



FINAL DRAFT Technical Report

ISO/DTR 14872

Health informatics — Identification of medicinal products — Core principles for maintenance of identifiers and terms

*Informatique de santé — Identification des médicaments —
Principes essentiels pour la mise à jour des identifiants et des
termes*

ISO/TC 215

Secretariat: **ANSI**

Voting begins on:
2025-05-16

Voting terminates on:
2025-07-11

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Published in Switzerland

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Foreword

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This document was prepared by Technical Committee ISO/TC 215, *Health informatics*.

This second edition cancels and replaces the first edition (ISO/TR 14872:2019), which has been technically revised.

The main changes are as follows:

- the focus on describing the current (2024) implementation experiences and processes has been strengthened;
- references to a “federated service delivery model” have been removed, since this remains inspirational and does not correspond to the current practice;
- information about operating models (governance) of existing maintenance organizations impacting IDMP on a global level has been included.

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