
Medicinski pripomočki - Navodilo za uporabo ISO 14971 - 2. del: Strojno učenje v umetni inteligenci (ISO/DTS 24971-2:2025)

Medical devices - Guidance on the application of ISO 14971 - Part 2: Machine learning in artificial intelligence (ISO/DTS 24971-2:2025)

Medizinprodukte- Leitfaden zur Anwendung von ISO 14971- Teil 2: Maschinelles Lernen in der künstlichen Intelligenz (ISO/DTS 24971-2:2025)

Dispositifs médicaux - Recommandations pour l'application de l'ISO 14971 - Partie 2: Apprentissage automatique dans le cadre de l'intelligence artificielle (ISO/DTS 24971-2:2025)

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Part 2:

Machine learning in artificial intelligence

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