



Standard Test Method for Linear Dimensional Stability of a Gasket Material to Moisture¹

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1. Scope

1.1 This test method covers a procedure to determine the stability of a gasket material to linear dimensional change due to hygroscopic expansion and contraction. It subjects a sample to extremes, that is, oven drying and complete immersion in water, that have shown good correlation to low and high relative humidities.²

1.2 The values stated in SI units are to be regarded as the standard. The values given in parentheses are for information only.

1.3 *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 and health practices and determine the applicability of regulatory limitations prior to use.*

2. Summary of Test Method

2.1 A series of samples are preconditioned to a stable starting point, measured, and then conditioned to a second exposure condition, either wet or dry. These changes are then determined and recorded, and the results presented as percent change.

3. Significance and Use

3.1 Gasket materials undergo several processing steps from point of manufacture to installation in a flange. Many applications require close control of dimensional change. An accurate test method for determining the relative stability of various materials is needed for design and quality assurance purposes. This test method is useful towards that end. It simulates the extreme storage conditions that a material may undergo prior to installation. Samples are allowed unrestricted expansion or contraction, and so this test method should not be used to

¹ This test method is under the jurisdiction of ASTM Committee F03 on Gaskets and is the direct responsibility of Subcommittee F03.20 on Mechanical Test Methods.

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² Since this test method takes a “worst case” approach, actual dimensional change due to atmospheric conditions would be expected to be less.

predict behavior clamped in a flange or other applications, or during specific processing steps.

3.2 This test method measures linear change, and may need to be modified if the test specimen is not flat, homogeneous, or free of voids.

4. Apparatus

4.1 This test method allows individual laboratories to select measuring devices of their own choice, but requires that the device be able to measure with a precision of ± 0.025 mm (0.001 in.).

5. Sampling

5.1 At least six test specimens should be taken for each sample material, three for the high humidity and three for the low humidity tests. The samples should be cut 2.54 cm (1.00 in.) wide, and between 20 and 30 cm (8 and 12 in.) in length. The long direction should be in the direction that experiences the greatest dimensional change, generally the cross machine or against the grain direction. If there is doubt, both directions should be sampled, and the results of the direction with the greatest change reported.

6. Conditioning

6.1 Test specimens should be preconditioned at least 20 h in a controlled environment maintained between 21 to 30°C (70 to 85°F) and 50 to 55 % relative humidity.

7. Procedure

7.1 Measure test specimens to ± 0.025 mm (± 0.001 in.) and record values as initial readings. If the test specimen is marked for identification or measurement, be certain that the mark is easily visible and will withstand exposure to heat and immersion in water.

7.2 *Testing for Dimensional Stability to Low Humidity*—Expose three prepared specimens in a forced hot-air oven set at $100 \pm 2^\circ\text{C}$ ($212 \pm 4^\circ\text{F}$) for 5 h. Remove specimens and allow to cool between 21 to 30°C (70 to 85°F) in a desiccator containing anhydrous-calcium chloride or suitable desiccant material. Remeasure and record measurements as final readings.

7.3 Testing for Dimensional Stability to High Humidity— Immerse three prepared specimens into a tray of deionized water to a depth of 1.2 cm (0.5 in.) for a 22 h period. For materials that are buoyant in water, a supported wire screen or expanded metal cover should be used with adequate weights to keep the specimens immersed at the specified depth. Be certain that the specimens are separated and able to expand in an unrestricted fashion. Remove specimens from water and lightly blot excess water from the surface of the specimens. Remeasure and record measurements as final readings.

8. Calculation of Results

8.1 Report the results as percent change to high or low humidity calculated from the following equations:

8.1.1 *High Humidity:*

$$L_{\Delta} = \frac{L_f - L_i}{L_i} \times 100 \quad (1)$$

8.1.2 *Low Humidity:*

$$L_{\Delta} = \frac{L_i - L_f}{L_i} \times 100 \quad (2)$$

where:

L_i = initial length,
 L_f = final length, and
 L_{Δ} = change in length, %.

8.2 Calculate and report the average of the three individual specimens. The total change can then be reported as the sum of the low humidity and high humidity changes.

TABLE 1 Change to Dry Condition

| Material | Average X_j | Repeat-ability (SR_j) | Reproduc-ability (SL_j) | Test Method Precision (SF_j) |
|----------|---------------|---------------------------|-----------------------------|----------------------------------|
| A | 0.02 | 0.010 | 0.009 | 0.014 |
| B | 1.02 | 0.081 | 0.235 | 0.248 |
| C | 0.45 | 0.034 | 0.096 | 0.244 |
| D | 0.52 | 0.043 | 0.121 | 0.128 |
| E | 0.35 | 0.042 | 0.124 | 0.131 |
| F | 0.11 | 0.023 | 0.019 | 0.030 |
| G | 0.21 | 0.058 | 0.124 | 0.137 |
| H | 1.74 | 0.039 | 0.130 | 0.136 |

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TABLE 2 Change to Wet Condition

| Material | Average X_j | Repeat-ability (SR_j) | Reproduc-ability (SL_j) | Test Method Precision (SF_j) |
|----------|---------------|---------------------------|-----------------------------|----------------------------------|
| A | 0.25 | 0.027 | 0.039 | 0.046 |
| B | 0.46 | 0.018 | 0.077 | 0.079 |
| C | 0.76 | 0.026 | 0.122 | 0.124 |
| D | 2.59 | 0.039 | 0.192 | 0.196 |
| E | 0.41 | 0.022 | 0.020 | 0.030 |
| F | 0.24 | 0.015 | 0.037 | 0.040 |
| G | 1.52 | 0.195 | 0.308 | 0.365 |
| H | 10.00 | 0.200 | 0.613 | 0.645 |

9. Report

9.1 Report the following information:

9.1.1 The complete identification of the sample material, grade, and caliper,

9.1.2 Date of test, and initial and final lengths for each specimen,

9.1.3 Type and precision of instrument used to determine length, and

9.1.4 Average percentage change in length (%) for low humidity, high humidity, and total.

10. Precision and Bias³

10.1 **Table 1** and **Table 2** present the results from which the repeatability within a laboratory, and the reproducibility between laboratories can be determined.

10.2 Since there is no accepted reference material suitable for determining the bias for the procedure in this test method for measuring the dimensional stability of gasket materials, no statement on bias is being made.

11. Keywords

11.1 linear dimensional stability; moisture; thickness

³ The precision data were derived from the results of an interlaboratory test program on materials A through H. Supporting data are available from ASTM International Headquarters. Request RR: F03-1011.