



Standard Test Method for Elapsed Time Calibration of Thermal Analyzers¹

This standard is issued under the fixed designation E1860; 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 test method describes the calibration or performance confirmation of the elapsed-time signal from thermal analyzers.

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

1.3 There is no ISO standard equivalent to this test method.

1.4 *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. Referenced Documents

2.1 *ASTM Standards:*²

[D3350](#) Specification for Polyethylene Plastics Pipe and Fittings Materials

[D3895](#) Test Method for Oxidative-Induction Time of Polyolefins by Differential Scanning Calorimetry

[D4565](#) Test Methods for Physical and Environmental Performance Properties of Insulations and Jackets for Telecommunications Wire and Cable

[D5483](#) Test Method for Oxidation Induction Time of Lubricating Greases by Pressure Differential Scanning Calorimetry

[E473](#) Terminology Relating to Thermal Analysis and Rheology

[E487](#) Test Method for Constant-Temperature Stability of Chemical Materials

[E691](#) Practice for Conducting an Interlaboratory Study to Determine the Precision of a Test Method

[E1142](#) Terminology Relating to Thermophysical Properties

[E1858](#) Test Method for Determining Oxidation Induction Time of Hydrocarbons by Differential Scanning Calorimetry

[E1868](#) Test Method for Loss-On-Drying by Thermogravimetry

[E2161](#) Terminology Relating to Performance Validation in Thermal Analysis

3. Terminology

3.1 *Definitions:*

3.1.1 The technical terms used in this test method are defined in Terminologies [E473](#), [E1142](#), and [E2161](#), including calibration, conformance, relative standard deviation, and thermal analysis.

4. Summary of Test Method

4.1 The elapsed time signal generated by a thermal analyzer is compared to a clock (or timer) whose performance is known and traceable to a national metrology institute. The thermal analyzer may be said to be in conformance, if the performance of the thermal analyzer is within established limits. Alternatively, the elapsed time signal may be calibrated using a two point calibration method.

5. Significance and Use

5.1 Most thermal analysis experiments are carried out under increasing temperature conditions where temperature is the independent parameter. Some experiments, however, are carried out under isothermal temperature conditions where the elapsed time to an event is measured as the independent parameter. Isothermal Kinetics (Test Methods [E2070](#)), Thermal Stability (Test Method [E487](#)), Oxidative Induction Time (OIT) (Test Methods [D3895](#), [D4565](#), [D5483](#), [E1858](#), and Specification [D3350](#) and Loss-on-Drying (Test Method [E1868](#)) are common examples of these kinds of experiments.

5.2 Modern scientific instruments, including thermal analyzers, usually measure elapsed time with excellent precision and accuracy. In such cases, it may only be necessary to confirm the performance of the instrument by comparison to a suitable reference. Only rarely will it may be required to correct the calibration of an instrument's elapsed time signal through the use of a calibration factor.

5.3 It is necessary to obtain elapsed time signal conformity only to 0.1 times the repeatability relative standard deviation

¹ This test method is under the jurisdiction of ASTM Committee [E37](#) on Thermal Measurements and is the direct responsibility of Subcommittee [E37.10](#) on Fundamental, Statistical and Mechanical Properties.

Current edition approved Sept. 15, 2013. Published September 2013. Originally approved in 1997. Last previous edition approved in 2007 as E1860–07. DOI: 10.1520/E1860-13.

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.

(standard deviation divided by the mean value) expressed as a percent for the test method in which the thermal analyzer is to be used. For those test methods listed in Section 2 this conformity is 0.1 %.

6. Apparatus

6.1 *Timer or Stopwatch*, with timing capacity of at least 3 h (10 800 s), a resolution of 0.1 s or better and an accuracy of 1.5 s per day which performance has been verified using standards and procedures traceable to a national metrology institute (such as the National Institute of Standards and Technology (NIST)). Such timers are available from most laboratory equipment suppliers.

7. Calibration

7.1 Perform any elapsed time signal calibration procedures recommended by the manufacturer of the thermal analyzer as described in the Operator's Manual.

8. Procedure

8.1 Obtain the instrument reaction time (I).

8.1.1 Reset the timer and the elapsed time signal for the thermal analyzer to zero elapsed time.

8.1.2 Simultaneously start the timer and the elapsed time signal for the thermal analyzer. Allow them to run for 6 to 10 s. Simultaneously stop the timer and the elapsed time signal for the thermal analyzer. Record the elapsed time from the timer as t_1 . Record the elapsed time from the thermal analyzer as t_2 .

