



Standard Test Method for Determination of Individual Components in Spark Ignition Engine Fuels by 100 Metre Capillary High Resolution Gas Chromatography¹

This standard is issued under the fixed designation D6729; 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 (ϵ) indicates an editorial change since the last revision or reapproval.

1. Scope*

1.1 This test method covers the determination of individual hydrocarbon components of spark-ignition engine fuels and their mixtures containing oxygenate blends (MTBE, ETBE, ethanol, and so forth) with boiling ranges up to 225 °C. Other light liquid hydrocarbon mixtures typically encountered in petroleum refining operations, such as blending stocks (naphthas, reformates, alkylates, and so forth) may also be analyzed; however, statistical data was obtained only with blended spark-ignition engine fuels.

1.2 Based on the cooperative study results, individual component concentrations and precision are determined in the range of 0.01 % mass to approximately 30 % mass. The procedure may be applicable to higher and lower concentrations for the individual components; however, the user must verify the accuracy if the procedure is used for components with concentrations outside the specified ranges.

1.3 The test method also determines methanol, ethanol, t-butanol, methyl t-butyl ether (MTBE), ethyl t-butyl ether (ETBE), t-amyl methyl ether (TAME) in spark ignition engine fuels in the concentration range of 1 % mass to 30 % mass. However, the cooperative study data provided sufficient statistical data for MTBE only.

1.4 Although a majority of the individual hydrocarbons present are determined, some co-elution of compounds is encountered. If this test method is utilized to estimate bulk hydrocarbon group-type composition (PONA) the user of such data should be cautioned that some error will be encountered due to co-elution and a lack of identification of all components present. Samples containing significant amounts of olefinic or naphthenic (for example, virgin naphthas), or both, constituents above *n*-octane may reflect significant errors in PONA type groupings. Based on the gasoline samples in the inter-

laboratory cooperative study, this procedure is applicable to samples containing less than 25 % mass of olefins. However, some interfering coelution with the olefins above C₇ is possible, particularly if blending components or their higher boiling cuts such as those derived from fluid catalytic cracking (FCC) are analyzed, and the total olefin content may not be accurate. Caution should also be exercised when analyzing olefin-free samples using this test method as some of the paraffins may be reported as olefins since analysis is based purely on retention times of the eluting components.

1.4.1 Total olefins in the samples may be obtained or confirmed, or both, if necessary, by Test Method **D1319** (percent volume) or other test methods, such as those based on multidimensional PONA type of instruments (Test Method **D6839**).

1.5 If water is or is suspected of being present, its concentration may be determined, if desired, by the use of Test Method **D1744**, or equivalent. Other compounds containing oxygen, sulfur, nitrogen, and so forth, may also be present, and may co-elute with the hydrocarbons. If determination of these specific compounds is required, it is recommended that test methods for these specific materials be used, such as Test Methods **D4815** and **D5599** for oxygenates, and **D5623** for sulfur compounds, or equivalent.

1.6 **Annex A1** of this test method compares results of the test procedure with other test methods for selected components, including olefins, and several group types for several interlaboratory cooperative study samples. Although benzene, toluene, and several oxygenates are determined, when doubtful as to the analytical results of these components, confirmatory analyses can be obtained by using specific test methods.

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

1.8 *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 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 **D02.04.0L** on Gas Chromatography Methods.

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

2. Referenced Documents

2.1 ASTM Standards:²

- D1319** Test Method for Hydrocarbon Types in Liquid Petroleum Products by Fluorescent Indicator Adsorption
- D1744** Test Method for Determination of Water in Liquid Petroleum Products by Karl Fischer Reagent
- D4815** Test Method for Determination of MTBE, ETBE, TAME, DIPE, tertiary-Amyl Alcohol and C₁ to C₄ Alcohols in Gasoline by Gas Chromatography
- D5599** Test Method for Determination of Oxygenates in Gasoline by Gas Chromatography and Oxygen Selective Flame Ionization Detection
- D5623** Test Method for Sulfur Compounds in Light Petroleum Liquids by Gas Chromatography and Sulfur Selective Detection
- D6839** Test Method for Hydrocarbon Types, Oxygenated Compounds, and Benzene in Spark Ignition Engine Fuels by Gas Chromatography
- E355** Practice for Gas Chromatography Terms and Relationships

3. Terminology

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

4. Summary of Test Method

4.1 Representative samples of the petroleum liquid are introduced into a gas chromatograph equipped with an open tubular (capillary) column coated with the specified stationary phase. Helium carrier gas transports the vaporized sample through the column, in which it is partitioned into individual components which are sensed with a flame ionization detector as they elute from the end of the column. The detector signal is recorded digitally by way of an integrator or integrating computer. Each eluting component is identified by comparing its retention time to that established by analyzing reference standards or samples under identical conditions. The concentration of each component in mass percent is determined by normalization of the peak areas after correction of selected components with detector response factors. The unknown components are reported individually and as a summary total.

5. Significance and Use

5.1 Knowledge of the specified individual component composition (speciation) of gasoline fuels and blending stocks is useful for refinery quality control and product specification. Process control and product specification compliance for many individual hydrocarbons may be determined through the use of this test method.

6. Apparatus

6.1 *Gas Chromatograph*, a gas chromatograph equipped with cryogenic column oven cooling and capable of producing

repeatable oven ramps from 0 °C to at least 300 °C is required. The following features are useful during the sample analysis phase: electronic flow readout, electronic sample split-ratio readout, and electronic pneumatic control of flow. Though their use is not required, careful review of this test method will demonstrate the usefulness of a gas chromatograph equipped with these features. These features will replace the need to carry out the manual calculations that must be performed as listed in **8.1** and **8.2**.

6.2 *Inlet*—a capillary split/splitless inlet system operated in the split mode is recommended. It must be operated in its linear range. Refer to **8.4** to determine the proper split ratio.

6.2.1 *Carrier Gas Pneumatic Control*—Constant carrier gas pressure control was used by all cooperative study participants. This may be either direct pressure to the inlet (injector) or by using a total flow/back pressure system.

6.2.2 *Pneumatic Operation of the Chromatograph*—The use of constant pressure was the mode of operating the gas chromatography used by the participants in the interlaboratory cooperative study. Other carrier gas control methods such as constant flow (pressure programming) may be used, but this may change the chromatography elution pattern unless the temperature programming profile is also adjusted to compensate for the flow differences.

6.2.3 *Temperature Control*—The injector operated in the split mode shall be heated by a separate heating zone and heated to temperatures of 200 °C to 275 °C.

6.3 *Column*, a fused silica capillary column, 100 m in length by 0.25 mm inside diameter, coated with a 0.5 μm film of bonded dimethylpolysiloxane. The column must meet the resolution requirements expressed in **8.3**. Columns from two different commercial sources were used in the interlaboratory cooperative study.

6.4 *Data System*, a computer based chromatography data system capable of accurately and repeatedly measuring the retention time and areas of eluting peaks. The system shall be able to acquire data at a rate of at least 10 Hz. Although it is not mandatory, a data system which calculates column resolution (R) is extremely useful as it will replace the need to carry out the manual calculations which must be performed as listed in **8.3**.

6.4.1 *Electronic Integrators*, shall be capable of storing up to 400 components in the peak table and shall be able to acquire the data at 10 Hz or faster speeds. They shall be capable of integrating peaks having peak widths at half height which are 1.0s wide. The integrator must be capable of displaying the integration mode of partially resolved peaks. In addition, these integrators should be able to download a commonly readable format of data (that is, ASCII) to a computer in order to facilitate data processing.

6.5 *Sample Introduction*—Sample introduction by way of a valve, automatic injection device, robotic arm or other automatic means is highly recommended. An automatic sample introduction device is essential to the reproducibility of the analysis. Manual injections are not recommended. All of the

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.

reproducibility data reported by this test method for the samples analyzed were gathered using automatic injection devices.

6.6 Flame Ionization Detector (FID)—The gas chromatograph should possess a FID having a sensitivity of 0.005 coulombs/g for *n*-butane. The linear dynamic range of the detector should be 10⁶ or better. The detector is heated to 300 °C.

7. Reagents and Materials

7.1 Calibrating Standard Mixture—A spark ignition engine fuel standard of known composition and concentration by mass can be used. In order to corroborate the identification of the sample, a typical chromatogram (Fig. 1) was obtained from reference sample ARC96OX.³

7.2 Gas Chromatograph Gases—All of the following gases shall have a purity of 99.999 % (V/V) or greater.

NOTE 1—Warning: Gases are compressed. Some are flammable and all gases are under high pressure.

7.2.1 Helium—The test data was developed with helium as the carrier gas. It is possible that other carrier gases may be used for this test method. At this time, no data is available from this test method with other carrier gases.

7.2.2 Air, Hydrogen and Make-up Gas (Helium or Nitrogen), shall have a purity of 99.999 % (V/V) or greater.

8. Instrument Check Out Prior to Analysis

8.1 Setting:

8.1.1 Linear Gas Velocity—If the gas chromatograph is equipped with an electronic flow readout device, set the flow to 1.8 mL/min. This is achieved by setting the carrier gas flow rate by injection of cm/s methane or natural gas at 35 °C. Ensure that the retention time is 7.00 min ± 0.05 min. This corresponds to a linear velocity of 25 cm/s to 26 cm/s. This is equivalent to retention times of methane at 0 °C ranging from 6.5 min to 6.8 min.

8.1.2 If the gas chromatograph is not equipped with an electronic flow readout device, calculate the linear gas velocity in cm/s using Eq 1.

$$\text{linear gas velocity} = V = \frac{\text{column length (cm)}}{\text{retention time of methane (s)}} \quad (1)$$

8.1.3 The typical retention times for methane and linear gas velocity for helium are 6.5 cm/s to 6.8 cm/s and 24 cm/s to 26 cm/s, respectively.

