-to-end encryption for all cross-border data exchanges, ensuring secure data transmission and storage.
- Adopt multi-factor authentication (MFA) and role-based access control for healthcare providers accessing patient data from other EU countries.
- Regularly audit cross-border data-sharing processes to identify vulnerabilities and ensure compliance with NIS2 Directive on cybersecurity.

7.4.3. BENEFITS

Interoperable and secure data sharing ensures that patients receive seamless healthcare, whether in Luxembourg or abroad. This approach promotes continuity of care while maintaining stringent data protection standards.

It is expected that this proposal is within the scope of the future EHDS implementation.

7.5. CREATE A COMPREHENSIVE TRAINING PROGRAM FOR STAKEHOLDERS

7.5.1. OBJECTIVE

Equip healthcare providers, administrative staff and IT personnel with the knowledge and skills required to manage patient data securely, comply with GDPR and utilize cross-border health data sharing systems effectively.

7.5.2. KEY ACTIONS

7.5.2.1 DEVELOP EHEALTH COMPLIANCE TRAINING MODULES

Lead: Health Professions Council and CNPD, in collaboration with Agence eSanté.

- Develop tailored training programs for different stakeholder groups (healthcare providers, insurers, IT staff) on topics such as:
 - GDPR compliance for health data.
 - Use of the centralized consent management system.
 - o Secure data-sharing practices, both nationally and cross-border.
 - o Cybersecurity protocols for protecting health data.

7.5.2.2. CONDUCT MANDATORY WORKSHOPS AND CERTIFICATIONS

- Ensure that all healthcare providers undergo mandatory training and certification on data privacy, consent management and cross-border data sharing.
- IT professionals responsible for maintaining eHealth platforms should receive specialized training on cybersecurity and data encryption technologies.

7.5.2.3. CONTINUOUS EDUCATION AND UPDATES

- Provide regular updates and refresher courses on emerging trends in data protection and new regulatory requirements (e.g., updates to EHDS or GDPR amendments).
- Collaborate with EU bodies to ensure alignment with new cross-border healthcare protocols.

7.5.3. BENEFITS

This ensures that all stakeholders are equipped with the necessary skills to manage patient data responsibly, comply with regulations and use digital health tools effectively, minimizing compliance risks. It is expected that this goes beyond the EHDS Implementation.



7.6. ESTABLISH CONTINUOUS MONITORING, AUDITING AND FEEDBACK MECHANISMS

7.6.1. OBJECTIVE

Set up continuous monitoring and auditing processes to ensure that data privacy, consent management and cross-border data-sharing protocols are followed consistently and adjusted as necessary.

7.6.2. KEY ACTIONS

7.6.2.1. IMPLEMENT A COMPLIANCE MONITORING SYSTEM

Lead: CNPD in collaboration with the Ministry of Health and Social Security and Agence eSanté.

- Develop a real-time monitoring system that tracks data access, consent updates and data-sharing activities to detect non-compliance or security risks.
- Set up automated alerts for unauthorized access attempts or breaches of GDPR rules.

7.6.2.2. CONDUCT REGULAR AUDITS AND ASSESSMENTS

- Perform annual audits of healthcare providers, insurers and eHealth platforms to ensure adherence to GDPR, EHDS and cross-border data-sharing regulations.
- Use audit findings to make continuous improvements to the consent management system, data security protocols and interoperability standards.

7.6.2.3. FEEDBACK LOOPS WITH STAKEHOLDERS

- Create regular feedback sessions with patients, healthcare providers and regulators to gather insights on the effectiveness of the compliance framework.
- Use feedback to refine data-sharing protocols, update consent management tools and address emerging compliance challenges.

7.6.3. BENEFITS

Continuous monitoring, auditing and feedback mechanisms ensure that the system remains compliant with evolving regulations and that stakeholders are consistently aligned with best practices in data privacy and security.

It is expected that over time, in line with the EHDS implementation, this will be addressed (to a certain extend).

7.7. CONCLUSION: A PATHWAY FOR HARMONIZING COMPLIANCE

By establishing a self-learning healthcare system and a centralized governance framework, implementing a centralized consent management system, ensuring interoperability and secure cross-border data sharing, providing comprehensive training and setting up monitoring mechanisms, Luxembourg can harmonize compliance across all stakeholders. This approach not only ensures GDPR and EHDS compliance but also fosters trust and collaboration among healthcare providers, patients and national authorities, enabling efficient and secure healthcare delivery.