NOTE 1—The elapsed time of the timer (t_1) is equal to the elapsed time of the thermal analyzer (t_2) plus the instrument reaction time (I). The instrument reaction time is that required for the thermal analyzer to initialize and terminate the thermal analysis experiment and may be up to several seconds. The instrument start up time does not affect the elapsed time of the thermal analysis experiment since the experiment is exclusive of this time.

NOTE 2—Data acquisition rate shall be set to the maximum available.

NOTE 3—Time measurements shall be recorded in seconds retaining all available digits.

8.1.3 Calculate the instrument reaction time I by Eq 2 (9.2).

8.2 Obtain the calibration constant (S).

8.2.1 Reset the timer and the elapsed time signal for the thermal analyzer to zero elapsed time.

8.2.2 Simultaneously start the timer and the elapsed time signal for the thermal analyzer. Allow them to run for a minimum of 10 000 s (= 167 min = 2.8 h = 2 h, 47 min). Simultaneously stop the timer and the elapsed time signal for the thermal analyzer (see Note 2, Note 3, and Note 4). Record the elapsed time from the timer as t_r . Record the elapsed time from the thermal analyzer as t_o .

8.2.3 Calculate the value for S using Eq 3 (see 9.3).

8.3 Using the values for I and S from 8.1.3 and 8.2.3, calculate the percent conformity (C) using Eq 4 or table of percent conformity values (see 9.4).

9. Calculation

9.1 For the purpose of these procedures, it is assumed that the relationship between observed elapsed time (t_o) and the actual elapsed time (t) is linear and is governed by Eq 1:

$$t = t_o S \quad (1)$$

where:

t = true experimental elapsed time (s),

t_o = thermal analyzer observed elapsed time (s), and

S = slope (nominal value = 1.00000).

9.2 Using the values for t_1 and t_2 from 8.1, the instrument reaction time (I) may be calculated by:

$$I = t_1 - t_2 \quad (2)$$

9.3 Using the values for t_r and t_o from 8.2, the calibration constant S may be calculated by:

$$S = (t_r - I)/t_o \quad (3)$$

where:

t_r = observed time of reference timer.

9.3.1 When performing this calculation, retain all available decimal places in the measured value and in the value of S .

9.4 Using the value for S from 9.3, the percent conformity of the instrument elapsed time indicator may be calculated as follows:

$$C = (1.00000 - S) \times 100 \% \quad (4)$$

NOTE 4—The percent conformity is usually a very small number and expressing it as a percent value may be inconsistent with SI metric notation. Because of its effect on the experiment and because of common use, it is expressed as a percent in this procedure.

9.4.1 Conformity may be estimated to one significant figure using the following criteria:

9.4.1.1 If S lies:

Between 0.9999 and 1.0001, then conformity is better than 0.01 %,

Between 0.9990 and 0.9999 or between 1.0001 and 1.0010, then conformity is better than 0.1 %, and

Between 0.9900 and 0.9990 or between 1.0010 and 1.0100, then conformity is better than 1 %.

9.5 Using the determined value for S , Eq 1 may be used to calculate the true elapsed time (t) from an observed elapsed time (t_o).

10. Report

10.1 Report the following information:

10.1.1 Model number and description of the Thermal Analyzer used,

10.1.2 The value of S as determined in 8.2.3 reported to at least five places to the right of the decimal point, and

10.1.3 Conformity as determined in 9.4.

11. Precision and Bias

11.1 An interlaboratory study of elapsed time calibration was conducted in 1996 that included participation by nine laboratories using instruments from four manufacturers. The results were treated by Practice E691.³

11.2 *Precision:*

11.2.1 The mean value for the calibration constant was $S = 0.999853$.

³ Supporting data have been filed at ASTM International Headquarters and may be obtained by requesting Research Report RR:E37-1019. Contact ASTM Customer Service at service@astm.org.

11.2.2 The repeatability (within laboratory) standard deviation for S was 0.000034.

11.2.2.1 Two values, each the mean of duplicate determinations within a single laboratory should be considered suspect if they differ by more than the 95 % repeatability limit $r = 0.000095$.

11.2.3 The reproducibility (between laboratory) standard deviation for S was 0.00024.

11.2.3.1 Two values, each the mean of duplicate determinations in differing laboratories, should be considered suspect, if they differ by more than the 95 % reproducibility limit $R = 0.00069$.

11.3 *Bias:*

11.3.1 The measurement of conformity in this test method is a comparison of the calibration constant S with the theoretical value of 1.000000 and provides an indication of bias.

11.3.2 The mean value for conformity was $C = 0.015 \%$.

11.3.3 Conformity was found to vary widely among instrument models but in no case exceeded $C = 0.05 \%$. This value is far better than the nominal conformity of 1 % required for most thermal analysis experiments.

12. Keywords

12.1 calibration; elapsed time; thermal analysis; time

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