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**ISO/IEEE
11073-10206**

Health informatics — Device interoperability —

Part 10206: Personal health device communication — Abstract content information model

Informatique de santé — Interopérabilité des dispositifs —

*Partie 10206: Communication entre dispositifs de santé
personnels — Modèle d'information de contenu abstrait*

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Institute of Electrical and Electronics Engineers, Inc
3 Park Avenue, New York
NY 10016-5997, USA

Email: stds.ipr@ieee.org
Website: www.ieee.org

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