

Standard Test Methods for Performance Characteristics of Metallic Bonded Resistance Strain Gages¹

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This standard has been approved for use by agencies of the Department of Defense.

INTRODUCTION

The Organization of International Legal Metrology is a treaty organization with approximately 75 member nations. In 1984, OIML issued International Recommendation No. 62, 'Performance Characteristics of Metallic Resistance Strain Gages.' Test Methods E 251 has been modified and expanded to be the United States of America's compliant test specification. Throughout this standard the terms "strain gage" and "gage" are to be understood to represent the longer, but more accurate, "metallic bonded resistance strain gages."

1. Scope

1.1 The purpose of this standard is to provide uniform test methods for the determination of strain gage performance characteristics. Suggested testing equipment designs are included.

1.2 Test Methods E 251 describes methods and procedures for determining five strain gage parameters:

	Section
Part I—General Requirements	7
Part II—Resistance at a Reference Temperature	8
Part III—Gage Factor at a Reference Temperature	9
Part IV—Temperature Coefficient of Gage Factor	10
Part V—Transverse Sensitivity	11
Part VI—Thermal Output	12

1.3 Strain gages are very sensitive devices with essentially infinite resolution. Their response to strain, however, is low and great care must be exercised in their use. The performance characteristics identified by these test methods must be known to an acceptable accuracy to obtain meaningful results in field applications.

1.3.1 Strain gage resistance is used to balance instrumentation circuits and to provide a reference value for measurements since all data are related to a change in the gage resistance from a known reference value.

1.3.2 Gage factor is the transfer function of a strain gage. It relates resistance change in the gage and strain to which it is

subjected. Accuracy of strain gage data can be no better than the precision of the gage factor.

1.3.3 Changes in gage factor as temperature varies also affect accuracy although to a much lesser degree since variations are usually small.

1.3.4 Transverse sensitivity is a measure of the strain gage's response to strains perpendicular to its measurement axis. Although transverse sensitivity is usually much less than 10 % of the gage factor, large errors can occur if the value is not known with reasonable precision.

1.3.5 Thermal output is the response of a strain gage to temperature changes. Thermal output is an additive (not multiplicative) error. Therefore, it can often be much larger than the gage output from structural loading. To correct for these effects, thermal output must be determined from gages bonded to specimens of the same material on which the tests are to run; often to the test structure itself.

1.4 Bonded resistance strain gages differ from extensioneters in that they measure average unit elongation ($\Delta L/L$) over a nominal gage length rather than total elongation between definite gage points. Practice E 83 is not applicable to these gages.

1.5 These test methods do not apply to transducers, such as load cells and extensioneters, that use bonded resistance strain gages as sensing elements.

1.6 Strain gages are part of a complex system that includes structure, adhesive, gage, leadwires, instrumentation, and (often) environmental protection. As a result, many things affect the performance of strain gages, including user technique. A

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further complication is that strain gages once installed normally cannot be reinstalled in another location. Therefore, gage characteristics can be stated only on a statistical basis.

1.7 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.

1.8 The values stated in SI units are to be regarded as the standard.

2. Referenced Documents

2.1 ASTM Standards:

E 83 Practice for Verification and Classification of Extensometers²

- E 228 Test Method for Linear Thermal Expansion of Solid Materials with a Vitreous Silica Dilatometer³
- E 289 Test Method for Linear Thermal Expansion of Rigid Solids with Interferometry³
- E 1237 Guide for Installing Bonded Resistance Strain Gages²

2.2 OIML International Recommendation No. 62:' Performance Characteristics of Metallic Resistance Strain Gages

3. Terminology

3.1 Definitions of Terms Specific to This Standard:

3.1.1 The vocabulary included herein has been chosen so that specialized terms in the strain gage field will be clearly defined. A typical strain gage nomenclature is provided in Appendix X1.

3.1.1.1 *batch*—a group of strain gages of the same type and lot, manufactured as a set (made at the same time and under the same conditions).

3.1.1.2 calibration apparatus-equipment for determining a characteristic of a bonded resistance strain gage by accurately producing the necessary strains, temperatures, and other conditions; and, by accurately measuring the resulting change of gage resistance.

3.1.1.3 error-strain gage-the value obtained by subtracting the actual value of the strain, determined from the calibration apparatus, from the indicated value of the strain given by the strain gage output. Errors attributable to measuring systems are excluded.

3.1.1.4 gage factor—the ratio between the unit change in strain gage resistance due to strain and the causing strain. The gage factor is dimensionless and is expressed as follows:

$$K = \frac{R - R_o}{R_o} / \frac{L - L_o}{L_o} = \frac{\Delta R}{R_o} / \epsilon$$
(1)

where:

K = the gage factor,

= the strain gage resistance at test strain, R

= the strain gage resistance at zero or reference strain, R_o

- L = the test structure length under the strain gage at test strain.
- = the test structure length under the strain gage at zero L_o or reference strain,
- ΔR = the change in strain gage resistance when strain is changed from zero (or reference strain to test strain),

= the mechanical strain $L - L_o / L_o$. ε

3.1.1.5 gage length (see Fig. 1)—the length of the strain sensitive section of a strain gage in the measurement axis direction. An approximation of this length is the distance between the inside of the strain gage end loops. Since the true gage length is not known, gage length may be measured by other geometries (such as the outside of the end loops) providing that the deviation is defined.

3.1.1.6 grid (see Fig. 1)—that portion of the strain-sensing material of the strain gage that is primarily responsible for resistance change due to strain.

3.1.1.7 *lot*—a group of strain gages with grid elements from a common melt, subjected to the same mechanical and thermal processes during manufacturing.

3.1.1.8 matrix (see Fig. 1)—an electrically nonconductive layer of material used to support a strain gage grid. The two main functions of a matrix are to act as an aid for bonding the strain gage to a structure and as an electrically insulating layer in cases where the structure is electrically conductive.

3.1.1.9 measurement axis (grid) (see Fig. 1)—that axis that is parallel with the grid lines.

3.1.1.10 strain gage, metallic, resistive, bonded (see Fig. 1)-a resistive element, with or without a matrix that is attached to a solid body by cementing, welding, or other suitable techniques so that the resistance of the element will vary as the surface to which it is attached is deformed. These test methods apply to gages where the instantaneous gage resistance, R, is given by the equation:

$$R = R_o (1 + \epsilon K) \tag{2}$$

where:

- R_o = element resistance at reference strain and temperature levels (frequently initial test or balanced circuit conditions),
- = linear strain of the surface in the direction of the ε strain-sensitive axis of the gage, and

K = a proportionality factor (see gage factor).

3.1.1.11 strain, linear-the unit elongation induced in a specimen either by a stress field (mechanical strain) or by a temperature change (thermal expansion).

3.1.1.12 temperature coefficient of gage factor-the ratio of the unit variation of gage factor to the temperature variation, expressed as follows:

$$\left(\frac{K_{t1} - K_{t0}}{K_{t0}}\right) \cdot \left(\frac{1}{T_1 - T_0}\right)$$
(3)

where:

- T_1 = the test temperature,

 T_0 = the reference temperature, K_{t1} = the gage factor at test temperature, and

 K_{t0} = the gage factor at reference temperature.

² Annual Book of ASTM Standards, Vol 03.01.

³ Annual Book of ASTM Standards, Vol 14.02.



FIG. 1 Typical Strain Gage

3.1.1.13 *thermal expansion*—the dimensional change of an unconstrained specimen subject to a change in temperature that is uniform throughout the material.

