



# GS1 EUDAMED Submission Guideline

How to implement UDI submissions for EUDAMED via GDSN

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## Document Summary

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1.0	January 2026	Initial creation and finalisation
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## Involved parties

There are several parties involved in the GS1 EUDAMED Submission.

- **GS1 Member Organisations:** your local contact point that offers you the GDSN Data pool for exchanging master data information of your medical devices. And the data pool where you enter the relevant information for GS1 UDI Connector. This data pool also offers automatic validations which helps you to make sure you have the required information for EUDAMED. At this moment **GS1 Switzerland, GS1 Denmark, GS1 Czech Republic** and **GS1 Netherlands** are offering a EUDAMED Submission service.
- **Trade Connectors:** The technical party behind the GDSN Data pool solutions that your local GS1 Member Organisation offer.
- **p36:** Technical service provider (access point) for the UDI connect interface. They deliver the integration between the GDSN Data pool and the UDI databases.

## Revocation (disclaimer)

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Whilst/although every care has been taken to ensure that the content of this document is correct, GS1 Member Organisation cannot be held responsible for errors or missing information in this publication. For questions regarding the contents of this publication, please contact GS1 Member Organisation.

<sup>1</sup> GS1 Member Organisations involved: GS1 Czech Republic, GS1 Denmark, GS1 Netherlands, GS1 Switzerland

## Table of contents

<b>1</b>	<b>Introduction - Preconditions</b>	<b>4</b>
1.1	Regulatory disclaimer	4
1.2	Document and project structure	5
<b>2</b>	<b>Phase 1 - Onboarding Process</b>	<b>6</b>
2.1	Access Point Configuration	7
<b>3</b>	<b>Phase 2 - GDSN Attribute Analysis for UDI Data</b>	<b>12</b>
<b>4</b>	<b>Phase 3 - Learn UDI Upload and Testing</b>	<b>14</b>
4.1	For Web-User Interface Users	14
4.2	For Machine-to-Machine Users	15
4.3	Registration Process for Regulation Devices	16
<b>5</b>	<b>Phase 4 - Data Capturing and Production Configuration</b>	<b>18</b>
5.1	For Web-User Interface Users	18
5.2	For Machine-to-Machine Users	19
<b>6</b>	<b>Phase 5 - Live Submissions and Error Handling</b>	<b>20</b>
6.1	Publication	20
6.2	Follow-up Submissions	21
6.3	Error Handling	22
6.4	Support	23
6.5	Discard	23
<b>7</b>	<b>Sources of Information</b>	<b>25</b>
7.1	Required Templates for Access Point Configuration	25
7.2	Data Pool Resources (e.g. User Manuals)	25
7.3	EUDAMED Information	25
7.4	Table of Abbreviations	26



# 1 Introduction - Preconditions

GS1 is an international not-for-profit organisation that offers a solution to share product information of medical devices between manufacturers, distributors and care institutions. This solution is called the Global Data Synchronisation Network (GDSN). Manufacturers also must upload part of this information in EUDAMED.

EUDAMED is the IT system established by Regulation (EU) 2017/745 on medical devices and Regulation (EU) 2017/746 on in vitro diagnosis medical devices. EUDAMED is an integral part of the implementation of the two Medical Devices Regulations.

To support manufacturers in this process the solution that GS1 offers also facilitates the uploading of data in EUDAMED.

This document serves as a guide for your initial configuration of the European database of medical devices – EUDAMED. It is focused on UDI (Unique Device Identification). The aim is to describe the process of user registration and the registration of an economic operator in the role of Legal Manufacturer (MF) and System/Procedure Pack Producer (PR), for the purpose of configuring Machine-to-Machine (M2M) Connector between EUDAMED and the GS1 GDSN for the UDI submissions.

At the same time, neither GS1 nor any Machine-to-Machine data upload alter or transfer any rights or obligations to other parties beyond those explicitly defined in the MDR (Medical Device Regulation) and IVDR (In Vitro Medical Device Regulation) regulations, which form the legal basis for the obligation to submit medical device data to EUDAMED by the Legal Manufacturer and Producer.

## 1.1 Regulatory disclaimer

GS1 provides participants with non-binding guidance aimed at applying GS1 standards. Each participating company is responsible for determining how such guidance is applied within its own organisation. GS1 is a neutral organisation providing voluntary recommendations. Participants remain independent entities and are free to define and follow their own procedures. GS1 employees have no authority to act on behalf of regulatory agencies, to interpret legal provisions in a legally binding manner, or to grant exemptions or approvals.

The information contained in this documentation has been developed and compiled by GS1 to the best of its knowledge. GS1 makes no representations or warranties, express or implied, regarding the completeness, accuracy or suitability of this information for any particular purpose.

To the extent permitted by applicable law, GS1 shall be liable only for damages caused by wilful misconduct or gross negligence. Liability for slight negligence is excluded, except where mandatory law provides otherwise.

Each participating company is solely responsible for ensuring compliance with all applicable legal and regulatory requirements.

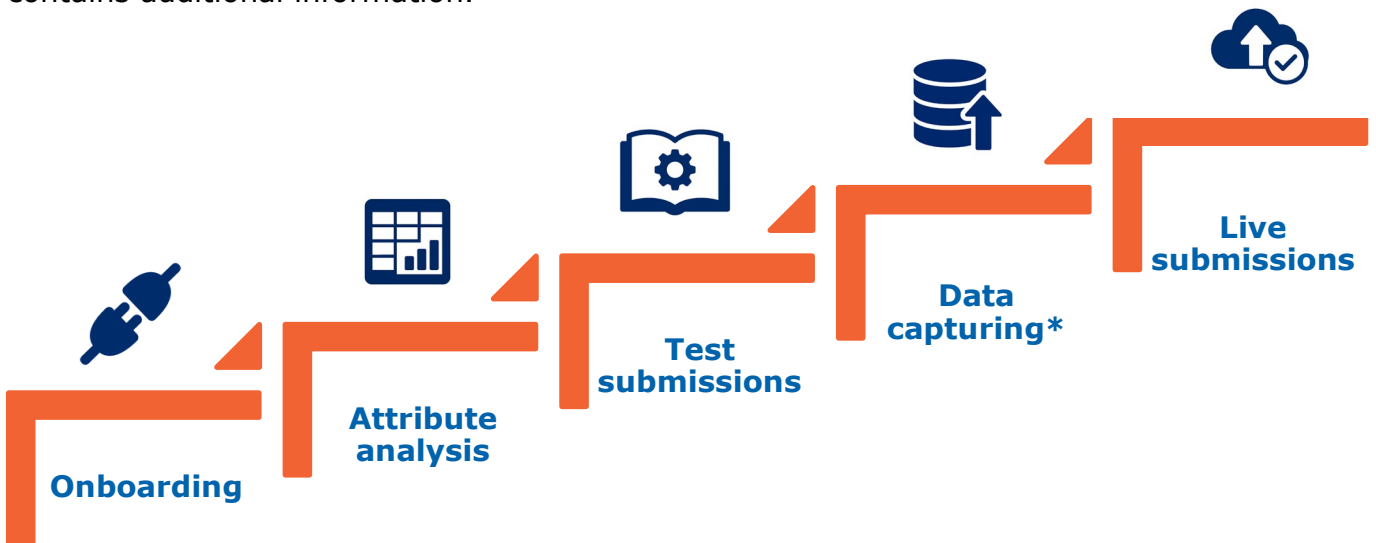
Any services provided by a local Member Organisation of GS1 shall be governed exclusively by the applicable general terms and conditions of the respective local Member Organisation, in their then-current version, as made available by such organisation (e.g. on its website or upon request). The participant alone is responsible for ensuring compliance with all applicable legal and regulatory requirements, including, MDR, IVDR, device registration and synchronization requirements.

