

## EUDAMED user guide Legacy Devices registration

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## **1** Introduction

This document contains the details of how Legacy Devices are identified in EUDAMED and how you can register a Legacy Device in EUDAMED.<sup>1</sup>

#### INFOGRAPHIC: Identifiers of a legacy device





<sup>1</sup>For a wider understanding on how to use the platform, including FAQs and process infographics, visit the EUDAMED Information Centre. For information specific to UDI, visit the UDI Helpdesk.

## **2** Identification details

To keep the same standard structure and identification elements for all Devices registered in EUDAMED, an identification element EUDAMED DI (the equivalent of the Basic UDI-DI) will be required, and a EUDAMED ID (the equivalent of the UDI-DI in case no UDI-DI has been assigned) will be generated from the EUDAMED DI.

#### 2.1 Generation of identification details for a Legacy Device when a UDI-DI already exists

When the manufacturer already has a UDI-DI assigned to a Legacy Device, the manufacturer will be able to use it as a device identifier for the Legacy Device. In this case, the EUDAMED DI is automatically generated from the UDI-DI value.

The Legacy Device will therefore have the following identification elements: a EUDAMED DI (equivalent of the Basic UDI-DI and generated based on the UDI-DI) and a UDI-DI (provided by the manufacturer).

In order to generate the EUDAMED DI identification element from the UDI-DI provided, EUDAMED will use a standard format, placing the characters **B-** in front of the UDI-DI.

UDI-DI: M991CVS12130NES2

Generated EUDAMED DI: B-M991CVS12130NES2

#### 2.2 Generation of identification details for a Legacy Device when a UDI-DI does not exist

When the manufacturer does not have a UDI-DI assigned for a Legacy Device, it will have to assign a EUDAMED DI and EUDAMED ID.

The EUDAMED DI will have a strict format, starting with **B**- as a prefix and continuing with a set of characters. For further information on the EUDAMED DI format please consult the Format of the Unique Device Identifiers for the Legacy Devices document.

The EUDAMED ID will have the same format and value as EUDAMED DI except the first prefix character. It will start with **D**- instead of **B**-.

EUDAMED DI: B-BEMF000000106CR023335WE

EUDAMED ID: D-BEMF000000106CR023335WE

## 3 Registering Legacy Devices

On the dashboard, click on Register a Legacy device:

Welcome to EUDAI	MED	
	d by the European Commission to implement es and Regulation (EU) 2017/746 on in vitro	See all the news
MDR EUDAMED is structured around 6 inte	erconnected modules and a public site.	
Tasks		
By module, consult, verify and/or manage y	our own and related data (managed by your actor), o	depending on your profile.
	UDI-DIs/Device	User management
My Actor data	Register a new Basic UDI-DI	Assess user access requests
	Register a legacy device	Manage your users
Manage your actor data	Manage your Basic UDI-DIs / EUDAMED DIs Manage your Devices details	
Manage your email notifications	manage your period dotails	

# 3.1 Step 1: EUDAMED DI identification information

1. Select the applicable legislation:

Legacy Device registration

anisation name:	Belgian MF A
N:	BE-MF-00000041
ddress:	Rue A, 1 1060 Brussels
elephone number:	-
mail:	public-contact@belgian-mf-a.be
Applicable Legislation	
VDD (Directive 98/79/EC on in v	tro Diagnostic Medical Devices)
MDD (Directive 93/42/EEC on M	dical Devices)
AIMOD (Directive 00/285/EEC	ctive Implantable Medical Devices)

 Select Yes or No to whether a UDI-DI is already assigned to the legacy device. If yes, enter the Issuing Entity and the UDI-DI code, and click Generate. EUDAMED will create a corresponding EUDAMED DI (the UDI-DI code with "B-" as prefix).

Basic UDI-DI main in	formation
* Is it a kit?	
🔾 Yes 💿 No	
Special device type	
Yes 🚺 No	Special device type is required unless you select the option - No
* Special device type:	
<ul> <li>Software</li> </ul>	

3. Non-EU manufacturers have to select the authorised representative (AR) for the current device from the options available.

If there is only one AR with an active Mandate with the manufacturer, it will be automatically retrieved:



4. Click on **Save & Next** to save it as a draft and continue with the following steps:



5. On the left you will see a summary of the device characteristics. Choose a *Risk class* from the list and select **Yes** or **No** for each of the options.