3.1.1.14 *thermal output*—the reversible part of the temperature induced indicated strain of a strain gage installed on an unrestrained test specimen when exposed to a change in temperature.

3.1.1.15 *transverse axis* (see Fig. 1)—the strain gage axis at 90° to the measurement axis.

3.1.1.16 *transverse sensitivity*—the ratio, expressed as a percentage, of the unit change of resistance of a strain gage mounted perpendicular to a uniaxial strain field (transverse gage) to the unit resistance change of a similar gage mounted parallel to the same strain field (longitudinal gage).

3.1.1.17 *type*—a group of strain gages that are nominally identical with respect to physical and manufacturing characteristics.

4. Significance and Use

4.1 Strain gages are the most widely used devices for the determination of materials, properties and for analyzing stresses in structures. However, performance parameters of strain gages are affected by both the materials from which they are made and their geometric design. These test methods detail the minimum information that must accompany strain gages if they are to be used with acceptable accuracy of measurement.

4.2 Most performance parameters of strain gages require mechanical testing that is destructive. Since test gages cannot be used again, it is necessary to treat data statistically and then apply values to the remaining population from the same lot or batch. Failure to acknowledge the resulting uncertainties can have serious repercussions. Resistance measurement is nondestructive and can be made for each gage. 4.3 Properly designed and manufactured strain gages, whose properties have been accurately determined and with appropriate uncertainties applied, represent powerful measurement tools. They can determine small dimensional changes in structures with excellent accuracy, far beyond that of other known devices. It is important to recognize, however, that individual strain gages cannot be calibrated. If calibration and traceability to a standard are required, strain gages should not be employed.

4.4 To be used, strain gages must be bonded to a structure. Good results depend heavily on the materials used to clean the bonding surface, to bond the gage, and to provide a protective coating. Skill of the installer is another major factor in success. Finally, instrumentation systems must be carefully designed to assure that they do not unduly degrade the performance of the gages. In many cases, it is impossible to achieve this goal. If so, allowance must be made when considering accuracy of data. Test conditions can, in some instances, be so severe that error signals from strain gage systems far exceed those from the structural deformations to be measured. Great care must be exercised in documenting magnitudes of error signals so that realistic values can be placed on associated uncertainties.

5. Interferences

5.1 In order to assure that strain gage test data are within a defined accuracy, the gages must be properly bonded and protected with acceptable materials. It is normally simple to ascertain that strain gages are not performing properly. The most common symptom is instability with time or temperature change. If strain gages do not return to their zero reading when the original conditions are repeated, or there is low or changing resistance to ground, the installation is suspect. Aids in strain gage installation and verification thereof can be found in Guide E 1237.

6. Hazards

6.1 In the specimen surface cleaning, gage bonding, and protection steps of strain gage installation, hazardous chemicals may be used. Users of these test methods are responsible for contacting manufacturers of these chemicals for applicable Material Safety Data Sheets and to adhere to the required precautions.

7. Test Requirements

7.1 General Environmental Requirements:

7.1.1 Ambient Conditions at Room Temperature—The nominal temperature and relative humidity shall be 23° C (73° F) and 50 %, respectively. In no case shall the temperature be less that 18° C (64° F) nor greater than 25° C (77° F) and the relative humidity less than 35 % nor more than 60 %. The fluctuations during any room temperature test of any gage shall not exceed \pm 2°C and \pm 5 % RH.

7.1.2 Ambient Conditions at Elevated and Lower Temperatures—The temperature adjustment error shall not exceed $\pm 2^{\circ}C$ ($\pm 3.6^{\circ}F$) or ± 2 % of the deviation from room temperature, whichever is greater. The total uncertainty of temperature shall not exceed $\pm 2^{\circ}C$ ($\pm 3.6^{\circ}F$), or ± 1 % of the deviation from room temperature, whichever is greater. At



FIG. 2 Wheatstone-Bridge Circuit

elevated temperatures the mixing ratio shall be constant, that means independent of temperature, at a nominal value of 0.009 g of water per 1 g of air at a pressure of 1 bar. This value corresponds to a relative humidity of 50 % at 23° C (73°F).

Note 1—This mixing ratio, independent of temperature, can be realized by a furnace that is well connected to an atmosphere meeting the conditions of 7.1.1.

7.2 Test Measurement Requirements:

7.2.1 Several methods are available for measuring the change of gage resistance with sufficient resolution and accuracy. In general, any of these methods that are convenient may be used after it has been shown that the particular combination of instruments or components used produce a system with the required accuracy.

7.2.2 Examples of potentially satisfactory methods are as follows:

7.2.2.1 Balanced Bridge Circuit—In this circuit, a change in gage resistance is matched by an equal unit resistance change in a calibrated arm of the bridge circuit so as to produce a balanced condition with zero electrical output. This circuit is not sensitive to excitation voltage changes except for self-heating effects. A sensitive null detector (galvanometer) is required to obtain adequate resolution. Direct-current excitation is usually, but not necessarily, used. Thermal emfs generated within the circuit and reactive changes in the circuit may cause errors. This circuit is shown in Fig. 2.

7.2.2.2 Unbalanced Bridge Circuit—This circuit is similar to the Wheatstone bridge except that the bridge components are not adjusted after a nearly balanced initial condition is obtained. The output voltage of an unbalanced bridge circuit in which one arm is varying, E_o , is given by the equation:

$$E_o = E_i [\Delta R / (4R_o + 2\Delta R)] \tag{4}$$

where:

 E_i = input voltage,

 R_o = resistance required for initial bridge balance, and

 ΔR = difference between the instantaneous resistance and R_o .

This circuit is readily adaptable to automatic recording of data. Either ac or dc excitation may be used, but errors due to thermal emfs and reactive changes are possible. Loading effects due to the impedance of the recording instruments may be significant and must be considered. To avoid the necessity of accurate absolute measurement of the input and output voltages, the readout (recording) system may be calibrated in terms of unit resistance change of a bridge arm by use of a calibrating



resistor that can be varied so that the total arm resistance changes in accurately known steps. This resistor should be in the opposite arm of the bridge circuit from the gage. This circuit is shown in Fig. 3.

7.2.2.3 Several types of instruments are available for obtaining strain data directly from a resistance strain gage. These instruments use various types of excitation and read-out systems. Such indicators may be used only after their resolution, accuracy, and stability have been verified by connecting a resistor that can be varied in accurately known increments in place of the gage and calibrating the strain indicator over the entire range for which it will be used. The calibrating resistor steps shall be accurate to 0.1 % of the resistance change or 2 ppm of the total resistance, whichever is greater. The effects of the following factors should be determined: thermal emf's within the bridge circuit and within the leads to the gage; reactive changes within the bridge and lead circuits; initial bridge unbalance; and, battery conditions or power line fluctuations.

7.3 Strain Gage Attachment:

7.3.1 The attachment conditions shall correspond exactly to the instructions published by the gage manufacturer.

8. Test Method for Determining Strain Gage Resistance at Ambient Conditions

8.1 The standard 23° C (73° F) temperature resistance of each unbonded strain gage shall be measured and stated. Alternatively, strain gages may be combined in sets (4, 5, or 10, for example) from the same batch that have close resistance values. All gages combined in sets shall fall within the stated nominal resistance value and uncertainty from all sources.

