
Vsadki (implantati) za srce in ožilje ter zunajtelesni pretočni sistemi - Žilni medicinski kombinirani proizvodi/zdravila - 1. del: Splošne zahteve (ISO/DIS 12417-1:2021)

Cardiovascular implants and extracorporeal systems - Vascular device-drug combination products - Part 1: General requirements (ISO/DIS 12417-1:2021)

Kardiovaskuläre Implantate und extrakorporale Systeme - Vaskuläre Medizinprodukt-Arzneimittel-Kombinationsprodukte - Teil 1: Allgemeine Anforderungen (ISO/DIS 12417-1:2021)

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Implants cardiovasculaires et circuits extra-corporels - Produits de combinaison médicament-dispositif vasculaire - Partie 1: Exigences générales (ISO/DIS 12417-1:2021)

Ta slovenski standard je istoveten z: prEN ISO 12417-1

ICS:

11.040.40	Implantanti za kirurgijo, protetiko in ortetiko	Implants for surgery, prosthetics and orthotics
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DRAFT INTERNATIONAL STANDARD

ISO/DIS 12417-1

ISO/TC 150/SC 2

Secretariat: ANSI

Voting begins on:
2021-10-25Voting terminates on:
2022-01-17

Cardiovascular implants and extracorporeal systems — Vascular device-drug combination products —

Part 1: General requirements

Implants cardiovasculaires et circuits extra-corporels — Produits de combinaison médicament-dispositif vasculaire —

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Reference number
ISO/DIS 12417-1:2021(E)

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Published in Switzerland

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Foreword

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The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

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For an explanation on the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the WTO principles in the Technical Barriers to Trade (TBT) see the following URL: Foreword - Supplementary information

The committee responsible for this document is ISO/TC 150, *Implants for surgery*, Subcommittee SC 2, *Cardiovascular implants and extracorporeal systems*.

ISO 12417 consists of the following parts under the general title, *Cardiovascular implants and extracorporeal systems — Vascular device-drug combination products*:

- Part 1: *General requirements*
- Part 2: *Local regulatory guidance*

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Introduction

This part of ISO 12417 was prepared in order to provide minimum requirements for vascular device-drug combination products (VDDCPs).

Only issues related to vascular devices combined with drug(s), wherein the drug serves an ancillary function of the VDDCP are covered by this part of ISO 12417.

Many vascular device-drug combination products have been shown to be safe and effective in clinical use. This revision is not intended to require additional evaluation of these products as the testing would not provide useful information regarding the expected clinical performance of the product. Manufacturers can rely on historical data gathered under the specifications of the previous edition. Similarly, for product modifications or changes in intended clinical use, this revision is not intended to require additional evaluation of any aspects of the product that are not expected to change clinical performance.

It was impossible, when writing this part of ISO 12417, to take into consideration all future and emerging technologies. VDDCPs using such technologies will need to be evaluated following the basic requirements of this International Standard. Testing beyond the scope of this part of ISO 12417 might also be necessary to characterize these future and emerging device systems. Consideration shall be given to the failure modes of the VDDCP and their effects on the performance in deciding what testing will be appropriate.

For issues related to the primary mode of action (PMOA) of the vascular VDDCP, the reader might find it useful to consider a number of other International Standards (see Bibliography).

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Cardiovascular implants and extracorporeal systems — Vascular device-drug combination products —

Part 1: General requirements

1 Scope

This part of ISO 12417 specifies requirements for vascular device-drug combination products (VDDCPs) based upon current technical and medical knowledge. VDDCPs are medical devices with various clinical indications for use in the human vascular blood system. A VDDCP incorporates, as an integral part, substance(s) which, if used separately, can be considered to be a medicinal substance or product (drug substance, drug product) but the action of the medicinal substance is ancillary to that of the device and supports the primary mode of action (PMOA) of the device. With regard to safety, this part of ISO 12417 outlines requirements for intended performance, design attributes, materials, design evaluation, manufacturing, sterilization, packaging, and information supplied by the manufacturer. For implanted products, this International Standard should be considered as a supplement to ISO 14630, which specifies general requirements for the performance of non-active surgical implants. This International Standard should also be considered as a supplement to relevant device-specific standards, such as the ISO 25539-series specifying requirements for endovascular devices. Requirements listed in this part of ISO 12417 also address VDDCPs that are not permanent implants.

