

Designation: D7807 – 20

# Standard Test Method for Determination of Boiling Range Distribution of Hydrocarbon and Sulfur Components of Petroleum Distillates by Gas Chromatography and Chemiluminescence Detection<sup>1</sup>

This standard is issued under the fixed designation D7807; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon ( $\varepsilon$ ) indicates an editorial change since the last revision or reapproval.

#### 1. Scope\*

1.1 This test method covers the determination of the boiling range distribution of petroleum products. The test method is applicable to petroleum products and fractions having a final boiling point of 538 °C (1000 °F) or lower at atmospheric pressure as measured by this test method. This test method is limited to samples having a boiling range greater than 55 °C (100 °F), and having a vapor pressure sufficiently low to permit sampling at ambient temperature.

1.1.1 The applicable sulfur concentration range will vary to some extent depending on the boiling point distribution of the sample and the instrumentation used; however, in most cases, the test method is applicable to samples containing levels of sulfur above 10 mg/kg.

1.2 This test method requires the use of both FID and SCD for detection. The hydrocarbon simulated distillation data obtained from the FID signal should be performed according to Test Method D2887 Procedure B.

1.3 The test method is not applicable for analysis of petroleum distillates containing low molecular weight components (for example, naphthas, reformates, gasolines, crude oils). Materials containing heterogeneous components (for example, alcohols, ethers, acids, or esters) or residue are not to be analyzed by this test method. See Test Methods D3710, D7096, D5307, D7169, or D7500.

1.4 This test method does not purport to identify all sulfur species in a sample. The detector response to sulfur is equimolar for all sulfur compounds within the scope (1.1) of this test method. Thus, unidentified sulfur compounds are determined with equal precision to that of identified substances. Total sulfur content is determined from the total area of the sulfur detector.

1.4.1 This test method uses the principles of simulated distillation methodology.

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

1.6 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, health, and environmental practices and determine the applicability of regulatory limitations prior to use.

1.7 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:<sup>2</sup>
- D86 Test Method for Distillation of Petroleum Products and Liquid Fuels at Atmospheric Pressure
- D1160 Test Method for Distillation of Petroleum Products at c- Reduced Pressure 0 ceda 1 b4/astm-d7807-20
- D2622 Test Method for Sulfur in Petroleum Products by Wavelength Dispersive X-ray Fluorescence Spectrometry
- D2887 Test Method for Boiling Range Distribution of Petroleum Fractions by Gas Chromatography
- D2892 Test Method for Distillation of Crude Petroleum (15-Theoretical Plate Column)
- D3120 Test Method for Trace Quantities of Sulfur in Light Liquid Petroleum Hydrocarbons by Oxidative Microcoulometry
- D3710 Test Method for Boiling Range Distribution of Gasoline and Gasoline Fractions by Gas Chromatography (Withdrawn 2014)<sup>3</sup>
- D4626 Practice for Calculation of Gas Chromatographic Response Factors

#### \*A Summary of Changes section appears at the end of this standard

<sup>&</sup>lt;sup>1</sup>This test method is under the jurisdiction of ASTM Committee D02 on Petroleum Products, Liquid Fuels, and Lubricants and is the direct responsibility of Subcommittee D02.04.0H on Chromatographic Distribution Methods.

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<sup>&</sup>lt;sup>2</sup> 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.

<sup>&</sup>lt;sup>3</sup> The last approved version of this historical standard is referenced on www.astm.org.

