

SLOVENSKI STANDARD

SIST EN ISO 11238:2013

01-februar-2013

Zdravstvena informatika - Identifikacija medicinskih izdelkov - Elementi in zgradba podatkov za enotno identifikacijo in izmenjavo predpisanih informacij o substancah (ISO 11238:2012)

Health informatics - Identification of medicinal products - Data elements and structures for the unique identification and exchange of regulated information on substances (ISO 11238:2012)

Medizinische Informatik - Identifikation von Arzneimitteln - Struktur und kontrollierte Vokabularen zur Identifikation und Beschreibung von Substanzen und Inhaltsstoffen (ISO 11238:2012)

iTeh STANDARD PREVIEW
(standards.iteh.ai)
SIST EN ISO 11238:2013
<https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-400240300000/iso-11238-2012>
Informatique de santé - Identification des médicaments - Éléments de données et structures pour l'identification unique et l'échange d'informations réglementées sur les substances (ISO 11238:2012)

Ta slovenski standard je istoveten z: EN ISO 11238:2012

ICS:

11.020	Medicinske vede in zdravstvenovarstveni pripomočki na splošno	Medical sciences and health care facilities in general
35.240.80	Uporabniške rešitve IT v zdravstveni tehniki	IT applications in health care technology

SIST EN ISO 11238:2013

en

iTeh STANDARD PREVIEW
(standards.iteh.ai)

SIST EN ISO 11238:2013

<https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-93a8a29a468f/sist-en-iso-11238-2013>

EUROPEAN STANDARD
NORME EUROPÉENNE
EUROPÄISCHE NORM

EN ISO 11238

November 2012

ICS 35.240.80

English Version

Health informatics - Identification of medicinal products - Data elements and structures for the unique identification and exchange of regulated information on substances (ISO 11238:2012)

Informatique de santé - Identification des médicaments -
Éléments de données et structures pour l'identification
unique et l'échange d'informations réglementées sur les
substances (ISO 11238:2012)

Medizinische Informatik - Identifikation von Arzneimitteln -
Struktur und kontrollierte Vokabularien zur Identifikation
und Beschreibung von Substanzen und Inhaltsstoffen (ISO
11238:2012)

This European Standard was approved by CEN on 24 May 2012.

CEN members are bound to comply with the CEN/CENELEC Internal Regulations which stipulate the conditions for giving this European Standard the status of a national standard without any alteration. Up-to-date lists and bibliographical references concerning such national standards may be obtained on application to the CEN-CENELEC Management Centre or to any CEN member.

This European Standard exists in three official versions (English, French, German). A version in any other language made by translation under the responsibility of a CEN member into its own language and notified to the CEN-CENELEC Management Centre has the same status as the official versions.

[SIST EN ISO 11238:2013](https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-9b3a2546c98f/iso-11238-2012)

CEN members are the national standards bodies of Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, Former Yugoslav Republic of Macedonia, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Romania, Slovakia, Slovenia, Spain, Sweden, Switzerland, Turkey and United Kingdom.



EUROPEAN COMMITTEE FOR STANDARDIZATION
COMITÉ EUROPÉEN DE NORMALISATION
EUROPÄISCHES KOMITEE FÜR NORMUNG

Management Centre: Avenue Marnix 17, B-1000 Brussels

Contents

Page

Foreword.....	3
---------------	---

iTeh STANDARD PREVIEW (standards.iteh.ai)

[SIST EN ISO 11238:2013](https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-93a8a29a468f/sist-en-iso-11238-2013)

<https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-93a8a29a468f/sist-en-iso-11238-2013>

Foreword

This document (EN ISO 11238:2012) has been prepared by Technical Committee ISO/TC 215 "Health informatics" in collaboration with Technical Committee CEN/TC 251 "Health informatics" the secretariat of which is held by NEN.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by May 2013, and conflicting national standards shall be withdrawn at the latest by May 2013.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. CEN [and/or CENELEC] shall not be held responsible for identifying any or all such patent rights.

According to the CEN/CENELEC Internal Regulations, the national standards organisations of the following countries are bound to implement this European Standard: Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, Former Yugoslav Republic of Macedonia, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Romania, Slovakia, Slovenia, Spain, Sweden, Switzerland, Turkey and the United Kingdom.

iTeh STANDARD PREVIEW **(standards.iteh.ai)**

Endorsement notice

The text of ISO 11238:2012 has been approved by CEN as a EN ISO 11238:2012 without any modification.