NOTE Due to variations in the design of combination products covered by this part of ISO 12417 and due to the relatively recent development of some of these combination products, acceptable standardized *in vitro* test results and clinical study results are not always available. As further scientific and clinical data become available, appropriate revision of this part of ISO 12417 might be necessary.

Delivery systems or parts of the delivery system are included in the scope of this part of ISO 12417, if they comprise an integral component of the vascular device and if they are drug-covered (e.g. drug-covered balloon catheters and drug-covered guidewires).

Devices whose PMOA is to provide a conduit for delivery of a drug, are excluded from the scope of this part of ISO 12417 (e.g. infusion catheters), unless they contain a drug component that is intended to have an ancillary action to the device part (e.g. antimicrobial coated infusion catheter).

Procedures and devices used prior to and following the introduction of the VDDCP (e.g. balloon angioplasty devices) are excluded from the scope of this part of ISO 12417 if they do not affect the drug-related aspects of the device.

This part of ISO 12417 is not comprehensive with respect to the pharmacological evaluation of VDDCPs. Some information on the requirements of different national and regional authorities is given in [Annex B](#).

Absorbable components of VDDCPs (e.g. coatings) are addressed by this part of ISO 12417 in their connection with drug-related aspects of the device. Degradation and other time-dependent aspects of absorbable implants and coatings are not completely addressed by this part of ISO 12417.

NOTE See also ISO/TS 17137 and ASTM F3036-13.

This part of ISO 12417 does not address issues associated with viable or non-viable biological materials such as tissues, cells, or proteins.

This part of ISO 12417 does not address issues associated with active surgical implants (i.e. implants that require power not generated by the human body or gravity).

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2 Normative references

The following documents, in whole or in part, are normatively referenced in this document and are indispensable for its application. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 10993-1, *Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process*

ISO 10993-2, *Biological evaluation of medical devices — Animal welfare requirements*

ISO 10993-7:2008, *Biological evaluation of medical devices — Part 7: Ethylene oxide sterilization residuals*

ISO 11070, *Sterile single-use intravascular introducers, dilators and guidewires*

ISO 11607-1, *Packaging for terminally sterilized medical devices — Part 1: Requirements for materials, sterile barrier systems and packaging systems*

ISO 14155, *Clinical investigation of medical devices for human subjects — Good clinical practice*

ISO 14630:2012, *Non-active surgical implants — General requirements*

ISO 14937, *Sterilization of health care products — General requirements for characterization of a sterilizing agent and the development, validation and routine control of a sterilization process for medical devices*

ISO 14971:2019, *Medical devices — Application of risk management to medical devices*

ISO 15223-1:2021, *Medical devices — Symbols to be used with information to be supplied by the manufacturer — Part 1: General requirements*

ISO 25539-2, *Cardiovascular implants — Endovascular devices — Part 2: Vascular stents*

NOTE See the Bibliography for additional device-specific and regional information about standards and guidance documents.

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 14630 and the following apply.

NOTE Potential clinical events are defined in [Annex A](#).

3.1 active pharmaceutical ingredient API

Drug substance

pharmacologically active (drug or medicinal) substance used as a raw material, which is coated on, bound to or incorporated into the device to achieve an ancillary device function (e.g. minimizing vascular restenosis)

3.2 batch

quantity of VDDCP at the final stage or pre-final stage of manufacture which has undergone the same manufacturing cycle, using the same components (e.g. same coating solution, same device size), and meets the same specifications

3.3 change

alteration to an activity or to the VDDCP to improve or maintain the composition or performance of a VDDCP

Note 1 to entry: This include small alterations to a VDDCP, a manufacturing process, or a test procedure, even if it is not necessarily captured by a Corrective Action/Preventative Action (CAPA) system, and may require reporting to local regional authorities

3.4 clinical event

complication, failure, or device-related observation that might be observed with clinical use of a VDDCP

Note 1 to entry: Such events might not have clinical significance and might not be attributable to the VDDCP.

