



Designation: F2528 – 06 (Reapproved 2023)

Standard Test Methods for Enteral Feeding Devices with a Retention Balloon¹

This standard is issued under the fixed designation F2528; 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 These test methods cover the establishment of performance requirements for the utilization of a single-use, enteral feeding device with a retention balloon, used by medical professionals for providing a means of nutrition and/or administration of medication to patients by means of natural orifice (nasal, oral, transluminal) and or a surgically created stoma. The product is manufactured in various sizes and materials such as silicone, urethane, and various polymers (as well as combinations of these) and is provided nonsterile for sterilization and sterile for single use only. Rationale for these test methods can be found in [Appendix X1](#).

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

[F623 Performance Specification for Foley Catheter](#)

2.2 *Other Standard:*

[Simulated Gastric Fluid, USP Official Compendia of Standards](#)³

3. Terminology

3.1 Definitions:

¹ These test methods are under the jurisdiction of ASTM Committee F04 on Medical and Surgical Materials and Devices and are the direct responsibility of Subcommittee F04.35 on GI Applications.

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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.

³ USP Official Compendia of Standards, available from U.S. Pharmacopeia (USP), 12601 Twinbrook Pkwy., Rockville, MD 20852.

3.1.1 *balloon integrity (resistance to rupture)*, n —volume of liquid that corresponds with balloon failure, or bursting.

3.1.2 *distal*, n —refers to the balloon end of the enteral feeding device

3.1.3 *enteral feeding device with retention balloon*, n —a two-way medical device intended to provide a means of nutrition or administration of medication, or both, to patients by means of natural orifice (nasal, oral, transluminal) or a surgically created stoma, or both, consisting of a drainage lumen and inflation lumen (see [Fig. 1](#)). Common balloon inflation sizes are 5 cm³, 15 cm³, and 20 cm³.

3.1.4 *French size (Fr)*, n —a scale used for denoting the size of catheters and other tubular instruments. The French size value is three times the outer diameter of the tube as measured in millimetres. For example, a diameter of 18 Fr indicates a diameter of 6 mm.

3.1.5 *inflation volume*, n —volume of liquid used to inflate the retention balloon of the enteral feeding device for proposed testing in this standard.

3.1.6 *rated volume*, n —stated volume of inflation of the retention balloon of the enteral feeding device in the manufacturer's labeling and instructions for use.

3.1.7 *simulated gastric fluid*, n —a solution consisting of hydrochloric acid, salt, and pepsin with a pH of approximately 1.2, per USP standard recipe.

3.1.8 *sterility*, n —the state of being free from viable microorganisms.

4. Specimen Preparation

4.1 All test specimens for test methods listed below shall consist of the manufacturer's new, finished, untested, unsterilized product. At the minimum, statistically valid samples of the smallest and the largest diameter of enteral feeding devices shall be tested.

5. Test Methods

PROCEDURE A: FLOW RATE THROUGH FEEDING LUMEN

5.1 *Scope*—This test method covers the determination of flow rates through the drainage lumen of the enteral feeding device with retention balloon.

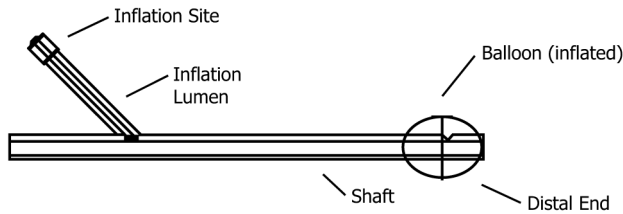


FIG. 1 Enteral Feeding Device with Retention Balloon

5.2 *Summary of Test Method*—The apparatus is set up as shown in Fig. 2. The flow rate is adjusted through the water inlet to a rate sufficient to maintain flow through the overflow outlet while each enteral feeding device is tested. A head pressure of 20 ± 1.0 cm of water (196 ± 10 kPa) above the tank bottom shall be maintained throughout the test to approximate actual physiological conditions. The overflow outlet should not be covered by water.

