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AMENDMENT 1
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**Medical devices — Hierarchical coding
structure for adverse events —**

**Part 1:
Event-type codes**

AMENDMENT 1

*Dispositifs médicaux — Structure de codage pour la cause et le type
d'événement défavorable —*

Partie 1: Codes de type d'événement

AMENDEMENT 1



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Foreword

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Amendment 1 to ISO/TS 19218-1:2011 was prepared by Technical Committee ISO/TC 210, *Quality management and corresponding general aspects for medical devices*.