

Patient Summary Evaluation Form for the Functional e2e testing MyHealth@EU Spring 2025 Test Session

Fields marked with * are mandatory.



My health @ EU
eHealth Digital Service Infrastructure
A service provided by the European Union

Evaluation Form for the Functional end-to-end Testing - Patient Summary

Introduction

The functional end-to-end testing validates, from the user point of view, the process and the information presented to health professionals using the MyHealth@EU services.

The evaluation is carried out for all the services in an environment that intends to emulate the normal operation as much as possible. For more details on the methodology of the testing, please check the dedicated page [here](#).

The tool used to collect the feedback from the health professionals and semantic experts participating in the testing is the Evaluation Form, which is provided using the European Commission online tool EUSurvey.

Thanks very much for your participation.

1 Information on the participants in the evaluation

* 1.1 Please, indicate the NCPeH you are representing

IE - Ireland

* 1.2 Role of the Healthcare Professional

- ☐ Doctor of medicine
☒ Nurse

- ☐ Dental practitioner
- ☐ Midwife
- ☐ Pharmacist
- ☐ Pharmacist technician
- ☐ Other

* 1.4 Professional role of the Semantic Expert

Technical Architect

2 Information on the Evaluation

* 2.1 Date of the evaluation

11/04/2025

* 2.2 Country of origin of the PS document being evaluated

LU - Luxembourg

3 Identification of the Patient Process

3.1 Identification of the Patient

3.1.1 Could the patient be identified using the fields provided by the country of origin?

(whether the returned information was sufficient to identify the patient)

- ☒ Yes
- ☐ No

3.1.3 Did you encounter any technical error during the Patient Search?

- ☐ Yes
- ☐ No

3.2 Identification of Authorized Third Party (Next of Kin)

(optional; Section only applicable when the identification of Authorized Third Party is supported by the implementations in both countries participating in the test)

3.2.1 Is your country implementing the identification of the Authorized Third Party (Next of Kin) operation?

- ☐ Yes
- ☒ No

4 Non-Functional Requirements

* 4.1 Was the system available during the performance of the test?

- ☒ Yes
☐ No

4.3 Response Time

	>15s	<15s	<10s
* To find a patient	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>
* To retrieve the document	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>

* 4.4 What is your perception of the response time?

- ☐ Good
☒ Acceptable
☐ Not acceptable

5 Patient Information

5.1 Patient name and other information

5.1.1 Family name of the patient

DOE-CALLA

5.1.2 Prefix of the patient, if present

NA

5.1.3 Given name of the patient

CELESTINA

5.1.4 Gender of the patient, if present

- ☐ Male
☒ Female
☐ Undifferentiated
☐ Null value

5.1.5 Date of birth of the patient

15/10/1968

5.2 Patient identifiers

5.2.1 Patient identifier

3843082788

5.2.2 Additional patient identifier

(1.3.182.2.4.2), 1968101545978 (1.3.182.4.4)

5.2.3 Please, introduce here any comment you consider relevant regarding patient identifiers

The document is not presented in English. We are also guessing the first name and surname of the patient.

6 Patient Summary List

6.1 Was only one Patient Summary provided for the patient?

(One Patient Summary considering that the same document is provided as CDA Level 1, i.e. with the original embedded document as a PDF file, as well as Level 3, i.e. with the coded entries translated into the Country of treatment language)

- ☒ Yes
☐ No

6.3 Was the initial Patient Summary from the country of origin of the patient provided (PDF file)?

- ☒ Yes
☐ No

6.5 Did you have any error during the process?

- ☐ Yes
☒ No

7 Document and Patient Information

7.1 Are the creation date of the document and the date of last update of the information present?

(please, note that, due to the way the Patient Summary is created, the date of last update of the information might be an earlier date than the date of creation of the document)

- ☒ Yes
☐ No

7.2 Please explain

The date headings are not in English

7.3 Does the patient's contact information (address, telecom) seem to be complete and usable?

- ☒ Yes
☐ No

7.5 Do you find the administrative information provided sufficient for the purpose of the service?

(this question refers to the information regarding patient's contacts, such as guardian or preferred contact, and about the author and legal authenticator of the document)

Yes - other than the headings not being Translated

8 Clinical Section: Allergies and adverse reactions Document

8.1 Is the Allergies and adverse reactions Document Section present?

- ☒ Yes
☐ No

8.2 Does the translated part of the section present the information in your national language?

- ☐ Yes
☒ No

8.3 Please explain

The content is partially in English

8.4 Please mark for which of the following data elements information is provided:

- ☒ Reaction Type
☐ Clinical Manifestation
☒ Agent
☒ Duration: From
☐ Duration: Until
☒ Severity
☐ Criticality
☐ Allergy status
☐ Certainty

8.5 Please introduce here any comment you consider relevant about the above data elements:

It's not translated, so not 100% sure what the patient is allergic to. Missing columns for the data elements.