For the service that is explained in this document, the general terms and conditions of GS1 are applicable. These are available at the local Member Organisation’s website or upon request.

In the MDR regulation the responsibilities regarding EUDAMED are described in Annex VI part A. <https://eur-lex.europa.eu/eli/reg/2017/745/oj/eng>

## 1.2 Document and project structure

There are 5 phases to successfully reach the EUDAMED submission. These are described in the picture below and in this document in the following 5 chapters. The last chapter contains additional information.



\* DRIFT = **Do it Right the First Time**

Figure 1 - UDI submission process

## 2 Phase 1 - Onboarding Process

EUDAMED offers to work with two environments. The first one is a EUDAMED Playground which is a test environment used for training and verifying system integration, especially for M2M data exchange. It allows users to simulate data submissions without any legal consequences.

In contrast, the second is the EUDAMED Production Environment representing the official live system where legally binding data must be submitted in compliance with MDR and IVDR regulations.

In this document the registration phases are explained in a logical and structured manner – EUDAMED Playground first and whenever certain steps differ in the EUDAMED Production, these differences are highlighted.

For the User and Actor registration follow these steps:

1. Follow this link to EUDAMED Playground:  
[https://webgate.training.ec.europa.eu/eudamed-play/landing-page#/.](https://webgate.training.ec.europa.eu/eudamed-play/landing-page#/)
2. Sign in with your EU Login or create one if not available.
3. Create a Test Actor Request, the role of *Legal Manufacturer (MF Actor)* or System or Procedure Pack Producer (PR Actor)
4. Please note that in EUDAMED production environment, all the applications get approved by Competent Authority but in Playground environment, they are approved by EUDAMED application support.
  - a. Contact [SANTE-EUDAMED-SUPPORT@ec.europa.eu](mailto:SANTE-EUDAMED-SUPPORT@ec.europa.eu).
  - b. Provide: country, application ID, manufacturer name, account email address
  - c. Kindly ask them with the approval of the test account

### European Commission (EC) EUDAMED website:

An official source of information is this [EC EUDAMED website](#) which provides a comprehensive information about EUDAMED in general and specific the UDI-module.

Refer to EUDAMED information centre ([Playground](#), [Production](#)) for more detailed information of the UDI-Module and its setting up and managing M2M data exchange between external systems and the EUDAMED database. This setup is taken care of by the service of GS1's UDI-connector.

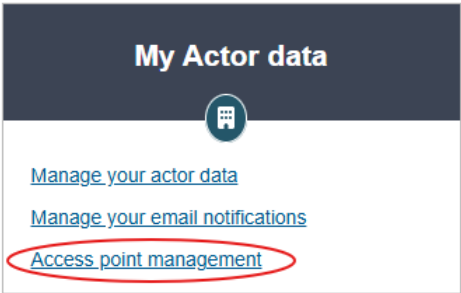
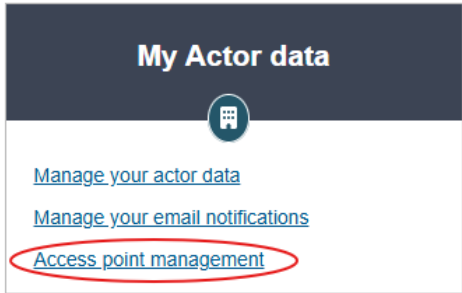
It also includes a video explaining the following relevant steps to use of an existing Access Point (AP) in EUDAMED for M2M data exchange with steps like entering the AP Party ID, uploading a 3<sup>rd</sup> Party Agreement, and submitting technical and legal contact details is found [here](#).

## 2.1 Access Point Configuration

This part provides information on configuring the M2M settings in EUDAMED.

**If you wish to activate your M2M services in EUDAMED Production environment you must first apply for an AP in EUDAMED Playground environment, complete the onboarding and test the service.** Please start in the EUDAMED Playground and later follow in the EUDAMED Production environment.

**NOTE:** In this part the details of 3<sup>rd</sup> Party Access Point provider p36 will be provided, as an example of such steps.

	EUDAMED PLAYGROUND environment	EUDAMED PRODUCTION environment
1.	<p>Log into <b>EUDAMED Playground</b> as a Local Actor Administrator (LAA). Under the <i>My Actor data</i> section click on the <i>Access point management</i> link:</p> 	<p>Log into <b>EUDAMED Production</b> as a Local Actor Administrator (LAA). Under the <i>My Actor data</i> section click on the <i>Access point management</i> link:</p> 
2.	Click on the <i>Request a new M2M access</i> button.	
3.	Agree to the disclaimer and click on the <i>Next</i> button	
4.	In the next screen select <u>Yes</u> to use an existing EUDAMED AP and click on the <i>Next</i> button.	
5.	<p>Enter the following Party ID and click on the <i>Validate AP</i> button: Example: p36 Access Point use "EUDAMED_0000018_ACC": Machine to Machine request access</p> <p><small>* Enter the Party Id of the Access Point you want to use:</small></p> <input type="text" value="EUDAMED_0000018_ACC"/> <p><b>Validate AP</b></p> <p><small>* 3rd Party Agreement:</small></p> <input type="button" value="Browse"/> <p><input type="button" value="Save &amp; Next"/> <input type="button" value="Cancel"/></p>	<p>When Proof-of-Testing is provided enter the following Party ID and click on the <i>Validate AP</i> button use 'EUDAMED_004861': Machine to Machine request access</p> <p><small>* Enter the Party Id of the Access Point you want to use:</small></p> <input type="text" value="EUDAMED_004861"/> <p><b>Validate AP</b></p> <p><small>* 3rd Party Agreement:</small></p> <input type="button" value="Browse"/> <p><input type="button" value="Save &amp; Next"/> <input type="button" value="Cancel"/></p>



