



Designation: ~~F2952--14~~ F2952 – 22

Standard Guide for Determining the Mean Darcy Permeability Coefficient for a Porous Tissue Scaffold¹

This standard is issued under the fixed designation F2952; 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 reappraisal. A superscript epsilon (ϵ) indicates an editorial change since the last revision or reappraisal.

1. Scope

1.1 This guide describes test methods suitable for determining the mean Darcy permeability coefficient for a porous tissue scaffold, which is a measure of the rate at which a fluid, typically air or water, flows through it in response to an applied pressure gradient. This information can be used to optimize the structure of tissue scaffolds, to develop a consistent manufacturing process, and for quality assurance purposes.

1.2 The method is generally ~~non-destructive~~ nondestructive and non-contaminating.

1.3 The method is not suitable for structures that are easily deformed or damaged. Some experimentation is usually required to assess the suitability of permeability testing for a particular material/structure and to optimize the experimental conditions.

1.4 Measures of permeability should not be considered as definitive metrics of the structure of porous tissue scaffolds and should complement measures obtained by other investigative techniques e.g., techniques, for example, scanning electron microscopy, gas flow porometry, and micro-computer ~~x-ray~~ X-ray tomography (~~ASTM~~ ASTM Guides F2450, F2603, and F3259).

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

1.6 *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

- ~~D4525~~ D737 Test Method for Air Permeability of ~~Roeks by Flowing Air~~ Textile Fabrics (Withdrawn 2022)
- D2434 Test Methods for Measurement of Hydraulic Conductivity of Coarse-Grained Soils
- F2450 Guide for Assessing Microstructure of Polymeric Scaffolds for Use in Tissue-Engineered Medical Products
- F2603 Guide for Interpreting Images of Polymeric Tissue Scaffolds
- F3259 Guide for Micro-computed Tomography of Tissue Engineered Scaffolds
- F3510 Guide for Characterizing Fiber-Based Constructs for Tissue-Engineered Medical Products

¹ This ~~test method~~ guide is under the jurisdiction of ASTM Committee F04 on Medical and Surgical Materials and Devices and is the direct responsibility of Subcommittee F04.42 on Biomaterials and Biomolecules for TEMPs.

Current edition approved ~~March 1, 2014~~ April 1, 2022. Published ~~April 2014~~ April 2022. Originally approved in 2014. Last previous edition approved in 2014 as F2952 – 14. DOI: ~~10.1520/F2952-14~~ 10.1520/F2952-22.

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.

2.2 American Petroleum Institute (API) Document.⁴

RP-27 Recommended Practice for Determining Permeability of Porous Media

3. Terminology

3.1 Definitions: Definitions of Terms Specific to This Standard:

3.1.1 *tortuosity, n*—the ratio of the actual path length through connected pores to the Euclidean distance (shortest linear distance).

4. Significance and Use

4.1 This document describes the basic principles that need to be followed to obtain a mean value of the Darcy permeability coefficient for structures that consist of a series of interconnected voids or pores. The coefficient is a measure of the permeability of the structure to fluid flowing through it that is driven by a pressure gradient created across it.

4.2 The technique is not sensitive to the presence of closed or blind-end pores (Fig. 1).

4.3 Values of the permeability coefficient can be used to compare the consistency of manufactured samples or to determine what the effect of changing one or more manufacturing settings has on permeability. They can also be used to assess the homogeneity and anisotropy of tissue scaffolds. Variability in the permeability coefficient can be also be indicative of:

4.3.1 Internal damage within the sample e.g., sample, for example, cracking or permanent deformation.

4.3.2 The presence of large voids, including trapped air bubbles, within the structure.

4.3.3 Surface effects such as a skin formed during manufacture.

4.3.4 Variable sample geometry.

4.4 This test method is based on the assumption that the flow rate through a given sample subjected to an applied pressure gradient is constant with time.

NOTE 1—If a ~~steady-state~~ steady-state flow condition isn't reached, then this could be due to structural damage (i.e., that is, crack formation or the porous structure deformed as a result of the force being placed upon it by the fluid flowing through it). Sample deformation in the form of stretching (bowing) can also occur for less resilient structures as a result of high fluid flow rates. This topic is discussed in more detail in Section 7.