8.2 Setting the Split Ratio—If the gas chromatograph is equipped with an electronic split-ratio readout device, set the split ratio to a sample split of 200:1. If the gas chromatograph is not equipped with an electronic split-ratio readout device, one must first calculate column flow rate and then proceed to calculating split ratio using Eq 2 and 3.

$$\text{column flow rate} = F = \frac{(60 \pi r^2) L (T_{ref}) 2(P_i - P_o)}{(T) 3(P_{ref})(P_i^2 - P_o^2) \mu} \quad (2)$$

where:

F = flow rate as calculated by using the equation,
r = column radius, cm,
L = column length, cm,
P_i = inlet pressure,
P_o = outlet pressure,
P_{ref} = reference pressure, 1 atm,
T = temperature of the column oven,
T_{ref} = temperature at the column outlet, and
μ = linear velocity, cm/s.

$$\text{split ratio} = S = \frac{\text{split vent flow} + F}{F} \quad (3)$$

8.2.1 The column flow rate is calculated by the use of Eq 2. Use the results obtained from Eq 3 to adjust the split flow until a split flow of approximately 200:1 is achieved.

8.3 Evaluation of Column Performance:

8.3.1 Prior to using the column described in Table 1, measure the resolution of the column under the conditions of Table 2. Check that the resolution for the following pairs of components is obtained using Eq 4 to calculate the resolution of a pair of components:

$$R = \frac{2(t_{R2} - t_{R1})}{1.699 (W_{h1} + W_{h2})} \quad (4)$$

where:

R = resolution,
t_{R2} = retention time of the first member of the pair,
t_{R1} = retention time of the second member of the pair,
W_{h1} = peak width at half height of the first member of the pair, and
W_{h2} = peak width at half height of the second member of the pair.

8.3.1.1 Column resolution should be checked frequently by examining the resolution of these compounds.

8.3.2 Evaluation of the Baseline—Carry out a blank baseline run utilizing no solvent injection, by setting the GC in accordance with the conditions of Table 1.

8.3.3 Subtract the baseline from a sample chromatogram and verify that the residual signal at the beginning of the chromatogram does not differ from the end of the chromatogram by more than 2 %.

8.4 Evaluation of Splitter Linearity—Using the reference gasoline sample, inject this sample according to the schedule listed in Table 3.

8.4.1 Select from the chromatogram about 10 to 15 components, which have concentrations in the range of 0.01 % mass to 30 % mass. Tabulate for each split ratio the concentrations of the 10 to 15 components. Verify that for each component selected, its concentration does not vary by more than 3 %.

9. Procedure

9.1 Set the operating conditions of the gas chromatograph as shown in Table 1. These conditions will elute all components up to and including pentadecane (nC₁₅).

³ Reference spark ignition sample No. ARC 960X obtained from the Alberta Research Council, Edmonton, Alberta, Canada. Other samples are available from suppliers.

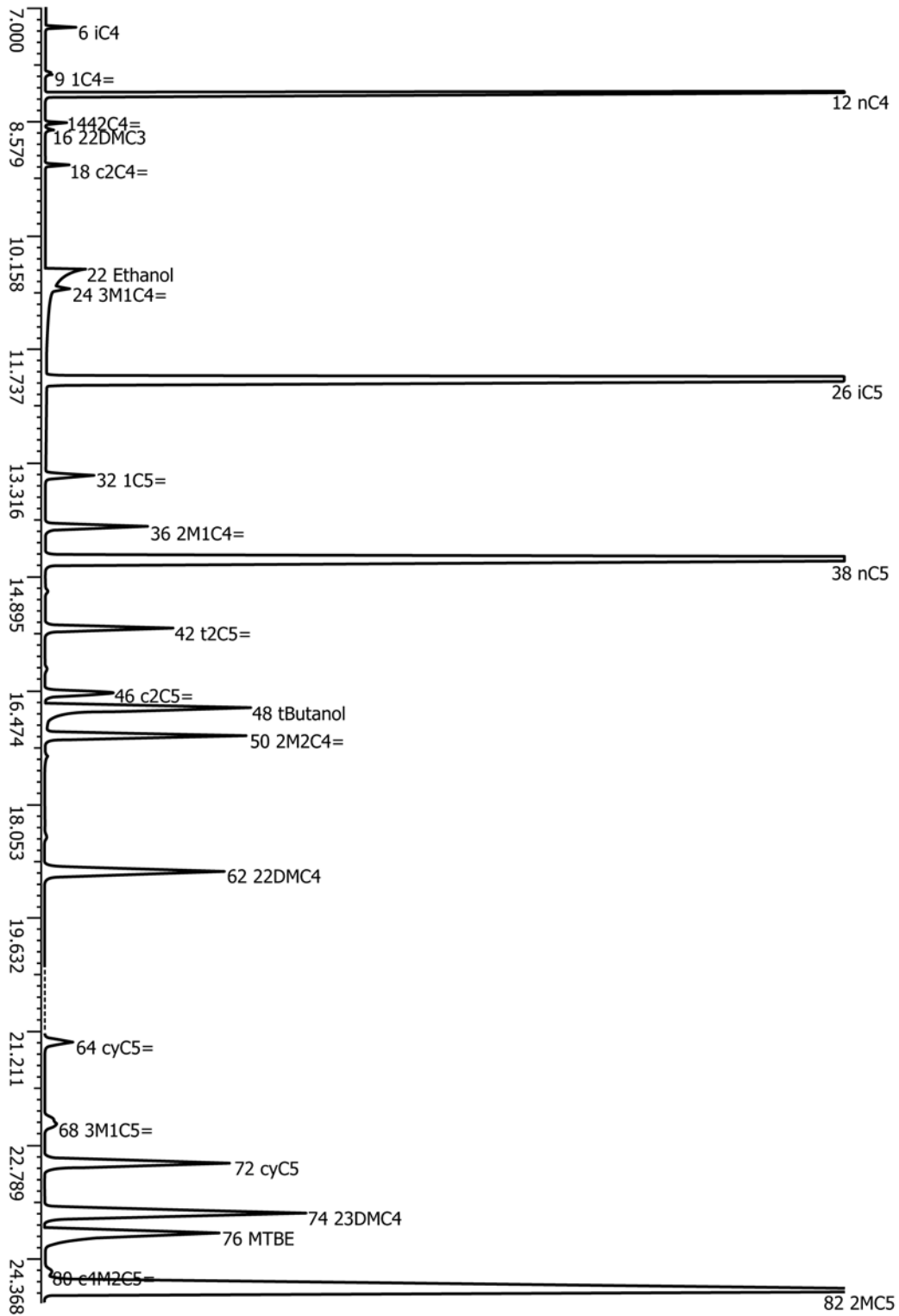


FIG. 1 Chromatogram for Reference Spiked Gasoline

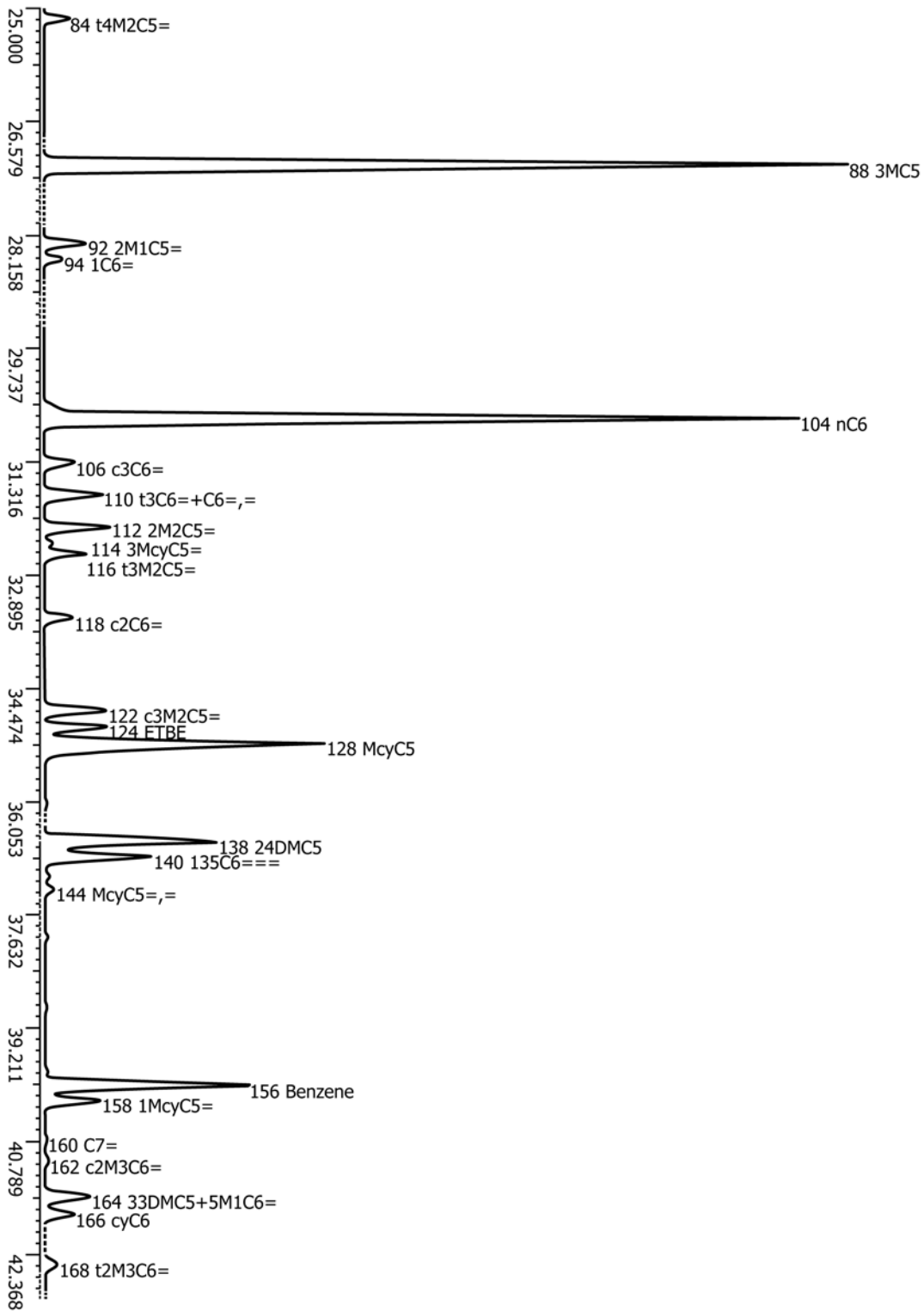


FIG. 1 Chromatogram for Reference Spiked Gasoline (continued)

9.2 All of the parameters in Table 1 can be marginally changed to optimize for sample types and optimize for characteristics of each gas chromatographic system. The final boiling point of samples should not exceed nC₁₅ and the

column resolution (R) performance requirements listed in Table 2 should not be compromised.