Switzerland



Czech Republic



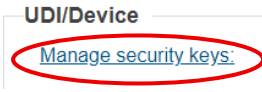
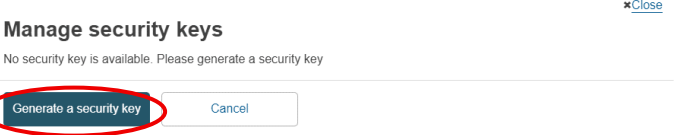
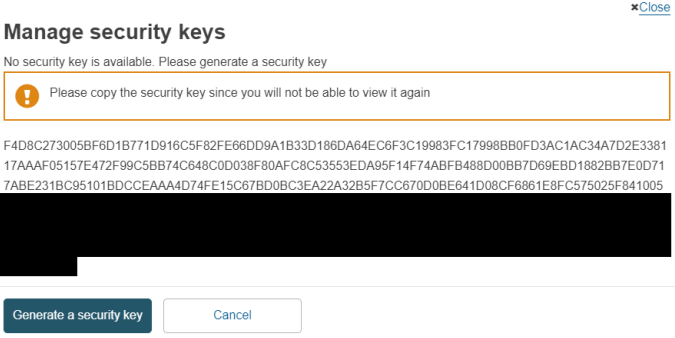
Netherlands



Denmark

6.	<p>You will be able to view information regarding your selected-Access Point. You will not be able to edit that information. Example:</p> <div data-bbox="255 470 774 929"> <p>Access Point Name</p> <p>Access Point Name: p36 Test</p> <p>Party ID: EUDAMED_0000018_ACC</p> <p>Access Point Country: Germany</p> <p>Access Point City: Frankfurt</p> <p>Organisation</p> <p>Organisation name: p36</p> <p>Street information, if applicable: No</p> <p>PO box: -</p> <p>City name: Bad Hersfeld</p> <p>Postal code: 36251</p> <p>Country: Germany</p> <p>Telephone: -</p> <p>Email: patrick.pfau@p36.io</p> </div>	<p>You will be able to view information regarding your selected-Access Point. You will not be able to edit that information. Example:</p> <div data-bbox="861 470 1380 1030"> <p>Access Point Name</p> <p>Access Point Name: p36</p> <p>Party ID: ΓUDAMED 004861</p> <p>Access Point Country: Germany</p> <p>Access Point City: Frankfurt</p> <p>Organisation</p> <p>Organisation name: p36 GmbH</p> <p>Street information, if applicable: Yes</p> <p>Street: Hof Meisebach</p> <p>Street number: -</p> <p>Address line 2: -</p> <p>PO box: -</p> <p>City name: Bad Hersfeld</p> <p>Postal code: 36251</p> <p>Country: Germany</p> <p>Telephone: +49 8621 7954500</p> <p>Email: eudamed@p36.io</p> </div>
7.	<p>Sign and upload the <b>3rd Party Agreement</b> as a PDF file and click on the <i>Save &amp; Next</i> button. A draft to be downloaded <a href="#">here</a>. For the signature contact your involved 3<sup>rd</sup> Party Provider. See also <b>Note #1</b> at the bottom of this section.</p>	<p>Sign and upload the <b>3rd Party Agreement</b> as a PDF file and click on the <i>Save &amp; Next</i> button. A draft to be downloaded <a href="#">here</a>. For the signature contact your involved 3<sup>rd</sup> Party Provider. See also <b>Note #1</b> at the bottom of this section.</p>
8.	<p>In the next screen fill in the <i>Technical Contact</i> details with the contact information of the Provider of the AP. Contact your involved 3<sup>rd</sup> Party Provider for these details OR see <b>Note #1</b> at the bottom of this section. Fill in also the <i>Legal Contact</i> details. This person must belong to <b>your organisation (Manufacturer)</b> and could be one of your Local Actor Admins in EUDAMED.</p>	
9.	<p>Upload your <b>Business justification</b> as PDF file. You will find the Business justification template <a href="#">here</a>. Contact your involved 3<sup>rd</sup> Party Provider OR see <b>Note #1</b> at the bottom of this section.</p>	<p>Upload your <b>Proof of testing</b> document as a PDF file which the 3<sup>rd</sup> Party Provider should have provided to you (EUDAMED Playground step #21) OR see <b>Note #1</b> at the bottom of this section.</p>
10.	<p>Select all services of the <i>UDI/Device</i> module and click on the <i>Submit</i> button.</p>	

	<div style="display: flex; justify-content: space-between;"> <div style="width: 45%;"> <p><b>Certificates/Notified Body</b></p> <ul style="list-style-type: none"> <li><input checked="" type="checkbox"/> SS(C)P Download</li> </ul> </div> <div style="width: 45%; border: 1px solid #ccc; padding: 5px;"> <p style="text-align: center; margin: 0;"><b>UDI/Device</b></p> <ul style="list-style-type: none"> <li><input checked="" type="checkbox"/> Update product original manufacturer</li> <li><input checked="" type="checkbox"/> Upload of Legacy / Regulation Device / SPP ( Basic UDI and UDI-DI / Master UDI-DI )</li> <li><input checked="" type="checkbox"/> Update Basic UDI</li> <li><input checked="" type="checkbox"/> Download of Legacy/ Regulation Device/SPP</li> <li><input checked="" type="checkbox"/> Upload of UDI-DI / Master UDI-DI for existing Basic UDI-DI</li> <li><input checked="" type="checkbox"/> Update of UDI-DI / Master UDI-DI</li> <li><input checked="" type="checkbox"/> Update container package</li> <li><input checked="" type="checkbox"/> Update market information</li> </ul> </div> </div> <div style="margin-top: 20px;"> <p><b>Actor</b></p> <ul style="list-style-type: none"> <li><input checked="" type="checkbox"/> Actor download</li> </ul> </div> <div style="margin-top: 20px; display: flex; justify-content: center; gap: 20px;"> <div style="border: 2px solid red; border-radius: 10px; padding: 5px 15px; background-color: #334d5d; color: white; font-weight: bold;">Submit</div> <div style="border: 1px solid #ccc; padding: 5px 15px;">Cancel</div> </div>
11.	Select Yes in the pop-up window to complete the Access Point registration process.
12.	In the next screen you can see a confirmation message and the Access Point Party ID.
13.	Click on the <i>Go back to Access Point Link dashboard</i> link to manage your AP. As soon as the request has been approved by EC and the AP has the status <b>"Active"</b> , you can continue. This can take a few days.
14.	<p>Under the <i>My Actor data</i> section click on the <i>Access point management</i> link:</p> <div style="border: 1px solid #ccc; padding: 10px; margin: 10px 0;"> <p style="text-align: center; background-color: #334d5d; color: white; padding: 5px;"><b>My Actor data</b></p> <div style="text-align: center; margin: 5px 0;"> </div> <p style="margin: 5px 0;"><a href="#">Manage your actor data</a></p> <p style="margin: 5px 0;"><a href="#">Manage your email notifications</a></p> <p style="margin: 5px 0; border: 2px solid red; border-radius: 10px; padding: 2px;"><a href="#">Access point management</a></p> </div>
15.	In the Access Point management page, you can view all your APs listed in the table. Make sure that the filter is set to <i>Status: Active</i> .
16.	Choose e.g. the p36 AP and click on the <i>Edit</i> link under the three dots to view the AP's settings
17.	In the displayed screen you can view and edit the details you entered regarding the p36 AP. Moreover, you can now change the services attached and manage