8.2 The unpackaged strain gages selected for testing should be stored under the ambient conditions described in 7.1.1 for at least 72 h before and during resistance measurement.

8.3 The uncertainty of the strain gage resistance measurement shall be less than \pm 0.1 %. Repeated measurements shall have a range no greater than \pm 0.04 % of the measured value. The influence of the measuring current on the strain gage shall not be greater than \pm 0.1 % of the resistance value.

8.4 For the resistance measurement no particular mechanical requirements are necessary. However, if the influence of the flatness of the strain gage on the resistance measurement exceeds \pm 0.1 % of the actual value, the gage must be held in contact with a substantially flat surface using a suitable pressing device. Care must be exercised to assure that the probes used to contact the tabs of gages without leads do not damage foil areas.



FIG. 4 Constant Bending-Moment Beam Method for Gage-Factor Determination

9. Test Methods for Determining the Gage Factor of Resistance Strain Gages at a Reference Temperature

9.1 These test methods describe procedures for the determination of the gage factor of bonded resistance strain gages. It is suggested that gage factor values be obtained for at least five gage installations of one type.

9.2 For gage factor determination, the uncertainty of the relative resistance change measurement shall not exceed ± 2 µohm/ohm or ± 0.1 % of the actual value, whichever is greater. Any of the test methods described in Section 7 may be used. In addition, special circuits designed to compare the gages being tested to a calibrated reference gage may be used if it is shown that equal accuracy is obtained.

9.3 Determination of the gage factor K requires mechanical equipment consisting of a test specimen and a loading device capable of producing a uniform uniaxial stress in the test specimen corresponding to nominal mean principal strain values of 0, \pm 1000 and \pm 1100 µm/m (µin./in.). The Poisson's ratio of the test specimen shall be 0.28 ± 0.01 or suitable corrections must be made. The mean principal strain shall be within \pm 50 µm/m (µin./in.) of the nominal value. The strain at the various gage stations shall differ by no more than \pm 0.5 % of the mean value and the strain within a gage station shall vary by no more than \pm 0.5 % of the nominal value. The uncertainty of the mean strain measurement shall be less than \pm 2 µm/m (µin./in.) or \pm 0.2 % of the actual value, whichever is greater. Any test apparatus that meets these criteria may be used for determination of gage factor.

9.4 To the extent possible, test specimens with attached strain gages for tests of the gage factor should be stored under the ambient conditions described in 7.1.1 for at least 72 h before being tested

9.5 For the determination of the gage factor, the strain gages under test should be prestrained three times with strain cycles similar to the ones used for the measurement, but with maximum strain levels about 10 % higher. That means that the loading cycle should nominally be:

- 0, + 1100 μ m/m (µin./in.), 1100 μ m/m (µin./in.),
 - + 1100 µm/m (µin./in.), 1100 µm/m (µin./in.),
 - + 1100 μm/m (μin./in.), 1100 μm/m (μin./in.),
- 0, + 1000 μ m/m (µin./in.), 0, 1000 μ m/m (µin./in.), 0. (5)

If possible, one half of the sample (group of gages to be tested) should be strained this way and the other half of the sample should be subjected to strains of the same magnitude but opposite sign. The gage factor is determined from the slope of the straight line between the measurement points at + 1000 μ m/m (µin./in.) and - 1000 µm/m (µin./in.). Although less desirable, it is permissible to use the strain cycles of:

for one half of the sample and strain cycles of:

0, - 1100 μm/m (μin./in.), 0, - 1100 μm/m (μin./in.)

 $0, -1100 \ \mu m/m \ (\mu in./in.), 0, -1000 \ \mu m/m \ (\mu in./in.), 0$ (7)

for the other half of the sample.

The gage factor is determined from the average of the slopes, of the straight lines between the measurement points at 0 and $+ 1000 \,\mu\text{m/m}$ (µin./in.) and 0 and $- 1000 \,\mu\text{m/m}$, (µin./in.).

9.6 As a guide, three separate test methods are described, the choice of the test method used being determined by the particular application and by the facilities that are available. These test methods do not classify strain gages according to accuracy or other performance characteristics. The three test methods that are described differ primarily in the manner of producing an accurately known surface strain, and they are thereby classified. These test methods are described in the following sections:

9.6.1 Constant Bending Moment Beam Test Method:

9.6.1.1 *Summary of Test Method*—This test method utilizes a strain on the surface of a test bar produced by loading it as a constant moment beam by the application of dead-weight loads.

9.6.1.2 *Mechanical System*—A typical mechanical system is shown in Fig. 4. The test beam may be of any suitable material that meets the requirements of 9.3, and shall have minimum dimensions of 19 by 25 by 760 mm (0.75 by 1 by 30 in.). The minimum distance between the pivot points on the supports shall be 2.45 m (96 in.). The beam assembly shall be symmetrical about a vertical line through its midpoint. The positions of the pivots and the weight values shall be adjusted to provide the required strains. The strain over the usable section of the beam shall vary by not more than 1 % of the strain at the reference point. The usable portion of the beam shall be at least one half of the exposed length.



FIG. 5 Constant-Stress Cantilever Beam Method for Gage-Factor Determination

9.6.1.3 Verification-The need for measuring calibration strain directly during each test is eliminated by maintaining a calibration of the system. Such a calibration is made by measuring with a Class A extensometer (see Practice E 83) the actual strain produced on the surface of the beam when it is loaded. Measurements shall be made with the extensometer centered over each station of the beam. At least three measurements shall be made at each station to verify the strain distribution over the width of the beam. The dimensions of the beam shall be checked at each station periodically. A change of 0.2 % in the thickness at any station shall disqualify that station. Other dimensional changes that would cause a change of surface strain of 0.2 % shall disqualify the beam. The strain at the reference station shall be determined each time the beam is used either with a Class A extensometer, or with a carefully selected, permanently mounted resistance strain gage that has been calibrated by spanning with a Class A extensometer. The response of this reference gage shall be verified periodically to assure compliance with specifications using a Class A extensometer. The beam shall be completely recalibrated after 50 applications or 6 months, whichever comes last.

9.6.1.4 *Procedures*—Mount test gages with any appropriate installation technique that will not change the characteristics of the test beam (for example, excessive cure temperatures could be damaging). Mount the gages at the stations on the beam where the strain level has been determined by the calibration procedure outlined in 9.6.1.3.

9.6.1.5 Install the test specimen bearing previously unstrained gages in the loading system and test environment. After temperature equilibrium has been attained, follow the loading sequence of 9.5. Take readings from the strain gages before applying the load, with the load applied, and after the load is removed for each loading cycle. Obtain compression loads by mounting the beam with the gaged surface up. Obtain tension loads by mounting the beam with the gaged surface down.

9.6.1.6 Calculate the gage factors.

9.6.2 Constant Stress Cantilever Beam Test Method:

9.6.2.1 *Summary of Test Method*—This test method produces strain on the surface of a cantilever beam that is designed to have a constant stress over the major portion of its length when loaded in the prescribed manner.

9.6.2.2 Mechanical System—A typical mechanical system is shown in Fig. 5 and detailed design of a beam that has been used satisfactorily is shown in Fig. 6 (Note 2). The size and arrangement of the equipment must be such that the beam may be bent sufficiently in either direction to produce a surface strain of at least 1100 µm/m (µin./in.). Two or more carefully selected strain gages, for use as reference standards, shall be permanently bonded to the constant-stress section of the beam as shown in Fig. 6. Great care must be taken to install these gages, using the best current techniques to ensure bonding integrity and long-time stability. These references gages shall be individually calibrated to determine their gage factor by placing a Class A extensometer (Practice E 83) so as to span the gage, bending the beam by means of the deflecting apparatus, and measuring the resulting change in gage resistance and strain. Readings shall be taken for the strain cycles stipulated in 9.5 and the gage factor computed (Note 3 and Note 4).