8.6 Can you safely understand the information present?

- ☐ Yes
☒ No

8.7 Please explain

We don't know what the agent is.

8.10 Is there any missing information or empty cell difficult to understand/interpret?

Note: whether it is clearly differentiated that there is "no information" or the "information is known to be absent", or other challenging circumstance

- ☒ Yes
☐ No

8.11 Please explain

As above, We don't know what the agent is and other data fields are missing.

8.12 Is there any other source of possible medical error?

- ☒ Yes
☐ No

8.13 Please explain

Incorrect medication could be given.

9 Clinical Section: Relevant diagnostic tests/laboratory data Narrative

9.1 Is the Relevant diagnostic tests/laboratory data Narrative Section present?

- ☐ Yes
☒ No

10 Clinical Section: History of Medication use Narrative

10.1 Is the History of Medication use Narrative Section present?

- ☒ Yes
☐ No

10.2 Does the translated part of the section present the information in your national language?

- ☐ Yes
☒ No

10.3 Please explain

Not presented in English

10.4 Please mark for which of the following data elements information is provided:

- ☒ Medicinal Product
☒ Active Ingredient
☒ Strength
☐ Dose Form
☒ Units per Intake
☒ Frequency of Intakes
☒ Route of Administration
☒ Duration of Treatment: From
☒ Duration of Treatment: Until

☐ Medication reason

10.5 Please introduce here any comment you consider relevant about the above data elements

The headings aren't translated nor the nor the medicinal data

10.6 Can you safely understand the information present?

- ☐ Yes
☒ No

10.7 Please explain

The information is not in English

10.10 Is there any missing information or empty cell difficult to understand/interpret?

Note: whether it is clearly differentiated that there is "no information" or the "information is known to be absent", or other challenging circumstance

- ☒ Yes
☐ No

10.11 Please explain

There is no reason for the medication prescribed/given

10.12 Is there any other source of possible medical error?

- ☒ Yes
☐ No

10.13 Please explain

One can make an incorrect assumption for the reason a medication was given

11 Medical problem

11.1 Clinical Section: Problem list - Reported

11.1.1 Is the Problem list - Reported Section present?

- ☒ Yes
☐ No

11.1.2 Does the translated part of the section present the information in your national language?

- ☐ Yes
☒ No

11.1.3 Please explain

The information is not in English

11.1.4 Please mark for which of the following data elements information is provided:

- ☒ Active Problem
- ☒ Onset Date
- ☐ Diagnosis Assertion Status
- ☒ Related Health Professional
- ☒ Related External Resource

11.1.5 Please introduce here any comment you consider relevant about the above data elements

The data elements are not in English

11.1.6 Can you safely understand the information present?

- ☐ Yes
- ☒ No

11.1.7 Please explain

The information isn't clear, not being in English

11.1.10 Is there any missing information or empty cell difficult to understand/interpret?

Note: whether it is clearly differentiated that there is "no information" or the "information is known to be absent", or other challenging circumstance

- ☒ Yes
- ☐ No

11.1.11 Please explain

The diagnosis assertion status isn't shown

11.1.12 Is there any other source of possible medical error?

- ☒ Yes
- ☐ No

11.1.13 Please explain

The information is not in English - one might need to guess, if not able to understand the original language

11.2 Clinical Section: History of medical device use

11.2.1 Is the History of medical device use Section present?

- ☒ Yes
- ☐ No

11.2.2 Does the translated part of the section present the information in your national language?

- ☐ Yes
☒ No

11.2.3 Please explain

The information is not in English

11.2.4 Please mark for which of the following data elements information is provided:

- ☒ Device/Implant
☒ Implant Date: From
☒ Implant Date: Until

11.2.5 Please introduce here any comment you consider relevant about the above data elements

The information is not in English.

11.2.6 Can you safely understand the information present?

- ☐ Yes
☒ No

11.2.7 Please explain

The information is not in English so one can't decide on the purpose of the implant

11.2.10 Is there any missing information or empty cell difficult to understand/interpret?

Note: whether it is clearly differentiated that there is "no information" or the "information is known to be absent", or other challenging circumstance

- ☒ Yes
☐ No

11.2.11 Please explain

It's not shown what the device was implanted for. The information is not in English

11.2.12 Is there any other source of possible medical error?

- ☒ Yes
☐ No

11.2.13 Please explain

One would need to know what the device is, else incorrect decisions can be made.