	<p>your <b>security keys</b><sup>2</sup> by clicking on the <i>Manage security keys</i> link in the edit mode:</p> 
18.	<p>In the pop-up window click on the <i>Generate a security key</i> button to generate the security key for the UDI/Device module (security keys are generated per module):</p> 
19.	<p>In the pop-up window you will be able to view the generated security key:</p>  <p><b>Please copy the security key. You will not be able to view your security key after closing the pop-up window.</b> If you lose your security key, you must regenerate it.</p> <div style="background-color: #f4a460; padding: 5px; border: 1px solid #ccc;"> <p><b>IMPORTANT</b> Please copy the regenerated security key. You will not be able to view your security key after closing the pop-up window. If you lose your security key, you must regenerate it.</p> </div>
20.	<p>Please forward the latest generated security key, Actor SRN &amp; and Actor company name:</p> <ol style="list-style-type: none"> <li>a. to a GS1 contact person.</li> <li>b. to the 3<sup>rd</sup> Party Provider p36, see <b>Note #1</b> at the bottom of this section.</li> </ol>
21.	<p>Continue and follow with the testing phase. Once this phase is successfully completed, ask for a <b>Proof of testing document (POT) in the PDF format only</b> for the EUDAMED Production environment setup:</p> <ol style="list-style-type: none"> <li>1. via a GS1 contact person.</li> </ol> <p style="text-align: right;">DONE</p>

<sup>2</sup>A security key (Token) is a crucial element for authorizing the connection between the legal entity (identified by its Single Registration Number - SRN) and the 3<sup>rd</sup> Party Provider, and it must be handed over to the provider to enable M2M communication for UDI registration on their behalf.

	2. via the 3 <sup>rd</sup> Party Provider p36, see <b>Note #1</b> at the bottom of this section.	
22.	The setup for <b>EUDAMED Playground</b> is done, go to the <a href="#">#1</a> of this Setup and follow the process referring the EUDAMED Production environment.	N/A

## Note #1

The process described above is very general and follows the mandatory steps in EUDAMED Playground/Production. Some AP providers may offer more support to efficiently manage the setup. For instance, the upcoming [p36 web tool](#) which digitalizes and half-automates the entire process from the AP Management. Screen shot below:

The screenshot displays the 'EUDAMED Actor Onboarding' interface. At the top, there's a 'UDH Hub Essential - Active' status. The main section shows four key metrics: Total Actors (2), In Progress (2), Playground Complete (0), and Production Ready (0). Below these metrics is a '+ Add Economic Actor' button and a search bar. The bottom section, titled 'EU Manufacturers & System and Procedure Pack Producers', lists two entities: 'P36 GmbH' (Manufacturer) and 'p36 GmbH' (SPP). Each entity has progress indicators for 'Playground' and 'Production' stages, with 'Continue Setup' links.

### 3 Phase 2 - GDSN Attribute Analysis for UDI Data

To ensure accurate and compliant data submission to EUDAMED, users must analyse the alignment between the **EUDAMED UDI Data Dictionary requirements** and the **mapped GDSN Profile**. For this purpose, the prepared document "GS1\_UDI\_Connector\_Profile\_Overview.xlsx", which you have received during your onboarding process, serves as the central reference, providing a comprehensive overview of attribute definitions and code list mappings.

To prepare successful UDI submissions the following tasks should be carried out by the responsible internal experts (e.g. Regulatory Affairs, Master Data Management or IT):

1. Define the scope of relevant device types (MDR, SPP, IVDR, MDD/AIMDD, IVDD).
2. Review the mandatory and optional fields per device type.
3. Compare these requirements with the mapped attributes in the GDSN Profile.
4. Assess the applicability of each attribute against the manufacturer's internal device data in source systems.
5. Identify potential gaps where internal data may not fully meet the requirements.
6. Repeat the same actions for the code list mapping.

For efficient execution, we recommend using the provided "GS1\_UDI\_Connector\_Profile\_Overview.xlsx". Add auxiliary columns (highlighted in green) in both tabs "EUDAMED\_swissdamed\_Attributes" and "UDID\_CodeLists" document internal notes, responsibilities, or gap resolutions.