- D5307 Test Method for Determination of Boiling Range Distribution of Crude Petroleum by Gas Chromatography (Withdrawn 2011)<sup>3</sup>
- D5504 Test Method for Determination of Sulfur Compounds in Natural Gas and Gaseous Fuels by Gas Chromatography and Chemiluminescence
- D5623 Test Method for Sulfur Compounds in Light Petroleum Liquids by Gas Chromatography and Sulfur Selective Detection
- D6299 Practice for Applying Statistical Quality Assurance and Control Charting Techniques to Evaluate Analytical Measurement System Performance
- D6300 Practice for Determination of Precision and Bias Data for Use in Test Methods for Petroleum Products, Liquid Fuels, and Lubricants
- D6352 Test Method for Boiling Range Distribution of Petroleum Distillates in Boiling Range from 174 °C to 700 °C by Gas Chromatography
- D7096 Test Method for Determination of the Boiling Range Distribution of Gasoline by Wide-Bore Capillary Gas Chromatography
- D7169 Test Method for Boiling Point Distribution of Samples with Residues Such as Crude Oils and Atmospheric and Vacuum Residues by High Temperature Gas Chromatography
- D7500 Test Method for Determination of Boiling Range Distribution of Distillates and Lubricating Base Oils—in Boiling Range from 100 °C to 735 °C by Gas Chromatography
- E178 Practice for Dealing With Outlying Observations
- E355 Practice for Gas Chromatography Terms and Relationships
- E594 Practice for Testing Flame Ionization Detectors Used in Gas or Supercritical Fluid Chromatography
- E1510 Practice for Installing Fused Silica Open Tubular Capillary Columns in Gas Chromatographs

### 3. Terminology

3.1 *Definitions*—This test method makes reference to many common gas chromatographic procedures, terms, and relationships. Detailed definitions of these can be found in Practices E355, E594, and E1510.

3.2 Definitions of Terms Specific to This Standard:

3.2.1 *area slice*, n—the area, resulting from the integration of the chromatographic detector signal, within a specified retention time interval. Peak detection parameters are bypassed and the detector signal integral is recorded as area slices of consecutive, fixed duration time intervals.

3.2.2 *corrected area slice*, *n*—an area slice corrected for baseline offset, by subtraction of the exactly corresponding area slice in a previously recorded blank (non-sample) analysis.

3.2.3 *cumulative corrected area*, *n*—the accumulated sum of corrected area slices from the beginning of the analysis through a given retention time, ignoring any non-sample area (for example, solvent).

3.2.4 *final boiling point (FBP), n*—the temperature (corresponding to the retention time) at which a cumulative corrected

area count equal to 99.5 % of the total sample area under the chromatogram is obtained.

3.2.5 *initial boiling point (IBP), n*—the temperature (corresponding to the retention time) at which a cumulative corrected area equal to 0.5 % of the total sample area under the chromatogram is obtained.

3.2.6 response factor (RF), *n*—the factor used in order to calculate the mg/kg sulfur recovery of the sample.

3.2.7 *slice rate, n*—the time interval used to integrate the continuous (analog) chromatographic detector response during an analysis. The slice rate is expressed in Hz (for example, integrations or slices per second).

3.2.8 slice time, n—the cumulative slice rate (analysis time) associated with each area slice throughout the chromatographic analysis. The slice rate is the time at the end of each contiguous area slice.

3.2.9 *total sample area, n*—the cumulative corrected area from the initial point to the final area point.

3.3 *Abbreviations*—A common abbreviation of hydrocarbon compounds is to designate the number of carbon atoms in the compound. A prefix is used to indicate the carbon chain form, while a subscripted suffix denotes the number of carbon atoms (for example, normal decane n-C10, iso-tetradecane = i-C14).

#### 4. Summary of Test Method

4.1 The boiling range distribution determination by distillation is simulated by the use of gas chromatography. A nonpolar open tubular (capillary) gas chromatographic column is used to elute the hydrocarbon components of the sample in order of increasing boiling point. The column temperature is raised at a reproducible linear rate and the area under the chromatogram is recorded throughout the analysis. Boiling points are assigned to the time axis from a calibration curve obtained under the same chromatographic conditions by analyzing a known mixture of hydrocarbons covering the boiling range expected in the sample. A quantitative standard is used to determine the SCD detector response factor. Finally, the sample solution is injected and with the use of the response factor, the amount of sample recovered is calculated. After converting the retention times of the sample slices to temperature, the boiling point distribution can be calculated up to the recovered amount. From these data, the boiling range distribution can be obtained.