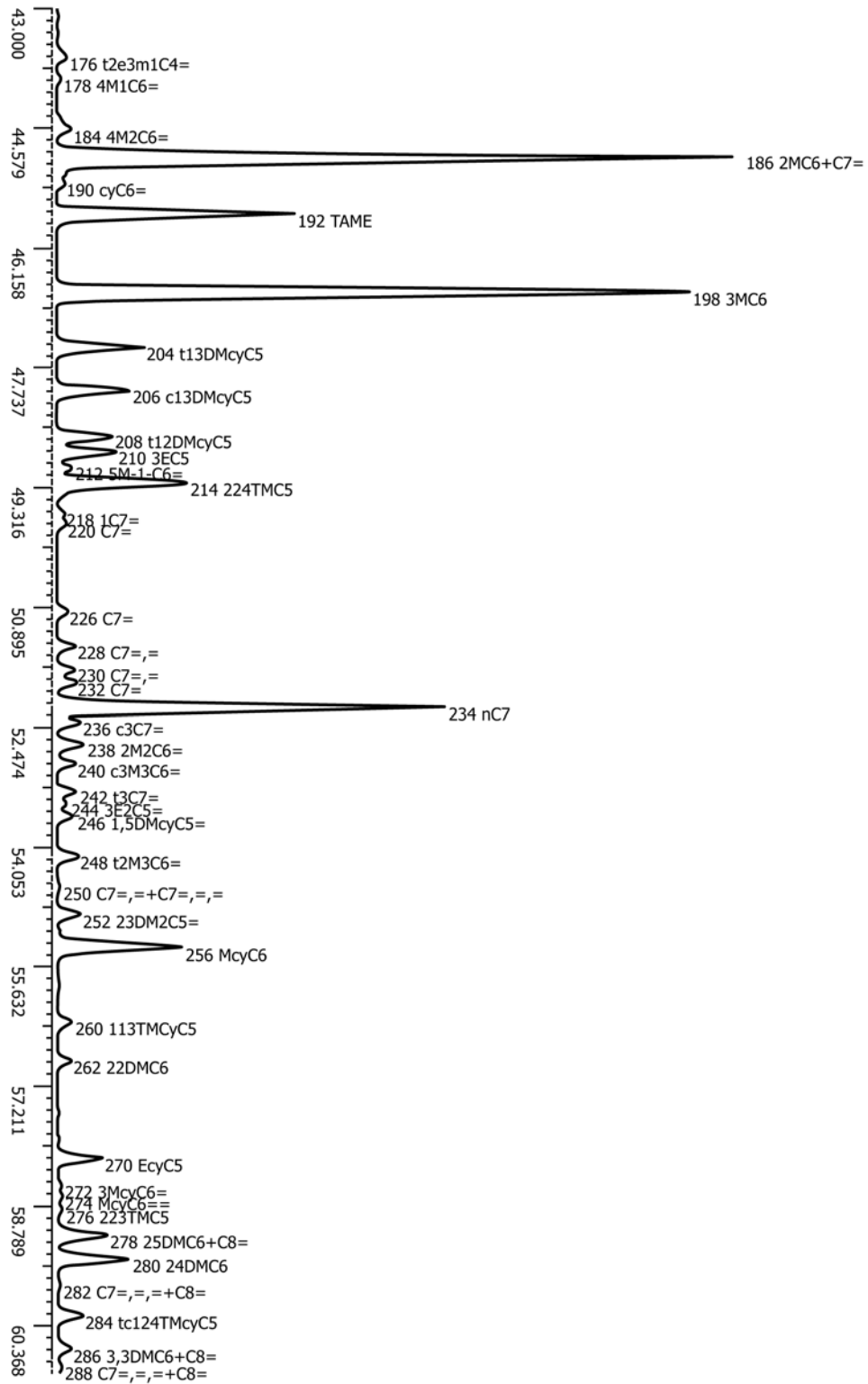


FIG. 1 Chromatogram for Reference Spiked Gasoline (continued)

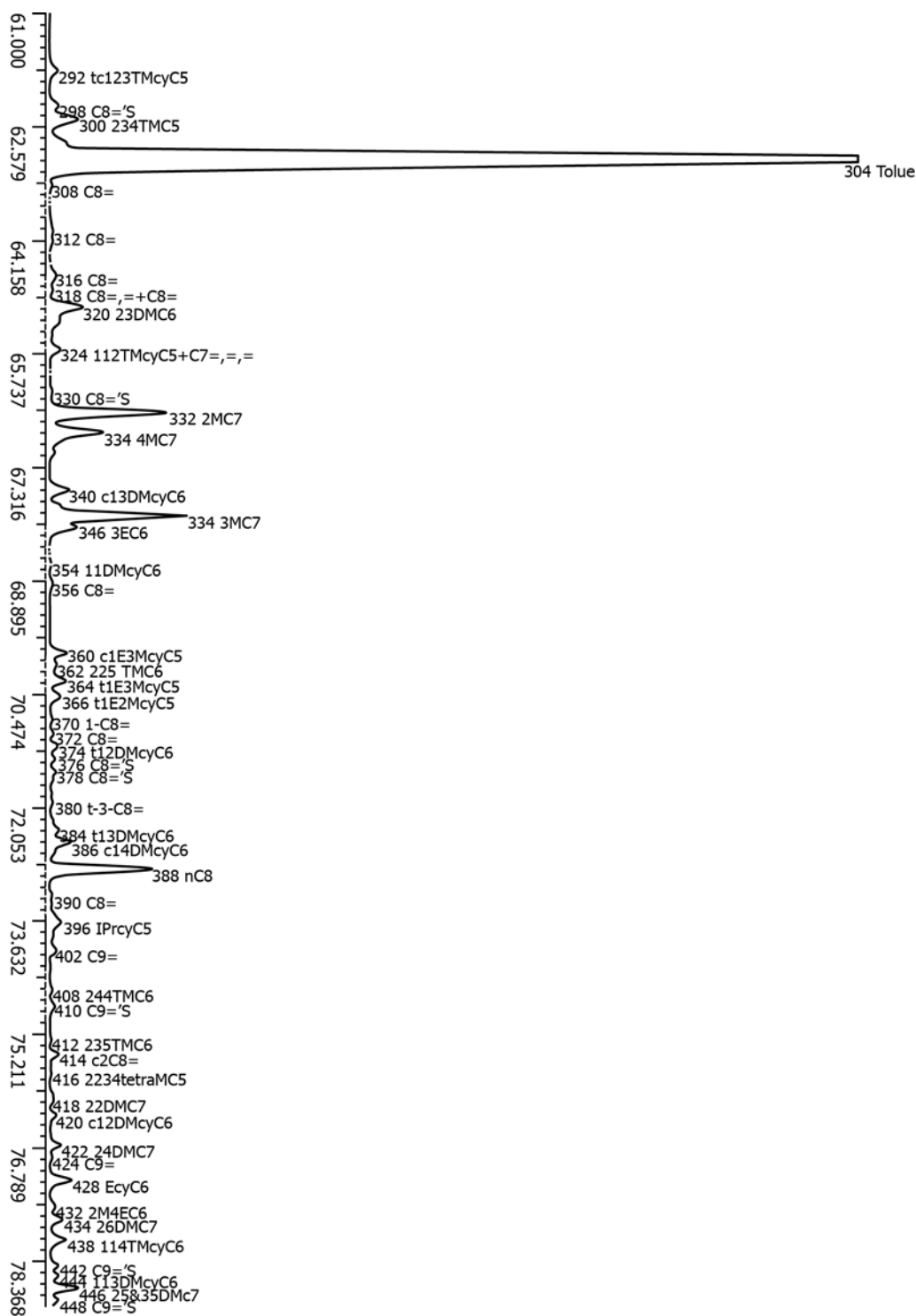


FIG. 1 Chromatogram for Reference Spiked Gasoline (continued)

9.3 Obtain a representative sample following the guidelines of Practice D4057 and any other applicable guidelines. Take precautions to minimize the loss of light ends from volatile samples. The sample container may be cooled prior to transfer of sample into it. Cool the sample to less than 4 °C, maintain at that approximate temperature until the autosampler is loaded and analysis begins.

9.4 Preparation/Storage:

9.4.1 Samples Stored in Vials—Cool the original sample to less than 4 °C prior to taking a sample aliquot or prior to filling the sample vials. The sample aliquot container, or the vial, or both, can also be cooled prior to the transfer of the original sample. Syringes may also be cooled along with the sample for manual injections.

