IVDR

Basic UDI-DI set of data in UDI database

Principle: Each UDI-DI inherits the attributes of its linked Basic UDI-DI and devices DI

Basic UDI-DI

Applicable legislation (IVDR) (*)

- •2. Basic UDI-DI value (*)
- •2b Basic UDI-DI Issuing entity (*);
- •6. Manufacturer SRN (*)
- •5. Name and address of manufacturer
- •7. Name and address and SRN of AR
- •9. Risk class (*)
- •A.2.14 Intended for self-testing (Y/N) (*)
- •A.2.14 Intended for near-patienttesting (Y/N) (*)
- •Companion diagnostics (Y/N) (*)
- •Intrument(Y/N) (*)
- •Reagent(Y/N) (*)
- •Professional testing (Y/N) (*)

•11. A. Name and/or, if applicable, device model that identifies the device(s) with this BASIC UDI-DI in the technical documentation and/or certificate or declaration of conformity (Name and/or model shall be provided)

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UDI-DIs

- •0. UDI-DI value (*)
- •0b. UDI-DI Issuing entity (*)
- •Secondary DI (value and issuing entity)
- •11.B. Reference, Article or Catalogue number (*)
- •Device with Direct marking (Y/N) (*)
- •Direct marking UDI-DI value (*)
- •Direct marking UDI-DI issuing entity (*)
- •1. Quantity of device(s) (*)
- •3. Type of UDI-PI (*)
- •4. Unit of use UDI-DI (*)
- •13. Storage/handling conditions
- •10-14. Name(s)/Trade name(s) (including languages)
- •12. Additional product description
- •19. URL for additional information
- •15. Labelled as single use (YN) (*)
- •16. Maximum number of reuse (*)
- •17. Device labelled sterile (Y/N) (*)
- •18. Need for sterilisation (Y/N) (*)
- •20. Critical warnings or contra-indications
- •8. Medical device nomenclature (CND) code (1)
- •21. Status
- •27 (A.2.10). In the case of devices designed and manufactured by another legal or natural person as referred in Article 10(14), the name, address and contact details of that natural/legal person

UDI-DIs (container package DI)

•0. UDI-DI value (*)
• 0b. Issuing entity (*)
•1. Quantity per package (*)
•21. Status

(1) Nomenclature decision: <u>https://ec.europa.eu/doc</u> <u>sroom/documents/34264</u>

(*) may not be changed
 Mandatory
 Mandatory if applicable
 Optional

IVDR



Other Device Data attributes

Basic UDI-DI

A.2.2 Certificate IDs (with NB, type .. Link);
A.2.11 SSP;
A.2.9 Performance study IDs (..link);
A.2.5 Presence of Human tissues/Cells (Y/N) (*);
A.2.6 Presence of Animal tissues/Cells (Y/N) (*)
A.2.7 Presence of Substances/cells of microbial origin (Y/N) (*);
Kit (Y/N) (*);

UDI-DIs

•A.2.13 New Device (Y/N) (*);
•A.2.3 Member State of the Placing on the EU Market of the Device (*);

•A.2.4 Member State(s) were the Device is made available in the Country;





(*) may not be changed Mandatory Mandatory if applicable Optional