4.1.1 By splitting the column effluent to FID and Sulfur Chemiluminescence Detector, simultaneous detection for hydrocarbon (FID) and sulfur (SCD) components boiling range distribution is obtained. The hydrocarbon simulated distillation data should be calculated according to Test Method D2887.

4.1.2 Alternatively, the FID may be used with the SCD detector superimposed over the FID and thus avoiding splitting the sample through the column exit. This type of arrangement will lower the sensitivity of the detector in the sulfur mode.

4.2 A sample aliquot is introduced into the chromatographic system. Sample vaporization is provided by separate heating of the point of injection or in conjunction with column oven heating.

4.3 The column oven temperature is raised at a reproducible linear rate to effect separation of the sample components in order of increasing boiling point. The elution of sample components is quantitatively determined using a flame ionization detector and a sulfur chemiluminescence detector. The detector signal integral is recorded as area slices for consecutive retention time intervals during the analysis.

4.4 Retention times of known normal paraffin hydrocarbons spanning the scope of this test method (C5- C44) are determined and correlated to their theoretical boiling point temperatures. The normalized cumulative corrected sample areas for each consecutive recorded time interval are used to calculate the boiling range distribution. The boiling point temperature at each reported percent off increment is calculated from the retention time calibration.

4.5 *Sulfur Chemiluminescence Detection*—As sulfur compounds elute from the gas chromatographic column, they are processed in a heated combustion zone. The products are collected and transferred to a sulfur chemiluminescence detector (SCD). This technique provides a sensitive, selective, linear response to volatile sulfur compounds and is used for the selective sulfur detection, while collecting hydrocarbon data from the FID.

4.6 Alternative Detectors—This test method is written for the sulfur chemiluminescence detector but other sulfur specific detectors can be used provided they have sufficient linearity, sensitivity, and have equimolar response to all eluted sulfur compounds, do not suffer from interferences, and satisfy quality assurance criteria. Regulatory agencies may require demonstration of equivalency of alternative detection systems to the SCD.

### 5. Significance and Use

5.1 The boiling range distribution of light and medium petroleum distillate fractions provides an insight into the composition of feed stocks and products related to petroleum refining processes. This gas chromatographic determination of boiling range can be used to replace conventional distillation methods for control of refining operations. This test method can be used for product specification testing with the mutual agreement of interested parties.

5.2 This test method extends the scope of Test Method D2887 (538 °C) boiling range determination by gas chromatography to include sulfur boiling range distribution in the petroleum distillate fractions. Knowledge of the amount of sulfur and its distribution in hydrocarbons is economically important in determining product value and in determining how best to process or refine intermediate products. Sulfur compounds are known to affect numerous properties of petroleum and petrochemical products. The corrosion of metals and poisoning of catalysts is of particular concern. In addition, the content of sulfur in various refined products may be subject to governmental regulations. Test Methods, such as, D2622, D3120, D5504 and D5623, are available to determine total sulfur content or content of individual sulfur compounds in petroleum and petroleum products. Test Methods, such as, D86, D1160, D2887, D3710, and D2892, are also available to determine the hydrocarbon boiling ranges of such samples. The gas chromatographic determination of the sulfur boiling range assists in process development, in treatment and control of refining operations and is useful for assessing product quality. This determination produces detailed information about the sulfur distribution in a sample that cannot be obtained by either total sulfur analysis or analysis of sulfur in discreet distillation cuts.

5.2.1 The hydrocarbon boiling range distributions obtained by Test Method D2887 are theoretically equivalent to those obtained by true boiling point (TBP) distillation (see Test Method D2892). They are not equivalent to results from low efficiency distillation such as those obtained with Test Method D86 or D1160.

