

Designation: F 745 – 00

Standard Specification for 18Chromium-12.5Nickel-2.5Molybdenum Stainless Steel for Cast and Solution-Annealed Surgical Implant Applications¹

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1. Scope

1.1 This specification covers the requirements for 18chromium-12.5nickel-2.5molybdenum stainless steel alloy shot, bar, or ingot used for the manufacture of cast and solution-annealed surgical implants.

1.2 The values stated in inch-pound units are to be regarded as the standard.

1.3 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

2.1 ASTM Standards:

- A 744/A 774M Specification for Castings, Iron-Chromium-Nickel, Corrosion Resistant, for Severe Service²
- E 8 Test Methods for Tension Testing of Metallic Materials³
- E 353 Test Methods for Chemical Analysis of Stainless, Heat-Resisting, Maraging, and Other Similar Chromium-Nickel-Iron Alloys⁴

2.2 American Society for Quality (ASQ) Standard:⁵

ASQC 1 Specification of General Requirements for a Quality Program

3. Ordering Information

3.1 Inquiries and orders for material under this specification shall include the following information:

3.1.1 Quantity (weight or number of pieces),

- 3.1.2 ASTM Designation,
- 3.1.3 Form (Section 4.1),
- 3.1.4 Special tests, and

² Annual Book of ASTM Standards, Vol 01.02.

³ Annual Book of ASTM Standards, Vol 03.01.

⁵ Available from American Society for Quality Control, 161 West Wisconsin Ave., Milwaukee, WI 53203.

3.1.5 Special requirements.

4. Materials and Manufacture

4.1 The base material furnished to the implant manufacturer for purposes of casting surgical implants shall be supplied in the form of bar, shot, or ingot.

5. Chemical Composition

5.1 The heat analysis shall conform to the chemical composition listed in Table 1. The manufacturer shall not ship material that is outside the limits specified in Table 1.

5.1.1 Requirements for the major and minor elemental constituents are listed in Table 1. Also listed are important residual elements. Analysis for elements not listed in Table 1 is not required to verify compliance with this specification.

5.2 *Product Analysis*—The product analysis is either for the purpose of verifying the composition of a heat or lot or to determine variations in the composition within the heat.

5.2.1 Acceptance or rejection of a heat or lot of material may be made by the purchaser on the basis of this product analysis.

5.2.2 Product analysis tolerances do not broaden the specified heat analysis requirements but cover variation between laboratories in the measurement of chemical content. Product analysis limits shall be as specified in Table 2.

5.3 For referee purposes, Test Methods E 353 shall be used.

6. Mechanical Properties

6.1 The material shall conform to the mechanical property requirements prescribed in Table 3.

6.2 Specimens for tension tests shall be cast from remelted material from each master heat by the same general procedures used in casting surgical implants.

6.2.1 Specimens may be cast, ground, or machined to final dimensions in accordance with the 0.25 in. (6.35 mm) diameter specimen in Fig. 8 of Test Methods E 8.

6.2.2 Specimens shall be solution annealed using the same procedures used to solution anneal surgical implants.

6.3 A minimum of two tension specimens shall be tested. If one specimen fails below the specified mechanical requirements, two additional specimens shall be tested and both must pass.

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⁴ Annual Book of ASTM Standards, Vol 03.05.