
Zdravstvena informatika - Zahteve za mednarodne strojno berljive kode za pakiranje zdravil (ISO/DIS 16791:2025)

Health informatics - Requirements for international machine-readable coding of medicinal product package identifiers (ISO/DIS 16791:2025)

Medizinische Informatik - Anforderungen für internationale maschinenlesbare Kodierungen von Identifikatoren für Arzneimittelpackungen (ISO/DIS 16791:2025)

Informatique de santé - Exigences relatives au codage international lisible par machine des identifiants de paquetage de médicaments (ISO/DIS 16791:2025)

Ta slovenski standard je istoveten z: prEN ISO 16791

[oSIST prEN ISO 16791:2025](https://standards.net/en/catalog/standards/sist/c36a0f56-1d00-4101-8a2c-03a0a1570c00/osist-pr-en-iso-16791-2025)

ICS:

35.040.50	Tehnike za samodejno razpoznavanje in zajem podatkov	Automatic identification and data capture techniques
35.240.80	Uporabniške rešitve IT v zdravstveni tehniki	IT applications in health care technology

oSIST prEN ISO 16791:2025

en,fr,de



DRAFT International Standard

ISO/DIS 16791

Health informatics — Requirements for international machine-readable coding of medicinal product package identifiers

*Informatique de santé — Exigences relatives au codage
international lisible par machine des identifiants de paquetage de
médicaments*

ICS: 35.240.80

ISO/TC 215

Secretariat: **ANSI**

Voting begins on:
2025-03-03

Voting terminates on:
2025-07-04

<https://standards.iteh.ai/catalog/standards/sist/e56a8f36-1d60-4401-8a3e-83af0af370e0/osist-pren-iso-16791-2025>

This document is circulated as received from the committee secretariat.

ISO/CEN PARALLEL PROCESSING

Reference number
ISO/DIS 16791:2025(en)

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Published in Switzerland

ISO/DIS 16791:2025(en)

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This document was prepared by Technical Committee ISO/TC 215, *Health informatics*, in collaboration with the European Committee for Standardization (CEN) Technical Committee CEN/TC 251, *Health informatics*, in accordance with the Agreement on technical cooperation between ISO and CEN (Vienna Agreement).

This third edition cancels and replaces the second edition (ISO/TS 16791:2020), which has been technically revised.

The main changes to the previous edition are as follows:

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The main changes to the previous edition are as follows:

- addition of a definition on ePL [3.1.11](#)
- Adjustment of [5.5.1](#) to reference ePL of ;
- Addition of [Annex F](#).

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