SIST EN ISO 11238:2013

<https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-93a8a29a468f/sist-en-iso-11238-2013>

iTeh STANDARD PREVIEW
(standards.iteh.ai)

SIST EN ISO 11238:2013

<https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-93a8a29a468f/sist-en-iso-11238-2013>

INTERNATIONAL STANDARD

ISO
11238

First edition
2012-11-01

Health informatics — Identification of medicinal products — Data elements and structures for the unique identification and exchange of regulated information on substances

*Informatique de santé — Identification des médicaments — Éléments
de données et structures pour l'identification unique et l'échange
d'informations réglementées concernant les substances*

iTeh STANDARD PREVIEW
(standards.iteh.ai)

SIST EN ISO 11238:2013

<https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-93a8a29a468f/sist-en-iso-11238-2013>



Reference number
ISO 11238:2012(E)

© ISO 2012

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 11238:2013

<https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-93a8a29a468f/sist-en-iso-11238-2013>



COPYRIGHT PROTECTED DOCUMENT

© ISO 2012

All rights reserved. Unless otherwise specified, no part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying and microfilm, without permission in writing from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office
Case postale 56 • CH-1211 Geneva 20
Tel. + 41 22 749 01 11
Fax + 41 22 749 09 47
E-mail copyright@iso.org
Web www.iso.org

Published in Switzerland

Contents

Page

Foreword	iv
Introduction	v
1 Scope	1
2 Terms, definitions, symbols and abbreviated terms	1
2.1 Terms and definitions	1
2.2 Symbols and abbreviated terms	8
3 Requirements	9
3.1 General	9
3.2 Concepts required for the unique identification and description of substances	9
3.3 Concepts required for the description of specified substances	11
3.4 Naming of substances	12
3.5 Requirements for unique identifiers	13
3.6 Types of substances	14
3.7 Defining specified substances	27
Annex A (informative) Existing identifiers and molecular structure representations	35
Bibliography	38

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 11238:2013

<https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-93a8a29a468f/sist-en-iso-11238-2013>

ISO 11238:2012(E)

Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting a vote.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights.

ISO 11238 was prepared by Technical Committee ISO/TC 215, *Health informatics*.

iTeh STANDARD PREVIEW
(standards.iteh.ai)

[SIST EN ISO 11238:2013](https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-93a8a29a468f/sist-en-iso-11238-2013)

<https://standards.iteh.ai/catalog/standards/sist/c4484857-a56e-4da5-b747-93a8a29a468f/sist-en-iso-11238-2013>

Introduction

This International Standard was developed in response to a worldwide demand for internationally harmonized specifications for medicinal products. It is one of a group of five standards which together provide the basis for the unique identification of medicinal products. The group of standards comprises:

ISO 11615, *Health informatics — Identification of medicinal products — Data elements and structures for the unique identification and exchange of regulated medicinal product information*;

ISO 11616, *Health informatics — Identification of medicinal products — Data elements and structures for the unique identification and exchange of regulated pharmaceutical product information*;

ISO 11238, *Health informatics — Identification of medicinal products — Data elements and structures for the unique identification and exchange of regulated information on substances*;

ISO 11239, *Health informatics — Identification of medicinal products — Data elements and structures for the unique identification and exchange of regulated information on pharmaceutical dose forms, units of presentation, routes of administration and packaging*;

ISO 11240, *Health informatics — Identification of medicinal products — Data elements and structures for the unique identification and exchange of units of measurement*.

These standards for the identification of medicinal products (IDMP) support the activities of medicines regulatory agencies worldwide by jurisdiction. These include a variety of regulatory activities related to development, registration and life cycle management of medicinal products, as well as pharmacovigilance and risk management.

To meet the primary objectives of the regulation of medicines and pharmacovigilance, it is necessary to reliably exchange medicinal product information in a robust and reliable manner. The IDMP standards therefore support the following interactions:

- between one medicine regulatory agency and another, e.g. European Medicines Agency to the US Food and Drug Administration (FDA); or vice versa;
- between pharmaceutical companies and medicine regulatory agencies, e.g. “Pharma Company A” to Health Canada;
- between the sponsor of a clinical trial to a medicine regulatory agency, e.g. “University X” to the Austrian Medicines Agency;
- between a medicine regulatory agency and other stakeholders, e.g. UK Medicines and Health Care Products Regulatory Agency (MHRA) to the National Health Service (NHS);
- between medicine regulatory agencies and worldwide-maintained data sources, e.g. the Pharmaceutical and Medical Device Agency (PMDA) and the organization responsible for assigning substance identifiers.

