

Designation: F 621 – 9702

Standard Specification for Stainless Steel Forgings for Surgical Implants¹

This standard is issued under the fixed designation F 621; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon (ϵ) indicates an editorial change since the last revision or reapproval.

1. Scope*

- 1.1 This specification covers the requirements of forged stainless steel for surgical implants when the material forged conforms to Specification F 138 (RUNS S31673), Specification F 1314 (UNS S21910), or Specification F 1586 (UNS S31675).
 - 1.2 The values stated in inch-pound units are to be regarded as the standard.

2. Referenced Documents

2.1 ASTM Standards:

¹ This specification is under the jurisdiction of ASTM Committee F-4 F04 on Medical and Surgical Materials and Devices and is the direct responsibility of Subcommittee F04.12 on Metallurgical Materials.

Current edition approved April Apr. 10, 1997; 2002. Published March 1998. June 2002. Originally published as F 621 – 79. Last previous edition F 621 – 927.



A 262 Practices for Detecting Susceptibility to Intergranular Attack in Austenitic Stainless Steels²

A 370 Test Methods and Definitions 473 Specification for Mechanical Testing of Steel Products²

A 473 Specification for Stainless and Heat-Resisting Steel Forgings²

E 8 Test Methods for Tension Testing of Metallic Materials³

E 18 Test Methods for Rockwell Hardness and Rockwell Superficial Hardness of Metallic Materials³

E 112 Test Methods for Determining Average Grain Size³

E 165 Practice for Liquid Penetrant Inspection Method⁴ Examination⁵

E 353 Test Methods for Chemical Analysis of Stainless, Heat-Resisting, Maraging, and Other Similar Chromium-Nickel-Iron Alloys⁴

² Annual Book of ASTM Standards, Vol 01.03.

³ Annual Book of ASTM Standards, Vol. 01.05. 03.01.

⁴ Annual Book of ASTM Standards, Vol 03.035.

⁵ Annual Book of ASTM Standards, Vol 03.0±3.



- F 138 Specification for Wrought-18Chromium-14Nickel-2.5Molybdenum Stainless Steel Bar and Wire for Surgical Implants (UNS S31673)⁶
- F-138 Specification 601 Practice for Stainless Steel Bars and Wire for Fluorescent Penetrant Inspection of Metallic Surgical Implants (Special Quality)⁷⁶
- F-601 Practice 1314 Specification for Fluorescent Penetrant Inspection of Metallic Wrought Nitrogen Strengthened-22Chromium-13Nickel-5Manganese-2.5Molybdenum Stainless Steel Alloy Bar and Wire for Surgical Implants (UNS S21910)⁶
- F 1314586 Specification for Wrought Nitrogen Strengthened-_221Chromium-12.50Nickel-53Manganese-2.5Molybdenum Stainless Steel Alloy Bar-and Wire for Surgical Implants
- F 1586 Specification for Wrought Nitrogen Strengthened–21 Chromium-10 Nickel-3 Manganese-2.5 Molybdenum Stainless Steel Bar for Surgical Implants⁷ (UNS S31675)⁶
- 2.2 ISO Standards:⁷

⁶ Annual Book of ASTM Standards, Vol 03.05. 13.01. Annual Book of ASTM

⁷ Available from American National Standards, Vol 13.01. Institute, 25 W. 43rd St., 4th Floor, New York, NY 10036.