3.5 compendial pharmaceutical reference standard

general term covering reference substances, reference preparations, and reference spectra that are recognized by a national pharmacopoeia

3.6 device part of the VDDCP DP

part of the VDDCP intended to treat vascular disease by temporary or long-term intervention or implantation that does not achieve its PMOA in or on the human body by pharmacological, immunological, or metabolic means, but might be assisted in its function by such means

3.7 Assay

biological or chemical method to determine the activity or potency of a substance

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3.8 drug product medicinal product

API, in its final form for administration to the patient (e.g. tablet, solution, spray), that is intended to prevent, diagnose, or treat disease and that achieves its principal intended action in or on the body by pharmacological, immunological, or metabolic means

3.9 drug-containing part of the VDDCP DCP

part of the VDDCP that consists of the API or matrix and associated device interfaces intended to assist in the PMOA of the device and/or diminish or ameliorate an unintended effect that placement of the device part might stimulate

Note 1 to entry: Some VDDCPs might have an incorporated medicinal or drug substance primarily intended to optimize the DP properties of the VDDCP.

3.10 DCP interface

common boundary or interconnection between the various components of the device part(s) and the drug-containing part(s) of a VDDCP

EXAMPLE

- a) the interface between the matrix containing the API and packaging materials with direct DCP contact;
- b) the device surface(s);
- c) the interface between the matrix and the API.

ISO/DIS 12417-1:2021(E)**3.11
delivery system**

transport device that physically or mechanically positions the VDDCP and/or the DCP at the intended anatomic location

EXAMPLE The delivery system of a drug-coated balloon, would position the balloon in the lumen of the lesion intended to be treated.

**3.12
drug content**

total labelled amount of API in a VDDCP

Note 1 to entry: Drug content might be expressed as µg/DCP of a certain size.

**3.13
drug delivery**

local interaction between the VDDCP drug and the *in vivo* environment, whether the drug is released from, eluted from, or remains bound to the VDDCP

**3.14
drug-related impurity**

any substance in the DCP of a VDDCP that is not the API or an excipient

Note 1 to entry: Drug-related impurities might include drug degradation products or degradants, drug-synthesis-related impurities, isomers of the drug, residual drug solvents, or biological contaminants (e.g. occurring with drugs derived from biological systems).

**3.15
drug release characterization**

in vitro characterization of the API released from the DCP of a VDDCP over time

EXAMPLE The release might be determined by a drug elution test, and could include a curve shape (or profile), a drug release rate, or both.

**3.16
durability**

ability to maintain adequate integrity and robustness during procedural (i.e. access, deployment, withdrawal), post-procedural, and long-term use (i.e. over time) in accordance with the design specifications

**3.17
efficacy
effectiveness**

ability of the VDDCP to achieve the planned and desired physiological result

**3.18
evaluate**

appraise or analyse qualitatively

**3.19
excipient**

additional material(s), other than the API, that are intentional components of the drug-containing part of a VDDCP

EXAMPLE Filler, extender, diluent, wetting agent, solvent, colorant, stabilizer, antioxidant, preservative, pH maintainer, polymers, adhesives.

**3.20
functionality**

ability of the VDDCP to perform physically, chemically, and/or mechanically, as designed

Note 1 to entry: Functionality does not include the physiological response to the VDDCP (i.e. efficacy).

3.21 matrix

any organic or inorganic material, other than living cells, intentionally applied by a manufacturer to a vascular device and designed for the purpose of drug storage, local drug activity at the surface and/or enabling, retarding, delaying, or modifying drug release

Note 1 to entry: The matrix might: be permanent or temporary (dissolvable, absorbable or degradable); include surface treatments such as primers; be a coating with or without an API, or consist of multiple excipients and/or multiple APIs.

3.22 particulates particles

mobile matter, other than gas bubbles, present on, or arising from the use of the VDDCP

3.23 pharmacokinetics

absorption, distribution, metabolism, and elimination of a drug *in vivo*

3.24 procedural fluids

blood and serum, saline, and contrast media that come into contact with a VDDCP

3.25 stability studies

tests undertaken according to a prescribed stability protocol to establish, support, or confirm the shelf life of a VDDCP

Note 1 to entry: Additional guidance on the drug-related aspects of the drug-containing part of the VDDCP can be found in International Conference on Harmonization Guideline ICH Q1A.

3.26 uniformity of drug content

comparison of the uniformity of the drug content between individual VDDCPs within each batch as compared to the labelled claim

3.27 vascular device-drug combination product VDDCP

vascular medical device that incorporates one or more APIs as an integral part (ancillary mode of action) to that of the device, but not necessarily to the VDDCP PMOA

Note 1 to entry: The VDDCP can be permanently deployed (i.e. it can be an implant like a drug-eluting stent) or temporarily deployed (i.e. it can be a drug-eluting balloon).