Manufacturer identification BE-MF-000000041, Belgian MF <u>A</u>	EUDAMED DI information	2 Certificate information	3 Device identification information	4 Device characteristics	5 Device information	
EUDAMED DI identification Applicable legislation: IVDD (Directive 98/79/EC on in	EUDAMED * Risk class:	DI informatio	on			
Ultro Diagnostic Medical Devices)  EUDAMED DI code: B-56909 Issuing Entity: EUDAMED	* Near-patient	testing				
Kit: No Special device type: Software	* Self-patient	-				
	* Companion O Yes O No	diagnostic				
	* Reagent O Yes O No					
	★ Instrument ○ Yes ○ No					

6. Select **Yes** or **No** if the device model is applicable and, if applicable, enter the Device model and enter a Device name if there is one, otherwise enter only a Device name:

Device model applicable Yes No	Device model is required by default unless you select the option - No
* Device model:	
Device Name:	

7. Click on **Save** to save your draft and complete it later, or **Save & Next** to save it as a draft and continue with the following steps:

Save	Save & Next >
Guile	

#### 3.2 Step 2: Certificate information

Select a certificate type, enter the NB number or name and click **Find**. Enter the *Certificate number* and *Expiry date*. If applicable, enter a *Revision number*.

<b>NOTE</b> <i>Annex - Legacy Device Certificate</i> Devices.	Types [16] lists the certificate types for Legacy

ertificate information	
em #1	•
Certificate Type:	
EC Certificate Full Quality Assurance System	~
	🖍 Change Notified Body
Organisation name: EVPU a.s.	
NB number: 1293	
Address:	
Telephone number: 421 42 44 03 600	
Email: hudak@evpu.sk	
Certificate number:	Revision number:
276898081	
Expiry date:	
2021-06-30	
YYY-MM-DD	
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# **3.3 Step 3: Device identification information**

EUDAMED will display the identifier of the Device (the previously provided UDI-DI or the EUDAMED **ID** generated based on the provided/generated EUDAMED **DI**. EUDAMED ID has the same code as the EUDAMED DI, except that it is with a "**D**-" prefix instead of the "**B**-" prefix):

ssuing Entity:		* EUDAMED ID code:	
UDAMED	~	D-LM100X3PL	

1. Enter the EMDN code. Click on Find and select the correct one:



2. If applicable, enter the trade name and select the language, otherwise select No:

Trade name:	* Select the language:
Trade_Name_01	_ I 🗸

3. Enter a reference/catalogue number and any additional information you might have:

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Additional product description:	Select the I	anguage:
		× •
	#	
Add additional product description in anoth	ner language	
21 for additional information (or alcohomic instru-	ctions for use):	
RL for additional information (as electronic instrue		
CL for additional information (as electronic instruc		
Device status:		

4. Choose the market status of the Device:

On the EU market	~	
On the EU market		
No longer placed on the EU market		

#### **3.4 Step 4: Device characteristics**

1. Select **Yes** or **No** for the first three options, then select **Yes** or **No** whether if Storage/ handling conditions are applicable:

* Labelled as single use
○ Yes ○ No
* Need for sterilisation before use
○ Yes ○ No
* Device labelled as sterile
○ Yes ○ No
Storage/handling conditions, if applicable
Yes No Storage/handling conditions are required unless you select the option - No
* Storage/handling conditions type: Description:
Add another storage/handling condition

2. If applicable, provide the correct values by selecting from the options provided and enter a description:

corage/handling conditions, if appli es No	Storage/handling conditions are required unless you select	at the option - No
Storage/handling conditions type:	Description:	
_		

3. Select **Yes** or **No** for Critical warnings or contra-indications and if Yes, enter the type and description. After completing, click on Save or Save & Next:

Critical warnings or contra-ind	ications, if applicable
Yes 🚺 No	Critical warning or contra-indications are required unless unless you select the option - No
* Critical warning type:	Description
	✓
Add critical warnings or con	<u>tra-indications</u>

#### 3.5 Step 5: Device information

1. Select **Yes** or **No** if the device was designed and manufactured by another legal or natural person.

If Yes, there are two different ways to find the *Product original manufacturer* of the device:

• Check the box *I know the Actor ID/SRN*, enter the Actor ID/SRN or name of the *Product original manufacturer* of the device and click **Check registry**:

Is the device designed and manufactured by another legal or natural person?				
Yes No				
I know the Actor ID/SRN				
* Enter Actor ID/SRN or name:				
	Q Check registry			



#### NOTE

Please ensure to check the box *I know the Actor ID/SRN* in order to search for an existing registered Manufacturer Actor either by SRN or by name.

