# INTERNATIONAL STANDARD

ISO 5832-1

Fifth edition 2016-07-15

## Implants for surgery — Metallic materials —

Part 1: Wrought stainless steel

Implants chirurgicaux — Produits à base de métaux —

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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 <a href="www.iso.org/directives">www.iso.org/directives</a>).

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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 World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT) see the following URL: <a href="https://www.iso.org/iso/foreword.html">www.iso.org/iso/foreword.html</a>.

The committee responsible for this document is ISO/TC 150, *Implants for surgery*, Subcommittee SC 1, *Materials*.

ISO 5832-1:2016

This fifth edition cancels and replaces the fourth edition (ISO 5832-1:2007), which has been technically revised. It also incorporates the Technical Corriger dum ISO 5832-1:2007/Cor 1:2008.

ISO 5832 consists of the following parts, under the general title *Implants for surgery — Metallic materials*:

- Part 1: Wrought stainless steel
- Part 2: Unalloyed titanium
- Part 3: Wrought titanium 6-aluminium 4-vanadium alloy
- Part 4: Cobalt-chromium-molybdenum casting alloy
- Part 5: Wrought cobalt-chromium-tungsten-nickel alloy
- Part 6: Wrought cobalt-nickel-chromium-molybdenum alloy
- Part 7: Forgeable and cold-formed cobalt-chromium-nickel-molybdenum-iron alloy
- Part 8: Wrought cobalt-nickel-chromium-molybdenum-tungsten-iron alloy
- Part 9: Wrought high nitrogen stainless steel
- Part 11: Wrought titanium 6-aluminium 7-niobium alloy
- Part 12: Wrought cobalt-chromium-molybdenum alloy
- Part 14: Wrought titanium 15-molybdenum 5-zirconium 3-aluminium alloy

## 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 this part of ISO 5832 has shown that an acceptable level of biological response can be expected when the material is used in appropriate applications.

The following definitions apply in understanding how to implement an ISO International Standard and other normative ISO deliverables (TS, PAS, IWA):

- "shall" indicates a requirement;
- "should" indicates a recommendation;
- "may" is used to indicate that something is permitted;
- "can" is used to indicate that something is possible, for example, that an organization or individual is able to do something.
- 3.3.1 of the ISO/IEC Directives, Part 2 (sixth edition, 2011) defines a requirement as an "expression in the content of a document conveying criteria to be fulfilled if compliance with the document is to be claimed and from which no deviation is permitted."
- 3.3.2 of the ISO/IEC Directives, Part 2 (sixth edition, 2011) defines a recommendation as an "expression in the content of a document conveying that among several possibilities one is recommended as particularly suitable, without mentioning of excluding others, or that a certain course of action is preferred but not necessarily required, or that (in the negative form) a certain possibility or course of action is deprecated but not prohibited." (Caros. 11e. 1.2)

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## Implants for surgery — Metallic materials —

## Part 1:

## Wrought stainless steel

### 1 Scope

This part of ISO 5832 specifies the characteristics of, and corresponding test methods for, wrought stainless steel for use in the manufacture of surgical implants.

NOTE 1 The mechanical properties of a sample obtained from a finished product made of this alloy can differ from those specified in this part of ISO 5832.

NOTE 2 The alloy described in this part of ISO 5832 corresponds to UNS S31673 referred to in ASTM F138/ASTM F139 and to alloy code 1.4441 given in the withdrawn DIN 17443.

#### 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.

 $ISO\ 377, Steel\ and\ steel\ products-Location\ and\ preparation\ of\ samples\ and\ test\ pieces\ for\ mechanical\ testing$ 

ISO 404, Steel and steel products — General technical delivery requirements

ISO 439, Steel and iron — Determination of total silicon content — Gravimetric method

ISO 629, Steel and cast iron — Determination of manganese content — Spectrophotometric method

ISO 643, Steels — Micrographic determination of the apparent grain size

ISO 671, Steel and cast iron — Determination of sulphur content — Combustion titrimetric method

 $ISO\ 4967:2013, Steel-Determination\ of\ content\ of\ non-metallic\ inclusions-Micrographic\ method\ using\ standard\ diagrams$ 

ISO 6892-1:2016, Metallic materials — Tensile testing — Part 1: Method of test at room temperature

ISO 10714, Steel and iron — Determination of phosphorus content — Phosphovanadomolybdate spectrophotometric method

### 3 Terms and definitions

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

#### 3.1

#### original gauge length

length between gauge length marks on the test piece measured at room temperature before the test

[SOURCE: ISO 6892-1:2016, 3.1.1]

## 4 Chemical composition

#### 4.1 Test samples

The selection of samples for analysis shall be carried out in accordance with ISO 377.

## 4.2 Cast analysis

The cast analysis of the steel when determined in accordance with <u>Clause 6</u> shall comply with the chemical composition specified in <u>Table 1</u>. The molybdenum and chromium contents shall be such that the *C* value obtained from <u>Formula (1)</u> is not less than 26.

$$C = 3.3 w_{\text{Mo}} + w_{\text{Cr}} \tag{1}$$

where

 $w_{\text{Mo}}$  is the molybdenum content, expressed as a percentage by mass;

 $w_{Cr}$  is the chromium content, expressed as a percentage by mass.

**Element Mass fraction** P%FVIFY Teh ST 0,030 max. Carbon Silicon Standards. Itel max Manganese 2,0 max.  $0,025 \, \text{max}.$ Phosphorus Sulfur 0,010 max. Nitrogen 0,10 max. Chromium 17,0 to 19,0 max. Molybdenum 2,25 to 3,00 Nickel 13,0 to 15,0 Copper 0,50 max. Iron Balance

Table 1 — Chemical composition

## 5 Microstructure in the fully annealed condition

#### 5.1 Grain size

The austenitic grain size, determined in accordance with <u>Clause 6</u>, shall not be coarser than grain size No. 5.

## 5.2 Microstructure

The steel shall have a structure free from delta ferrite, chi or sigma phase, when examined in accordance with <u>Clause 6</u>.

#### 5.3 Inclusion content

The non-metallic inclusion content of the steel, determined at finished size after a hot-rolling process stage and in accordance with <u>Clause 6</u>, shall not exceed the limits given in <u>Table 2</u>.

NOTE It can be necessary to use vacuum or electroslag melting to produce a steel complying with these cleanliness requirements.

Inclusion content reference number Type of inclusion Thin Thick A - Sulfides 1,5 1 B - Aluminates 1,5 1 1 C – Silicates 1,5 D - Oxides, globular 1,5 1

Table 2 — Inclusion content limits

## 6 Mechanical properties

### 6.1 Test pieces

The selection and preparation of samples and test pieces for tensile testing shall be in accordance with ISO 377. **Teh STANDARD PREVIEW** 

## 6.2 Tensile test (standards.iteh.ai)

The tensile properties of the steel in the form of bars, wires, and sheet and strip, when tested in accordance with <u>Clause 6</u>, shall comply with the values specified in <u>Tables 3</u>, 4 and 5, respectively.

Should any of the test pieces not meet the specified requirements or break outside the gauge limits, retests shall be carried out in accordance with ISO 404.

#### 6.3 Gauge length

Original gauge length  $l_0$  shall be either  $5,65 \times \sqrt{S_0}$  or 50 mm, where  $S_0$  is defined as the original cross-sectional area in square millimetres. The gauge length chosen for testing shall be reported with the test results.

#### 7 Test methods

The test methods to be used in determining compliance with the requirements of this part of ISO 5832 shall be those given in Table 6.