The necessary messaging specifications are included as an integral part of the IDMP standards to secure the interactions listed above.

Unique identifiers produced in conformance with the IDMP standards will support applications for which it is necessary to reliably identify and trace the use of medicinal products and the materials within medicinal products.

This International Standard provides a structure that enables the assignment and maintenance of unique identifiers for all substances in medicinal products or in packaging materials in which medicinal products are contained. This International Standard sets out the general rules for defining and distinguishing substances, and provides a high-level model that structures substances and specified substances for the organization and capturing of data.

This International Standard has been developed using HL7's Common Product Model, and detailed modelling of substances and specified substances has been undertaken in that domain. It is anticipated that implementation will use the HL7 substances implementation guide and messaging to deliver a strong, non-semantic unique identifier for every substance present in a medicinal product. It is anticipated that a single organization will be

ISO 11238:2012(E)

responsible for the generation of identifiers for every substance and that such an organization would retain the defining elements upon which the substance identifier was based. At the specified substance level, a more regional approach may be necessary because of the proprietary nature of much of the information.

The use of the identifier is essential for the description of substances in medicinal products on a global scale. This International Standard does not involve developing nomenclature for substances or specified substances, but common and official substance names in current use can be mapped to each identifier.

Materials used in medicinal products range from simple chemicals to gene-modified cells to animal tissues. To unambiguously define these substances is particularly challenging. This International Standard defines substances based on their scientific identity (i.e. what they are) rather than on their use or method of production. Molecular structure or other immutable properties, such as taxonomic, anatomical and/or fractionation information, are used to define substances. This International Standard contains five groups of elements that are sufficient to define all substances. Although it is certainly possible to define or classify substances in other ways, this International Standard uses a minimalist structured scientific concept approach focusing on the critical elements necessary to distinguish two substances from one another. There are frequently interactions between substances when they are mixed together, but this International Standard has intentionally not included these supramolecular interactions at the substance level because of the variable nature and strength of such interactions. This International Standard also allows for the capture of multiple terms which refer to a given substance and a variety of reference information that could be used to classify substances or relate one substance to another.

In addition to the substance level, this International Standard also provides elements for the capture of further information on substances, such as grade, manufacturer, manufacturing specifications, and also to capture information on substances that are frequently combined together in commerce but are not strictly a medicinal product. At the specified substance level, four groups of elements provide information essential to the tracking and description of substances in medicinal products.

The basic concepts in the regulatory and pharmaceutical standards development domain use a wide variety of terms in various contexts. The information models presented in this International Standard depict elements and the relationship between elements that are necessary to define substances. The terms and definitions described in this International Standard are to be applied for the concepts that are required to uniquely identify, characterize and exchange information on substances in regulated medicinal products.

The terms and definitions adopted in this International Standard are intended to facilitate the interpretation and application of legal and regulatory requirements, but they are without prejudice to any legally binding document. In case of doubt or potential conflict, the terms and definitions contained in legally binding documents prevail.

Health informatics — Identification of medicinal products — Data elements and structures for the unique identification and exchange of regulated information on substances

1 Scope

This International Standard provides an information model to define and identify substances within medicinal products or substances used for medicinal purposes, including dietary supplements, foods and cosmetics. Other standards and external terminological resources are referenced that are applicable to this International Standard.

2 Terms, definitions, symbols and abbreviated terms

2.1 Terms and definitions

For the purposes of this document, the following terms, definitions and abbreviations apply.

2.1.1

active marker

constituent, or groups of constituents, of an herbal substance, herbal preparation or herbal medicinal product which are of interest for control purposes and are generally accepted to contribute to therapeutic activity

NOTE Active markers are not equivalent to analytical or signature markers that serve solely for identification or control purposes.

2.1.2

analytical data

set of elements to describe and capture methods and reference material used to determine purity, potency or identity in a specified substance

2.1.3

chemical bond

condition that occurs when forces acting between two atoms or groups of atoms lead to the formation of a stable discrete molecular entity

2.1.4

chemical substance

type of substance that can be described as a stoichiometric or non-stoichiometric single molecular entity and is not a protein or nucleic acid substance

NOTE Chemical substances are generally considered “small” molecules which have associated salts, solvates or ions and may be described using a single definitive or representative structure.