8. CONCLUSION

This concluding chapter summarizes the key outcomes, opportunities and proposed approaches to overcome challenges, which will ensure the successful implementation of the DS4H project and *may* lay the grounds for the future EHDS implementation.

The DS4H project represents a significant step forward in Luxembourg's journey towards a more integrated, efficient and secure healthcare system. By leveraging the EHDS and Gaia-X initiatives, the project aims to create a harmonized Health Data Space that facilitates secure data exchange, enhances healthcare delivery and fosters medical innovation.

The EHDS and Gaia-X have common goals, notably in regards to data sovereignty, security and privacy and interoperability aspects. EHDS *could* rely on Gaia-X to be the technical backbone by providing the technical infrastructure for storing, processing and exchanging health data across borders.

In the context of Luxembourg, both in regards to the digital health infrastructure as well as relevant stakeholders, there is a large variety of entities, both public and private, with different roles and responsibilities, collaborative needs and balancing aspects. The limited size of the country allows for shorter lines of communication (and constructive collaboration), making Luxembourg uniquely positioned to pioneer in creating the Health Data Space.

Even though there are numerous European- and Luxembourgish regulations that need to be adhered to, also through the upcoming EHDS, there is a significant paradigm shift to in a secure, sensible, ethical and meaningful way create value from usage of health data with the aim to ultimately improve patient care.

Identified challenges include barriers to data protection and privacy, interoperability and data exchange, balancing data security and accessibility, cross-border data exchange and sovereignty as well as healthcare professionals within the digital transformation.

Approaches to overcome these challenges consist of establishing a self-learning health system as well as a multi-stakeholder governance framework, developing a national standard for consent management, ensuring national interoperability and secure cross-border health data sharing, creating a comprehensive training program for stakeholders and finally by establishing continuous monitoring, auditing and feedback mechanisms.

In conclusion, there is a large long-tern potential for Luxembourg('s healthcare system) by creating a Health Data Space in Luxembourg with a clear governance and by adhering to all relevant regulations. The Health Data Space stimulates and allows for constructive collaboration between both care institutions amongst each other (and thus primary use of health data), between care- and research institutions (and thus secondary use of health data) and finally returning medical innovation from research- to care institutions (and thus closing the loop between primary- and secondary use of health data). This not only has the potential to ultimately improve patient care, but also to foster innovation and thus allows economic growth and attracting investment.