NOTE 2—In order for the beam to fulfill the requirements of a constant-stress beam, the drive rod must be attached to the beam at the apex of the angle formed by the sides of the beam. The ratio of the free length of the beam to width at the base should not be less than 9.1.

NOTE 3—For the reference gage, the gage factor for compression strains may differ from the gage factor for tension strains and it must be determined for both directions of loading.

NOTE 4—It may be convenient to obtain strain of the beam surface as a function of the deflection of the end of the beam as measured by a dial gage while the strain gages are being calibrated.

9.6.2.3 Verification of Beam—The constant-stress area of the beam shall be explored with a Class A extensometer to determine the area where the strain is the same as that experienced by the reference gages. The gage length of the extensometer shall not exceed 25 mm (1 in.). Only areas of the beam where differences between the strains indicated by the extensometer and the reference gage do not exceed 10 μ m/m (μ in./in.) at a strain of 1000 μ m/m (μ in./in.) are acceptable for testing gages. The beam shall be verified after each 50 uses or 6 months, whichever comes last.

9.6.2.4 *Procedure*—Install the gages to be tested on the beam in the areas that have been found to be satisfactory; connect them to instruments for measuring their change of resistance. The active axes of the gages shall be parallel to the



FIG. 6 Constant Stress Cantilever Beam

center line of the beam. A selector switch may be used to connect several gages into the measuring circuits if it is shown that repeated switchings do not change indicated strain readings by more than 2 μ m/m (μ in./in.).

9.6.2.5 Follow the loading schedule of 9.5 and calculate gage factors.

9.6.3 Direct Tension or Compression Test Method:

9.6.3.1 *Summary of Test Method*—This test method produces strain in a test bar by applying direct tensile or compressive loads to the bar.

9.6.3.2 *Mechanical System*—A typical mechanical system is shown in Fig. 7. In this system the test bar is strained directly in tension or compression by a testing machine or other device capable of applying an axial load to the specimen. The horizontal position of the bar is convenient for mounting the reference extensometer, but it is not necessary. The load may be applied by hydraulic, mechanical, or other means, but care must be taken to prevent any twisting or bending of the bar. Twisting in the mechanical system of Fig. 7 is prevented by the torque arm. Fig. 8 shows a test bar that has been used

successfully for both tension and compression loading. The strain gage under test shall be mounted at the center of the reduced section; and a Class A extensometer shall be mounted so as to span the gage. The extensometer should have a gage length as near that of the gage as possible in order to minimize the effect of nonuniform strain along the length of the bar.

9.6.3.3 Verification—Since the calibration strain is measured during each test, no calibration of the system is necessary. The thickness and width of the test bar must be uniform within ± 0.25 % of their average values over a length extending 13 mm (0.5 in) beyond the extensometer gage points in each direction. The absence of twisting and bending of the test bar must be verified.

9.6.3.4 *Procedure*—Mount a test gage by any appropriate technique so that the center of its sensitive portion coincides with the center line of the bar. Mount the bar in the loading device taking care to avoid bending or loading of the bar. Connect the gage electrically to the resistance-measuring circuit, and mount the reference extensometer so as to span the gage. Follow the loading cycle in 9.5 (plus or minus strains

E 251 - 92 (2003)



FIG. 7 Testing Machine for Gage-Factor Measurements



FIG. 8 Test Bar for Gage Factor Test

only) except that preload, not exceeding 5 % of the maximum load, may be applied to align the bar in the machine, to remove backlash, etc. Take readings simultaneously from the electrical circuit and the extensometer. Calculate gage factors. Repeat for strains in the opposite direction.

10. Test Methods for Determining the Temperature **Coefficient of Gage Factor of Resistance Strain Gages**

10.1 These test methods describe procedures for the determination of temperature coefficient of gage factors of bonded resistance strain gages.

10.2 For temperature coefficient of gage factor determination, the uncertainty of the relative resistance change measurement shall not exceed \pm 5 µohm/ohm or \pm 0.1 % of the actual value, whichever is greater.

10.3 If convenient, strain gages may be tested in tension/ compression half bridges (one gage in tension, the other in compression) by mounting two gages opposite each other and connecting them in a half bridge. This practice helps to eliminate errors from drift and leadwires. If gages are tested individually, a three-lead wiring arrangement is used (see Fig. 2 and Fig. 3).

10.4 To determine the temperature coefficient of gage factor, it is necessary to have equipment consisting of a test specimen, a loading device, and a furnace for producing the temperatures needed. It must be possible to adjust the strain in the specimen to mean values of 0 and + 1000 μ m/m (μ in./in.). It is desirable that a strain of $-1000 \ \mu m/m$ ($\mu in./in.$) may be produced.

Instead of the reference strain of zero, a small prestrain of between 20 and 100 µm/m (µin./in.) may be used. The adjustment error shall be no more than \pm 50 µm/m (µin./in.). The uncertainty of the mean strain should be less than \pm 5 µm/m (µin./in.). The strain at the various gage stations shall differ by no more than ± 2 % of the actual strain and the strain within a gage station shall vary by no more than ± 2 % of the nominal value.

10.5 Two test methods for determining the temperature coefficient of gage factor of bonded resistance strain gages are given, a static method and a dynamic method. The choice of test method will be determined by the temperature range, ultimate user needs, and the number of tests to be conducted. The two test methods differ in the manner in which the strain is produced, one test method making use of measurements made under static strain and static temperature conditions, and the other test method making use of measurements made under dynamic strain and transient temperature conditions.

10.5.1 Static Test Method:

10.5.1.1 Summary of Test Method-This test method⁴ utilizes a constant-stress cantilever beam that is forcibly deflected in a series of fixed, accumulative steps that can be accurately repeated at various temperatures of interest.

10.5.1.2 Typical equipment used to produce the strain and a typical test beam are shown in Fig. 9. The beam is designed to have a considerable area of uniform stress that is directly proportional to the deflection of the end point (the apex of the angle formed by the sides of the beam) of the beam. The frame is designed to hold the base of the beam rigidly and provide a base for the sliding-stepped block. The rider on the beam is attached at the apex of the angle formed by the beam sides. The frame must be much more rigid than the beam to prevent errors due to bending of the frame. The stepped block can provide several deflection steps, as shown in Fig. 9. However, it is sufficient that the maximum deflection produces a surface

⁴ This test method is based on apparatus and techniques proposed by McClintock, R.M., "Strain Gage Calibration Device for Extreme Temperatures," Review of Scientific Instruments, Vol 30, No. 8, 1959, p. 715.



FIG. 9 Apparatus for Static Determination of Gage-Factor Variation Versus Temperature

strain on the beam of $1000 \pm 50 \,\mu\text{m/m}$ (µin./in.). The stepped surfaces must be parallel to each other and to the opposite sliding surface of the block. The apparatus must be designed so the beam end is deflected about 2 % of its total planned deflection when the rider is in contact with the lowest step of the sliding block. This is to ensure that contact is always maintained between the beam and the rider. To avoid differential expansion problems, all parts of the test rig, and the specimen, should be made from the same material, selected to assert proper operation over the entire temperature span to be encountered.