2.1.5

chiral substance

substance whose molecular structure is not superimposable on its mirror image

2.1.6

component

intended constituent of a specified substance

EXAMPLE Dimethicone and silicon dioxide are components of simethicone. Human insulin protamine and zinc are the components in human insulin isophane.

NOTE Components are used to describe the substances and specified substances that form a multi-substance material.

ISO 11238:2012(E)

2.1.7

composition stoichiometry

quantitative relationships between the chemical elements or moieties that make up a substance

EXAMPLE Sodium phosphate dibasic heptahydrate and sodium phosphate dibasic dihydrate are defined as different substances.

2.1.8

constituent

substance present within a specified substance

NOTE Constituents can be impurities, degradants, active markers or signature substances, or single substances mixed together to form a product. Constituents shall have an associated role and amount. Constituent specifications shall be used to describe components as well as limits on impurities or related substances for a given material.

2.1.9

controlled vocabulary

finite set of values that represent the only allowed values for a data item

NOTE 1 The allowed values can be codes, text or numeric.

NOTE 2 Adapted from CDISC Clinical Research Glossary V8.0, 2009.

2.1.10

copolymer

polymer with more than one type of structural repeating unit linked through covalent bonds

NOTE Copolymers are obtained by copolymerization or sequential polymerization of two or more monomers. Copolymers can be random, alternating, block or graft.

2.1.11

critical process step

manufacturing step necessary for production of a specified substance

2.1.12

degree of polymerization

number of structural repeating units in a polymeric block or chain

NOTE Applies to both homopolymers and block copolymers where it refers to the degree of polymerization within a block.

2.1.13

diverse origin

substances that are not isolated together or the result of the same chemical synthetic process

NOTE Material containing multiple substances is defined either as a mixture substance or a multi-substance (group 1) specified substance based on origin. Two substances brought together that do not undergo a chemical reaction resulting in the formation or breakage of specific chemical bonds would be defined as separate substances, even if there are non-bonding interactions between the substances.

2.1.14

enhancer

cis-acting sequence of DNA that increases the utilization of some eukaryotic promoters and which can function in either orientation and in any location (upstream or downstream) relative to the promoter

2.1.15

fraction

distinct portion of material derived from a complex matrix, the composition of which differs from antecedent material

NOTE This concept is used to describe source material and is recursive in that a subsequent fraction can be derived from an antecedent fraction, which is implied in the order of listing.

EXAMPLE Serum immunoglobulins to polyclonal IgG is an example of recursive fractionation.

2.1.16**gene**

basic unit of hereditary material that encodes and controls the expression of a protein or protein subunit

2.1.17**gene element**

individual element within a gene such as a promoter, enhancer, silencer or coding sequence

2.1.18**glycosylation**

enzymatic process that links saccharides or oligosaccharides to proteins, lipids or other organic molecules

2.1.19**glycosylation type**

significant differences in glycosylation between different types of organisms

NOTE This distinguishes the pattern of glycosylation across organism types, e.g. human, mammalian and avian. The glycosylation type is a defining element when polydisperse organism-based glycosylation exists in a substance.

2.1.20**grade**

set of specifications indicating the quality of a specified substance

2.1.21**homopolymer**

polymer containing a single structural repeating unit

2.1.22**isotope**

variants of a chemical element that differ by molecular mass

NOTE Radionuclides or nuclides with a non-natural isotopic ratio are shown in the structural representation with the nuclide number displayed. Natural abundance isotopes are represented by an elemental symbol without a nuclide number.

EXAMPLE ^{13}C refers to a carbon atom that has an atomic mass of 13.

2.1.23**manufacturing**

process of production for a substance or medicinal product from the acquisition of all materials through all processing stages

NOTE The critical process, starting and processing materials, and critical production parameters are included.

2.1.24**material**

any entity that has mass, occupies space and consists of one or more substances

2.1.25**medicinal product**

any substance or combination of substances which may be administered to human beings or animals for treating or preventing disease, with the view to making a medical diagnosis or to restore, correct or modify physiological functions

NOTE 1 A medicinal product may contain one or more manufactured items and one or more pharmaceutical products.

NOTE 2 In certain jurisdictions, a medicinal product may also be defined as any substance or combination of substances which may be used to make a medical diagnosis.