ANNEX I

Who	Role	Responsibility	Collaboration Needs	Balancing aspects
Ministry of Health and Social Security	The central governmental authority responsible for setting health policy, regulating the healthcare system, overseeing	To supervises hospitals, healthcare professionals, medical products and public health campaigns. It also leads the response to health crises (e.g., pandemics) and ensures the	The Ministry must provide clear leadership and policy direction while ensuring that regulations related to data privacy, security and public health are enforced. It works with CNS, Agence eSanté and Healthcare Providers to integrate these policies into day-to-day healthcare operations. The Ministry also needs to collaborate with Professional- and Patient Associations and the Pharmaceutical Industry to ensure that the needs of all stakeholders are considered when establishing eHealth policies.	The Ministry balances regulatory compliance with innovation, ensuring that Luxembourg's healthcare system embraces new technologies without compromising on quality and safety.
CNS	To manage healthcare reimbursements for medical services, prescriptions, hospital stays and other healthcare-related expenses. The CNS also negotiates tariffs with healthcare providers and ensures that medical services remain accessible and affordable. CNS needs to work with Healthcare Providers, the Ministry of Health and Soc Security and the UCM to streamline health data exchange for accurate billing and reimbursements. Collaboration with Agence eSanté is necessary to exproviders and ensures that medical services remain accessible and affordable. CNS needs to work with Healthcare Providers, the Ministry of Health and Soc Security and the UCM to streamline health data exchange for accurate billing and reimbursements. Collaboration with Agence eSanté is necessary to expression and clinical data are synchronized. CNS also interacts with Patients' Association in prove access to digital tools for managing insurance claims.		CNS must balance operational efficiency in healthcare reimbursements with maintaining affordability and accessibility for patients, while aligning with regulatory frameworks for data protection.	
Agence eSanté	Agence eSanté must collaborate with all stakeholders, particularly CNS, To oversee the secure exchange of patient health data, facilitate interoperability between healthcare systems, ensure data privacy and support eHealth initiatives like telemedicine. In regards to the EHDS, Agence eSanté is responsible for the primary use of data. Agence eSanté must collaborate with all stakeholders, particularly CNS, Healthcare Providers and the Ministry of Health and Social Security, to build and maintain a robust, secure and interoperable eHealth platform. It ensures that different systems (clinical, financial and administrative) can communicate with each other. Close cooperation with Professional Associations and Health Professions Council is also necessary to ensure healthcare providers are trained in and using the eHealth systems effectively.		Agence eSanté must balance technical demands for system's interoperability and data security with the operational needs of healthcare providers and regulatory compliance.	
	Public and private hospitals, specialized centres To provide secondary and tertiary care services. To act as the first point of contact for patients. To play a critical			
		role in diagnosing and referring patients to specialists or hospitals. To provide secondary care either within hospitals or in private	Healthcare providers must collaborate with the Ministry of Health and Social Security, CNS and Agence eSanté to ensure that health data is accurately	
	Specialists	practices. To dispense medications, provide advice on drug use and	recorded, securely shared and accessible for patient care. Providers work closely with Professional- and Patient Associations to ensure the medical community's	, Providers must balance patient care with compliance to eHealth
Key healthcare providers	Pharmacies	contribute to public health efforts such as vaccinations. To take and analyse clinical specimens to obtain information	interests are aligned with patient safety and satisfaction. Cooperation with Pharmaceutical Industry is also necessary to ensure the smooth digital	protocols, using technology to improve service delivery while safeguarding patient data.
	Private and public laboratories	about the health of a patient to aid in diagnosis, treatment, and prevention of disease.	integration of prescription and medication data. Finally, healthcare providers collaborate with the Health Professions Counsil for licensing and qualifications	sareguarding patient data.
	Different institutions (elderly care, disabilities etc.)	To assist with activities of daily living, such as bathing, dressing, and medication management.	and with the Health Scientific Council for good medical practices.	
	Other healthcare professionals	To carry out various tasks to contribute to the healthcare system.		