4.5 Care should be taken to ensure that hydrophobic materials are fully wetted out when using water or other aqueous-based liquids as permeants.

4.6 Conventionally, the pressure differential created across a sample is measured as a function of both increasing and decreasing flow rates. An alternative approach, which may be practically easier to create, is to apply a range of different pressure differentials across the sample and measure the resultant flow of fluid through it. The hysteresis that occurs during a complete cycle of

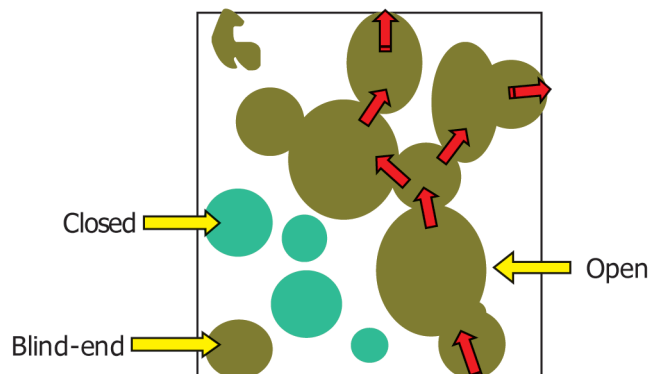


FIG. 1 Schematic of the Different Pores Types Found in Tissue Scaffolds. Fluid Flow through the Structure is via the Open Pores

increasing flow rate followed by a progressive decrease in flow rate can provide an excellent measure of the behavioural consistency of the matrix. Significant hysteresis in the measured pressure differential during increasing and decreasing flow rates can indicate the existence of induced damage in the structure, the fact that the material is behaving viscoelastically, or is suffering from permanent plastic deformation. Some guidance on how to identify which of these factors are responsible for hysteresis is provided in Section 7.

4.7 It is assumed that Darcy’s law is valid. This can be established by plotting the volume flow through the specimen against the differential pressure drop across the specimen. This plot should be linear for Darcy’s law to apply and a least-squares fit to the data should pass through the origin. It is not uncommon for such plots to be non-linear which may indicate that the structure does not obey Darcy’s law or that the range of pressures applied is too broad. This topic is further discussed in Section 7.

5. Characterisation/Characterization and the Structural Features of Tissue Scaffolds

5.1 Porous tissue scaffolds are typically manufactured from polymers and ceramics and consist of a network of connected voids through which cells, macromolecules such as growth factors, and small molecules such as nutrients and dissolved gases can move (1).³ In most cases, the material used to create the scaffold will disappear over time, either as a result of enzyme activity or some other degradation processes (e.g., for example, hydrolysis). The time-dependent permeability of tissue scaffolds to dissolved gases and solutes is critical to their function, particularly for high levels of cell occupancy due to the demands for oxygen and nutrients as well as the need to remove waste products.

5.2 There are many methods available for characterizing the structural features of scaffolds (ASTM(Guides F2450-10), and F3510), but these can be time-consuming, expensive to use, and can result in permanent damage or contamination to the scaffold.

5.3 Most investigators report some measure of pore size and an estimate of the scaffold porosity (2, 3). However, there are significant practical issues associated with these measurements. Techniques such as mercury porosimetry and gas flow porometry are used to estimate pore size distributions which typically differ by an order of magnitude due to differences in the underlying physics of the techniques (ASTM(Guide F2450). Despite the shortfalls of these techniques, both can be used to infer a useful amount of information regarding the structure of the scaffold. Both porosimetry and porometry represent the scaffold structure as a distribution of differently sized parallel-sided pores i.e., pores, that is, the model assumes a simple structure that is equivalent to the more complicated structures usually manufactured where the pores are not parallel-sided and not of uniform diameter.