10.5.1.3 A furnace or cryostat capable of producing the desired temperature conditions is required but not shown.

10.5.1.4 Mount the gage or gages to be tested on the beam so they are symmetrically centered on the constant-stress area and aligned with the longitudinal center line of the beam. Mount temperature sensors as near the gage(s) as practicable and at each end of the constant-stress area. Mount the beam in the frame, and connect the gages electrically to the read-out instruments.

10.5.1.5 With the loading apparatus in the furnace or cryostat and the gage connected to its read-out instrumentation, allow the beam to come to temperature equilibrium at the reference temperature (usually room temperature). With the rider resting on the lowest step of the block, take a measurement of the gage output. Then move the sliding block so as to increase the beam deflection and take gage output readings at each step. Again take readings as the deflection is decreased in steps. Repeat this procedure to obtain three sets of readings. Take the gage output due to strain for each step as the average of the differences from the value at the lowest step for all loading cycles.

10.5.1.6 Bring the temperature of the test fixture and beam to each of the preselected temperatures of interest and repeat the procedure. Take care to ensure that the temperature has

stabilized. Make tests at a minimum of five nearly equally spaced temperatures over the temperature range of interest, compute the temperature coefficient of gage factor (see section 3.12).

10.5.2 Dynamic Test Method:

10.5.2.1 *Summary of Test Method*—This test method depends upon the output voltage from a bridge circuit composed of stable resistors and one or more resistance strain gages:

$$E_0 \approx E_1 \, K(N/4)\epsilon \tag{8}$$

where:

 E_0 = output voltage from bridge circuit,

 E_1 = input voltage to bridge circuit,

$$K =$$
 gage factor of the gages,

 ϵ = strain to which the gages are subjected, and

N = number of active gages.

If such a bridge circuit is connected to a constant d-c voltage source and the gages are subjected to a sinusoidal strain of constant amplitude, the change in the alternating output voltage will be a measure of the change of gage factor.

10.5.2.2 This test method requires a means of vibrating a constant-stress cantilever beam at a constant amplitude; varying the temperature of the beam at a nearly uniform rate; and measuring the output voltage, or change of output voltage, of the bridge circuit as a function of temperature. These operations must be done simultaneously.

10.5.2.3 The beam vibration may be conveniently produced by a motor-driven cam or by an electromechanical vibrator. If the vibrator is used, a method of maintaining the amplitude of vibration constant is required. Monitoring the vibration amplitude by means of a velocity sensing pick-up may not be satisfactory because of changes in the vibration frequency.

10.5.2.4 The temperature environment is conveniently produced by radient heaters of the tungsten filament quartz tube type. Power may be supplied to these heaters by a temperature



FIG. 10 Dynamic Apparatus for Determining Variation of Gage Factor

programming unit or by manual control with an autotransformer. In order to maintain a nearly uniform temperature over the length of the beam, supplemental heat must be supplied to the clamped end of the beam. This may be done by resistancewire heating elements built into the clamping fixture.

10.5.2.5 Equipment for producing the vibratory motion, by means of a cam, and temperature environment is shown in Fig. 10. The control units for the heating elements are not shown. Care must be taken in the design of the apparatus to prevent changes in the rigidity of the beam support and clamping with time or temperature. The design of the beam is shown in Fig. 11.

10.5.2.6 Measuring the ac output of the strain gage circuit and obtaining changes by taking differences of measured values will not usually be satisfactory because of the small differences of large values involved. However, the change of ac voltage may be measured directly by use of circuits such as those shown in Fig. 12 and Fig. 13. The input circuit, Fig. 12, provides a selected constant voltage of 4 to 12 V to the gage circuit, and also provides means for varying this input voltage over a range of \pm 10 % of the nominal value in known steps. After the ac output voltage from the gage circuit has been amplified to about 5 V and filtered to remove all signals except that of the vibration frequency, it becomes the input signal to the output circuit, Fig. 13. The signal is rectified, filtered to remove ripple, and suppressed by a bucking voltage from a stable dc voltage source. The difference between the rectified signal and the suppressing voltage is recorded as a function of test-beam temperature. The dc voltage input to the gage circuit must be constant during the test.

10.5.2.7 Mount two resistance strain gages on opposite sides of the constant-stress cantilever beam as shown in Fig. 11. Clamp the wide end of the beam firmly to the rigid mount, and connect the narrow end to equipment for producing sinusoidal deflections of constant amplitude. Make the connection to this equipment at the apex of the angle made by the sides of the main portion of the beam. Connect the gages as

adjacent arms of a bridge circuit, the other arms being stable resistors of approximately the same resistance as the gages and chosen so that the bridge circuit is nearly balanced when the beam is in a neutral position. With the input terminals of the bridge circuit connected to a constant-voltage source, vibrate the beam to produce a strain of about \pm 500 µm/m (µin./in.). Adjust the suppressing voltage to give zero output to the recorder. Obtain the recorder sensitivity in terms of change of gage-circuit in known steps. The change of output voltage due to a change in input voltage is the same as would be caused by the same percentage change of gage factor, the strain amplitude remaining constant.

10.5.2.8 After the recorder sensitivity has been determined, return the input voltage to its nominal value, and increase the temperature of the beam at a uniform rate to the maximum desired temperature. During this time record the difference between the rectified gage circuit output and the suppressing voltage as a function of the test beam temperature. Obtain the temperature from a temperature sensor mounted as near the gage installation as practicable. A heating rate of 10°C (20°F)/ min has been used satisfactorily. During the test keep the temperature gradient over the area of the beam near the gages small. Measure this temperature gradient by the difference between two temperature sensors, one mounted near the clamping fixture and the other mounted an equal distance the other side of the gage installation. The temperature difference between these points should not exceed $3^{\circ}C$ ($5^{\circ}F$) or 1 % of the beam temperature. The power to the heaters in the clamping fixture can be adjusted to minimize this temperature difference. Calculate the temperature coefficient of gage factor.

10.5.2.9 Since both the length and thickness of the beam change with temperature, the recorded output should be corrected for the resulting change in strain. Applying this correction gives:

Gage factor change,
$$\% = \Delta E_0 / (1t\alpha_b t)$$
 (9)



FIG. 11 Beam Used for Gage Factor Variation Test



where:

- ΔE_0 = recorded change in gage circuit output, percent,
- $\alpha_b = \text{coefficient of linear expansion of the beam material.}$ (See Test Methods E 228 and E 289 for methods of determining α_b), and
- Δt = difference between the instantaneous beam temperature and the initial temperature.

11. Test Method for Determining the Transverse Sensitivity of Resistance Strain Gages at a Reference Temperature

11.1 Strain gage transverse sensitivity results in an undesired signal induced by strains along directions other than the one to be measured. In two-dimensional stress analysis the errors resulting from transverse sensitivity will depend upon the stress distribution in gaged areas, and for accurate results, strain-gage readings should be corrected for this effect.

11.2 For transverse sensitivity determination, the uncertainty of the relative resistance change measurements shall not exceed \pm 5 µohm/ohm or \pm 0.1 % of the actual value, whichever is greater.