EUDAMED UDI v2.22.0				swissdamed			GDSN v3.1.33 Mappi			Manufacturer's Analysis			
Scope	Field ID	Field Label	Entity Name	MDR Status	Overlap with EUDAMED?	Attribute Name	Example Value	Applicable?	Sample Value	Attribute name in EXP / PIM	Remark		
No	Basic-UDI / EUDI Attributes			x	-	Global model information							
Yes	FLD-UID-14	Basic UDI- DI code	BasicUIDData	M	Yes	Global Model Number (GMN)	7612345GOLDENtest19JG	Yes	7612345GOLDENtest19JG	fmm			
Auto-popul	FLD-UID-01	Issuing Entity Basic UDI-DI	BasicUIDData	M	Yes			Yes	MDR	Legislation			
Yes	FLD-UID-11	Applicable Legislation / Regulation	BasicUIDData	M	Yes	Regulatory act	MDR	default=EU					
Yes	FLD-UID-11	Applicable Legislation / Regulation	BasicUIDData	M	Yes	Regulatory agency	EU	Yes	CLASS_IIB	Risk Class			
Yes	FLD-UID-16	Risk Class	BasicUIDData	M	Yes	Additional trade item	EU_CLASS_IIB	default=76					
Yes	FLD-UID-16	Risk Class	BasicUIDData	M	Yes	Additional trade item	76	default=76					
Yes	FLD-UID-10	Legal Manufacturer SRN	BasicUIDData	M	Yes	Additional party identification	SRN / CH-MF-000023141	Yes	DK-MF-000021581	EU Actor			
Yes	FLD-UID-10	Legal Manufacturer SRN	BasicUIDData	M	Yes	Contact type code	EMA	default=EMA					
Yes		Legal manufacturer CHRN	CHRN (MF)	M	swissdamed only	Additional party identification	CHRN / CHRN-MF-20010136	No					
Yes		Legal manufacturer CHRN	CHRN (MF)	M	swissdamed only	Contact type code	EMA	No					
Yes		Authorised representative CHRN	CHRN (AR)	CM	swissdamed only	Additional party identification	CHRN / CHRN-AR-20010004	Yes	CHRN-AR-20010004-MF-00004	CH Mandate			
Yes		Authorised representative CHRN	CHRN (AR)	CM	swissdamed only	Contact type code	EAR	default=EMA					
Yes	FLD-UID-20	Device Model	BasicUIDData	CM	Yes	Additional trade item	MODEL_NUMBER /						
Yes	FLD-UID-22	Device Name	BasicUIDData	CM	Yes	Global model description	Compl_MDR_IIB_name						
Yes	FLD-UID-12	Is it a System which is a Device in	BasicUIDData	M	Yes	Multi component device type	PROCEDURE_PACK						
Yes	FLD-UID-13	Special Device Type	BasicUIDData	O	Yes	Special device type code	SOFTWARE						
Yes	FLD-UID-28	Active Device	BasicUIDData	M	Yes	Is active device	false						
Yes	FLD-UID-29	Device Intended to administer and/	BasicUIDData	M	Yes	Is device intended to administer	true						

This analysis forms the foundation for a structured implementation, ensuring that all relevant device data is correctly captured, validated, and ready for submission. By systematically comparing EUDAMED field requirements with internal data structures, organizations can proactively address compliance issues and secure efficient UDI registration.



## Note #2

If you have any specific technical or attribute-related questions, please contact your local GS1 Member Organisation.

## 4 Phase 3 - Learn UDI Upload and Testing

When [Phase 2 - GDSN Attribute Analysis for UDI Data](#) is completed, it is time to test upload of your data. It is important to start with a record for one product first and then scale up with the rest of your records afterwards.

Even when having a M2M solution, submitting your data in EUDAMED can be complicated, because of the requirements from EUDAMED. For that reason, it is extra important to have the **Do it Right the First Time** approach (DRIFT). Be very sure the data information is submitted with the right content, by having internal quality assurance steps, all those steps that are possible.

The intention of testing is to make sure, that your M2M solution supports all the scenarios and attributes relevant to your submission of data to EUDAMED.

The recommended number of test submission depends on your range of products. It is recommended to do a test submission for each class in MDR and IVDR that you have in your range of products.

These steps are recommended to follow.

### 4.1 For Web-User Interface Users

1. Use Editor (Web-UI) to create your first record with all the attributes required for your product. Implemented validation rules will directly control (DRIFT) if your record is affected by any consistency mistake, business rule dependency or error appearance. The screenshot below shows you what that looks like. Publish first UDI when the product has all required attributes.

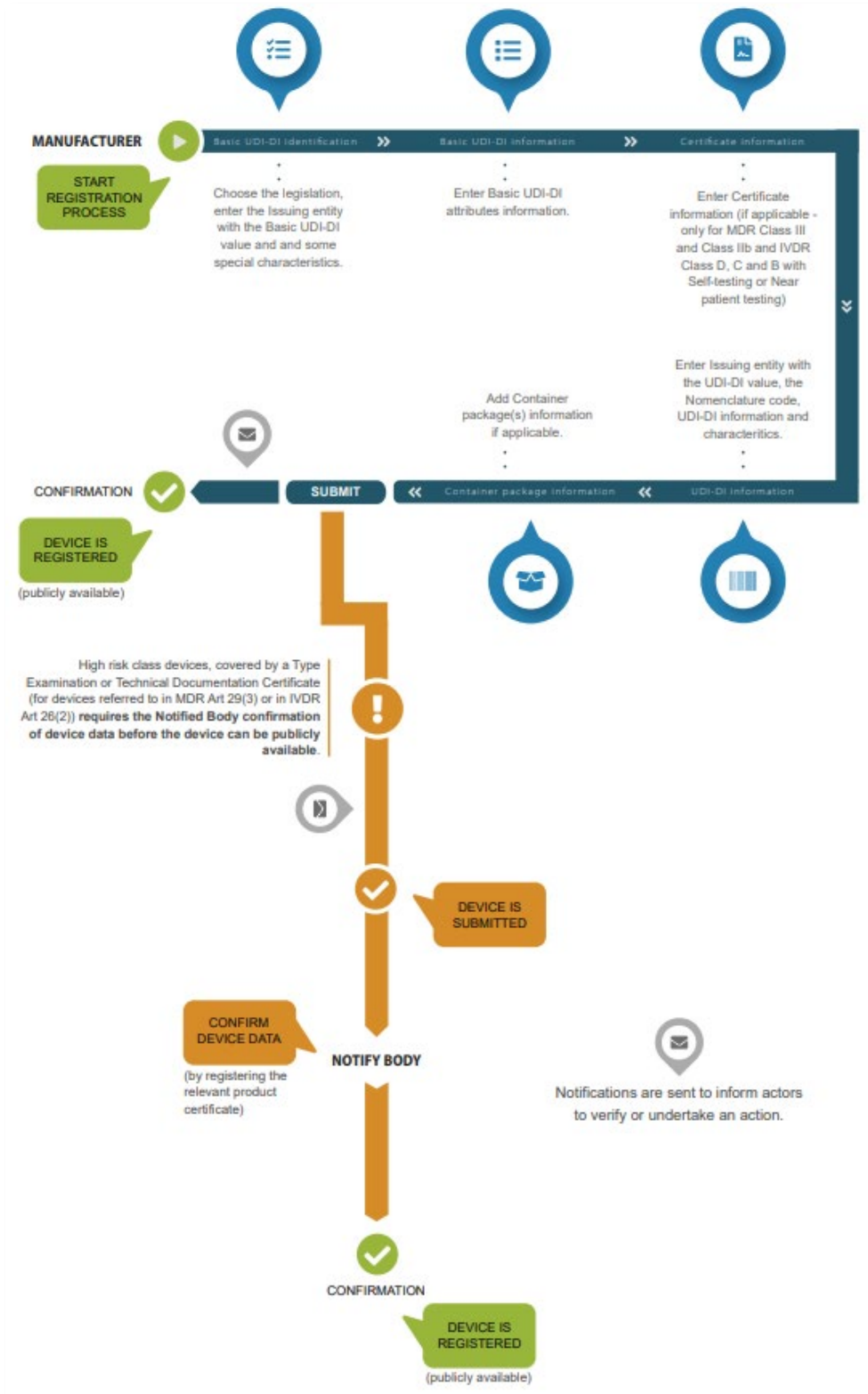
The screenshot displays a web-based data entry form for 'Additional Trade Item Classification'. It includes fields for 'Additional product classification type code', 'Additional Trade Item Classification System Code', 'Additional Trade Item Classification Value', 'Additional product classification value', and 'Additional Trade Item Classification Version'. Two entries are shown: one for 'EU Regulation (MDR/IVDR) Risk class (76)' and another for 'GMDN - Global Medical Device Nomenclature (35)'. On the right side, a red 'Errors' panel lists two issues: 'G541 Error: For all attributes for which the Trade Item Standard and associated extensions identifies a code list, only the values in that code list are valid in the GDS network.' and '097.007 EUDAMED: At least one iteration of additionalTradeItemClassification (EMDN value) must be used where the corresponding additionalTradeItemClassificationSystemCode must equal '88''. Below each error is a snippet of the corresponding code list.

