

Extension of the IFA database as of 15.12.2025: Information for supplier



Foreword

The EU Deforestation Regulation – Regulation (EU) 2023/1115 (EUDR) – is expected to enter into force on 30 December 2025. The IFA database will contain two new data fields on the publication date of 15 December 2025.

The [Guidelines for Notifying Product and Address Data](#) have been expanded to include the new features.

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1. New data fields

1.1 EUDR – EU-Entwaldungsverordnung – EU Deforestation Regulation

It must be indicated whether the product belongs to a product group that is subject to Regulation (EU) 2023/1115 (EUDR). The supplier identifies whether a product is subject to EUDR in a two-step process, whereby (I) and (II) both must be fulfilled:

- (I) A product is generally subject to the EUDR if its HS code is listed in Annex I, right-hand column (*'Relevant products'*) of the EUDR. This is a conclusive list: a product whose HS code is not listed is not subject to the EUDR. For example, medicinal products are not listed here as relevant products.
- (II) If a product is in principle subject to the EUDR according to (I), the relevant commodity must also be included in accordance with Annex I, left column (*'Relevant commodity'*) of the EUDR in order to determine the final EUDR obligation of the product. However, if the product does not contain the relevant commodity, it is not subject to the EUDR – even if (I) is fulfilled.

Suppliers who sell products subject to the EUDR must comply with the due diligence obligations of the EUDR and submit a due diligence statement to the competent authorities. Without prior submission, a supplier shall not place any products subject to the EUDR on the market in Germany.

Acceptable values	0	no
	1	yes, product subjects to EUDR

Please indicate in the EAD file in the appendix whether your products are subject to EUDR or not.

- Medicinal products are not relevant products according to Annex I of the EUDR, so they are assigned the *value 0 = no* in the IFA database.
- Medical devices and other pharmacy-typical products must be identified according to the two-step procedure (I) and (II) described above and labelled accordingly.

1.2 HS-Code – HS code

The HS code under which all products are classified, must be specified. The nomenclature of Harmonized Commodity Description and Coding System, commonly known as the 'HS nomenclature', is an international multi-purpose nomenclature. It was developed by the World Customs Organisation (WCO), assigns six-digit codes for the classification of products into specific product groups and is valid worldwide.

An HS code is already required for customs clearance for every import into the EU. If this is not yet available, it can be determined, for example, via the information system of the German Generalzolldirektion [EZT-Online](#).

Acceptable values	6-digit numeric data field
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Please indicate the applicable HS codes (first six-digits of the customs tariff number) for all products (medicinal products, medical devices and other pharmacy-typical products) **in the EAD file attached**.