



Designation: F601 – 23

Standard Practice for Fluorescent Penetrant Inspection of Metallic Surgical Implants¹

This standard is issued under the fixed designation F601; 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 practice is intended as a standard for fluorescent penetrant inspection of metallic surgical implants.

1.2 *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, health, and environmental practices and determine the applicability of regulatory limitations prior to use.*

1.3 *This international standard was developed in accordance with internationally recognized principles on standardization established in the Decision on Principles for the Development of International Standards, Guides and Recommendations issued by the World Trade Organization Technical Barriers to Trade (TBT) Committee.*

2. Referenced Documents

2.1 *ASTM Standards:*²

E1417/E1417M Practice for Liquid Penetrant Testing

3. Significance and Use

3.1 This practice is intended to confirm the method of obtaining and evaluating the fluorescent penetrant indications on metallic surgical implants.

4. Fluorescent Penetrant Method

4.1 Perform fluorescent penetrant inspection of metallic surgical implants in accordance with Practice E1417/E1417M.

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

Current edition approved Sept. 1, 2023. Published September 2023. Originally approved in 1978. Last previous edition approved in 2018 as F601 – 18. DOI: 10.1520/F0601-23.

² For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

4.2 The penetrant system used shall be Type I, Method A, minimum of Level 3 sensitivity, in accordance with Practice E1417/E1417M.

4.3 All penetrant materials shall be compatible with each other in accordance with Practice E1417/E1417M.

5. Preparation for Testing

5.1 Pre- and post-cleaning requirements are to be agreed upon between the purchaser and supplier.

5.2 If sand/grit blasting is used for pre-cleaning, care shall be taken to ensure that defects are not masked or peened over.

6. Penetrant Method Materials and Equipment Controls

6.1 The penetrant method materials deteriorate in usefulness through contamination and age. System performance and process control checks shall be performed in accordance with Practice E1417/E1417M.

7. Evaluation

7.1 The product acceptance and rejection criteria shall be as agreed upon between the purchaser and supplier.

7.2 Indication verification per Practice E1417/E1417M is permitted. If the purchaser has unique requirements, these shall take precedence.

8. Personnel Certification

8.1 If specified in the contractual agreement, personnel performing examination to this practice shall be qualified in accordance with a nationally or internationally recognized nondestructive testing (NDT) personnel qualification practice or standard and certified by the employer or certifying agency, as applicable.

9. Keywords

9.1 fluorescent; penetrant inspection; testing methods; surgical implants

*A Summary of Changes section appears at the end of this standard