2. After one successful registered record, that record can be exported in excel. The exported excel file can then be used to fill out other test submissions and be imported into your GDSN service.
3. Then you can publish all test records in EUDAMED. Then check EUDAMED Response Feedback (CIC - Catalogue Item Conformation) for the records send. Complete the *step 1* again if any mistakes occurred.

## 4.2 For Machine-to-Machine Users

1. Send your first UDI record through your implemented M2M interface into our UDI Connector service via GDSN or the data validation. Complete this step for all your chosen test scenarios.
2. Then you can publish all test records in EUDAMED. Then check EUDAMED Response Feedback (CIC) for the records send. If there are errors mentioned in the CIC look at [Phase 5 - Error Handling](#) for your guidance. Complete [step 1](#) again if any mistakes occurred.

### 4.3 Registration Process for Regulation Devices



Source: [https://health.ec.europa.eu/document/download/c3231845-228e-437a-8d77-510ecc3a548b\\_cs?filename=md\\_eudamed-udi-registration-process\\_en.pdf](https://health.ec.europa.eu/document/download/c3231845-228e-437a-8d77-510ecc3a548b_cs?filename=md_eudamed-udi-registration-process_en.pdf)

The steps in this part of the process depend on the type of risk class for your products. There will be two levels of risk class:

- Low risk class (for MDR: I, IIA, IIB non-implants, for IVDR: D, C, and B with Self-testing or Near patient testing).
- and High risk class (for MDR: IIB implants, III, for IVDR: A or B without Self-testing or Near patient testing).

For Low risk class, when you send your data to EUDAMED, it will be with no involvement of Notified Bodies. The data gets the status "REGISTERED", and you will get a CIC receipt with the status "SYNCHRONISED", and your registration is completed.

For High risk class (MDR: IIB implants, III) you are only able to send data with status "SUBMITTED" and you will get a CIC receipt with the status "RECIEVED", which means you as MF has a valid submission, but the Notified Body will have to "confirm" the device data. That will happen when Notified Body register the certificate. After approval, status will change to "REGISTERED" and you will get a CIC receipt with the status "SYNCHRONISED", and your registration is completed. In the Playground environment this will not happen if no Notified Body is involved for collaboration testing. This is not required for a Proof of testing.

After testing is completed, it is important to get the **Proof of Test (POT)** file in PDF format and store it in our documentation. The POT file must include the used Actor Code (SRN), DI Codes and successful XML messages from EUDAMED. You can get your POT file from one of the following options:

- a) Contact your 3<sup>rd</sup> Party Provider for POT file.
- b) Generate the POT file from p36 "Authority Onboarding" (see [Phase 1 - Onboarding Process Note #1](#))

By choosing method b), you will learn more about the service you use, and how it works. Both methods will work fine though.

If you have any questions related to this preparation step, contact your GS1 Member Organisation for support and training.

## 5 Phase 4 - Data Capturing and Production Configuration

For Production configuration lookup the steps of [Phase 1 - Onboarding Process](#).

Once you have your configuration in production ready, you can start adding product data for the products you want to register in EUDAMED. It is recommended that you identify two user roles to enable working with a 4-eye principle: one person as "Editor", this person adds the relevant data to your product. The other person is "Access control Editor (Publisher)", this person checks the data and publishes the data to EUDAMED. You can identify these roles in the 'users' tab in your GDSN Data pool.

First collect the required UDI data internally, see [Phase 2 - GDSN Attribute Analysis for UDI Data](#). If available, you can also reuse data that was successfully submitted in the TEST environment.

Add the data of one product in your GDSN Data pool. Look for help on the working of the GDSN Data pool here on the website of your local GS1 Member Organisation. The following steps are recommended.

### 5.1 For Web-User Interface Users

1. Use the Editor role to create a first product (UDI-DI) under a Basic UDI-DI and enter all the attributes required for that product. As mentioned in the previous step under testing, it's recommended to follow DRIFT. Implemented validation rules (errors) will support that and directly control if your record is affected by any missing mandatory, consistency mistake or other business rule dependency. The screenshot below shows you what that looks like.

The screenshot displays the EUDAMED web interface for product configuration. It shows two sections for 'Additional Trade Item Classification'. Each section includes a dropdown for 'Additional product classification type code/Additional Trade Item Classification System Code' and a text input for 'Additional Trade Item Classification Value'. The first section has 'EU Regulation (MDR/IVDR) Risk class (76)' selected, and the second has 'GMDN - Global Medical Device Nomenclature (35)'. The 'Additional product classification value' field contains 'EU\_CLASS\_1' and '12345' respectively. On the right side, an 'Errors' panel lists two error messages: 'G541 Error: For all attributes for which the Trade Item Standard and associated extensions identifies a code list, only the values in that code list are valid in the GDS network.' and '097.007 EUDAMED: At least one iteration of additionalTradeItemClassification (EMDN value) must be used where the corresponding additionalTradeItemClassificationSystemCode must equal '88''. Below each error message are two examples of the 'Additional product classification type code/Additional Trade Item Classification System Code' field with arrows pointing to the error.