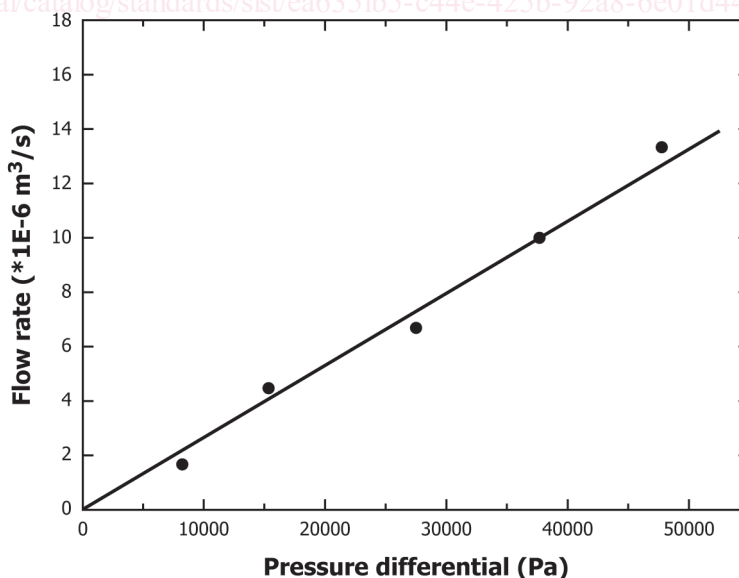


FIG. 2 Example of a Plot of Flow Rate versus Pressure Differential

³ The last approved version of this historical standard is referenced on www.astm.org.

⁴ Available from American Petroleum Institute (API), 1220 L St., NW, Washington, DC 20005-4070, <http://www.api.org>.

³ The boldface numbers in parentheses refer to the list of references at the end of this standard.

5.4 Electron and other microscopies are extensively used to image scaffolds, but the data that these techniques produce is often challenging to interpret without some undefinable level of uncertainty (i.e., that is, quantifying the dimensions of typically irregularly shaped and sized structural features). The same arguments apply to tomographic methods such as magnetic resonance imaging and micro-computer tomography (~~(μ CT)~~); for example, calculations based on the analysis of a series of scaffold images obtained from a tomographical method such as μ CT will depend on how well the boundaries of the voids or pores can be defined, on the instrument resolution in the x, y and z planes, and the methodology used to obtain dimensional information. Nevertheless, many groups have pursued quantitative analysis of pore size distributions in polymeric (3) and bioceramic (4) matrices in recognition of the important correlation between this parameter and tissue ingrowth.

5.5 The pores in a tissue scaffold typically consist of a series of irregularly shaped voids⁴ that can be connected to each other both by partial fusion and connecting channels (connects). Through pores provide a path through the scaffold from one side to the other, and are the primary routes for fluid penetration into the scaffold. The dimensions of a given pore can be difficult to define due to, for example, merging of adjacent cavities that result in fenestrations or “windows” forming in the void walls. Blind-end and closed-pores, although not contributing to measures of fluid permeability, play an important role in gas diffusion through the structure.

6. The Darcy Permeability Coefficient

6.1 The Darcy permeability coefficient is a measure of the resistance of a porous material to flow of a fluid through it that is governed by the dimensions and density of open (or through) pores and by the tortuosity of the structure.

6.2 In its simplest form, the permeability coefficient, k , of the scaffold can be determined by measuring the flow of fluid through the material in a given time under a known pressure gradient using Darcy’s law (5), i.e., that is,

$$Q = \frac{-kA(P_b - P_a)}{\mu L} \quad (1)$$

which states that the flow rate (Q , (m^3/s)) through the material is directly proportional to the cross-sectional area (A , (m^2)) and the pressure drop ($P_b - P_a$, (Pa)) and inversely proportional to the viscosity of fluid (μ , (Pa.s)) and the length (L , (m)) over which the pressure drop occurs.

6.3 The permeability coefficient, k , is then derived from the slope of a linear plot of flow rate versus pressure drop where the slope is forced to pass through the origin (see Fig. 2).