# 6. Apparatus

6.1 *Chromatograph*—Any gas chromatograph, with hardware necessary for interfacing to a chemiluminescence detector and containing all features necessary for the intended application(s) can be used. The gas chromatographic system used shall have the following performance characteristics:

6.2 Column Temperature Programmer—The chromatograph must be capable of linear programmed temperature operation over a range sufficient to elute compounds up to a boiling temperature of 538 °C (1000 °F) before reaching the upper end of the temperature program. The programming rate must be sufficiently reproducible to obtain retention time repeatability of 0.01 min (0.6 s, corresponding to approximately 0.5 °C) for each component in the calibration mixture described in 7.7.

6.3 *Detectors*—This test method requires a Flame Ionization Detector (FID) and a Sulfur Chemiluminescence Detector (SCD).

6.3.1 *FID*—The FID shall meet or exceed the following specifications in accordance with Practice E594. Check the detector according to the instrument manufacturer's instructions.

6.3.2 *SCD*—The sulfur chemiluminescence detector shall meet or exceed the following specifications: (1) greater than  $10^3$  linearity, (2) less than 1 pg S/s sensitivity, (3) greater than  $10^6$  selectivity for sulfur compounds over hydrocarbons, (4) no quenching of sulfur compound response from co-eluting hydrocarbons when the same volume of sample is injected as for regular analysis, and (5) equimolar response (<±10 %) on a sulfur basis.

6.3.2.1 For the purpose of boiling point calibration, the system must be capable of measuring sulfur compounds and hydrocarbons simultaneously from a single column and injection, for example, flame ionization detector with splitting the column effluent prior to the sulfur chemiluminescence detector. Alternatively, a combined FID/SCD can also be used in order to obtain simultaneous sulfur and FID chromatogram.

6.3.2.2 Sulfur compounds eluting from the chromatographic column are processed in a heated hydrogen-rich combustion zone fitted to the end of the column. Products are transferred under reduced pressure to the reaction chamber of the chemiluminescence detector. An excess of ozone present in the chamber reacts with the sulfur combustion product(s) to liberate blue (480 nm) and ultraviolet light (260 nm).

6.3.3 *Detector Split Requirements*—To ensure the low levels of sulfur are detected properly, the system must be capable to detect the components in the system sulfur test mixture (see 8.9) with signal to noise (peak-to-peak) ratio of at least 100. Connections of the column to the detector shall be such that no temperature below the column temperature exists. Refer to Practice E1510 for proper installation and conditioning of the capillary column.

6.4 *Sample Inlet System*—Any sample inlet system capable of meeting the performance specification in 8.7 may be used. Programmed temperature vaporization (PTV) and programmable cool on-column injection systems have been used successfully.

6.5 *Carrier Gas Flow Control*—The chromatograph shall be equipped with carrier gas pressure or flow control capable of maintaining constant carrier gas flow control through the column throughout the column temperature program cycle.

6.6 *Micro Syringe*—A micro syringe with a 23 gauge or smaller stainless steel needle is used for sample introduction. Syringes of 0.1  $\mu$ L to 10  $\mu$ L capacity are commercially available. Automatic syringe injection is recommended.

6.7 *Column*—This test method is limited to the use of non-polar wall coated open tubular (WCOT) columns.

6.7.1 Any column and conditions may be used that provide separation of typical petroleum hydrocarbons in order of increasing boiling point and meet the column performance requirements of 8.7.1 and 9.3.1.1.

6.7.2 Glass, fused silica, and stainless steel columns, with a 0.53 mm diameter have been successfully used. Cross-linked and bonded 100 % dimethyl-polysiloxane stationary phases with film thickness of 0.5  $\mu$ m to 2.65  $\mu$ m have been used. The column length and liquid phase film thickness shall allow the elution of at least C44 n-paraffin (BP = 545 °C).

6.8 Data Acquisition System—Use of an electronic integrating device or computer is mandatory for determining the detector response and for boiling point calibration. The device must have the following capabilities: (1) graphic presentation of the chromatogram, (2) digital display of chromatographic peak areas, (3) measurement of area and time intervals, (4) calculation and use of response factors in accordance with Practice D4626, for example, external standardization, and (5) the maximum area measured must be within the linear range of the measuring system used.