- ISO 5832-1 Implants for Surgery—Metallic Materials Part 1: Wrought Stainless Steel
- ISO 5832-9 Implants for Surgery—Metallic Materials Part 9: Wrought High Nitrogen Stainless Steel
- 2.3 American Society for Q\(\infty\) uality Standard:
- Cl Specifications of General Requirements for a Quality Control Program⁸

3. Ordering Information

- 3.1 Inquiries and orders for forgings under this specification shall include the following information:
- 3.1.1 Quality; number of pieces,
- 3.1.1 Quantity,
- 3.1.2 ASTM designation; and date of issue,
- 3.1.3 ASTM material-grade,
- 3.1.3 Condition,
- 3.1.4 Mechanical properties (other than those specified herein),
- 3.1.5 Finish,
- 3.1.6 Applicable (alloy) standard and date of issue,
- 3.1.4 Condition,
- 3.1.5 Mechanical properties,
- 3.1.6 Finish,
- 3.1.7 Applicable dimensions or print drawing number,
- 3.1.78 Special tests, if any, and
- 3.1.89 SOther special requirements.

4. General Requirements for Delivery

- 4.1 Material furnished to this specification shall conform to the applicable requirements in the current edition of Specification A 473.
 - 4.2 In the case where a conflict exists between this specification and that listed in 4.1, this specification shall take precedence.

5. Materials and Manufacture

- 5.1 Material for forgings shall be bars or wire fabricated in accordance with Specifications F 138, F 1314, or F 1586, generally in the unannealed condition with a finish suitable for forging.
- 5.2 The material shall be forged by hammering, pressing, rolling, extruding, or upsetting, and shall be processed, if practicable, so as to cause metal flow during the hot-working operation to be in the direction most favorable direction for resisting stresses encountered in service, as may be indicated to the fabricator supplier by the implant manufacturer. purchaser.
- 5.3 Forgings shall be free of splits, scale, cracks, inequalities, flaws, and other imperfections not consistent with good commercial practice (see Note 1). Offset or mismatch allowance, dependent upon part size and configuration, shall be within standard forging tolerances.
 - Note 1—Compliance to these requirements may be verified by Practices E 165 or F 601 or other suitable methods.
- 5.4 After all hot working-operations, the forgings shall receive an annealing treatment, when necessary, by heating the parts to an appropriate elevated temperature for a specified dwell time followed by rapid cooling to meet the applicable metallurgical requirements specified herein. by the purchaser.
- 5.5 Optional identification marks, including the <u>manufacturer's purchaser's</u> logo, material designation, heat code number, and impression number, may be placed upon each forging, the method and location of which shall be as specified by the purchaser.

6. Chemical Composition

- 6.1 When specified by the implant manufacturer, the chemical compositions of either the raw forging bars or the completed forgings
- 6.1 The stainless steel forgings shall be determined and confirmed by conform to the forger, and shall meet chemical requirements prescribed in the product analysis limits of Specification F 138, F 1314, applicable alloy specification: F 138, F 1314, or F 1586.
 - 6.2 For referee purposes, Test Methods E 353 shall be used.

7. Mechanical Properties Requirements

7.1 The mechanical properties of stainless steel forgings produced from the three alloys of concern in this specification depend on a number of factors, including the specific alloy from which the forgings are produced and the fabrication practice. Different combinations of specific alloy and fabrication practice will provide substantially different mechanical properties.

⁸ Available from American Society for Quality Control, 161 W. Wisconsin Quality, 600 N. Plankinton Ave., Milwaukee, WI 53203.



- 7.2 When specified by the implant manufacturer, the mechanical properties of forgings shall be tested by the forger and shall comply with the minimum mechanical properties as specified by the implant manufacturer.
 - 7.3 If tension testing is required, test in Specifications F 138, F 1314, or F 1586.
- <u>7.1.1 Test</u> specimens-should shall be taken from a representative forging; if possible, or from a specially representative forged test bar, only if the configuration does not lend itself to yielding the required specimen. Any specially forged test bar must be in the same condition as the forgings it represents.
- 7.42 When desired, Rockwell hardness may be specified-on by the implant manufacturer's purchase order or print purchaser and shall be determined in accordance with Test Methods E 18.
 - 7.53 The mechanical properties shall be determined in accordance with Test Methods-A 370.-E 8.