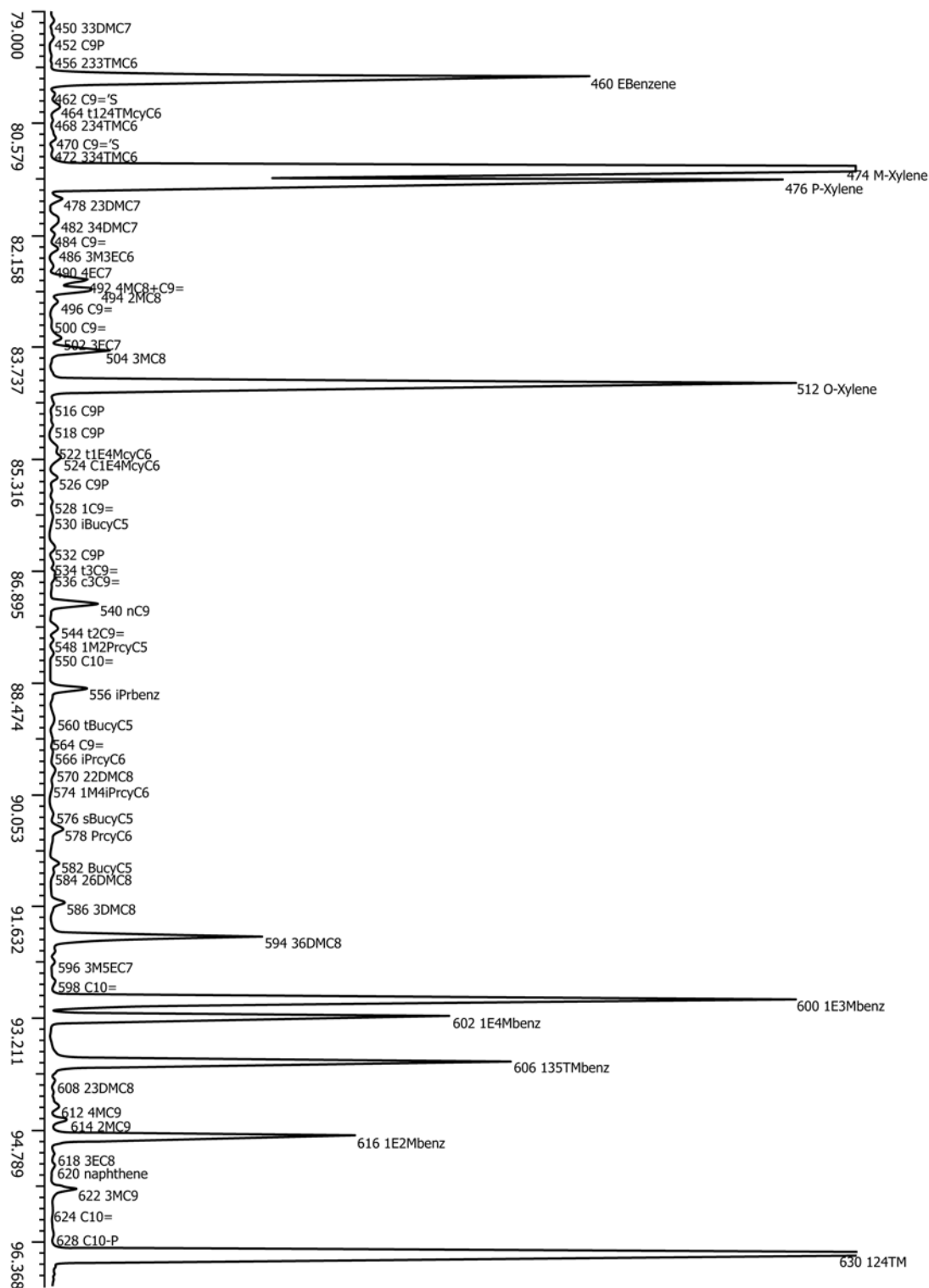


FIG. 1 Chromatogram for Reference Spiked Gasoline (continued)

9.4.2 Samples Stored in Pressurized Containers—It is recommended that they be kept away from direct heat or light. No other sample preparations are necessary for samples stored in pressurized containers. Avoid storage at temperatures greater than 25 °C. Store pressure containers in accordance with the manufacturer’s instructions.

9.5 It is recommended that a quality assurance (QA) sample similar to the reference material gasoline sample be run at regular intervals (see Fig. 1). An interval of once per week or after every 15 samples is recommended. The quantitation results use statistical quality control charts can track benzene. Other components of interest in the reference sample can be

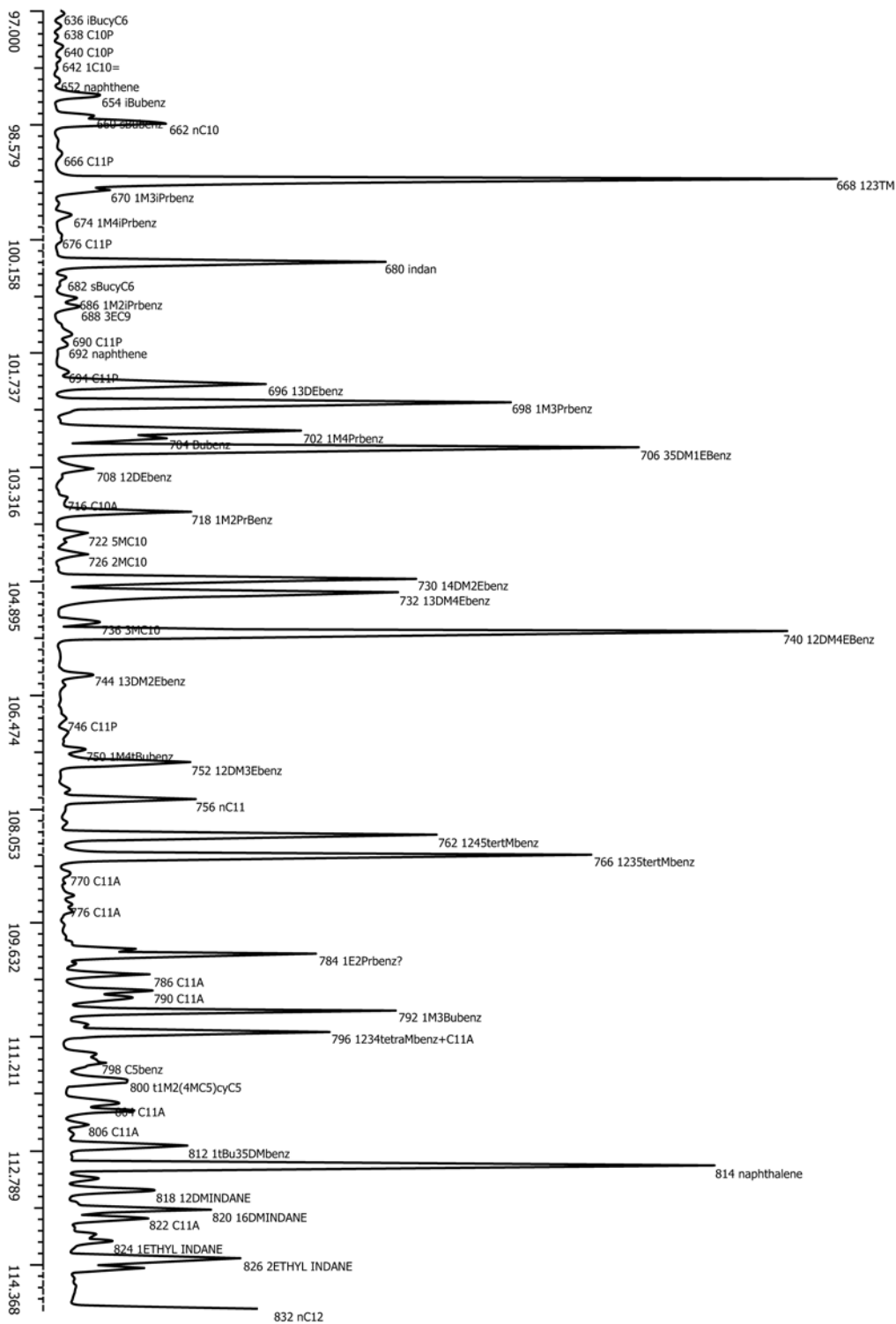


FIG. 1 Chromatogram for Reference Spiked Gasoline (continued)

tracked in a similar manner. By monitoring these components over an extended period of time, the performance of the column and the chromatographic system can be determined.