11.3 To determine the transverse sensitivity; mechanical equipment is required for producing uniform uniaxial strains of 0, 1000, and 1100 μ m/m (μ in./in.) in the longitudinal direction of the specimen. The adjustment error shall be less than \pm 50 μ m/m (μ in./in.). The strain at the various gage stations shall differ by no more than \pm 2% of the actual strain in the longitudinal direction of the specimen and no more than \pm 5 μ m/m (μ in./in.) in the transverse direction. The uncertainty of the strain shall be no more than \pm 10 μ m/m (μ in./in.) in the



longitudinal direction of the specimen and $\pm 1 \,\mu\text{m/m}$ (µin./in.) in the transverse direction.

11.4 This test method describes a procedure for determining transverse sensitivity of bonded resistance strain gages including those with wire and foil strain-sensitive elements. It involves mounting gages on a test specimen that can be loaded so as to produce a nearly uniaxial strain field. The transverse sensitivity is determined from the relative response of nearly identical gages mounted parallel to a perpendicular to the direction of principal strain.

11.5 Apparatus:

11.5.1 *Electrical Measurements*—Any of the test methods described in Section 7 may be used.

11.5.2 *Mechanical System*—Typical equipment used to produce the strain and a suggested test beam is shown in Fig. 14. The control gage may be either a Class-A extensometer or permanently installed and waterproofed bonded resistance strain gages that are temperature compensated for the beam material and have been calibrated in place with a Class-A extensometer.

11.5.3 Verification—The test beam shall be lightly scribed to define 16 stations as shown in Fig. 15. The difference between the strain measured by the control gage and the actual strain at each of these stations, both parallel and perpendicular to the direction of principal strain, shall be determined. The strains at the test station shall be measured with a Class-A extensometer centered over the station. The reference strain may be measured with a Class-A extensometer or with the previously calibrated control gage of the bonded resistance type. The strain perpendicular to the direction of maximum principal strain must be less than 4 μ m/m (μ in./in.) or 0.5 % of the maximum principal strain with a maximum principal strain of $1000 \pm 50 \,\mu\text{m/m}$ (µin./in.). A record of the station-by-station calibration must be maintained, and data must be corrected to within 5 µm/m (µin./in.). The behavior of the beam surface shall be verified after 50 tests or 6 months, whichever comes last.

NOTE 5—Tests are usually made for tension loading only, using the upper surface of the beam. If compression tests are desired utilizing the

lower surface of the beam, station-to-station calibration must also be obtained for this surface.

11.6 Procedure:

11.6.1 A test requires a minimum of five strain gages of one type. Install these gages on the test beam in a manner similar to that shown in Fig. 15, using manufacturer recommended installation techniques. Mount at least three gages perpendicular to the direction of maximum principal strain and at least two gages parallel to the direction of maximum principal strain.

11.6.2 For the determination of the transverse sensitivity, the strain gages under test should be prestrained three times to a maximum transverse strain of 1100 μ m/m (μ in./in.) and then the test strain of 1000 μ m/m (μ in./in.) should be applied. The transverse sensitivity must be determined from the slope of the straight line between the measurement points at 0 and 1000 μ m/m (μ in./in.).

11.6.3 Compute the transverse sensitivity of each of the gages mounted perpendicular to the direction of principal strain as follows:

transverse sensitivity =
$$[(\Delta R_t / R_{to}) / (\Delta R_l / R_{lo})] \times 100$$
 (10)

where:

- $\Delta R_{f}/R_{io}$ = unit change of resistance of the transverse gage, and
- $\Delta R_{l}/R_{lo}$ = average unit change of resistance of the gages mounted parallel to the direction of principal strain.

11.6.4 Report the transverse sensitivity of the gage type as the average of all values obtained. Also report the range of values obtained.

12. Test Method for Determining the Thermal Output of Bonded Resistance Strain Gages

12.1 *Scope*:

12.1.1 The thermal output of a strain gage installed on a given material may be expressed in at least three ways:

12.1.1.1 A complete record of the unit change of gage resistance as a function of temperature,



FIG. 14 Transverse-Sensitivity Test Rig



FIG. 15 Testing Stations and Gage Arrangement for Transverse Sensitivity Test

12.1.1.2 The unit change of gage resistance between two given temperatures, a chord function, and

12.1.1.3 The slope of a tangent to the unit change of resistance-versus-temperature curve at a given temperature.

12.1.2 The values of 12.1.1.2 and 12.1.1.3 can be obtained from 12.1.1.1. The test method described is primarily useful when a temperature range greater than 50°C (90°F) is involved. This test method requires heating uniformly a small test strip with a strain gage installed upon it while the unit change of gage resistance is recorded as a function of test strip temperature. It depends upon the test strip being able to expand freely with temperature changes; upon the test strip not being distorted or otherwise affected by external loads, residual stresses, etc; upon the temperature of the test strip being uniform at any given time; and upon changes of resistance of the gage due to other factors being insignificant during the test.

12.2 For thermal output determination, the uncertainty of the relative resistance change measurements shall not exceed \pm 5 µohm/ohm or \pm 0.1 % of the actual value, whichever is greater.

12.3 For determination of the thermal output (1.3.5), a test specimen is required. It must be properly heat-treated and free of residual stresses. Its dimensions, especially the thickness, must be sufficient to prevent any distortion and thermal hysteresis greater than \pm 5 µm/m (\pm µin./in.) for any temperature difference of 100°C (180°F). This applies to the whole temperature range for which the thermal output has to be determined. If this temperature range is less than 100°C (180°F), the requirements applied should be those for a minimum of 100°C (180°F). The thermal expansion of the specimen must be known over the full temperature range with an uncertainty of less than \pm 0.2 µm/m/°C (\pm 0.1 µin./in./°F). If necessary, the thermal expansion must be determined using a dilatometer and the specimen material.

Note 6—A test strip 25 by 150 by 1.3 mm (1 by 6 by 0.050 in.) has been used. The test strip may be of any suitable material.

12.4 Apparatus:

12.4.1 *Electrical Measurements*—The unit change of gage resistance is conveniently recorded as a function of temperature by the circuit shown in Fig. 3. The output voltage from the bridge circuit is recorded on the y-axis of a sensitive X-Y



FIG. 16 Apparatus for Determining Temperature Sensitivity

recorder, the x-axis of which records the temperature indicated by a thermocouple mounted on the test strip as near the gage as possible.

12.4.2 *Thermal System*—A system that has been used to produce satisfactory thermal environments at temperatures up to 800°C (1475°C) is shown in Fig. 16. The heating elements are eight tungsten filament quartz tube radiant-heat lamps rated at 1000W each. The test strip is supported at the center of the heater configuration by porcelain tubes that also serve to carry electrical leads and thermocouples to the gaged area. Air from the cooling coils surrounding the reflecting cylinder prevents the reflecting surface from overheating during a test series.

12.5 Procedure:

12.5.1 Install the strain gage at the center of the test strip using manufacturer recommended installation procedures. Attach the test strip to its holder and mount in the furnace. Connect the leads from the gage and thermocouples to the measuring circuit and recorder. Calibrate the recorder by varying the calibrating resistor in at least five nearly equal increments that cover the expected gage resistance change and noting the recorder response at each step. With the bridge circuit again balanced, supply power to the heaters.