6.4 The SI units of the coefficient are m^2 .

6.5 Permeability coefficients are routinely used in assessing soils, filters, and other porous materials (ASTM (Test Methods D4525D737-08 and D2434RP-27)) and have also been used to characterize polymeric scaffolds and hard tissues e.g., tissues, for example, cancellous bone (6-911).

7. Methodology

7.1 Obtaining reliable values for the permeability coefficient involves a degree of experimental optimization to ensure that a range of flow rates and pressure differentials can be measured. Clearly, it is advantageous to measure a range of flow rates and pressure differentials to improve the reliability of the Darcy coefficient, but this can produce non-linear plots for reasons that are discussed in Section 8. This will require some experimentation to optimize the sample geometry and to select the most appropriate fluid, typically air or water, for a given structure/sample geometry and material type. Sections 7.3 and 7.4 describe the features that are required in an experimental system in order to obtain robust estimates of the coefficient.

7.2 Reliably determining the pressure differential across the scaffold and measuring the flow rate through it are fundamental aspects of permeability testing. In practice, the sensitivity of the apparatus used to measure pressure will limit the magnitude of the pressure gradient that can be used for a given sample geometry.

⁴ The terminology for scaffold structure is not well defined. The term pore “pore” is widely used to mean a void, a window in a void, or a conduit connecting two or more voids together.

7.3 *Gas-based Gas-Based Systems:*

7.3.1 Fig. 3 shows a schematic representation of apparatus that can be used to measure the flow of gas, in this case compressed air, through a disc-like sample mounted in a commercially available filter holder that can be purchased in a range of sizes.

7.3.2 The rate of flow through the sample is measured by a gas flow meter. These devices are commercially available for different ranges of flow rate. Care should be taken to ensure that the flow meter used is appropriate for the flow rates used to avoid potential measurement inaccuracies. The pressure upstream of the sample, P_b , is measured and used together with a measured value for atmospheric pressure (P_a) to determine the pressure gradient ($P_b - P_a$) required by Eq 1.

7.4 *Liquid-based Liquid-Based Systems:*

7.4.1 Fig. 4 shows an experimental configuration that measures the flow of a liquid, such as water, through a porous tubular scaffold sample. The apparatus consists of a circulating pump, which is used to generate an internal pressure within the circuit, P_b . P_a is the measured value of atmospheric pressure. The internal pressure that develops within the circuit is very dependent on the permeability of the scaffold and its geometry, but is usually sufficiently high that any changes in pressure along the length of a vertically mounted sample due to differences in height can be ignored. However, the user is advised to check that this assumption is valid for the sample and sample geometry that is being investigated.

7.4.2 The water that flows through the walls of the specimen and out through the overflow is collected at given time intervals, weighed, and converted into a flow rate. The fluid reservoir replenishes the fluid lost from the system via the overflow.

7.4.3 Alternative sample geometries can be used (i.e., (that is, a disc of material sandwiched between ‘O’ rings—O-rings in a commercially available filter holder), as used for gas-based systems. In both cases the practical considerations are the same: how to apply a progressively increasing pressure gradient without significantly deforming the sample or letting fluid flow around it.

8. Practical Considerations

8.1 There are many experimental configurations that can be used to generate the flow rate and pressure differential measurements required to determine Darcy’s permeability coefficient. It is not uncommon to observe a degree of non-linearity in plots of flow rate versus differential pressure, particularly when investigating a new sample, whether it is manufactured from a different material or produced using different processing conditions. The following considerations should be taken into account in designing the apparatus and defining the sample geometry to ensure that the relationship between flow rate and differential pressure is attributed to the structure and not an experimental artifact.