Note 1—Some gas chromatographs have an algorithm built into their operating software that allows a mathematical model of the baseline profile to be stored in memory. This profile is automatically subtracted from the detector signal on subsequent sample runs to compensate for the column bleed. Some integration systems also store and automatically subtract a blank analysis from subsequent analytical determinations.

### 7. Reagents and Materials

7.1 *Carrier Gas*—Helium, of at least 99.999 % (v/v) purity (**Warning**—Helium is a compressed gas under high pressure). Any oxygen present is removed by a chemical resin filter (**Warning**—Follow the safety instructions from the filter supplier.) The total amount of impurities should not exceed 10 mL/m<sup>3</sup>. Helium or Nitrogen (at least 99.999 % (v/v)) can also be used as detector make-up gas.

TABLE I Typical das enfoliatographic contations					
Instrument	A gas chromatograph equipped with an on-column or temperature programmable vaporizing injector (PTV)				
Column	Capillary 10 m, 0.53 mm ID, 0.88 $\mu m$ 100 % dimethylpolysiloxane stationary phase				
Flow Conditions	25 mL/min He carrier (constant flow)				
Inlet	Temperature: Programmed 100 ℃ to 350 ℃ at 25 ℃/min				
Detector	Split FID – SCD or SCD with FID Adapter				
FID	Temperature: 350 °C 35 mL/min H2, 350 mL/min Air				
SCD	Temperature: 950 °C 10 mL/min O <sub>2</sub> , 90 mL/min H <sub>2</sub> 100 mL/min to 150 mL/min He or N <sub>2</sub> make-up				
Oven	35 °C to 350 °C at 25 °C /min				
Sample Injection	0.1 µL neat				

TABLE 1 Typical Gas Chromatographic Conditions

7.1.1 Additional purification is recommended by the use of molecular sieves or other suitable agents to remove water, oxygen, and hydrocarbons. Available pressure shall be sufficient to ensure a constant carrier gas flow rate.

**7.2** *Hydrogen*—Hydrogen of at least 99.999 % (v/v) purity (suitable for the flame ionization detector (FID). (**Warning**— Hydrogen is an extremely flammable gas under high pressure.)

7.3 *Air*—High purity (for example, hydrocarbon-free) compressed air is used as the oxidant for the flame ionization detector (FID). (**Warning**—Compressed air is a gas under high pressure and supports combustion.)

7.4 Oxygen—High purity (for example, hydrocarbon-free) compressed oxygen is used as the oxidant for the sulfur chemiluminescence detector (SCD). (Warning—Compressed oxygen is a gas under high pressure and supports combustion.)

Note 2—Some SCD detectors allow the use of air instead of oxygen: contact the SCD manufacturer for information on the use of air as oxidant.

7.5 *Solvents*—Unless otherwise indicated, it is intended that all solvents conform to the specifications of the committee on analytical Reagents of the American Chemical Society<sup>4</sup> where such specifications are available.

7.5.1 Other grades may be used provided it is first ascertained that the solvent is of sufficiently high purity to permit its use without lessening the accuracy of the determination.

7.6 Cyclohexane  $(C_6H_{12})$ , (99+ % pure) may be used as a viscosity reducing solvent. It is miscible with asphaltic hydrocarbons, however, it responds well to the FID. The quality (hydrocarbon content) should be determined by this test method prior to use as a sample diluent. (Warning—Cyclohexane is flammable.)

<sup>&</sup>lt;sup>4</sup> ACS Reagent Chemicals, Specifications and Procedures for Reagents and Standard-Grade Reference Materials, American Chemical Society, Washington, DC. For suggestions on the testing of reagents not listed by the American Chemical Society, see Analar Standards for Laboratory Chemicals, BDH Ltd., Poole, Dorset, U.K., and the United States Pharmacopeia and National Formulary, U.S. Pharmacopeial Convention, Inc. (USPC), Rockville, MD.