исм	To coordinate the different sickness funds in Luxembourg.		UCM coordinates with CNS, Healthcare Providers and the Ministry of Health and Social Security to standardize health insurance policies, reimbursement procedures and rates. This collaboration ensures consistency across different insurance funds and smooth integration with the broader eHealth infrastructure. UCM must also work with Patients' Associations to ensure that reimbursement and insurance systems are patient-friendly.	UCM must balance the financial sustainability of the insurance system with ensuring equitable access to healthcare for all citizens.
IGSS	To oversee the functioning of the social security system, including healthcare. To monitor the financial stability and performance of the healthcare with national including healthcare. The social security system, ensuring compliance with national functioning efficiently. Collaboration with Healthcare Providers and		IGSS works with CNS, UCM and the Ministry of Health and Social Security to ensure financial oversight and that healthcare and social security systems are functioning efficiently. Collaboration with Healthcare Providers and Agence eSanté ensures that accurate data is collected for reporting, analysis and policy- making.	IGSS must balance financial oversight with operational flexibility to adapt to new healthcare models, while ensuring the long-term sustainability of the social security system.
LNDS	To implement Luxembourg's strategies in research, innovation, and digitalization.	To enable value creation from secondary use of data, for public and private partners and support the sharing and re-use of public sector data, in a trustable manner. In regards to the EHDS, LNDS has a leading role for the country's implementation in terms of secondary use of data.	LNDS collaborates closely with government, (healthcare) industry, research institutions and citizens to create impactful data projects.	LNDS must balance the need for open and accessible data with the requirement to protect personal data and ensure privacy. This involves implementing robust data protection measures and adhering to regulations such as the GDPR.
CNER	The national ethics body for health studies.	To protect persons participating in a health study by providing an opinion concerning the ethical acceptability of projects (such as clinical trials) submitted to it.	The CNER collaborates with the CNPD, as well as with Research Organizations and Key Healthcare Providers (who act as Principal Investigator within clinical trial studies).	The CNER needs to balance ethical considerations by protecting individuals who are participating in health studies, while allowing medical research to advance.
OSIS	OSIS was created by the Ministry of Health within a national cybersecurity framework. Its goal is to facilitate discussions on cybersecurity and the various entities involved.	To define both general- and specific cybersecurity guidelines for hospitals and "transversal" structures (such as LUXITH or eSanté).	OSIS works closely with other government agencies, such as the Ministry of Health, CNS, eSanté, FHL and LUXITH, to coordinate cybersecurity efforts and share best practices.	OSIS must balance the need for stringent data protection measures with the requirement to keep health information accessible to authorized personnel. This involves implementing security protocols that do not hinder the day-to-day operations of healthcare providers.
	AMMD To represents doctors and dentists and involved in negotiations regarding tariffs and working conditions. To represents the interests of hospitals and healthcare institutions in Luxembourg, Healthcare Workers' Unions To represent various healthcare professionals, advocating for		Professional associations and Trade Unions must work with Healthcare Providers, the Ministry of Health and Social Security and Agence eSanté to	
Professional Associations and			ensure that the needs and concerns of healthcare professionals are reflected in	These associations must balance advocacy for professionals' rights with supporting the integration of eHealth innovations to
Trade Unions			eHealth policies. They play a crucial role in training healthcare professionals to use eHealth systems effectively and in advocating for working conditions that	improve healthcare outcomes.
		their working conditions, salaries and professional development. At the centre of the healthcare system, with rights to access high-	support eHealth adoption.	
	Patients	quality care, reimbursement for medical services and involvement in healthcare decisions. To represent patients with specific health conditions, advocating	Patients and their associations need to work with the Ministry of Health and Social Security, CNS and Healthcare Providers to ensure patient rights, privacy	Patients' associations balance advocacy for better healthcare services with supporting digital transformation efforts that improve convenience and efficiency for patients.
Patient (Associations)	Patient Associations	for better care, support and research.	and safety are protected. They also provide feedback on eHealth tools, such as patient portals and digital records and collaborate with Agence eSanté to	
	National Health Information and Mediation Service	To provide information, advice, as well as conflict prevention and resolution between healthcare providers and patients.	improve user-friendliness and accessibility.	