8.2 *Sample Characteristics:*

8.2.1 The samples will need to have sufficient stiffness to ensure that they are able to withstand a pressure gradient without

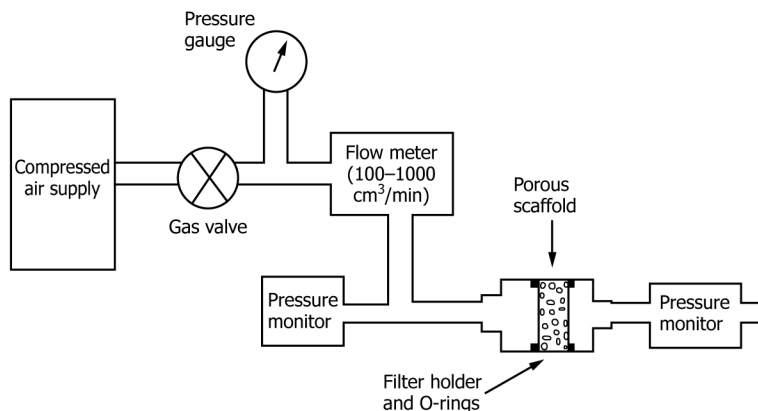


FIG. 3 Measuring the Pressure Differential Across a Disc of a Porous Scaffold Produced by the Controlled Flow of Gas Through the Disc

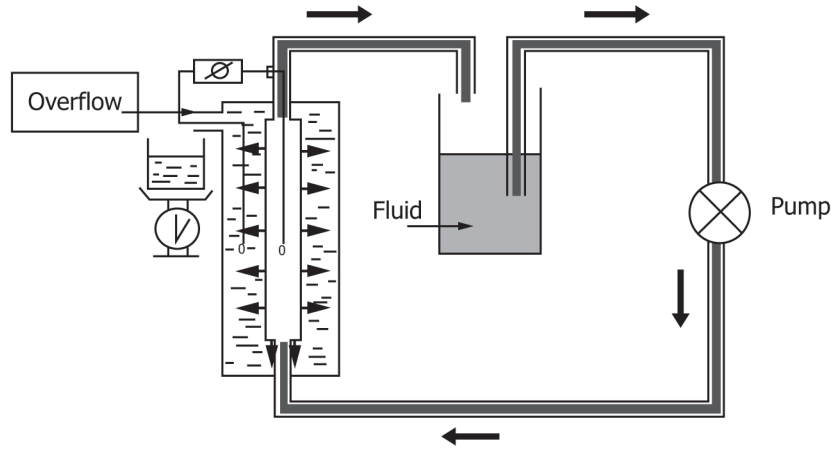


FIG. 4 Passage of Fluid Through the Wall of a Porous Tubular Scaffold in a Closed Loop Pumped System can be Determined by Weighing Fluid Samples at Defined Time Intervals

incurring damage or deforming significantly e.g., significantly, for example, bowing. Unfortunately the suitability of permeability testing as an experimental method for a given material/structure/sample geometry will need to be established by experimentation using the guidance given in [Table 1](#).

8.2.2 It is recommended that measurements of flow rate and pressure differential be made cyclically to assess potential hysteresis of the system. This can be the simple approach of measuring the flow rates at increasing pressure differentials followed by a progressive decrease back to the start point. A time lag should be allowed before a measurement is made at a given pressure differential to allow the experimental system to reach a steady-state condition.

8.2.3 A significant difference in flow rate after one or more cycles of pressurizing/depressurizing the sample is indicative of a structural change having occurred within a sample. This may be due to viscoelastic effects in a polymer-based scaffold, or may be indicative of either structural damage or permanent plastic deformation. Differentiating between potentially permanent changes in the sample structure from viscoelastic effects is usually straightforward if the tests are repeated after a period of several hours as viscoelastic effects are reversible.

ASTM F2952-22

NOTE 2—The sample must be left *in situ* between successive tests in an unloaded condition if this approach is followed.

8.3 Sample Geometry:

8.3.1 The value of the mean permeability coefficient obtained by experimentation will be influenced by errors in the true sample area and thickness.

8.3.2 Care should be taken to minimize thickness variations across a specimen.

TABLE 1 Potential Solutions to Commonly Encountered Problems

Problem	Potential Solution
Insufficient flow rate	Increase the sample area Reduce the sample thickness Reduce the viscosity of the fluid used (e.g., substitute air for water)
Pressure gradient too small	Increase the sample thickness Reduce the sample area Increase the viscosity of the fluid used (e.g., substitute water for air)
Sample deforms during the experiment	Increase the sample thickness Reduce the sample area Reduce the viscosity of the fluid used (e.g., substitute air for water)

8.3.3 The sample dimensions should be much larger than the dimensions of the voids/pores that they contain to avoid the scenario where the presence of one or more large voids significantly reduces the effective sample thickness.