2. After completion publish this first UDI-DI with its Basic-UDI data to your targeted "UDI Connector" GLN (Publish to GLNs see [Phase 5 - Live Submissions and Error Handling, Publication section](#)). Make sure that this first publication of a Basic UDI-DI with its first UDI-DI succeeds with CIC SYNCHRONISED or RECEIVED.



## 6 Phase 5 - Live Submissions and Error Handling

### 6.1 Publication

When valid draft items are prepared, created or uploaded in the GDSN Data pool and the approval (review by 4 eye-principle) is done, only then **publication** of the first live submissions can follow. To avoid additional error handling, it's recommended to go step by step with the following approach. It is the same procedure as in the test environment (UAT instance) of your GDSN Data pool.

For MDD/AIMDD/IVDD records:

- the publication can be done per each single UDI-DI (GTIN) submission separately or in bulk.
- Multiple UDI-DI's can be sent in parallel.

For MDR/IVDR/SPP records:

1. it is required to start with the initial publication of the first UDI-DI belonging to the new Basic UDI-DI.
2. after this first submission has succeeded with CIC SYNCHRONISED (low risk class) or CIC RECEIVED (high risk class) the follow-up UDI-DIs of the same Basic UDI-DI can be published too.
3. repeat this sequencing order of 1+2 above for every first UDI-DI belonging to a Basic UDI-DI and its follow-up UDI-DIs

Use the following Publish to GLNs from your dedicated GDSN Data pool service:

GDSN Data Pool	Member Organisation	Publish To GLN
Synfony	GS1 Czech Republic	8594182509823 Synfony UDI Connector
GS1Trade Sync	GS1 Denmark	7609999484711 GS1 UDI Link
firstbase	GS1 Switzerland	4399902421386 firstbase UDI Connector
GS1 Data Source	GS1 Netherlands	7609999484728 GS1 Data Source UDI Connector

### Note #4

Please make sure, that your GDSN Data pool support has correctly setup the subscription for your participating Information Provider GLN by the relevant UDI Connector service. Only with a valid match between 'Publication' and 'Subscription' ("Pub-Sub-Match) the synchronisation between both partners can be proceeded. Normally this should be the case during the setup and onboarding ([Phase 1 - Onboarding Process](#)). After the publication is done, the relevant active subscription can be checked under 'subscriptions', for example:

GLN	Company name	Recipient data pool
4399902421386	firstbase UDI Connector	7612345000343 GS1 Switzerland - firstbase

## 6.2 Follow-up Submissions

After the publication is done, for any UDI submission EUDAMED gives direct responses like other UDI databases. As described in [Phase 3 - Learn UDI Upload and Testing](#) the CIC will be receipt with the status "SYNCHRONISED" (=Registered in EUDAMED) or "RECEIVED" (=Submitted in EUDAMED) will return. It is recommended to wait for this initial submission per each Basic UDI-DI. Only if this initial submission was successful, follow-up submissions for other UDI-DIs shall be sent. **Update submissions** for already registered records can be **only** sent if the CIC SYNCHRONISED (=Registered in EUDAMED) was provided.

### Note #5



**ATTENTION:** Please be aware that in EUDAMED Production the Notified Body Approval can take some time. High risk class devices cannot be updated before the "Registered" state is achieved.

### 6.3 Error Handling

When a CIC REVIEW has been returned, and EUDAMED has responded with an error. This can be the case for several reasons, for instance:

- Issues regarding the access point configuration (Actor settings) e.g. UDI Token is not valid.
- Technical problems in the M2M interface between the UDI Connector service (e.g. p36) and EUDAMED because of a service downtime.
- Content error, when the GDSN Data pool cannot validate against specific data e.g. an invalid Authorised Representative Actor Code (SRN).

In any case click on CIC status "View on Synclist" link and open the detailed description of the CIC error message.

<input type="checkbox"/>	Publication status	CIC Status	GTIN	Description short	Target market	Unit descriptor	Who should see this?
<input type="checkbox"/>	Published	<b>Review</b> <a href="#">View on Synclist</a>	07612345779171	7612345GOLDENtest40J7	EU	Pack or Inner Pack	Restricted <a href="#">View details</a>
<input type="checkbox"/>	Live		07612345779164	7612345GOLDENtest40J7	EU	Base Unit or Each	Restricted <a href="#">View details</a>

Last change date	Status	GTIN	Target market	Data source	Scope	Data recipient								
07/11/2025 16:32:35	<b>Review</b>	07612345779133	EU	7612345000435 UDI manufacturer POC/MVP	Public	4399902421386 firebase UDI Connector								
<table border="1"> <thead> <tr> <th>Status code</th> <th>Status code detail</th> <th>Description</th> <th>Corrective action code</th> </tr> </thead> <tbody> <tr> <td>CIC999</td> <td>Error reported by P36 UDI connect</td> <td>: elementReport&gt; &lt;message:operationErrorCode&gt;ERR-DTX-EUD-403.01 &lt;/message:operationErrorCode&gt; &lt;message:operationErrorDetail&gt;Only the owner can update their entities. DE-MF-000005765 is not the owner of 7612345GOLDENtest38JL GS1.&lt;/message:operationErrorDetail&gt; &lt;/message:elementReport</td> <td>ACTION_NEEDED</td> </tr> </tbody> </table>							Status code	Status code detail	Description	Corrective action code	CIC999	Error reported by P36 UDI connect	: elementReport> <message:operationErrorCode>ERR-DTX-EUD-403.01 </message:operationErrorCode> <message:operationErrorDetail>Only the owner can update their entities. DE-MF-000005765 is not the owner of 7612345GOLDENtest38JL GS1.</message:operationErrorDetail> </message:elementReport	ACTION_NEEDED
Status code	Status code detail	Description	Corrective action code											
CIC999	Error reported by P36 UDI connect	: elementReport> <message:operationErrorCode>ERR-DTX-EUD-403.01 </message:operationErrorCode> <message:operationErrorDetail>Only the owner can update their entities. DE-MF-000005765 is not the owner of 7612345GOLDENtest38JL GS1.</message:operationErrorDetail> </message:elementReport	ACTION_NEEDED											

**Important:** It is good to know that the communication in the GDSN is always related to the top level of trade item packaging hierarchies. Also, here the EUDAMED response message (CIC) is related to the highest level in the packaging hierarchy, e.g. Case unit. If you click on "Open in Editor", it brings you to the relevant trade item hierarchy, where you can revise the data for error correction.