8. Special Requirements Tests

- 8.1 *Corrosion Tests*—Forgings furnished to this specification shall be capable of passing the test for intergranular corrosion susceptibility in accordance with the current edition Practice E of Practices A 262, Practice E. A 262.
- 8.2 *Grain Size*—On the cross section examined, the grain size shall be predominately ASTM No. 4 or finer. No regions exhibiting grain size larger than ASTM No. 3 shall be allowed. Test procedures shall be in accordance with Test Methods E 112.
- 8.3 When specified by the purchaser, fluorescent penetrant inspection shall be performed on forgings. Penetrant inspections shall be performed in accordance with Practice E 165 or Practice F 601.
 - 8.4 Other special requirements shall be as specified on by the purchase order or print. purchaser.

9. Certification

9.1 The <u>fabricator's supplier's</u> certification that the material was manufactured and tested in accordance with this specification, together with a report of the test results, shall be furnished to the <u>implant manufacturer</u> purchaser with each shipment.

10. Quality Program Requirements

- 10.1 The <u>producer of supplier and any other processor that supplies</u> material in accordance with this specification shall maintain a quality program as defined in <u>ASQC C1 or equivalent</u>.
- 10.2 The manufacturer of surgical implants or medical appliances shall be assured of or may audit the material producer's quality program for conformance to the intent of ASOC ASO C1 or equivalent.

11. Keywords

11.1 forgings—surgical implants; metals (for surgical implants)—stainless steel; stainless steel—surgical applications

APPENDIXES

(Nonmandatory Information)

X1. RATIONALE

- X1.1 The purpose of this specification is to characterize the chemical, mechanical, and metallurgical properties of wrought stainless steel forgings for surgical implants.
- X1.2 The microstructural requirements contained in this specification represent the current general consensus with respect to optimization of mechanical properties for implant applications.
- X1.3 This specification has been expanded to cover forgings of three specific alloys; each UNS designation has been included for clarification. A Biocompatibility section has been added as an appendix.
- X1.4 ISO standards are listed for reference only. Although the ISO standards listed in 2.2 are similar to the corresponding ASTM standards, they may not be identical. Use of an ISO standard in addition to or instead of a preferred ASTM standard may be negotiated between purchaser and supplier. In this specification, ISO 5832-1 composition D is similar to ASTM Specification F 138 and the composition of ISO 5832-9 is similar to ASTM Specification F 1586.



X2. BIOCOMPATIBILITY

- X2.1 The material compositions covered by this-standard specification have been employed successfully in human implant applications in contact with soft tissue and bone for over a decade.
 - X2.2 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 has shown an acceptable level of biological response can be expected, if these materials are used in appropriate applications.

SUMMARY OF CHANGES

Committee F04 has identified the location of selected changes to this standard since the last issue (F 621 - 97) that may impact the use of this standard.

- (1) No significant changes were made to this specification.
- (2) UNS numbers were updated, titles were corrected, and wording was revised to be in accordance with template language and other specifications.
- (3) ISO Standards were added for reference.

ASTM International takes no position respecting the validity of any patent rights asserted in connection with any item mentioned in this standard. Users of this standard are expressly advised that determination of the validity of any such patent rights, and the risk of infringement of such rights, are entirely their own responsibility.

This standard is subject to revision at any time by the responsible technical committee and must be reviewed every five years and if not revised, either reapproved or withdrawn. Your comments are invited either for revision of this standard or for additional standards and should be addressed to ASTM International Headquarters. Your comments will receive careful consideration at a meeting of the responsible technical committee, which you may attend. If you feel that your comments have not received a fair hearing you should make your views known to the ASTM Committee on Standards, at the address shown below.

This standard is copyrighted by ASTM International, 100 Barr Harbor Drive, PO Box C700, West Conshohocken, PA 19428-2959, United States. Individual reprints (single or multiple copies) of this standard may be obtained by contacting ASTM at the above address or at 610-832-9585 (phone), 610-832-9555 (fax), or service@astm.org (e-mail); or through the ASTM website (www.astm.org).