EC REP

european authorized representative



german office

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The Medical Device Regulations (MDR (EU) 2017/745 & IVDR (EU) 2017/746) require the designation of an **Authorised Representative (EAR)** for those legal manufacturers who are based outside the European Union. The EAR must be based within the EU. The relevant legal requirements are described in both regulations in Art. 11.

Art. 2 (32) MDR/Art. 2 (25) IVDR

'authorised representative' means any natural or legal person established within the Union who has received and accepted a written mandate from a manufacturer, located outside the Union, to act on the manufacturer's behalf in relation to specified tasks with regard to the latter's obligations under this regulation;

Art. 11 (1) MDR/IVDR

Where the manufacturer of a device is not established in a Member State, the device may only be placed on the Union market if the manufacturer designates a sole authorised representative.



MT Promedt Consulting GmbH, a Germany based organisation is acting since 1995 as a reliable European Authorised Representative and is providing qualified and professional services in all EU Member States under the new regulations.



us office

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The EAR is a registered economic operator. The EAR services under the new regulations will include:

- / provision of an authorised European address / verification activities regarding EU declaration of conformity, technical documentation and appropriate conformity assessment procedure
- / keeping available a copy of the technical documentation, the EU declaration of conformity and, if applicable, a copy of the relevant certificate, including any amendments and supplements, issued by the notified body / verification of compliance with all registration obligations for the EAR and manufacturer
- / interaction with competent authorities providing all relevant information and documentation necessary to demonstrate the conformity of a device as well as product samples on demand

- / liaison between competent authorities, manufacturer, importer, distributor in vigilance aspects
- / cooperation with the competent authorities on any preventive or corrective action taken to eliminate or mitigate the risks posed by devices;
- / information of the manufacturer about complaints and reports from healthcare professionals, patients and users about suspected product related incidents

Our EAR services also include

/ access to free sales certificate/ EAR under other specific regulations for non-medical devices / in-house and online trainings

All clients have permanent and safe access to our document server system. Make use of our personalised EAR services. Our EAR services meet the requirements of the new regulations and provide you full flexibility in your European distribution network.