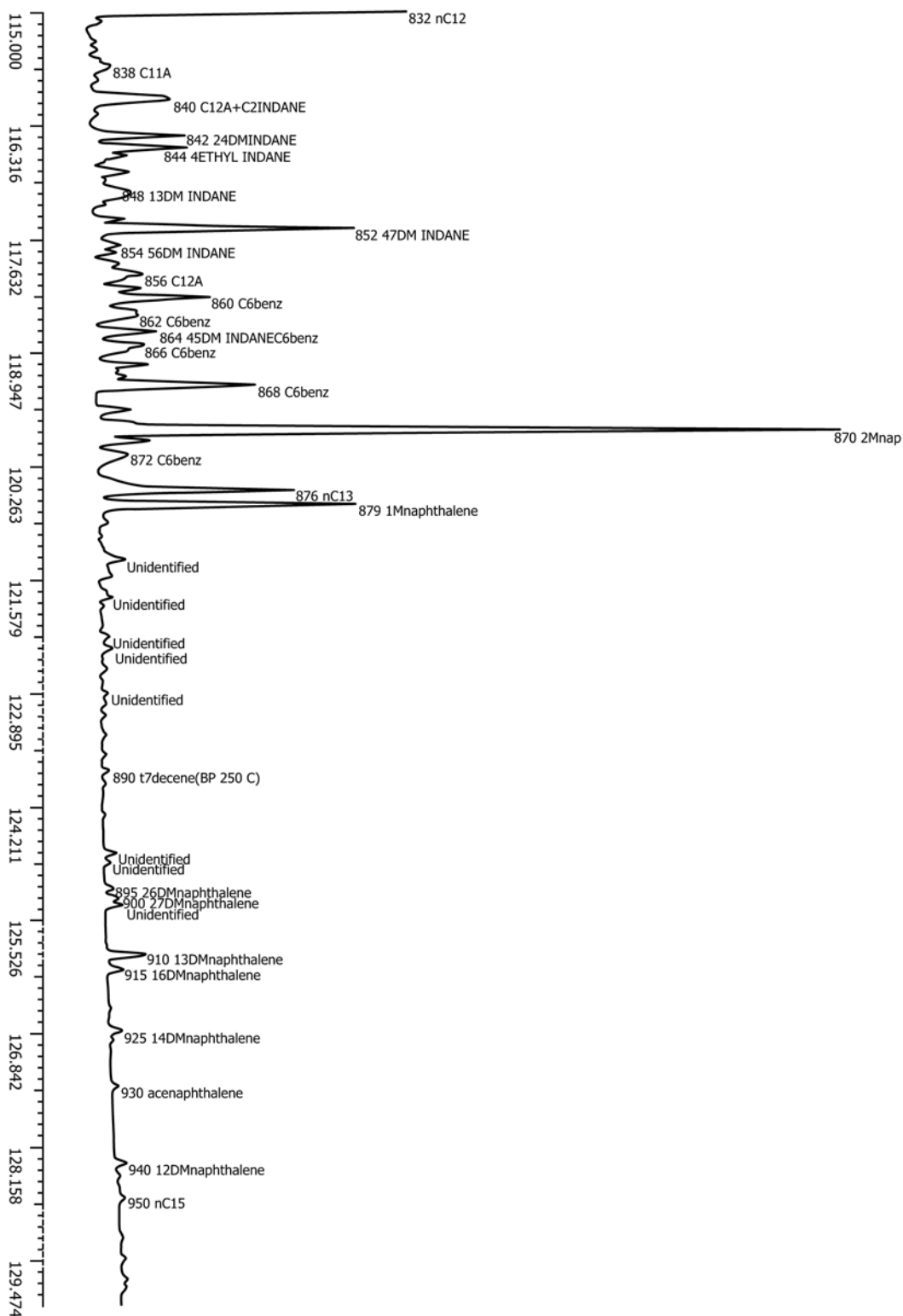


FIG. 1 Chromatogram for Reference Spiked Gasoline (continued)

10. Data Analysis

10.1 *Compound Identification*—Prepare a table listing all of the retention times of the components in the sample. Compare the retention time of each peak with that of the reference

gasoline. Pay particular attention to the fact that columns can be overloaded, and peaks can shift in retention time. Observe the peak pattern so that proper identification is made by comparison with the reference material.

TABLE 1 Chromatographic Operating Conditions, Column Requirements and Data Acquisition Requirements

Chromatographic Conditions	Requirements
Injector settings	
Injector temperature, °C	250
Split ratio	175:1 - 275:1
Liner	deactivated glass
Injection volume, µL	0.2–0.5
Detector settings	
FID detector temperature, °C ^A	300–350
Gas flows	
Hydrogen, mL/min ^B	30–40
Air, mL/min	300–450
Nitrogen make up, mL/min	30
Column oven settings	
Initial temperature, °C	0
Initial time, min	15
1st ramp rate, °C/min	1
Final temperature, °C	50
Final time, min	0
2nd ramp rate, °C/min	2
Final temperature, °C	130
Final time, min	0
3rd ramp rate, °C/min	4
Final temperature, °C	270
Final time, min ^C	0
Column Requirements	
Length, m	100
Inside diameter, mm	0.25
Liquid phase	100 % dimethylpolysiloxane
Film thickness, µ	0.5
Pressure, psig	40–50
Flow, mL/min	1.7–2.0
Linear gas velocity, cm/s	24.5
Data acquisition, Hz	10–20
Total analysis time, min	140–150

^A Set to 25 °C to 50°C above the highest column temperature.

^B Values to be set as recommended by instrument manufacturer.

^C Final temperature or time may be adjusted to ensure complete elution of the sample components.

TABLE 2 Resolution Performance Requirements

Component Pair	Minimum Resolution	Concentration of Each Component, W/W
Benzene 1-Methyl-cyclopentene	1.0	0.5 %–0.5 %
m-Xylene p-Xylene	0.4	2.0 %–2.0 %
n-Tridecane 1-Methylnaphthalene	1.0	0.5 %–0.5 %

TABLE 3 Injection Schedule

Split Ratio	Injection Volume, µL	Injection Temperature, °C
100:1	0.1	250
200:1	0.5	250
300:1	1.0	250

10.2 Consistency in peak identification can be achieved by using software (data handling software, spreadsheet software, and so forth). Alternatively, a retention index system can be used.

$$(R1)_i = 100n + 100 \left[\frac{\log(T_i) - \log(T_n)}{\log(TN) - \log(T_n)} \right] \quad (5)$$

where:

$(RI)_i$ = retention index of component I bracketed by the *N*-paraffin, *n* in its lower boundary and *N*-paraffin *N* in its upper boundary,

T_i = adjusted retention time of component *i* (retention time of component *i* minus the retention time of methane),

T_n = adjusted retention time of *N*-paraffin *n*, and

TN = retention time of *N*-paraffin *N*.

10.3 Determine the hydrocarbon response factors by using the following equation.⁴

$$RRF_{CH4} = \frac{MW_i}{N_c} \times \frac{1}{MW_{CH4}} \quad (6)$$

where:

RRF_{CH4} = relative response factor of each component with respect to methane ($RRF_{CH4} = 1.000$),

MW_i = molecular weight of the component, *i*,

N_c = number of carbon atoms in the molecule, and

MW_{CH4} = molecular weight of methane (16.04276).

10.4 Convert the acquired areas to corrected areas by multiplying each area by its corresponding relative response factor as indicated in the following equation.

⁴ Sevcik, J., *Detectors in Gas Chromatography*, Elsevier, NY, 1976, p. 94.

TABLE 4 Predominant Compounds and Identified Coeluting Compounds^A

NOTE 1—The response factor of the predominant compound will be used for the analyte and this analyte will be used for the calculations.

Peak Number (from Annex A1)	Predominant Compound	Coeluting Compound(s)
164	3,3-dimethylpentane	5-methyl-1-hexene
186	2-methylhexane	C ₇ -olefin
278	2,5-dimethylhexane	C ₈ -olefin
286	3,3-dimethylhexane	C ₈ -olefin
304	toluene	2,3,3-trimethylpentane ^B
324	1,1,2-trimethylcyclopentane	C ₇ -triolefin
326	C ₈ -diolefin	C ₈ -paraffin
492	4-methyloctane	C ₉ -olefin
796	1,2,3,4-tetramethylbenzene	C ₁₁ -aromatic

^A This is not an exhaustive list. Due to the possibility of coeluting peaks in other areas, the user is cautioned in the interpretation of the data.

^B In most alkylated gasolines, a split may occur between toluene and 233 TMC5.

TABLE 5 Response Factors of Oxygenated Compounds

Analytes	Relative Response Factors	
	RRF C ₇ = 1.000	RRF CH ₄ = 1.000
Methanol	2.996	2.672
Ethanol	2.087	1.862
t-Butanol (TBA)	1.302	1.161
Methyl-t-butyl ether (MTBE)	1.577	1.407
Ethyl-t-butyl ether (ETBE)	1.407	1.255
t-Amyl methyl ether (TAME)	1.356	1.210

$$A_{c_i} = (A)_i (RRF)_i \quad (7)$$

where:

$(A_{c_i})_i$ = corrected area,

A_i = acquired area for an individual component, and

RRF = relative response factor (weight basis).

10.4.1 The percent mass (% W) is calculated as follows:

$$\% W_i = \frac{(A_{c_i})_i}{\sum_{i=1}^{i=n} A_{c_i}} \times 100 \quad (8)$$

where:

$\% W$ = percent mass of the component i in the mixture, and

$i=n$ = summation of all the corrected areas for the components analyzed.

10.4.1.1 The subscript i indicates that the operations are carried out for each individual component in the matrix.

10.5 In the case of unidentified components, utilize a relative response factor of 0.800 (relative to methane).

11. Oxygenates

11.1 A cooperative study for linearity was performed for methanol, ethanol, t-butanol, methyl-t-butyl ether (MTBE), ethyl-t-butyl ether (ETBE), and t-amyl methyl ether (TAME) in concentration ranges from 1.0 % mass up to 30 % mass (Annex A2). The average relative response factors for the oxygenates were calculated from the study and are listed in Table A2.1. They have been incorporated into the IHA Method. The percent standard deviation of these relative response values was as high as 7 %. MTBE was the only oxygenate that was

present in sufficient number of samples to meet the ASTM requirements for round robin testing in accordance with RR:D02-1007. Therefore the statistical data for MTBE should be taken from Table A1.2.

12. Expression of Results

12.1 Report the concentration of each components as percent (m/m) to the nearest 0.001 % mass.

12.2 The data for individual components may be grouped by summing the concentration of compounds in each particular group type such as paraffin, isoparaffin, olefin, aromatic, naphthene, oxygenates, and unknowns. Commercially available software may be used to provide this function, as well as the calculation of other properties of petroleum liquids.

13. Precision and Bias⁵

13.1 The repeatability and reproducibility precision estimates are quoted in Annex A1.

13.2 *Precision Statement Outline*—(> Analyte Qualification Process):

13.2.1 For each analyte to qualify for a precision statement, it must be present in at least six samples, and detected by at least six laboratories, at least once, in accordance with RR:D02-1007⁶ requirements.

