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Device Week, 13 October 2021 – Spotlight On The Latest Medtech Industry News From The EU

by Amanda Maxwell

In this week's podcast, EU regulatory editor Amanda Maxwell brings listeners up to speed on the latest news from the European Union. Topics include discussion of ISO 13485, ongoing industry concerns with MDR implementation, and the Eudamed database expansion.

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Medtech Insight articles addressing topics discussed in this episode:

- European Regulatory Roundup, September 2021: EN ISO 13485 Updated And More Key Implementation Developments
- EU Classification 'Bible' Published To Support Risk Class Decision-Making
- <u>Two New Modules Added To Eudamed For Use On Voluntary Basis</u>
- EN ISO 13485 and EN ISO 14971: Next Wave Of EU Standards Now Due "By December"