Conseil Supérieur des Professions de Santé	Regulating the licensing, qualifications and ethical practices of healthcare professionals, including doctors, nurses and pharmacists.	To ensure that healthcare providers meet the required standards for professional practice and patient safety.	The Health Professions Council collaborates with the Ministry of Health and Social Security, Healthcare Providers and Professional Associations to ensure that healthcare professionals meet training standards for using eHealth systems It also ensures that regulations governing the use of digital tools by healthcare professionals are in place.	The Council balances the need for high standards in healthcare with the adoption of new technologies that require ongoing professional development.
Conseil scientifique du domaine de la santé	To develop and contribute to the implementation of standards of good medical practice.	To promote high-quality care, to guide healthcare professionals in the development of good practices and to make optimal use of available resources.	Collaboration with various government departments and agencies to ensure policies are evidence-based, working with healthcare providers to implement best practices as well as partnering with research institutions to foster innovation and scientific advancements.	Ensuring that new medical technologies and treatments are safe and effective before widespread use.
Division de l'Inspection Sanitaire	Responsible for inspecting and ensuring the hygiene, safety and compliance of healthcare institutions.	To oversee the licensing of healthcare facilities, investigate complaints and enforce public health laws.	Health Inspection works with Healthcare Providers, the Ministry of Health and Social Security and Agence eSanté to ensure that health facilities comply with eHealth regulations, data security standards and hygiene protocols. It also ensures that any newly introduced eHealth solution meets public health standards.	It must balance regulatory enforcement with supporting the adoption of eHealth innovations in clinical settings.
L'Observatoire national de la santé	To guide health decisions and policies and assess their impact by networking data.	To evaluate the population health status, publish and disseminate these findings and to propose improvements to the population's health status and the health system.	Collaboration with health ministries and other government agencies to ensure data is used effectively in policymaking, working with healthcare providers to gather accurate and timely data and partnering with research institutions to conduct studies and analyses.	Balancing the need for comprehensive data analysis with the need for timely reporting as well as ensuring that both local and national health trends are adequately addressed.
The Pharmaceutical Industry	To provide medications and treatments to healthcare providers in Luxembourg, Involvement in research, development, manufacturing and the distribution of pharmaceutical products, as well as negotiating pricing and reimbursement conditions with the CNS.		The pharmaceutical industry collaborates with Healthcare Providers, CNS and Agence eSanté to integrate digital solutions such as ePrescriptions into the healthcare system. It must also work with the Ministry of Health and Social Security and regulatory bodies to ensure compliance with drug safety and digital health regulations.	The industry must balance innovation in digital healthcare (e.g., digital drug tracking) with ensuring safety and regulatory compliance in a highly sensitive industry.
Academic and Research To contribute to medical research, healthcare innovation and the Institutions training of healthcare professionals.		health data and medical science.	Research institutions collaborate with the Ministry of Health and Social Security, Healthcare Providers and Agence eSanté to develop and test new digital health technologies and analyse health data. They work with Healthcare Providers and the population directly to obtain health data. They collaborate with CNER to obtain ethical approval. They also provide the educational support necessary for healthcare workers to adopt eHealth solutions effectively.	Institutions must balance innovation and research with practical
Direction de la Santé Publique	To define public health objectives and contribute to the national health strategy.	To develop and manage national health plans and interact with all healthcare system partners. To provide awareness, screening, and surveillance services. To ensure access, quality, and safety of healthcare. To guarantees compliance with applicable laws, regulations, and standards. To support the development of effective public health policies.	Public health bodies work with the Ministry of Health and Social Security, CNS and Healthcare Providers to gather data from eHealth systems for public health monitoring, disease surveillance and crisis response. These bodies ensure that eHealth solutions are used effectively for public health purposes, such as managing pandemics.	These bodies balance real-time health surveillance with protecting patient privacy and data security.



