



Designation: E788 – 97 (Reapproved 2019)

Standard Specification for Pipet, Blood Diluting¹

This standard is issued under the fixed designation E788; 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 requirements for glass reusable blood diluting pipets that are used for performing red and white cell corpuscle determinations.

1.2 The values stated in SI units are to be regarded as standard. No other units of measurement are included in this standard.

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

E694 Specification for Laboratory Glass Volumetric Apparatus

E920 Specification for Commercially Packaged Laboratory Apparatus

E921 Specification for Export Packaged Laboratory Apparatus

E1133 Practice for Performance Testing of Packaged Laboratory Apparatus for United States Government Procurements

E1157 Specification for Sampling and Testing of Reusable Laboratory Glassware

3. Classification

3.1 This specification covers two different pipet designs and permissible alternatives.

3.1.1 Red and white cell blood diluting pipets (see Fig. 1).

3.1.2 Permissible alternative designs (see Fig. 2).

¹ This specification is under the jurisdiction of ASTM Committee E41 on Laboratory Apparatus and is the direct responsibility of Subcommittee E41.01 on Laboratory Ware and Supplies.

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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.

4. Materials

4.1 The pipets shall be made of common spirit bore white back tubing or clear glass with a white stripe applied to the outer surface of the tubing.

4.2 The beads shall be made of glass or ceramic composition and shall be of red, clear, or white coloring.

5. Dimensions and Permissible Variations

5.1 *Design*—The red and white cell blood dilution pipets shall consist of a small uniform bore glass tube which shall have a bulb of proper size near the proximal end (see Fig. 1). As an alternative, the pipets may be constructed of three pieces of glass that are fused together to form a one-piece pipet (see Fig. 2). The bulb shall contain a nonspherical glass or ceramic bead sufficiently large enough to prevent its being impacted in the constriction portion of the bulb. The cross section of the pipet at any point shall be circular. The distal end of the white cell pipet shall be pulled to a point and then ground and polished to a tapered tip. The distal end of the red cell pipet shall be ground and polished to a tapered tip. The inside diameter (ID) of the pipet bore at the tip end of the white cell pipet shall be 0.2 to 0.5 mm. The external diameter of the ground and polished tip shall not exceed 2.0 mm. The overall length of both pipets shall be 104 to 121 mm. The proximal end may be of funnel design, with an exterior taper, or ground and polished with a taper of sufficient angle to permit application of a rubber suction tube or other suction device (see Fig. 2).

5.2 *Capacity*—The capacity of the red cell pipet bulb shall be 0.8 to 1.2 cm³ and the capacity of the white cell bulb shall be 0.2 to 0.4 cm³. The capacity of the red cell pipet stem (bulb to distal end) shall be 0.008 to 0.012 cm³ and the capacity of the white cell pipet stem shall be 0.02 to 0.04 cm³.

5.3 *Capacity Markings*:

5.3.1 *Pipet Bulb Markings*—The red and white cell pipet bulb capacity shall be confined to and indicated by calibration lines on the pipet. These lines shall be located on both sides of the bulb and shall be within 2 mm to 6 mm above the bulb (proximal end) and within 3 mm to 6 mm below the bulb (distal end). The exact manner for measuring these line placements is specified in Fig. 1.

5.3.2 *Pipet Stem Markings*—The graduation lines on the stem of the red and white cell pipets shall be equally divided into two or ten divisions. If there are two divisions, the two