13.2.2 The (repeatability standard deviation)/mean value for each analyte/sample combination must be less than or equal to

⁵ Supporting data describing the interlaboratory cooperative study to determine precision and bias has been filed at ASTM International Headquarters and may be obtained by requesting RR: RR:D02-1519.

⁶ Supporting data have been filed at ASTM International Headquarters and may be obtained by requesting Research Report RR:D02-1007.

0.1, in accordance with LOQ requirements which, while not a standard, is what CS94 is recommending.

13.3 A brief explanation of headers in **Table A1.2** follows:

13.3.1 ID: self explanatory,

13.3.2 r_{\min} : lower 95 % confidence limit of r_{est} ,

13.3.3 r_{est} : repeatability estimate in percentage of concentration,

13.3.4 r_{\max} : upper 95 % confidence limit of r_{est} ,

13.3.5 R_{\min} , R_{est} , R_{\max} : same as above except for reproducibility,

13.3.6 C_{\min} : lower concentration limit that rest, R_{est} is applicable, and

13.3.7 C_{\max} : upper concentration limit that rest, R_{est} is applicable.

13.4 The summaries for the paraffins, isoparaffins, C_2 benzene, and oxygenates follow the same procedure that was used for the analytes and are listed in **Table A1.3**.

13.5 *Bias*—The bias of this test method cannot be determined since an appropriate standard reference material is not available.

14. Keywords

14.1 gas chromatograph; gasoline; individual hydrocarbon analysis; oxygenated fuels; spark-ignition engine fuels

ANNEXES

(Mandatory Information)

A1. HYDROCARBON DATA

A1.1 **Table A1.1** presents the component retention times and properties.

A1.2 **Table A1.2** represents the repeatability and reproducibility precision estimates prepared by statisticians of CS94 in accordance with RR:D02-1007. The analyte qualification process for precision statements is outlined as follows:

A1.2.1 For each analyte to qualify for a precision statement, it must be present in at least six samples, and detected by at least six laboratories, at least once, in accordance with RR:D02-1007 requirements.

A1.2.2 The (repeatability standard deviation)/mean value for each analyte/sample combination must be less than or equal to 0.1, in accordance with LOQ requirements which, while not a standard, is what CS94 is recommending.

A1.3 *Summary for Oxygenates: Warning*—The statistical data could be done on the oxygenates but there was not an equal number of all oxygenates in the round robin. MTBE was the largest contributor to the statistical results. The number of samples that contained each oxygenate is as follows:

Oxygenate Type	No. of Samples	Approximate Concentration Range
Ethanol	2	1 %, and 12 %
t-butanol	2	0.20 %, and 1.0 %
MTBE	6	1, 2, 4, 8 and 16 %
ETBE	1	0.50 %
TAME	1	15.00 %

A1.4 The precision statement for the olefins and cycloparaffins is determined by taking the square root of the value determined in the summary; multiply by the coefficient (r_{coef}) for repeatability and the coefficient (R_{coef}) for the reproducibility.

Name	r_{\min}	r_{coef}	r_{\max}	R_{\min}	R_{coef}	R_{\max}	C_{\min}	C_{\max}
Cyclo-paraffins	0.0726	0.08	0.098	0.286	0.384	0.586	2	10
Olefins	0.1555	0.18	0.21	0.382	0.555	1.012	2	25

A1.5 The precision for the aromatics does not depend on level and is stated below in mass percent.

Name	r_{\min}	r %	r_{\max}	R_{\min}	R %	R_{\max}	C_{\min}	C_{\max}
Aromatics	0.8549	0.98	1.155	2.151	2.706	3.651	15	50

A1.6 The summaries for the paraffins, isoparaffins, C_2 benzene and oxygenates follow the same procedure that was used for the analytes. The statistics for the grouping are shown in **Table A1.3** as an indication of reproducibility and repeatability of reporting the results as a group summary. However, there is a possibility that significant error could occur due to co-elution of peaks, the presence of significant amounts of olefinic or naphthenic constituents, or both, above octane and the percent unknown in the sample. If more accurate summary results are needed that are not covered by the above precision statement, for some or all of the above families of components, please consider another ASTM test method.

TABLE A1.1 Component Retention Times and Properties

NOTE 1—The names used are from several other tables and changes have been made where the GCMS did not agree with the peak name or its retention time.

NOTE 2—*n*-propanol will coelute with 3M-1-C5=.

NOTE 3—MTBE will coelute with 23DN-1C4=.

NOTE 4—MSBE will coelute with 1-hexene.

NOTE 5—ETBE will coelute with 23DM-13C4=.

NOTE 6—*isobutanol* will coelute with 44DM-1-c5=.

NOTE 7—233TM pentane will coelute with toluene when the ratio with toluene is greater than 5.0:1.

NOTE 8—The coeluting olefins in Notes 2-6 will usually be below 1000 ppm.

NOTE 9—In some instances the chemical group is known, but the chemical structure is not known (for example, C₆-olefin; the position of the double bond is not known).

NOTE 10—Relative response factors for six of the major oxygenated compounds have been determined by using the average results from seven laboratories analyzing six samples in duplicate. These same samples were used to determine linearity of methanol, ethanol, *t*-butanol, MTBE, ETBE and TAME from a concentration level ranging from 1 % mass up to 30 % mass.

Peak No.	New Name	Retention Time	Molecular Mass, MWt	Theoretical Mass, Rf, (C1)
1	Methane	6.74	16.04	1.000
2	Ethene	7.10	28.05	0.874
3	Ethane	7.21	30.07	0.937
4	Propene	7.41	42.05	0.874
5	Propane	7.87	44.11	0.916
6	Isobutane	8.26	58.12	0.906
7	Methanol	8.64	32.03	2.672
8	Isobutene	8.95	56.11	0.874
9	1-butene	8.99	56.11	0.874
10	1,3-butadiene	9.17	54.09	0.843
12	<i>N</i> -butane	9.28	58.12	0.906
14	Trans-2-butene	9.70	56.11	0.874
16	2,2-dimethylpropane	9.82	72.15	0.899
18	Cis-2-butene	10.33	56.11	0.874
20	1,2-butadiene	10.88	54.09	0.843
22	Ethanol	11.39	46.07	1.862
24	3-methyl-1-butene	12.21	70.13	0.874
26	Isopentane	13.57	72.15	0.899
28	1,4-pentadiene	14.25	68.12	0.849
30	2-Butyne (dimethylacetylene)	14.57	54.09	0.843
32	1-pentene	15.03	70.13	0.874
34	Isopropanol	15.28	60.11	1.950
36	2-methyl-1-butene	15.76	70.13	0.874
38	<i>N</i> -pentane	16.24	72.15	0.899
40	2-methyl-1,3-butadiene	16.73	68.12	0.849
42	Trans-2-pentene	17.23	70.13	0.874
44	3,3-dimethyl-1-butene	17.86	84.16	0.874
46	Cis-2-pentene	18.17	70.13	0.874
48	Tert-butanol (TBA)	18.51	74.12	1.161
50	2-methyl-2-butene	18.76	70.13	0.874
52	Trans-1,3-pentadiene	19.12	68.12	0.849
54	3-methyl-1,2-butadiene	19.48	68.12	0.849
56	Cyclopentadiene	19.76	67.10	0.824
58	Cis-1,3-pentadiene	20.25	68.12	0.849
60	1,2-pentadiene	20.51	68.12	0.849
62	2,2-dimethylbutane	20.69	86.18	0.895
64	Cyclopentene	23.16	68.12	0.849
66	4-methyl-1-pentene	24.30	84.16	0.874
68	3-methyl-1-pentene	24.38	84.16	0.874
70	<i>n</i> -propanol	24.68	60.11	1.770
72	Cyclopentane	24.86	70.13	0.874
74	2,3-dimethylbutane	25.57	86.18	0.895
76	2,3-dimethyl-1-butene	25.99	84.16	0.874
78	Methyl tert-butyl ether (MTBE)	26.18	88.09	1.407
80	Cis-4-methyl-2-pentene	26.48	84.16	0.874
82	2-methylpentane	26.66	86.18	0.895
84	Trans-4-methyl-2-pentene	72.09	84.16	0.874
86	Methyl ethyl ketone (MEK)	28.00	72.06	1.570
88	3-methylpentane	29.15	86.18	0.895
90	C ₆ -olefin	29.61	84.16	0.874
92	2-methyl-1-pentene	30.29	84.16	0.874
94	1-hexene	30.52	84.16	0.874
96	Methyl sec-butyl ether (MSBE)	30.66	88.09	1.550