8.3.4 It can be practically challenging to determine the effective sample area, particularly when the sample is sandwiched between ‘O-rings-O-rings. It may be easier to determine this after the measurements have been made as the ‘O-rings-O-rings will leave an impression in the sample that serves to define the maximum diameter of the area exposed to fluid flow. Errors in determining the true sample diameter can be more significant for smaller sample areas.

8.3.5 While apparently excellent Darcy plots can be constructed for samples of ~~non-uniform~~nonuniform thickness, the obtained value of the coefficient is not reliable. This is an obvious issue for poorly prepared samples or those that are difficult to machine. It is also important to consider the thickness of the sample used in permeability measurements as shown in Fig. 5. If the ratio between the measured sample thickness and mean pore diameter is too low, then the Darcy coefficient obtained will not be representative of the structure as a whole. It is therefore recommended that the sample thickness be at least ten times that of the mean pore diameter to overcome this limitation.

8.4 Dealing with Hydrophobic Materials:

8.4.1 Commonly used scaffold materials, such as poly(lactic-glycolic acid) copolymer (PLGA), poly(lactic acid) (PLA) and polycaprolactone (PCL) which are hydrophobic, can be difficult to work with using water as the permeant as a result of incomplete wetting out. Hydrophobic materials are also susceptible to persistent trapped air bubbles remaining in the structure. A strategy for dealing with these problems is to add a few drops of a wetting agent, such as a surfactant or ethanol to the water used to ‘wet’“wet” out the sample prior to measurements being made. A very weak ethanol/water mixture is usually sufficient to overcome the surface tension that prevents wetting out of the structure. A weak vacuum can also be used to induce wetting out of hydrophobic structures for those materials that can craze in the presence of ethanol (environmental stress cracking). Care should be taken so that the vacuum applied does not cause any damage to the structure.

8.5 Mounting the Sample in a Holder:

8.5.1 Disc-like samples can be easily mounted in commercially available filter holders. These clamp the sample between two ‘O-rings-O-rings. When mounting the sample, care should be taken to:

8.5.1.1 Ensure that the seal created between the clamp and the sample is sufficiently good to avoid fluid leakage during the experiment.

8.5.1.2 Ensure that any clamping pressure applied does not damage or significantly distort the sample. This consideration is particularly relevant for small diameter (<7 (<7 mm diameter) disc-like samples clamped between two ‘O-rings-O-rings where the sample will bulge if the applied clamping pressure is too high, thereby effectively increasing the sample thickness.

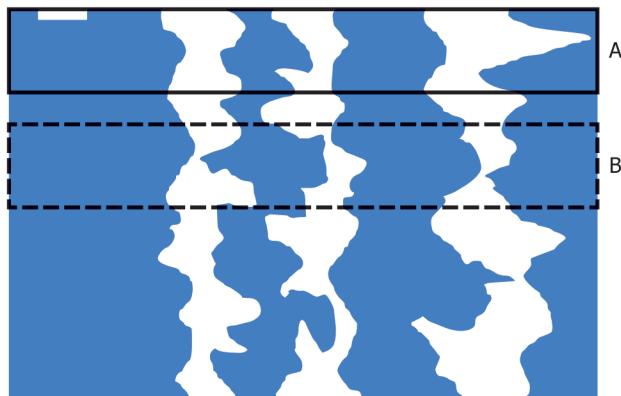


FIG. 5 A Representative Pore Distribution representative pore distribution must be Present present in Scaffold Samples scaffold samples in Order order to get a Good Measurement good measurement of Darcy’s Coefficient coefficient. Permeability through the Portion portion of the Scaffold Enclosed scaffold enclosed by Box A will be Higher higher than Permeability Measured permeability measured through the Portion portion of the Scaffold Enclosed scaffold enclosed by Box B. Box B has much Tighter Constriction tighter constrictions than Box A.