Last change date	Status	GTIN	Target market	Data source	Scope	Data recipient										
08/12/2025 15:01:07	<b>Review</b>	07612345779713	EU	7612345000435 UDI manufacturer POC/MVP	Public	4399902421386 firebase UDI Connector										
<table border="1"> <thead> <tr> <th>Status code</th> <th>Status code detail</th> <th>Description</th> <th>Corrective action code</th> <th>Corrective action</th> </tr> </thead> <tbody> <tr> <td>CIC999</td> <td>Validation at p36 failed.</td> <td>Submission blocked due to business rules Entity 7612345GOLDENtest46JK is not submittable because of its current state (In Submission)</td> <td>ACTION_NEEDED</td> <td>ACTION_NEEDED /</td> </tr> </tbody> </table>							Status code	Status code detail	Description	Corrective action code	Corrective action	CIC999	Validation at p36 failed.	Submission blocked due to business rules Entity 7612345GOLDENtest46JK is not submittable because of its current state (In Submission)	ACTION_NEEDED	ACTION_NEEDED /
Status code	Status code detail	Description	Corrective action code	Corrective action												
CIC999	Validation at p36 failed.	Submission blocked due to business rules Entity 7612345GOLDENtest46JK is not submittable because of its current state (In Submission)	ACTION_NEEDED	ACTION_NEEDED /												

## Note #6

Usually, the error message must be interpreted by the user, as EUDAMED responds with specific references like error code, SRN, Basic UDI-DI Code (GMN) or UDI-DI Code (GTIN) and the related error detail. The detail includes the attribute name or code value in EUDAMED language. If the reader does not find the **error root cause** directly using the Web UI, they could look-up the information in the current “*GS1\_UDI\_Connector\_Profile\_Overview.xlsx*”, which helps to map against the GDSN attribute definitions and code list mappings. If you have any specific error questions, please contact your local data pool support team.

After fixing the error, the affected item must be **re-published**.

## 6.4 Support

If the error is misunderstood or for any kind of user question in your GDSN Data pool, please don't hesitate to contact your **local GS1 Member Organisation for support** or training. See [Data Pool Resources](#) section for links to relevant information.

If the error, its root cause or the user question is related to the EUDAMED system itself, please contact [SANTE-EUDAMED-SUPPORT@ec.europa.eu](mailto:SANTE-EUDAMED-SUPPORT@ec.europa.eu). Please make sure that you respect their required information.

## 6.5 Discard

For specific exceptions EUDAMED allows to withdrawal and discard existing UDI registrations. This “escape-out” should be only used, when the submitted data is incorrect and non-updateable fields cannot be changed after successful registration. As an example, a Basic UDI-DI codes were wrongly entered and linked to its UDI-DI (GTIN).



**ATTENTION:** The discard functionality should be only used in exceptional circumstances, because in EUDAMED it can be only done manually per each single UDI-DI.

It is very important to follow the discard process in the following three interfaces:

1. **EUDAMED Web-UI** – please refer to the official documentation from <https://webgate.ec.europa.eu/eudamed-help/en/documentation/user-guides-and-templates.html> and [Devices: UDI Devices](#): Section 5.2.7 Discard registered UDI-DIs/EUDAMED IDs (and their Basic UDI-DI/EUDAMED DI)
2. Discard the submission under the **3<sup>rd</sup> Party Provider's** (e.g. p36) **interface**. steps to follow will be detailed here very soon, probably later in 2026.
3. Or use the item withdrawal process of **GDSN Data pool**, e.g. by using the button **“Withdraw”** to be clicked for the published item – but only in these kinds of exceptional situations:



Items

New Item ▾ View Edit **Withdraw** Templates ▾ Export ▾

Filters applied: Data source 7612345000435 - UDI manufacturer POC/MVP ✕

<input type="checkbox"/>	Publication status	CIC Status	GTIN ↕	Description short	Target market	Unit descriptor	Who should see this?
<input checked="" type="checkbox"/>	<b>Published</b>	<b>Accepted</b> <a href="#">View on Synclist</a>	07612345779201	7612345GOLDENtest41J9	EU	Case	Restricted <a href="#">View details</a>

## 7 Sources of Information

### 7.1 Required Templates for Access Point Configuration

EUDAMED Playground: [Business justification](#), [Third-party agreement](#)

EUDAMED Production: [Third-party agreement](#)

### 7.2 Data Pool Resources (e.g. User Manuals)

p36 web tool: <https://app.udiconnect.io/>

GS1 Czech Republic Synfony: <https://synfony.cz/>

GS1Trade Sync: [GS1Trade Sync | Product data exchange made easy](#)

GS1 Netherlands Data Source: <https://www.gs1.nl/producten-services/data-exchange/gs1-data-source/gezondheidszorg/>

GS1 Switzerland firstbase: <https://www.firstbase.ch/de/support>

### 7.3 EUDAMED Information

EC's EUDAMED Website: [https://health.ec.europa.eu/medical-devices-eudamed\\_en?prefLang=de](https://health.ec.europa.eu/medical-devices-eudamed_en?prefLang=de)

Information Centre - EUDAMED Playground: <https://webgate.ec.europa.eu/eudamed-play-help/en/welcome-to-the-eudamed-information-centre.html>

Information Centre - EUDAMED Production: <https://webgate.ec.europa.eu/eudamed-help/en/welcome-to-the-eudamed-information-centre.html>

## 7.4 Table of Abbreviations

AIMDD	Active Implantable Medical Device Directive 90/385/EEC
AP	Access Point
CIC	Catalogue Item Conformation
DRIFT	Do it Right the First-Time approach
EC	European Commission
EU	European Union
EUDAMED	European Database on Medical Devices
GDSN	Global Data Synchronisation Network
GLN	Global Location Number
GMN	Global Model Number
GTIN	Global Trade Item Number
GUDID	Global Unique Device Identification Database
IFU	Instructions for Use
IVDD	In Vitro Medical Device Directive 98/79/EC
IVDR	In Vitro Diagnostic Regulation (EU) 2017/746
LAA	Local Actor Administration
M2M	Machine-to-Machine
MDD	Medical Device Directive 93/42/EEC
MDM	Master Data Manager
MDR	Medical Device Regulation (EU) 2017/745
MF	Legal Manufacturer (as Economic Operator in EUDAMED)
N/A	Not Applicable
POT	Proof of Testing
PR	System/Procedure Pack Producer (as Economic Operator in EUDAMED)
PROD	Production (Environment)
SPP	System or Procedure Pack
SRN	Single Registration Number
swissdamed	Swiss Database on Medical Devices
UAT	User Acceptance Testing
UDI	Unique Device Identification
UDI-DI	Unique Device Identification - Device Identifier
UI	User Interface
Web-UI	Web User Interface
XML	eXtensible Markup Language