
**Implants for surgery —
Hydroxyapatite —**

**Part 2:
Thermally sprayed coatings of
hydroxyapatite**

Implants chirurgicaux — Hydroxyapatite —

Partie 2: Revêtements à base d'hydroxyapatite, obtenus par projection thermique



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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.

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).

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. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see www.iso.org/patents).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

For an explanation on the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT) see the following URL: www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 150, *Implants for surgery*, Subcommittee SC 1, *Materials*.

This third edition cancels and replaces the second edition (ISO 13779-2:2008), which has been technically revised.

A list of all parts in the ISO 13779 series can be found on the ISO website.

Any feedback or questions on this document should be directed to the user's national standards body. A complete listing of these bodies can be found at www.iso.org/members.html.

Introduction

No known surgical implant material has ever been shown to be completely free of adverse reactions in the human body. However, long-term clinical experience of the use of the material referred to in ISO 13779 has shown that an acceptable level of biological response can be expected, if the material is used in appropriate applications.

The biological response to hydroxyapatite coatings has been demonstrated by a history of clinical use and by laboratory studies (see References [1], [2], [3], [4], [5], [6]).

Implants for surgery — Hydroxyapatite —

Part 2:

Thermally sprayed coatings of hydroxyapatite

1 Scope

This document specifies requirements for single layer thermally sprayed hydroxyapatite coatings applied to metallic surgical implants.

These requirements are intended to describe properties of the materials and to communicate these between organizations. These requirements are not written with the objective of replacing a company's internal operational and assessment requirements although they could be used as such.

NOTE 1 For thin coatings with a thickness of less than 50 μm , some of the test methods described in this document might be difficult to apply without modification.

NOTE 2 The requirements of the hydroxyapatite layer of dual-layer coatings (consisting of a lower layer of metallic coating and an upper layer of hydroxyapatite coating) can follow this document; however, testing methods referred to in this document cannot be applied to dual layer coatings. If this document is taken in reference for the requirements of the hydroxyapatite layer of dual layer coatings, a rationale on how the single-layer tested coupons are representative of the dual-layer coated implant might be considered necessary.

This document does not cover coatings made from glasses, glass ceramics, alpha- and beta-tricalcium phosphate, biphasic calcium phosphate or other forms of calcium phosphate.

NOTE 3 While the requirements in this document are intended to be used as specifications of a thermally sprayed coating of hydroxyapatite, it might be necessary to establish routine control procedures specifying control tests and their time intervals to make sure the characteristics of the coating stay within specified limits.

NOTE 4 This document was developed with a focus on plasma sprayed coating of hydroxyapatite. It might also be used to characterize other thermally sprayed coatings of hydroxyapatite. However, thermally sprayed coatings that do not have a history of clinical use might present different risks and might need additional characterizations beyond those identified in this document.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 4288, *Geometrical Product Specifications (GPS) — Surface texture: Profile method — Rules and procedures for the assessment of surface texture*

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

ISO 13779-3, *Implants for surgery — Hydroxyapatite — Part 3: Chemical analysis and characterization of crystallinity ratio and phase purity*

ISO 13779-4, *Implants for surgery — Hydroxyapatite — Part 4: Determination of coating adhesion strength*

ISO 13779-6, *Implants for surgery — Hydroxyapatite — Part 6: Powders*

ASTM F1044, *Standard Test Method for Shear Testing of Calcium Phosphate Coatings and Metallic Coatings*

ASTM F1854, *Standard Test Method for Stereological Evaluation of Porous Coatings on Medical Implants*

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 13779-3 and the following apply.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at <https://www.iso.org/obp>
- IEC Electropedia: available at <http://www.electropedia.org/>

3.1 thermally sprayed hydroxyapatite coating

coating formed by thermal spraying of hydroxyapatite powders

Note 1 to entry: For the purpose of this document, the terms “coating” and “hydroxyapatite coating” both mean “thermally sprayed hydroxyapatite coating”

4 Coating preparation

The powder used for thermal spraying of the hydroxyapatite coating shall be in accordance with ISO 13779-6.

Unless documented and justified by the manufacturer, all test specimens shall be prepared using the same production methods of regular implant components, including initial hydroxyapatite powder, substrate material, production installations, substrate surface preparation process, coating process parameters, cleaning and sterilization.

5 Requirements

5.1 General

The minimum requirements for the hydroxyapatite coating are established in 5.2 to 5.8.

Other characterization tests, such as those described in A.2, A.3 and A.4, might also be requested (e.g. in order to satisfy applicable national or regional regulation).

NOTE 1 In addition to the tests described in 5.2 to 5.8, some regulatory bodies can request the tests given in Annex A to characterize the hydroxyapatite coating.

5.2 Calcium to phosphorus ratio ($Ca:P$)

Coating shall be scraped from the substrate before testing.

The calcium to phosphorus ratio, $Ca:P$, of the hydroxyapatite ceramic coating shall be determined in accordance with ISO 13779-3.

The calcium to phosphorus ratio, $Ca:P$, shall have a value in the range of 1,61 to 1,76 for the atomic ratio.

NOTE Calcium to phosphorous ratio is usually not influenced by the thickness of the coating.

5.3 Trace elements

Coating shall be scraped from the substrate before testing. The amount of sample required for chemical analysis is dependent on the chemical analysis technique used. The amount of sample used shall be sufficient to achieve adequate quantification limits. The technique used to remove the coating shall