7.7 *Calibration Mixture*—An accurately weighed mixture of approximately equal mass quantities of n-hydrocarbons dissolved in a suitable solvent. The mixture shall cover the boiling range from n-C5 to n-C44, but does not need to include every carbon number, but at least sufficient number of points to generate a reliable calibration curve and the C16 and C18.

7.7.1 At least one compound in the mixture must have a boiling point lower than the IBP of the sample and at least one compound in the mixture must have a boiling point higher than the FBP of the sample. Boiling points of n-paraffins are listed in Table 2.

7.7.2 If necessary, for the calibration mixture to have a compound with a boiling point below the IBP of the sample, propane or butane can be added to the calibration mixture, non-quantitatively, by bubbling the gaseous compound into the calibration mixture in a septum sealed vial using a gas syringe.

7.7.3 *Reference Material*—A reference sample that has been analyzed by laboratories participating in the test method cooperative study. Consensus values for the boiling range distribution of this sample are being determined.

#### 8. Preparation of Apparatus

#### 8.1 Gas Chromatograph Setup:

8.2 Place the gas chromatograph and ancillary equipment into operation in accordance with the manufacturer's instructions. Recommended operating conditions are shown in Table 1.

8.3 When attaching the column to the detector inlet, ensure that the end of the column terminates as close as possible to the FID jet. Follow the instructions in Practice E1510.

8.4 The FID should be periodically inspected and, if necessary, remove any foreign deposits formed in the detector from combustion of silicone liquid phase or other materials. Such deposits will change the response characteristics of the detector.

8.5 *SCD*—Place in service in accordance with the manufacturer's instructions. Optimization of the oxidant/fuel ratio is critical for ensuring complete combustion of hydrocarbon components in a sample. A flame or combustion zone that is too hydrogen rich will result in incomplete combustion. Matrix interference is occasionally observed when changing sample size. When matrix interference is indicated, samples may be analyzed by dilution or application of other mitigation efforts.

8.5.1 The typical flameless/combustion zone interface contains ceramic tubes in its construction. The performance of these tubes is critical to performance of the SCD system. Compromised ceramic tubes are susceptible to matrix effects. Compromised tubes may allow for reproducible duplicate sample analysis but will fail QA procedures such as matrix dilution and spike analyses. Poorly functioning tubes can also result in severe instrument drift, loss of equimolar response, and general response instability. Compromised tubes must be replaced to restore nominal instrument function.

8.6 *Column Conditioning*—A new column will require conditioning at the upper test method operating temperature to reduce or eliminate significant liquid phase bleed, resulting in a stable chromatographic baseline. Follow the guidelines outlined in Practice E1510.

**8.7** System Performance Specification:

Carbon No.	Boiling Point, °C	Boiling Point, F	Carbon No.	Boiling Point, °C	Boiling Point, °F		
https://sta	ndards -162 ai/catalog	standard -259 t/foal a	152-6a <b>23</b> -4a84	-a498-( <b>380</b> :50ceda1)	4/astm-d78716_20		
2	-89	-127	24	391	736		
3	-42	-44	25	402	755		
4	0	31	26	412	774		
5	36	97	27	422	791		
6	69	156	28	431	808		
7	98	209	29	440	825		
8	126	258	30	449	840		
9	151	303	31	458	856		
10	174	345	32	466	870		
11	196	385	33	474	885		
12	216	421	34	481	898		
13	235	456	35	489	912		
14	254	488	36	496	925		
15	271	519	37	503	937		
16	287	548	38	509	948		
17	302	576	39	516	961		
18	316	601	40	522	972		
19	330	626	41	528	982		
20	344	651	42	534	993		
21	356	674	43	540	1004		
22	369	695	44	545	1013		

TABLE 2 Boiling Points of Normal Paraffins<sup>A,B</sup>

<sup>A</sup> API Project 44, October 31, 1972 is believed to have provided the original normal paraffin boiling point data that are listed in Table 1. However, over the years some of the data contained in both API Project 44 (Thermodynamics Research Center Hydrocarbon Project) and Test Method D6352 have changed and they are no longer equivalent. Table 1 represents the current normal paraffin boiling point values accepted by Subcommittee D02.04 and found in all test methods under the jurisdiction of Section D02.04.0H.