12.5.2 For the determination of the thermal output, the test specimens with the attached strain gages should be heated to the upper end of the temperature range. As the temperature decreases, in steps or continuously but eventually with sufficient heat balance and without perceptible oxidation, the strain indication versus temperature is measured. At least five measurements should be taken over the temperature range:

12.5.2.1 Between 0 and 100°C (32 and 212°F), at temperature intervals of about 20°C (35°F) with a maximum of 30°C (55°F) (roughly 5 measurements per 100°C (180°F)),

12.5.2.2 Above 100°C (212°F) and below 0°C (32°F), at temperature intervals of about 25°C (45°F) with a maximum of 40°C (75°F) (at least 5 measurements per 200°C (350°F)). More measurements should be made if the thermal output changes rapidly.

12.5.3 The test specimen material shall be particularly specified; the linear coefficient of expansion and the nonlinear part of the thermal expansion must be known. If the strain

gages are of the self-compensated type, the thermal coefficient of expansion for which the strain gages are compensated shall be specified.

13. Report

13.1 Data Reduction and Statistics—Since strain gages used to determine values of performance characteristics are generally not reusable, the data obtained from a sample of such gages are used to predict values of these characteristics for all other gages of the same batch. Well established statistical methods can be used to make such predictions of values and the uncertainties associated with those values. These methods require that tests be made upon a sample taken at random from the batch, and generally assume that the test results will have a normal, that is, Gaussian, distribution. For purposes of verification of reported values, it will be assumed that these reported values are equivalent to those obtained from a sample of thirty gages and that systematic errors in these reported values are negligible or have been corrected for. A numerical example is given in Appendix X3.

13.1.1 For each characteristic to be verified, a sample of strain gages shall be tested using the procedures described. The control sample shall consist of at least five strain gages although a sample of ten or more gages would be preferred. The results of the several tests made on one gage shall be averaged, and these average values shall be used in the following computations to determine if the average performance or variability of the control sample differs from the reported values at the 95 percent confidence level.

13.1.2 A two-sided t-test is used to compare the manufacturer's statement of the value for a particular characteristic, V_R , shown on the gage package, and the average value from the control tests, V_C . The standard deviation of the reported value, S_R , is taken as $T_R/2$, where T_R is the tolerance given on the package. The standard deviation of the control value, S_C , is calculated from the control test data using the relationship:

$$S_{C} = \sqrt{\frac{\sum_{i=1}^{n} (X_{i} - \bar{X})^{2}}{n-1}}$$
(11)

TABLE 1	Values Used	to Compare	Reported a	and Control	Test
Valu	es of Average	Gage Perfo	rmance and	d Variability	

Number of gages in control sample,	Factor for comparing average values, ^A	Values for comparing variability, ^B	
п	t	А	В
5	2.036	0.306	8.47
10	2.025	0.386	3.57
15	2.018	0.424	2.74
20	2.013	0.448	2.40
30	2.002	0.476	2.10

^AObtained by interpolation for *t* (.975) from a table of percentiles of the *t* distribution assuming that S_R is based upon a sample of 30.

^BObtained by interpolation from a table of percentiles of the *F* distribution for *F* (.975) assuming that S_R is based upon a sample of 30.

where:

 X_i = the average value for i^{th} gage,

 \vec{X} = the average value for all gages in the control sample, and

n = the number of gages in control sample.

The manufacturer's reported value, V_R , is considered to be verified if:

$$|V_{R} - V_{C}| \le t - \sqrt{\left[\frac{29S_{R}^{2} + (N-1)S_{C}^{2}}{28+n}\right] \times \left[\frac{3+n}{3n}\right]} + e_{s}$$
(12)

where:

t = the value from Table 1 corresponding to n, and

 e_s = the estimate of systematic error in the control value, V_C .

13.1.3 An "equal tail" test is used to compare the variability reported with the gages and the variability of the control sample. The reported tolerance, T_R , shall be considered verified if:

$$A \le (S_R^{-2}/S_C^{-2}) \le B \tag{13}$$

where:

 $S_R = T_R/2,$

 S_C = the standard deviation of control sample, and

A and B = the values from Table 1 corresponding to the number of gages in the control sample, n.

13.1.4 An example is presented in Appendix X2.

13.2 Content of Report:

13.2.1 The report shall include the following:

13.2.1.1 Method of test used,

13.2.1.2 Statement of the accuracy of the apparatus used, including the latest date of verification and by whom verified,

13.2.1.3 The uncertainty from all sources, of the gage characteristics at a probability level of at least 95.5 % (that is to say, no less than two standard deviations).

13.2.1.4 Type of electrical circuit used, including the type of excitation (ac or dc, constant voltage or constant current), and the voltage across or current through the gage,

13.2.1.5 Temperature and humidity in the test area,

13.2.1.6 Number of gages included in the tests,

13.2.1.7 Complete description of the gages tested including, type, lot number, manufacturer, size, and method of attachment, including curing cycle,

13.2.1.8 Results from all gages tested, and

13.2.1.9 Average of all values of the characteristics determined and the uncertainties. Results may be presented in the form of a table or a graph (see Appendix X2 for samples). In the case of gage factor, state whether the values are for tension and compression load on each gage, tension on part of the sample and compression on the remainder, or tension or compression only.

13.3 Presentation of Gage Characteristics:

13.3.1 Supply the following information:

13.3.1.1 Gage type,

13.3.1.2 Lot number,

13.3.1.3 Batch number,

13.3.1.4 Gage resistance with uncertainty,

13.3.1.5 Gage factor with uncertainty,

13.3.1.6 Transverse sensitivity with uncertainty,

13.3.1.7 Temperature coefficient of gage factor, over recommended temperature range, expressed either graphically or numerically with uncertainty,

13.3.1.8 Thermal output versus temperature to be expressed either graphically or numerically with uncertainty. The thermal output data or graph must include a statement concerning whether the test was run at fixed stable temperature steps or with continuous but uniform temperature changes and recording of data.

14. Precision and Bias

14.1 *Precision*—No precision statement based on interlaboratory testing is possible at this time. Committee E-28 is not aware of sufficient laboratories to conduct an appropriate program of testing but seeks cooperation of any and all laboratories concerned with implementing these test methods. The Committee is willing to coordinate such an activity.

14.2 *Bias*—Since there are no reference or true values of strain gage performance characteristics, no bias estimates can be measured.



APPENDIXES

(Nonmandatory Information)

X1. GAGE DESIGNS (Reproduced from Reference Cited in 2.3)



FIG. X1.2 Typical Strain Gage Installation Displacement of Grid from Specimen Surface: H_d = H_a + H_m + ½ H_g



X2. PRESENTING THE INFORMATION

X2.1 A format for presenting the information either numerically or graphically, is given as follows:

> EXAMPLE: PACKAGE INFORMATION GAGE TYPE: RLX-25PB-8764 LOT: 36D91L BATCH: 6143

> > VALUE

PERFORMANCE CHARACTERISTIC	VALUE
1. Gage resistance at 24°C and 50 % RH	350 Ω \pm 0.2 %
2. Gage factor at 24°C and 50 % RH	2.05 ± 1 %
3. Transverse sensitivity at 24°C and 50 %	(-0.6 ± 0.2) %
RH	

 Temperature coefficient of gage factor (if expressed numerically)



^A See Fig. X2.1 for graphical presentation.