TABLE A1.1 *Continued*

Peak No.	New Name	Retention Time	Molecular Mass, MWt	Theoretical Mass, Rf, (C1)
98	C ₆ -olefin	30.94	84.16	0.874
100	2-butanol	31.56	74.12	1.600
102	2ethyl-1-butene	32.47	84.16	0.874
104	<i>N</i> -hexane	32.75	86.18	0.895
106	Cis-3-hexene	33.41	84.16	0.874
108	Di-isopropyl ether (DIPE)	33.58	102.00	1.600
110	Trans-3-hexene+hexadiene	33.86	84.16	0.874
112	2-methyl-2-pentene	34.33	84.16	0.874
114	3-methylcyclopentene	34.57	82.10	0.853
116	Trans-3-methyl-2-pentene	34.71	84.16	0.874
118	Cis-2-hexene	35.62	84.16	0.874
120	3,3-dimethyl-1-pentene	36.04	98.19	0.874
122	Cis-3-methyl-2-pentene	36.92	84.16	0.874
124	Ethyl tert-butyl ether (ETBE)	37.07	102.18	1.255
126	2,3-dimethyl-1,3-butadiene	37.19	82.10	0.853
128	Methylcyclopentane	37.40	84.16	0.874
130	2,2-dimethylpentane	37.60	100.21	0.892
132	4,4-dimethyl-1-pentene	37.91	98.19	0.874
134	Isobutanol	38.06	74.12	1.500
136	2,3-dimethyl-2-butene	38.30	84.16	0.874
138	2,4-dimethylpentane	38.99	100.21	0.892
140	1,3,5-hexatriene	39.31	80.06	0.832
142	2,2,3-trimethylbutane	39.48	100.21	0.892
144	Methylcyclopentadiene	40.17	80.06	0.832
146	C ₇ -olefin	40.30	98.19	0.874
148	C ₇ -olefin	40.68	98.19	0.874
150	C ₇ -diolefin	41.20	96.18	0.856
152	4-methylcyclopentene	41.44	82.10	0.853
154	Methylenecyclopentane	42.08	82.10	0.853
156	Benzene	42.30	78.05	0.812
158	1-methyl-1-cyclopentene	42.46	82.10	0.853
160	C ₇ -olefin	43.06	98.19	0.874
162	C ₂ -methyl-3-hexene	43.37	98.19	0.874
164	3,3-dimethylpentane+5-methyl-1-hexene	43.81	100.21	0.892
166	Cyclohexane	44.07	84.16	0.874
168	Trans-2methyl-3-hexene	44.82	98.19	0.874
170	3,3-dimethyl-1,4-pentadiene	45.44	96.18	0.856
172	<i>N</i> -butanol	45.58	74.12	1.500
174	Dimethylcyclopentadiene	45.69	94.17	0.838
176	t,2-ethyl-3-methyl-1-butene	45.97	98.19	0.874
178	4-methyl-1-hexene	46.27	98.19	0.874
180	C ₇ -olefin	46.55	98.19	0.874
182	3-methyl-1-hexene	46.78	98.19	0.874
184	4-methyl-2-hexene	46.92	98.19	0.874
186	2-methylhexane+C ₇ -olefin	47.29	100.21	0.892
188	2,3-dimethylpentane	47.51	100.21	0.892
190	Cyclohexene	47.65	82.10	0.853
192	Tert-amyl methyl ether (TAME)	48.10	102.18	1.210
194	C ₇ -olefin	48.46	98.19	0.874
196	C ₇ -olefin	48.64	98.19	0.874
198	3-methylhexane	49.05	100.21	0.892
200	C ₇ -olefin	49.47	98.19	0.874
202	C ₇ -olefin	49.62	98.19	0.874
204	Trans-1,3-dimethylcyclopentane	49.83	98.19	0.874
206	Cis-1,3-dimethylcyclopentane	50.40	98.19	0.874
208	Trans-1,2-dimethylcyclopentane	51.01	98.19	0.874
210	3-ethylpentane	51.21	100.10	0.892
212	C ₇ -olefin	51.43	98.19	0.874
214	2,2,4-trimethylpentane	51.61	114.23	0.890
216	C ₇ -olefin	51.75	98.19	0.874
218	1-heptene	52.05	98.19	0.874
220	C ₇ -olefin	52.18	98.19	0.874
222	2,3-dimethyl-1,3-pentadiene	52.69	96.18	0.874
224	C ₇ -diolefin	53.00	96.18	0.856
226	C ₇ -olefin	53.36	98.19	0.874
228	C ₇ -diolefin	53.81	96.18	0.856
230	C ₇ -diolefin	54.13	96.18	0.856
232	C ₇ -olefin	54.28	98.19	0.874
234	<i>N</i> -heptane	54.59	100.21	0.892
236	Cis-3-heptene	54.81	98.19	0.874
238	2-methyl-2-hexene	55.10	98.19	0.874
240	Cis-methyl-3-hexene	55.35	98.19	0.874
242	Trans-3-heptene	55.72	98.19	0.874
244	3-ethyl-2-pentene	55.88	96.18	0.856

TABLE A1.1 *Continued*

Peak No.	New Name	Retention Time	Molecular Mass, MWt	Theoretical Mass, Rf, (C1)
246	1,5-dimethylcyclopentene	56.06	96.18	0.856
248	Trans-2-methyl-3-hexene	56.58	98.19	0.874
250	C ₇ -diolefin+C ₇ -triolefin	57.01	96.18	0.856
252	2,3-dimethyl-2-pentene	57.35	98.19	0.874
254	3-ethylpentene	57.57	98.19	0.874
256	Methylcyclohexane	57.79	98.19	0.874
258	C ₇ -olefin	58.28	98.19	0.874
260	1,1,3-trimethylcyclopentane	58.79	112.22	0.874
262	2,2-dimethylhexane	59.29	114.10	0.890
264	2,3,4-trimethyl-1,4-pentadiene	59.45	110.21	0.859
266	3,3-dimethyl-1,5-hexadiene	59.79	110.21	0.859
268	C ₈ -olefin	60.12	98.19	0.874
270	Ethylcyclopentane	60.60	98.19	0.874
272	3-methylcyclohexene	60.99	96.18	0.856
274	Methylcyclohexadiene	61.14	94.17	0.838
276	2,2,3-trimethylpentane	61.22	114.10	0.890
278	2,5-dimethylhexane+C ₈ -olefin	61.59	114.23	0.890
280	2,4-dimethylhexane	61.91	114.23	0.890
282	C ₇ -triolefin+C ₈ -olefin	62.28	112.24	0.856
284	Trans,cis-1,2,4-trimethylcyclopentane	62.68	112.22	0.874
286	3,3-dimethylhexane+C ₈ -olefin	63.13	114.23	0.890
288	C ₇ -triolefin+C ₈ -olefin	63.39	112.24	0.856
290	C ₈ -olefin	63.69	112.22	0.874
292	Trans,cis-1,2,3-trimethylcyclopentane	64.27	112.22	0.874
294	C ₈ -olefin	64.52	112.22	0.874
296	C ₈ -olefin	64.73	112.22	0.874
298	C ₈ -olefin	64.82	112.22	0.874
300	2,3,4-trimethylpentane	64.94	114.23	0.890
302	C ₇ -diolefin	65.25	96.18	0.856
304	Toluene	65.50	92.06	0.821
306	2,3,3-trimethylpentane	65.76	114.23	0.890
308	C ₈ -olefin	65.90	112.22	0.874
310	C ₈ -diolefin	66.12	110.21	0.859
312	C ₈ -olefin	66.48	112.22	0.874
314	C ₈ -olefin	66.65	112.22	0.874
316	C ₈ -olefin	67.08	112.22	0.874
318	C ₈ -diolefin+C ₈ -olefin	67.30	110.21	0.859
320	2,3-dimethylhexane	67.47	114.23	0.890
322	2-methyl-3-ethylpentane	67.71	114.23	0.890
324	1,1,2-trimethylcyclopentane+C ₇ -triolefin	68.04	112.22	0.874
326	C ₈ -diolefin+C ₈ -paraffin	68.31	114.23	0.859
328	C ₈ -olefin	68.41	112.22	0.874
330	C ₈ -olefin	68.64	112.22	0.874
332	2-methylheptane	68.86	114.23	0.890
334	4-methylheptane	69.11	114.23	0.890
336	C ₈ -diolefin+C ₇ -olefin	69.41	112.22	0.874
338	C ₈ -olefin	69.70	112.22	0.874
340	Cis-1,3-dimethylcyclohexane	69.91	112.22	0.874
342	Trans-1,4-dimethylcyclohexane	70.01	112.22	0.874
344	3-methylheptane	70.23	114.23	0.890
346	3-ethylhexane	70.38	114.23	0.890
348	C ₈ -diolefin	70.51	110.21	0.874
350	C ₈ -olefin	70.72	112.22	0.874
352	C ₈ -olefin	70.92	112.22	0.874
354	1,1-dimethylcyclohexane	71.18	112.22	0.874
356	C ₈ -olefin	71.43	112.22	0.874
358	C ₈ -olefin	71.70	112.22	0.874
360	Cis-1-ethyl-3-methylcyclopentane	72.10	112.22	0.874
362	2,2,5-trimethylhexane	72.23	128.26	0.888
364	Trans-1-ethyl-3-methylcyclopentane	72.46	112.22	0.874
366	Trans-1-ethyl-2-methylcyclopentane	72.68	112.22	0.874
368	1-methyl-1-ethylcyclopentane	72.96	112.22	0.874
370	1-octene	73.16	112.22	0.874
372	C ₈ -olefin	73.26	112.22	0.874
374	Trans-1,2-dimethylcyclohexane	73.36	112.22	0.874
376	C ₈ -olefin	73.48	112.22	0.874
378	C ₈ -olefin	73.68	112.22	0.874
380	Trans-3-C ₈ -Olefin	74.08	112.11	0.874
382	C ₈ -olefins	74.45	112.22	0.874

