KCI Prepares for European Union Medical Device Regulation Transition

KCI, an Acelity Company, is preparing to meet the requirements outlined in the European Union Medical Devices Regulation (MDR). During this transitional period, KCI is actively updating its technical documentation and processes to ensure compliance with MDR requirements when it comes into force on 26 May 2020.

Notwithstanding the effective date of 26 May 2020, the MDR provides a period of grace for compliance certification issued by Notified Bodies and medical devices placed on the market before the effective date.

KCI is confident in its ability to meet all MDR requirements and deadlines ensuring our customers benefit from continued and compliant access to our technologies in Europe. As always, our focus is on providing best-in-class therapies.

Specific enquiries relating to KCI’s transition to MDR compliance can be submitted by email to EUMDR@Acelity.com.

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