ANNEX II

Level	What	Brief description	Key Requirements/aspects	Relevance for eHealth
European	EHDS - European Health Data Space, Regulation (EU) 2025/327	To create a framework for the secure and efficient sharing of health data within and across EU member states. It is designed to facilitate both primary use (clinical care) and secondary use (research, policymaking and innovation) of health data, while ensuring high levels of privacy and security.	Empowerment of Individuals (access to portable personal health data), Improved Healthcare Delivery (cross-border healthcare and interoperability standards), Data for Research and Innovation (secondary use of health data and data-driven health solutions) and Privacy and Security (GDPR compliance and data sovereignty)	The EHDS plays a pivotal role in advancing eHealth by creating a secure, interoperable and patient-centric framework for health data exchange across the EU.
European	GDPR - General Data Protection Regulation, (EU) 2016/679	The cornerstone of EU data protection law. It applies to all organizations processing personal data within the EU, including health data	Lawfulness, Fairness and Transparency, Data Minimization, Consent and Legal Basis, Data Security and Confidentiality, Right of Access, Rectification and Erasure, Data Portability, Data Sovereignty and Local Laws and requirement of an Data Protection Officer	To ensure that patients' personal health data is processed securely and transparently. To emphasize obtaining explicit consent from patients for the use of their data, particularly when sharing data across healthcare providers or borders
European	NIS2 - Network and Information Security 2 Directive, (EU) 2022/2555	Directive to strengthen cybersecurity in key sectors, including healthcare	Cybersecurity Measures, Incident Reporting and Risk Management and Governance	The NIS2 Directive ensures that the digital infrastructure supporting eHealth is secure, minimizing the risk of cyberattacks that could compromise sensitive health data or disrupt healthcare services.
European		Directive focused on privacy and electronic communications, complementing the GDPR by specifically addressing the confidentiality of communications and the tracking of user behaviour online	Explicit User Consent for cookie usage, email marketing, data minimization and other aspects of data privacy	To emphasize the confidentiality of communications, requiring consent for cookie usage, promoting data minimization and ensuring transparency in data processing. It also aligns with the GDPR to provide a robust framework for data protection in the EU.
European	Patients' Rights in Cross-Border Healthcare, Directive 2011/24/EU	Directive to facilitate patients' access to cross-border healthcare and to reinforce patients' rights in relation to receiving healthcare services in other EU countries	Access to Health Data Across Borders and reimbursements for cross-border healthcare services	The directive supports the exchange of medical data and the continuity of care for patients traveling or living in multiple EU countries. It is closely connected with the goals of the EHDS and GDPR in terms of data portability and security.
European	DGA - Data Governance Act, Regulation (EU) 2022/868	Regulation that complements the GDPR by promoting trustworthy mechanisms for data sharing within and across sectors, including healthcare	Data Intermediaries and Data Sovereignty	The DGA supports the governance of health data exchange, particularly for research and innovation, while ensuring that data protection principles are maintained.
European	Al Act, Regulation (EU) 2024/1689	A comprehensive legal framework introduced by the EU to regulate the development and use of AI systems. It aims to foster innovation while protecting individuals from the potential harms of AI.	Risk Classification, Transparency and Disclosure, Risk Management, Conformity Assessments, Prohibited Practices, AI Literacy and Governance and Compliance	In relation to the healthcare sector, the AI Act has significations implications, including the regulation of high-risk AI systems, the need for transparency and patient consent, and the integration with existing regulations. It aims to promote innovation while ensuring the safe and ethical use of AI in healthcare.
European	MDR - Medical Device Regulation, (EU) 2017/745	A comprehensive framework introduced by the EU to ensure the safety and efficacy of medical devices.	Classification of Devices, Unique Device Identifier, General Safety and Performance Requirements, Clinical Evaluation and Post-Market Clinical Follow-Up, Quality Management System, Notified Bodies, Technical Documentation, Vigilance and Post- Market surveillance, EUDAMED Database, Transparency and Traceability and Increased Scrutiny on High-Risk Devices	The introduction of the MDR resulted in strict regulatory and approval processes, classification and regulatory control, quality and safety regulations, innovation and compliance, a global regulatory landscape, post-market surveillance and interdisciplinary collaboration. These aspects aim to enhance the safety, efficacy, effectiveness and quality of medical devices, ensuring that patients and healthcare providers have access to high-quality, reliable medical devices.
European	IVDR - In Vitro Diagnostic	The In Vitro Diagnostic Medical Devices Regulation (EU) 2017/746 (IVDR) governs the placing on the market and use of in vitro diagnostic devices within the European Union. It replaces Directive 98/79/EC and aims to enhance patient safety by introducing stricter requirements for clinical evidence, risk classification, transparency, and postmarket surveillance.	The IVDR introduces a risk-based classification system, stricter conformity assessments, and reinforced clinical performance evaluation. It requires EUDAMED registration and post-market surveillance. In-house devices are allowed under specific conditions if no CE-marked equivalent exists.	The IVDR applies to diagnostic software, including AI tools. Hospital-developed digital tests fall under Article 5(5). The regulation promotes traceability, security, and interoperability with health IT systems,
National	Luxembourg Data Protection Law (Law of 1 August 2018)	Law that complements the GDPR and adapts specific provisions for Luxembourg's legal context, providing additional national-level enforcement and guidance.	The law includes derogations for the processing of health data for research, public health and archival purposes under strict conditions. Healthcare institutions must ensure that DPOs are appointed to oversee compliance and regularly audit data-handling procedures. The CNPD has the authority to impose significant fines for non-compliance. Penalties can be severe, especially in the case of mishandling sensitive health data.	
National	and Data Security (Law of 30 May 1	Law that ensures that all electronic communication involving health data (e.g., telemedicine, remote monitoring) maintains strict confidentiality.	Providers offering telehealth or digital consultations must adhere to secure communication standards to protect sensitive health information during remote exchanges.	
National	Law on Hospitals and Medical Establishments (Law of 8 March 2018)	Law that regulates the organization, operations and inspection of hospitals and medical institutions. It ensures that hospitals have appropriate digital systems in place to manage patient data securely and comply with Luxembourg's data protection standards.	Hospitals must integrate with national eHealth systems like the DSP to ensure that patient data can be shared securely across the healthcare network for better coordination of care. The law furthermore mandates that hospitals adopt standardized EHR systems that are interoperable with national eHealth infrastructure. This ensures seamless sharing of patient records between healthcare professionals	
	Law on Patient Rights and Obligations (Law of 24 July 2014)	Law that allows patients to access their personal health records, including electronic records. This applies to data stored in national systems like the DSP or in individual hospitals. Patients can request that healthcare providers provide information on how their health data is being used, ensuring transparency	This law reinforces the principle that patients must give informed consent before their health data is processed or shared, except in emergency situations where patient consent cannot be obtained. The law also mandates that patients are informed about their rights regarding their data, including how to request access or correction	