<sup>*B*</sup> Used *n*-paraffin boiling points are traditionally rounded to the nearest whole degree for calibration. The boiling points listed in Table 1 are correct to the nearest whole number in both degrees Celsius and degrees Fahrenheit. However, if a conversion is made from one unit to the other and then rounded to a whole number, the results will not agree with the table values for a few carbon numbers. For example, the boiling point of n-heptane is 98.425 °C, which is correctly rounded to 98 °C in the table. However, converting 98.425 °C gives 209.165 °F, which rounds to 209 °F, while converting 98 °C gives 208.4 °F, which rounds to 208 °F. Carbon numbers 2, 4, 7, 8, 9, 13, 14, 15, 16, 25, 27, and 32 are affected by rounding.

8.7.1 Column Resolution-The column resolution, influenced by both the column physical parameters and operating conditions, affects the overall determination of boiling range distribution. Resolution is therefore specified to maintain equivalence between different systems (laboratories) employing this test method. Resolution is determined using Eq 1 and the C16 and C18 paraffins from the calibration mixture analysis (see 7.7), and is illustrated in Fig. 1. Resolution (R) should be at least three, using the identical conditions employed for sample analyses:

$$R = 2(t_2 - t_1) / (1.699 (w_2 + w_1))$$
(1)

where:

R = resolution,

= time(s) for the n-C16 peak maximum,  $t_1$ 

= time(s) for the *n*-C18 peak maximum,  $t_2$ 

 $w_{I}$ = peak width(s), at half height, of the n-C16 peak, and

= peak width(s), at half height, of the n-C18 peak.  $W_2$ 

8.7.2 Column Elution Characteristics-The recommended column liquid phase is a non-polar phase such as 100 % dimethyl-polysiloxane.

8.8 Sulfur Standard—A petroleum sample with a known total Sulfur content and a known boiling point distribution of the sulfur (Warning-Sulfur compounds can be flammable and harmful or fatal when ingested or inhaled.).

8.8.1 Calculate the relative response factor for the Sulfur Standard (see 8.8).

8.8.2 Inject and analyze a suitable amount of the sulfur standard (8.8). Relative response factors shall be calculated for the Sulfur standard in accordance with Practice D4626 or Eq 2.

$$R_m = \frac{C_n \times A_r}{C_r \times A_n} \tag{2}$$

where:

 $R_m$  = relative response factor for the Sulfur standard,

= concentration of the sulfur compound as sulfur,  $C_n$ 

= peak area of sulfur compound,

 $A_n$  $C_r$ = concentration of referenced sulfur standard as sulfur, and

 $A_r$ = peak area of the referenced sulfur standard.

8.9 Linearity Check—A linearity check should be performed after installation of the instrument or whenever maintenance is performed. It is recommended to analyze known standards with different levels of sulfur; or, dilute a known sulfur standard with a sulfur-free standard with a similar boiling point distribution.

8.10 *Recovery Check*—To check the stability of the analysis system, a QC-sample analysis should be made at least every ten analyses. The total area for the sulfur components should not deviate more than 10 % from the value obtained in the previous calibration run.

#### 9. Procedure

9.1 Analysis Sequence Protocol—Define and use a predetermined schedule of analysis events designed to achieve maximum reproducibility for these determinations. The schedule will include cooling the column oven and injector to the initial starting temperature, equilibration time, sample injection and system start, analysis, and final temperature hold time.

9.1.1 After chromatographic conditions have been set to meet performance requirements, program the column temperature upward to the maximum temperature to be used and hold that temperature for the selected time. Following the analysis sequence protocol, cool the column to the initial starting temperature.

