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INTERNATIONAL STANDARD

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**Medical electrical equipment -
Part 2-89: Particular requirements for the basic safety and essential performance
of medical beds for children**

**Appareils électromédicaux -
Partie 2-89: Exigences particulières pour la sécurité de base et les performances
essentiels des lits médicaux pour enfants**

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Medical electrical equipment - Part 2-89: Particular requirements for the basic safety and essential performance of medical beds for children

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This publication is published as a double logo standard.

The text of this International Standard is based on the following documents of IEC:

Draft	Report on voting
62D/2239/FDIS	62D/2272/RVD

Full information on the voting for its approval can be found in the report on voting indicated in the above table. In ISO, the document was approved by XXX P members out of YYY having cast a vote.

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