5.3 *Significance and Use*—The flow rate is measured in reverse flow for ease in testing, since differences in the flow rate as a result of flow direction are theoretically insignificant.

5.4 Apparatus:

5.4.1 *Water Reservoir*, capable of maintaining 20 ± 1.0 cm (7.9 ± 0.4 in.) of water (196 ± 10 kPa) above the tip of the enteral feeding device connection throughout the test as shown in Fig. 2. (See Performance Specification F623.)

5.4.2 *Graduated Cylinder*, calibrated for suitable measurement of the effluent.

5.4.3 *Syringe*, with appropriate tip for inflation of enteral feeding device balloon.

5.5 Hazards:

5.5.1 Overflow should not be covered. Head pressure must be kept constant; water should always be exiting through the overflow outlet.

5.5.2 Establish equilibrium before testing.

5.5.3 Flow rates through all fittings must exceed that of the enteral feeding device being tested.

5.6 Procedure:

5.6.1 Test at 23 ± 4 °C (73.4 ± 7 °F).

5.6.2 Inflate the retention balloon of the test specimen with water to labeled volume.

5.6.3 Connect the enteral feeding device to enteral feeding device connector and open the stopcock. The tip of the enteral feeding device connection at the junction of enteral feeding device on-off valve should be level with the bottom of the tank ± 1 cm and it should deliver fluid at 20 ± 1 cm (196 ± 10 kPa) head pressure at that junction.

5.6.4 Establish flow equilibrium before taking test measurements.

5.6.5 Record the amount of fluid through the device feeding lumen in 30 s.

5.7 *Interpretation of Results*—Flow rates for enteral feeding devices tested must meet or exceed $9 \text{ cm}^3/\text{min}$.

5.8 *Precision and Bias*—To be determined within five years.

PROCEDURE B: BALLOON BURST VOLUME

5.9 *Scope*—This test method covers the determination of balloon integrity of enteral feeding devices with retention balloon.

5.10 *Summary of Test Method*—The enteral feeding device with retention balloon is submerged in a small container filled with water. The balloon is then inflated with water until rupture, which enables the volume at which the balloon bursts to be observed.

5.11 *Significance and Use*—The balloon burst volume is measured to quantify the resistance to rupture of the enteral feeding device with retention balloon member.

5.12 *Apparatus*—The testing apparatus is set up as shown in Fig. 3.

5.12.1 *System Reservoir*.

5.12.2 *Syringe*.

5.12.3 *Water*.

5.13 *Hazards*—Water should be emptied from system reservoir through purge valve when fill marked is reached.

5.14 Procedure:

5.14.1 Test at 23 ± 4 °C (73.4 ± 7 °F).

5.14.2 Insert uninflated enteral feeding device into test orifice in system reservoir per Fig. 3.

5.14.3 Close orifice so that it is positioned proximal to the enteral feeding device with retention balloon member. The device is not to be immersed in water within the reservoir per Fig. 3.

5.14.4 Fill syringe with amount of water greater than that listed in Table 1 for the desired French size. Attach tip of syringe to enteral feeding device inflation valve.

5.14.5 Inflate retention balloon at $1 \text{ cm}^3/\text{s}$ with water until balloon bursts. Record amount of water injected into balloon at time of burst.

5.15 *Interpretation of Results*—Burst volumes for enteral feeding devices tested must meet or exceed those listed in Table 1.

5.16 *Precision and Bias*—To be determined within five years.

PROCEDURE C: BALLOON VOLUME MAINTENANCE

5.17 *Scope*—This test method is applicable for enteral feeding devices with retention balloon to test the integrity of the inflation system to maintain balloon volume.

5.18 *Summary of Test Method*—The balloon retention device of the enteral feeding device is inflated with a test liquid. This test liquid contains a colorant which enables a leak of this fluid to be observed. If no leak is observed, the integrity of the inflation system is upheld, therefore maintaining the balloon volume.

5.19 *Significance and Use*—This test method establishes a standard test method for determining the functional integrity of the inflation system of the enteral feeding device with retention balloon by observing the consistency of volume of the balloon after it is filled with test liquid. Additionally, since it is the