Select the Actor from the list:

Select manufacturer	×Clo
Actor ID/SRN 1	Organisation name It
NL-MF-00000041	Medical Device Manufacturer
AU-MF-000004268	Trusted NonEUMF
AS-MF-000004249	Non_EU_MF_R3.3_Shriya
BE-MF-000004247	Bel_MF_R3.3_Shriya
US-MF-000003888	The Americans
US-MF-000004107	Ohio Pharmaceuticals
CO-MF-000004129	Non_EU_MF_3.2_Shriya
BE-MF-000004128	MF_BE_R3.2_Shriya
EL-MF-000004067	VIANEX S.A.
AI-MF-000004047	AR Aguilla Ionut 2nd
+ Previous	1 2 19 Next →
Close	

• Enter the name of the *Product original manufacturer organisation name* and click on **Check registry**:



Select manufacturer	× <u>Close</u>
Organisation name 11	
PDasOrg (3)	
PDasOrg (2)	
MANUF-1(1)	
Select the relevant result above. If there are no results or they are not applicable, please select the option data manually'	'Enter
Enter data manually Cancel	

If the Organisation name is not in the list, click on **Enter data manually** and fill in the required fields with the details on the *Product original manufacturer* of the device:

* Name (Manufacturer Name):	Change manufacture
PoMasOrg	
Street information, if applicable	
Yes 🚺 No 🚯 Street informa	ation is required unless you select the option - No
* Street:	Street number:
Via de Rosso	10
Address line 2:	
PO box: * City name:	* Postal code:
Milan	
* Country:	
× •	
Telephone:	
Telephone format example: +32 x xxx xx xx	
* Email:	

2. Select **Yes** or **No** to provide the Clinical Investigation reference(s):



3. Select **Yes** or **No** for the three following options on Tissues and cells:

Yes 🔿 No			
	tissues or cells, or the	ir derivatives:	
Yes 🔿 No			
Presence of cells	substances of microb	ial origin:	
) Yes 💿 No			

4. Select a Member State from the drop-down list where the device has been placed on the EU market, and click on **Submit** to submit it:

* Member State	e where the Device is to or has bee	n first placed on the EU market:
France	~	
Member States	where the device is or is to be man	de available on the market:
Finland	From YYYY-MM-DD	To To TYYYY-MM-DD
France	From YYYY-MM-DD	To H
Select one or m	nore countries >	

5. Once you confirm your submission, you will be brought to a page confirming the submission of your Legacy device:



# 4 Management of Regulation Devices and Legacy Devices

For further information on how to manage Regulation and Legacy Devices please consult the following chapters of the dedicated UDI Devices - user guide:

- · Manage your device Basic UDI-DI/EUDAMED DI details
- · Manage your device UDI-DI/EUDAMED ID details
- · Search & View Devices, Systems and/or Procedure Packs

# 5 Linking the Regulation Devices to Legacy Devices

EUDAMED will allow linking a Regulation Device with the correspondent Legacy Device and it will perform this linking automatically when both have the same UDI-DI. The link will be made at the level of the UDI-DI.



#### NOTE

See Sections Link a registered Regulation Device to a registered Legacy Device and Delete the link between a Regulation Device and a Legacy Device of the UDI Devices user guide.

# 6 Annex – Legacy Device certificate types

This Annex presents the certificate types that can be used when registering a Legacy Device.

Certificate types depend on the applicable legislation of the Device.

Applicable	Certificate Type
Legislation	
MDD	Directive 93/42/EEC Annex II excluding section 4
	Directive 93/42/EEC Annex II section 4
	Directive 93/42/EEC Annex III
	Directive 93/42/EEC Annex IV
	Directive 93/42/EEC Annex V
	Directive 93/42/EEC Annex VI
AIMDD	Directive 90/385/EEC Annex 2 excluding section 4
	Directive 90/385/EEC Annex 2 section 4
	Directive 90/385/EEC Annex 3
	Directive 90/385/EEC Annex 4
	Directive 90/385/EEC Annex 5
IVDD	Directive 98/79/EC Annex III section 6
	Directive 98/79/EC Annex IV excl. section 4 and 6
	Directive 98/79/EC Annex IV section 4
	Directive 98/79/EC Annex IV section 6
	Directive 98/79/EC Annex V
	Directive 98/79/EC Annex VI
	Directive 98/79/EC Annex VII excluding section 5
	Directive 98/79/EC Annex VII section 5