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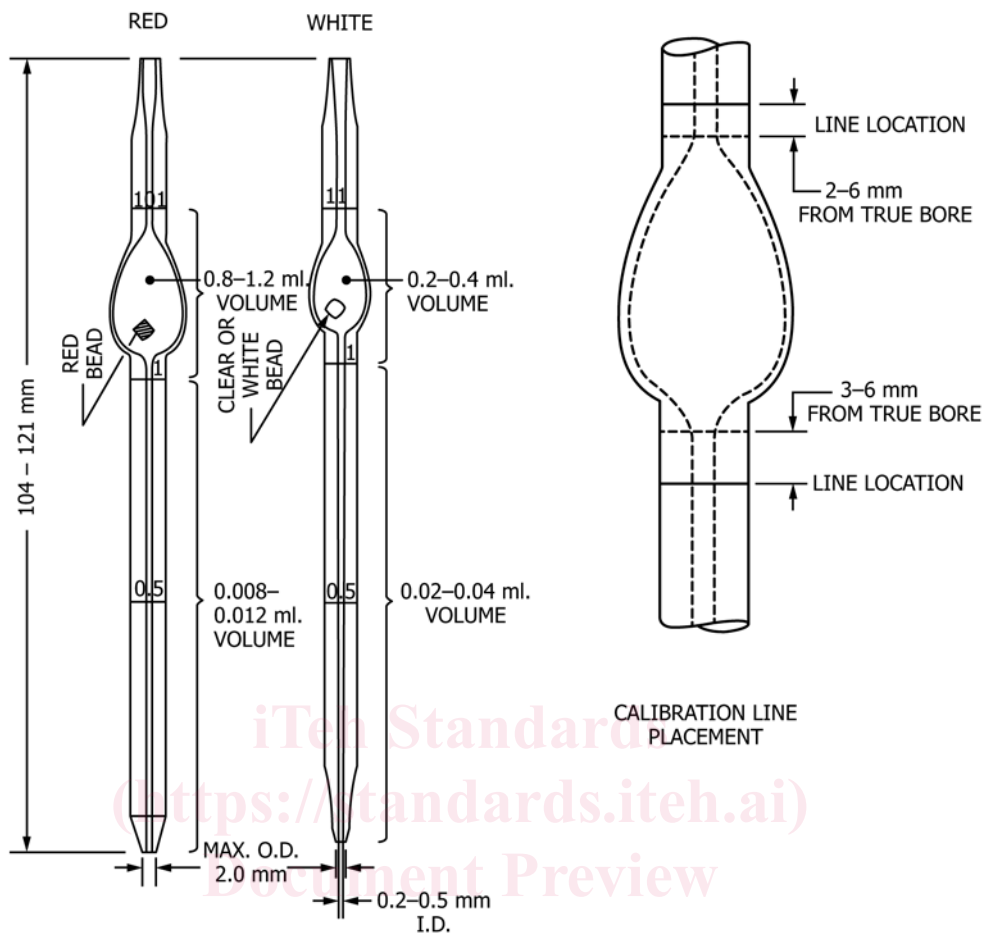


FIG. 1 Pipet Dimensions

calibration lines on the stem will be numbered 0.5 and 1. If there are ten divisions, the ten calibration lines will be numbered 0.5 on the fifth line and 1 on the tenth line respectively. The top line (above bulb at the proximal end) shall be numbered 101 on the red cell pipet and 11 on the white cell pipet, representing stem and bulb volume collectively. At the option of the manufacturer, pipet stems may be graduated with split lines or partial length lines (see Fig. 2).

5.4 *Graduation Lines*—All graduation lines on the red and white cell pipets shall be at right angles to the pipet axis and parallel to each other. The thickness of the graduation lines shall not exceed 0.4 mm.

5.5 *Accuracy*—The red and white cell pipet bulb volume shall be a volumetric ratio to the stem volume. The red cell pipet bulb to stem ratio shall be 100:1 and the white cell pipet bulb to stem ratio shall be 10:1 when tested in accordance with 6.1 and 6.2.

5.6 *Workmanship:*

5.6.1 The pipets shall be free of defects that detract from their appearance or impair their serviceability.

5.6.2 Construction shall be such that the mechanical strength is provided to withstand the rigors of normal use. The pipets shall be free of strain when tested in accordance with 6.4.

5.7 *Identification*—Each pipet shall have the name or registered trademark of the manufacturer. These markings are to be located on the stem of the pipet directly opposite the graduation lines. The manufacturer has the option to state the pipet volumetric tolerances on the pipet. These tolerances may be located on the stem or the proximal end of the pipet.

5.7.1 The white cell pipet shall have a clear or white glass or ceramic bead sealed within the bulb to identify it readily as a white cell pipet.

5.7.2 The red cell pipet shall have a red glass or ceramic bead sealed within the bulb to identify it readily as a red cell pipet.

5.8 *Pigmentation*—All markings shall be permanently fused in or on the pipet. The markings shall be dark amber or black in color. When tested in accordance with 6.3, the pigmentation shall not discolor the solution. The appearance of the markings,