



FINAL DRAFT Amendment

ISO 10993-17:2023/ FDAM 1

Biological evaluation of medical devices —

Part 17:

Toxicological risk assessment of medical device constituents

AMENDMENT 1

Évaluation biologique des dispositifs médicaux —

Partie 17: Appréciation du risque toxicologique des constituants

des dispositifs médicaux

AMENDEMENT 1

ISO/TC 194

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