3.28 VDDCP deployment

physical or mechanical positioning of the VDDCP so that the drug-containing part is in contact with the intended anatomic treatment site

Note 1 to entry: The VDDCP might be permanently deployed (e.g. a drug-eluting stent) or temporarily deployed (e.g. a drug-eluting balloon).

3.29 VDDCP specification

required list of test procedures and appropriate acceptance criteria which are numerical limits, ranges, or other criteria for the tests described

Note 1 to entry: A specification is a critical quality standard. It establishes the set of criteria to which a VDDCP has to conform.

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Note 2 to entry: Additional guidance on the drug-related aspects of the drug-containing part of the VDDCP can be found in International Conference on Harmonization Guideline ICH Q6A.

3.30

Primary mode of action (PMOA)

single mode of action of a combination product that provides the most important therapeutic action of the combination product. The most important therapeutic action is the mode of action expected to make the greatest contribution to the overall intended therapeutic effects of the combination product

Note 1 to entry: Additional guidance on the drug-related aspects of the drug-containing part of the VDDCP can be found in International Conference on Harmonization Guideline ICH Q1A.

4 Intended performance

4.1 General

The requirements of ISO 14630:2012, Clause 4, shall apply.

4.2 Classification

A VDDCP is a product that is considered to be a medical device but which incorporates, as an integral part, substances which, if used separately, can be considered to be a medicinal product or drug product. It is classified as a medical device, provided that the action of the medicinal or drug substance is ancillary to that of the device, as reflected in the product claim and as supported by the scientific data provided by the manufacturer of the device.

4.3 Intended clinical location

The intended clinical location shall be identified as one or more of the following:

- a) abdominal aorta;
- b) arterio-venous shunt for vascular access;
- c) carotid;
- d) coronary;
- e) femoral;
- f) iliac;
- g) popliteal;
- h) intracerebral;
- i) renal;
- j) thoracic aorta;
- k) thoraco-abdominal aorta;
- l) tibial;
- m) other arterial or venous vessels to be specified.

5 Design attributes

5.1 General

The design attributes to meet the intended performance of the VDDCP shall take into account at least the following:

- a) ability of the device part of the VDDCP (i.e. the device without the API and matrix) to fulfill all product-specific requirements for the PMOA (e.g. the mechanical function), which are defined in the device-related standards;
- b) ability of the drug-containing part of the VDDCP to fulfill the drug-specific function and requirements of the VDDCP as defined in 5.2.
- c) ability of the VDDCP to meet defined chemical, physical, mechanical or compatibility specifications after interaction with the DCP/matrix and device or manufacturing processes.
- d) ability of the VDDCP to meet applicable interactional (ergonomic, connections, coupling) requirements, unless justified.

5.2 Drug-containing part of the VDDCP (DCP)

5.2.1 General

The design attributes of the VDDCP to meet the intended performance of the DCP shall additionally take into account at least the following:

- a) ability of the DCP to be consistently, accurately, and safely brought into contact with the intended anatomic treatment site;
- b) appropriate physical and chemical compatibility of the DCP interfaces (i.e. the device, the drug, the matrix, and any packaging with direct DCP contact);
- c) appropriately justified/conducted biocompatibility of the DCP;
- d) conformance of the DCP to VDDCP specifications at the time of manufacture and after storage;
- e) ability of the DCP to deliver or maintain the intended amount of drug safely at the target site in accordance with the specification of the VDDCP at product release and for the duration of the labelled shelf life;
- f) appropriate interaction between the VDDCP and procedural fluids.

5.2.2 Matrix

The design attributes of the VDDCP to meet the intended performance of the matrix shall additionally take into account at least the following:

- a) ability of the matrix to maintain adequate integrity during procedural use in accordance with the design specifications (e.g. freedom from significant delamination, flaps, and bare spots); and over time as applicable for the VDDCP.
- b) ability of the matrix to maintain adequate resistance to unintended generation of particles;
- c) conformance of the matrix to VDDCP specifications at the time of manufacture and after storage;
- d) conformance of the matrix dimensions, physical and chemical properties, and other matrix parameters (e.g. porosity, mass, density, distribution, glass transition temperature, melting temperature, fragmentation point) to the design requirements;