11.3 Clinical Section: History of Procedures Document

11.3.1 Is the History of Procedures Document Section present?

- ☒ Yes

☐ No

11.3.2 Does the translated part of the section present the information in your national language?

☐ Yes

☒ No

11.3.3 Please explain

The information is not in English

11.3.4 Please mark for which of the following data elements information is provided:

☒ Procedure

☒ Body site

☒ Procedure Date

11.3.5 Please introduce here any comment you consider relevant about the above data elements

The information is not in English

11.3.6 Can you safely understand the information present?

☐ Yes

☒ No

11.3.7 Please explain

The information is not in English

11.3.10 Is there any missing information or empty cell difficult to understand/interpret?

Note: whether it is clearly differentiated that there is "no information" or the "information is known to be absent", or other challenging circumstance

☒ Yes

☐ No

11.3.11 Please explain

The information is not in English. The not about what the procedure was for, is good, but it's not translated

11.3.12 Is there any other source of possible medical error?

☒ Yes

☐ No

11.3.13 Please explain

One can't understand what the procedure was.

11.4 Clinical Section: Functional status assessment note

11.4.1 Is the Functional status assessment note Section present?

- ☐ Yes
☒ No

12 Medical history

12.1 Clinical Section: History of Past Illness Narrative

12.1.1 Is the History of Past Illness Narrative Section present?

- ☐ Yes
☒ No

12.2 Clinical Section: History of Immunization Narrative

12.2.1 Is the History of Immunization Narrative Section present?

- ☒ Yes
☐ No

12.2.2 Does the translated part of the section present the information in your national language?

- ☐ Yes
☒ No

12.2.3 Please explain

The information is not in English

12.2.4 Please mark for which of the following data elements information is provided:

- ☒ Vaccination
☒ Brand Name
☒ Vaccination Date
☐ Agent
☒ Marketing Authorization Holder
☐ Dose number in series
☒ Batch/lot number
☐ Administering Center
☐ Health Professional Identification
☐ Country of Vaccination
☒ Administered

12.2.5 Please introduce here any comment you consider relevant about the above data elements

A number of data elements are missing. The information needs to be translated.

12.2.6 Can you safely understand the medical information communicated?

- ☐ Yes
☒ No

12.2.7 Please explain

The information is not in English

12.2.10 Is there any missing information or empty cell difficult to understand/interpret?

Note: whether it is clearly differentiated that there is "no information" or the "information is known to be absent", or other challenging circumstance

- ☒ Yes
☐ No

12.2.11 Please explain

A number of data elements are missing. The information needs to be translated.

12.2.12 Is there any other source of possible medical error?

- ☒ Yes
☐ No

12.2.13 Please explain

The information is not in English

13 Clinical Sections: Social History Narrative

13.1 Is the Social History Narrative Section present?

- ☐ Yes
☒ No

14 Clinical Section: History of Pregnancies Narrative

14.1 Is the History of Pregnancies Narrative Section present?

- ☐ Yes
☒ No

15 Clinical Section: Vital signs

15.1 Is the Vital signs Section present?

- ☐ Yes
☒ No

16 Clinical Section: Others

16.1 Are the following Sections present?

- ☐ Plan of Care
- ☐ Advance Directives

17 About the Document

17.1 Is the structure of the document logical and easy to follow?

- ☒ Yes
- ☐ No

17.3 Is the data provided generally medically coherent?

- ☐ Yes
- ☒ No

17.4 Please explain

The information is not in English

17.5 Is the original content in the PDF from Country A provided in English?

- ☐ Yes
- ☒ No

17.7 Considering the information provided as a whole, do you find the service useful for the care you intend to provide?

- ☐ Yes
- ☒ No

17.8 Please introduce here any comment you consider relevant regarding the question above

The information is not in English and can result in errors being made

18 Additional Feedback

Please, upload here additional feedback about the evaluation: ideally, the CDA document (XML file) that was displayed to you and the rendering of the CDA Display Tool (PDF file). That information will greatly help Solution Provider when evaluating the submission.

The maximum upload limit is 1 MB.

18.1 Please upload your file here:

The maximum file size is 1 MB

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df354c97-4a35-4a11-8c01-8dd69230d7cb/LU_L3_PS_Summary_Friendly_04_25.htm

Contact

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