9.1.2 Inject either the calibration mixture, solvent, or sample into the chromatograph; or make no injection (baseline blank). At the time of injection, start the chromatograph time cycle and the integrator/computer data acquisition. Follow the analysis protocol for all subsequent repetitive analyses or calibrations. Since complete resolution of sample peaks is not expected, do not change the sensitivity setting during the analysis.

9.2 Baseline Blank-Perform a blank analysis (baseline blank) at least once per day. The blank analysis may be without injection or by injection of an equivalent solvent volume as used with sample injections, depending upon the subsequent data handling capabilities for baseline/solvent compensation. The blank analysis is typically performed prior to sample analyses, but may be useful if determined between samples or



FIG. 1 Column Resolution Parameters

at the end of a sample sequence to provide additional data regarding instrument operation or residual sample carryover from previous sample analyses.

NOTE 3—If automatic baseline correction (see Note 1) is provided by the gas chromatograph, further correction of area slices may not be required. However, if an electronic offset is added to the signal after baseline compensation, additional area slice correction may be required in the form of offset subtraction. Consult the specific instrumentation instructions to determine if an offset is applied to the signal. If the algorithm used is unclear, the slice area data can be examined to determine if further correction is necessary. Determine if any offset has been added to the compensated signal by examining the corrected area slices of those time slices which precede the elution of any chromatographic unretained substance. If these corrected area slices (representing the true baseline) deviate from zero, subtract the average of these corrected area slices from each corrected area slice in the analysis.

9.3 Retention Time versus Boiling Point Calibration—If this is the first time that an analysis is carried out, prepare the sequence to include the retention time calibration standard, the Reference Gas Oil and a blank which is necessary to calculate the Boiling Point Distribution of the Reference Gas Oil as well as for subsequent samples analysis. Calibration should be performed weekly when the instrument is in use, or whenever maintenance is performed and as dictated by the lab on-site precision and/or Quality Control protocol. Inject an appropriate aliquot of the calibration mixture (see 7.7) into the chromatograph, using the analysis sequence protocol. Obtain a normal (peak detection) data record in order to determine the peak retention times and the peak areas for each component. Collect a time slice area record if a boiling range distribution report is desired. Fig. 2 illustrates a graphical plot of a calibration analysis.

9.3.1 Inspect the chromatogram of the calibration mixture for evidence of skewed (non-Gaussian shaped) peaks. Skewness is often an indication of overloading the sample capacity of the column, which will result in displacement of the peak apex relative to non-overloaded peaks. Distortion in retention time measurement and hence errors in boiling point temperature calibration will be likely if column overloading occurs. The column liquid phase loading has a direct bearing on acceptable sample size. Reanalyze the calibration mixture using a smaller sample size or a more dilute solution to avoid peak distortion.

9.3.1.1 Skewness Calculation—Calculate the ratio A/B on specified peaks in the calibration mixture as indicated by the designations in Fig. 3. A is the width in seconds of the portion of the peak eluting prior to the time of the peak apex and measured at 10 % of peak height (0.10-H), and B is the width in seconds of the portion of the peak eluting after the time of the peak apex at 10 % of peak height (0.10-H). The skewness of all peaks shall not be less than 0.5 nor more than 2.

9.3.2 Prepare a calibration table based upon the results of the analysis of the calibration mixture by recording the time of each peak maximum and the boiling point temperature in degrees Celsius (or Fahrenheit) for every component in the mixture. Normal paraffin boiling point temperatures (atmospheric equivalent temperatures) are listed in Table 2.

9.3.3 It is recommended to perform a calibration at the start and end of the sequence of analysis.

9.4 Sample Preparation—The amount of sample injected must not overload the column stationary phase capacity nor exceed the detector linear range. A narrow boiling range sample will require a smaller amount injected than a wider boiling range sample. Samples that are of low enough viscosity to be sampled with a syringe at ambient temperature may be injected neat. This type of sample may also be diluted with an appropriate solvent (for example, cyclohexane) to control the amount of sample injected.

<u>9.4.1</u> Samples that are too viscous or waxy to sample with a syringe may be diluted with an appropriate solvent.



