Extension of the MDR Transitional Period and Removal of the "Sell Off" Periods - Q&A on Practical Aspects Related to the Implementation of Regulation (EU) 2023/607, Apr-2023

The following document of the European Commission provides the Q&A document that is intended to facilitate the application of the "Regulation (EU) 2023/608 of the European Parliament and of the Council Amending Regulations (EU) 2017/745 and (EU) 2017/746 as regards the Transitional Provisions for Certain Medical Devices and In Vitro Diagnostic Medical Devices, 25-Apr-2023".

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