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Health informatics - Requirements for international machine-readable coding of medicinal product package identifiers (ISO/TS 1 6791:2020)

Contents	Page
European foreword	3
Foreword	4
Introduction	5
1 Scope	6
2 Normative references	6
3 Terms, definitions and abbreviated terms	6
3.1 Terms and definitions	6
3.2 Abbreviated terms	11
4 Procedural background	11
4.1 General	11
4.2 Identification	11
4.3 International machine-readable coding	12
4.4 Medicinal product	12
4.5 Labelling	13
4.6 Package identifier	13
4.7 Serialization	14
5 Usage requirements	14
5.1 General	14
5.2 Traceability	15
5.2.1 Principles	15
5.2.2 Guidelines	16
5.3 Measures to combat falsification of medicines	17
5.3.1 Principles	17
5.3.2 Guidelines for both approaches	18
5.3.3 Product authentication	18
5.3.4 Supply chain integrity	19
5.4 Improving patient safety at point of care	19
5.4.1 Principles	19
5.4.2 Guidelines	19
5.5 Support of healthcare systems	20
5.5.1 Principles	20
5.5.2 Guidelines	21
5.6 Procurement and stock management	21
5.6.1 Principles	21
5.6.2 Guidelines	22
5.7 Overview of guidelines	22
6 Economic aspects	22
6.1 General	22
6.2 Manufacturer perspective	23
6.3 Healthcare provider perspective	23
Annex A (informative) Relationship between PhPID and MPID	24
Annex B (informative) Packaging hierarchy, relationship between MPID, PCID and GTIN®	26
Annex C (informative) Identification of trade items and logistic units	28
Annex D (informative) Examples for Package Identifier	29
Annex E (informative) Personalized Medicine	37
Bibliography	38