FIG. X2.1 Items 4 and 5 (Presented Graphically)

 $\begin{array}{l}(+0.9\pm 0.2)\ \%/100^{\circ}C^{A}\\ -45.1+3.32\ T-6.76\cdot10^{-2}\\ T^{2}\\ +3.20\cdot10^{-4}\ T^{3}\\ -2.86\cdot10^{-7}\ T^{4}\mu\text{m/m}\\ \pm\ 0.22\ (\mu\text{m/m})\cdot^{\circ}C^{-1A}\end{array}$

X2.2 The gage type designation used shall be unique in nature so that, from published literature, the following can be determined:

X2.2.1 strain gage geometry,

X2.2.2 type of strain sensing alloy,

X2.2.3 type of matrix material, if used,

X2.2.4 approximate matrix size,

X2.2.5 type of leads, terminals, solder dots or other optional features, if included, and

X2.2.6 the coefficient of expansion of the material for which the strain gage is to provide compensated minimum thermal output, if applicable.

X3. COMPARISON OF REPORTED AND CONTROL VALUES

🎢 E 251 – 92 (2003)

X3.1 Example—The following example is given to show how values and tolerances furnished with a shipment of strain gages can be compared to values obtained from a sample tested by a control laboratory to determine whether the reported values properly describe the supplied gages. The gage factor Kis the characteristic used in this example, but the process would be the same for other characteristics.

X3.1.1 Assume that the gage factor value furnished with the gages is 2.024 \pm 0.5 %, reported in accordance with these test methods.

X3.1.2 Select a sample of ten gages at random from the shipment.

X3.1.3 Test the gages of the sample using the procedures described in point four. Each gage is tested in both tension and compression. The test plan (Table X3.1) gives the same number of data points for each direction of loading.

X3.1.4 The results of the tests are tabulated as shown in Table X3.1. Note that the values given in this table were obtained using random number techniques and may not be representative of actual data from strain gages.

X3.1.5 An average gage factor K_i is calculated for each gage tested. Average values are used in the subsequent calculations since gage performance and not the effects of the test procedure are of interest. The average value is the best estimate of the gage factor of that particular gage. These average values are shown in Table X3.1.

X3.1.6 An average gage factor for the sample, K_C , is calculated:

$$K_{C} = \frac{1}{n} \sum_{i=1}^{n} K_{i}$$
 (X3.1)

This value is shown in Table X3.1.

X3.1.7 An estimate of the standard deviation for the sample, S_C , is calculated as:

$$S_{C} = \sqrt{\frac{\sum_{i=1}^{n} (K_{i} - K_{C})^{2}}{n-1}}$$
(X3.2)

This value is shown in Table X3.1.

X3.1.8 An estimate of the systematic error in K_c is made from the characteristics of instruments, test apparatus, test procedures, operators, etc. This estimate is influenced by the experience of the laboratory. Note that this does not include systematic effects for which corrections have been made, for example, instrument calibration corrections. For this example, the systematic error *e*, is estimated to be 0.05 %.

X3.1.9 The gage factor value, K_C , obtained from (1), is compared to the value furnished with the gages, K_R , using the equation:

$$|K_{R} - K_{C}| \le t \quad \sqrt{\left[\frac{29 S_{R}^{2} + (n-1) S_{C}^{2}}{28 + n}\right] \left[\frac{30 + n}{30 n}\right]} + e_{s}$$
(X3.3)

Using values from Table X3.1 in this equation gives:

$$2.024 - 2.020 \le 2.025$$

$$\sqrt{\left[\frac{29 \times 0.0051^2 + 9 \times 0.0064^2}{38}\right]\left[\frac{40}{300}\right]} + 0.001$$

$$\le 0.005$$
(X3.4)

Since this statement is true, there is no reason to believe that the two values are different at the 95 percent confidence level, and the value of K is therefore verified.

X3.1.10 The variability of the sample is compared to the tolerance furnished with the gages using Eq 13 in 13.1.3:

🖽 E 251 – 92 (2003)

TABLE X3.1 Test Plan

Odd numbered gages—Runs 1, 2 and 3 in tension; Runs 4 and 5 in compression Even numbered gages—Runs 1, 2 and 3 in compression; Runs 4 and 5 in tension

Gage No	Run 1	Run 2	Run 3	Run 4	Run 5	Avg. K _i	$K_i - K_C$	$(K_i - K_C)^2$
1	2.039	2.040	2.012	2.040	2.005	2.027	+ .0068	$4.624 imes10^{-5}$
2	2.008	2.015	2.013	2.022	2.019	2.015	0052	$2.704 imes10^{-5}$
3	2.038	2.039	2.001	2.040	2.000	2.024	+ .0038	$1.444 imes10^{-5}$
4	2.002	2.014	2.036	2.003	2.005	2.012	0082	$6.724 imes10^{-5}$
5	2.013	2.022	2.027	2.037	2.048	2.029	+ .0088	$7.744 imes10^{-5}$
6	2.000	2.021	2.046	2.008	2.034	2.022	+ .0018	$0.324 imes10^{-5}$
7	2.007	2.027	2.004	2.039	2.011	2.018	0022	$0.484 imes10^{-5}$
8	2.047	2.005	2.047	2.001	2.028	2.026	+ .0058	$3.364 imes10^{-5}$
9	2.013	2.008	2.007	2.015	2.011	2.011	0092	$8.464 imes10^{-5}$
10	2.005	2.010	2.036	2.003	2.034	2.018	0022	$0.484 imes10^{-5}$
						20 202		36.36×10^{-5}

Control Values (Calculated): $K_c = \Sigma K_i / n = 20.202 / 10 = 2.0202$ $S_c = \sqrt{\Sigma (K_i - K_c)^2 / n - 1} = \sqrt{3.636 \cdot 10^{-4} / 9} = 0.0064$

$$e_s = 0.0005 K_c = 0.0010$$

$$A \leq (S_R^2 / S_C^2) \leq B \tag{X3.5}$$

Using values from Table 1 in this equation gives:

$$0.386 \le \left[\frac{0.0051^2}{0.0064^2} = 0.635\right] \le 3.57 \tag{X3.6}$$

Reported values (from package):

$$S_R = 2.024$$

 $S_R = T_R/2 = 0.005 \cdot 2.024/2 = 0.0051$

Since this statement is true, there is no reason to believe that the variabilities are different at the 95 % confidence level, and the value of T_R is therefore verified.

REFERENCES

- (1) "Bonded Electric Resistance Strain Gauges with Metallic Measurement Grids—Characteristics and Testing Conditions"; VDE/VDI— Richtlinien NR 2365 August, 1974. Verein Deutscher Ingenieure, VDE/VDI—Fachgruppe Messtechnik, 4000 Dusseldorf 1, Graf-Recke-Str. 84, F. R. of Germany.
- (2) "Calibration of Measuring Instruments—Force and Strain Measuring Extensioneters"; Draft Recommendation, Rc. Aero 803 21. Bureao National de l'Aeronautique et de l'Espace, Boulogne 92100, France (undated).
- (3) "Measurements and Uncertainties"; by P. Giacomo, International Bureau of Weights and Measures. OIML Bulletin No. 65, 1976.
- (4) "Methods for Calibration of Bonded Electric Resistance Strain Gauges"; Draft for Development 6: 1972. British Standards Institution, 2 Park Street, London W1A 2BS, England.
- (5) "Strain Gages, Bonded Resistance"; Classification Specification NAS 942 (National Aerospace Standard 942) 1963. Aerospace Industries Association of America, Inc. Published by National Standards Association, Inc., 1315 Fourteenth Street, N.W., Washington, D.C., USA.
- (6) "Performance Characteristics of Metallic Resistance Strain Gauges"; International Recommendation No. 62, 1985. Organisation Internationale de Metrologie Legale, 11, Rue Turgot, 75009 Paris, France.

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