

ISO/TR 14872:2025-12 (E)

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

Contents		Page
Foreword		iv
Introduction		v
1	Scope	1
2	Normative references	1
3	Terms and definitions	1
4	Abbreviated terms	2
5	Maintenance of International Standards on IDMP -- description of existing relationships between regulators and other standards development organizations	2
6	IDMP terminology maintenance	3
6.1	IDMP maintenance organizations at global level	3
6.2	Core data within IDMP and data that is "derived"	3
7	Current maintenance processes	3
7.1	Generic description of governance and maintenance process	3
7.2	Governance and maintenance: substances and Global Pharmaceutical Product Identifier (PhPID)	4
7.3	Governance and maintenance: pharmaceutical dose forms	5
7.4	Governance and maintenance: units of measurement	5
7.5	Governance for SNOMED interactions with IDMP	6
7.6	Governance for MedDRA	7
7.7	Governance for Anatomical Therapeutic Chemical (ATC) classification system and Defined Daily Dose (DDD)	8
7.8	Governance for International Nonproprietary Names (INN)	9
7.9	Governance for GS1	10
8	IDMP core maintenance processes	11
8.1	General	11
8.2	General processes	11
8.3	Terminology processes	11
8.4	Unambiguity	12
9	International mapping and language translations	12
10	Integration of IDMP core with data sources that are considered "derived"	12
Bibliography		14