TABLE A1.1 *Continued*

Peak No.	New Name	Retention Time	Molecular Mass, MWt	Theoretical Mass, Rf, (C1)
384	Trans-1,3-dimethylcyclohexane	74.66	112.22	0.874
386	Cis-1,4-dimethylcyclohexane	74.79	112.22	0.874
388	<i>N</i> -octane	74.98	114.23	0.890
390	C ₈ -olefin	75.33	112.22	0.874
392	C ₈ -olefin	75.49	112.22	0.874
394	Trans-2-octene	75.62	112.22	0.874
396	Isopropylcyclopentane	75.72	112.22	0.874
398	C ₉ -olefin	75.85	126.24	0.874
400	C ₉ -olefin	75.89	126.24	0.874
402	C ₉ -olefin	75.90	126.24	0.874
404	C ₉ -olefin	76.08	126.24	0.874
406	2,2,4-trimethylhexane	76.31	128.26	0.888
408	2,4,4-trimethylhexane	76.62	128.26	0.888
410	C ₉ -olefins	76.86	126.24	0.874
412	2,3,5-trimethylhexane	77.29	128.26	0.888
414	Cis-2-octene	77.53	112.22	0.874
416	2,2,3,4-tetramethylpentane	77.77	128.26	0.888
418	2,2-dimethylheptane	78.02	128.26	0.888
420	Cis-1,2-dimethylcyclohexane	78.36	112.22	0.874
422	2,4-dimethylheptane	78.74	128.26	0.888
424	C ₉ -olefin	78.90	126.24	0.874
426	C ₉ -olefin	79.08	126.24	0.874
428	Ethylcyclohexane	79.24	112.22	0.874
430	Propylcyclopentane	79.39	112.22	0.874
432	2-methyl-4-ethylhexane	79.59	128.26	0.888
434	2,6-dimethylheptane	79.74	128.26	0.874
436	C ₉ -olefin	79.85	126.24	0.874
438	1,1,4-trimethylcyclohexane	80.05	126.24	0.874
440	C ₉ -olefin	80.28	126.24	0.874
442	C ₉ -olefin	80.38	126.24	0.874
444	1,1,3-trimethylcyclohexane	80.52	126.24	0.874
446	2,5 & 3,5-dimethylheptane	80.69	128.26	0.888
448	C ₉ -olefin	80.88	126.24	0.874
450	3,3-DMheptane	81.00	128.26	0.888
452	C ₉ -paraffin	81.13	128.26	0.888
454	C ₉ -olefin	81.34	126.24	0.874
456	2,3,3-trimethylhexane	81.56	128.26	0.888
458	C ₉ -olefin	81.68	126.24	0.874
460	Ethylbenzene	81.96	106.08	0.827
462	C ₉ -olefin	82.00	126.24	0.874
464	Trans-1,2,4-trimethylcyclohexane	82.31	126.24	0.874
466	C ₉ -olefin	82.33	126.24	0.874
468	2,3,4-trimethylhexane	82.63	128.26	0.888
470	C ₉ -olefin	82.73	126.24	0.874
472	3,3,4-trimethylhexane	82.89	128.26	0.888
474	<i>M</i> -Xylene	83.30	106.08	0.827
476	<i>P</i> -Xylene	83.43	106.08	0.827
478	2,3-dimethylheptane	83.57	128.26	0.888
480	3,5-dimethylheptane	83.83	128.26	0.888
482	3,4-dimethylheptane	83.91	128.26	0.888
484	C ₉ -olefin	84.08	126.24	0.874
486	3-methyl-3-ethylhexane	84.26	128.26	0.888
488	C ₉ -olefin	84.41	126.24	0.874
490	4-ethylheptane	84.52	128.26	0.888
492	4-methyloctane+C ₉ -olefin	84.70	128.26	0.888
494	2-methyloctane	84.84	128.26	0.888
496	C ₉ -olefin	85.01	126.24	0.874
498	C ₉ -paraffin	85.18	128.26	0.888
500	C ₉ -olefin	85.36	126.24	0.874
502	3-ethylheptane	85.51	128.26	0.888
504	3-methyloctane	85.69	128.26	0.888
506	C ₉ -paraffin	85.87	126.24	0.874
508	Cis-1,2,4-trimethylcyclohexane	85.91	126.24	0.874
510	1,1,2-trimethylcyclohexane	86.05	126.24	0.874
512	<i>O</i> -Xylene	86.27	106.08	0.827
514	C ₉ -olefin	86.47	126.24	0.874
516	C ₉ -paraffin	86.57	128.26	0.888
518	C ₉ -paraffin	86.75	128.26	0.888
520	C ₉ -olefin	86.90	126.24	0.874
522	Trans-1-ethyl-4-methylcyclohexane	87.08	126.24	0.874
524	Cis-1-ethyl-4-methylcyclohexane	87.23	126.24	0.874
526	C ₉ -paraffin	87.49	128.26	0.888

TABLE A1.1 *Continued*

Peak No.	New Name	Retention Time	Molecular Mass, MWt	Theoretical Mass, Rf, (C1)
528	1-nonene	87.79	126.24	0.874
530	Isobutylcyclopentane	88.00	126.24	0.874
532	C ₉ -paraffin	88.45	128.26	0.888
534	Trans-3-nonene	88.65	126.24	0.874
536	Cis-3-nonene	88.82	126.24	0.874
538	C ₉ -paraffin	89.09	128.26	0.888
540	<i>n</i> -nonane	89.24	128.26	0.888
542	C ₁₀ -olefin	89.41	140.27	0.874
544	Trans-2-nonene	89.74	126.24	0.874
546	1-methyl-1-ethylcyclohexane	89.61	126.24	0.874
548	1-methyl-2-propylcyclopentane	89.96	126.24	0.874
550	C ₁₀ -olefin	90.09	140.27	0.874
552	C ₁₀ -paraffin	90.18	142.28	0.887
554	C ₁₀ -paraffin	90.29	142.28	0.887
556	Isopropylbenzene	90.46	118.08	0.832
558	Cis-2-nonene	90.78	126.24	0.874
560	Tert-butylcyclopentane	90.80	126.24	0.874
562	C ₉ -olefins	90.88	126.24	0.874
564	Nonene	91.16	126.24	0.874
566	Isopropylcyclohexane	91.32	126.24	0.874
568	3,3,5-trimethylheptane	91.44	142.28	0.887
570	2,2-dimethyloctane	91.60	142.28	0.887
572	2,4-dimethyloctane	91.67	142.28	0.887
574	1-methyl-4-isopropylcyclohexane	91.82	140.27	0.874
576	Sec-butylcyclopentane	92.20	126.24	0.874
578	Propylcyclohexane	92.40	126.24	0.874
580	2,5-dimethyloctane	92.59	142.28	0.887
582	Butylcyclopentane	92.89	126.24	0.874
584	2,6-dimethyloctane	93.04	142.28	0.887
586	3,6-dimethyloctane	93.43	142.28	0.887
588	1-methyl-2-ethylcyclohexane	93.59	126.24	0.874
590	C ₁₀ -olefin	93.79	140.27	0.874
592	Propylbenzene	93.96	120.20	0.832
594	3,3-dimethyloctane	94.27	142.28	0.887
596	3-methyl-5-ethylheptane	94.54	142.28	0.887
598	C ₁₀ -olefin	94.66	140.27	0.874
600	1-ethyl-3-methylbenzene	94.88	120.20	0.832
602	1-ethyl-4-methylbenzene	95.09	120.20	0.832
604	Naphthene	95.30	140.27	0.874
606	1,3,5-trimethylbenzene	95.73	120.20	0.832
608	2,3-dimethyloctane	95.94	142.28	0.887
610	5-methylnonane	96.13	142.28	0.887
612	4-methylnonane	96.29	142.28	0.887
614	2-methylnonane	96.49	142.28	0.887
616	1-ethyl-2-methylbenzene	96.77	120.20	0.832
618	3-ethyloctane	97.01	142.28	0.887
620	Naphthene	97.14	140.27	0.874
622	3-methylnonane	97.47	142.28	0.887
624	C ₁₀ -olefin	97.69	140.27	0.874
626	C ₁₀ -paraffin	97.83	142.28	0.887
628	C ₁₀ -paraffin	98.16	142.28	0.887
630	1,2,4-trimethylbenzene	98.49	120.20	0.832
632	C ₁₀ -paraffin	98.74	142.28	0.997
634	C ₁₀ -paraffin	98.90	142.28	0.887
636	Isobutylcyclohexane	99.10	140.27	0.874
638	C ₁₀ -paraffin	99.09	142.28	0.887
640	C ₁₀ -paraffin	99.22	142.37	0.887
642	1-decene	99.52	140.27	0.874
644	C ₁₀ -paraffin	99.66	142.28	0.887
646	C ₁₀ -paraffin	99.70	142.28	0.887
648	C ₁₀ -aromatic	99.75	134.22	0.837
650	C ₁₀ -paraffin	99.82	142.28	0.887
652	Naphthene	99.93	140.27	0.874
654	Isobutylbenzene	100.06	134.22	0.837
656	Trans-1-methyl-2-propylcyclohexane	100.09	140.27	0.874
658	C ₁₀ -paraffin	100.19	142.28	0.887
660	Sec-butylbenzene	100.28	134.22	0.837
662	<i>n</i> -decane	100.40	142.28	0.887
664	C ₁₁ -paraffin	100.67	156.32	0.886
666	C ₁₁ -paraffin	100.85	156.32	0.886
668	1,2,3-trimethylbenzene	101.28	120.20	0.832
670	1-methyl-3-isopropylbenzene	101.40	134.22	0.837
672	C ₁₁ -paraffin	101.55	156.32	0.886

TABLE A1.1 *Continued*

Peak No.	New Name	Retention Time	Molecular Mass, MWt	Theoretical Mass, Rf, (C1)
674	1-methyl-4-isopropylbenzene	101.73	134.22	0.837
676	C ₁₁ -paraffin	102.06	156.32	0.886
678	C ₁₁ -paraffin	102.05	156.32	0.886
680	2,3-dihydroindene	102.42	118.17	0.819
682	Sec-butylcyclohexane	102.57	140.27	0.874
684	C ₁₁ -paraffin	102.87	156.32	0.886
686	1-methyl-2-isopropylbenzene	103.03	134.22	0.837
688	3-ethylnonane	103.26	156.32	0.886
690	C ₁₁ -paraffin	103.37	156.32	0.886
692	Naphthene	103.55	140.27	0.874
694	C ₁₁ -paraffin	103.88	126.19	0.886
696	1,3-diethylbenzene	104.08	134.22	0.837
698	1-methyl-3-propylbenzene	104.35	134.22	0.837
700	1,4-diethylbenzene	104.57	134.22	0.837
702	1-methyl-4-propylbenzene	104.73	134.22	0.837
704	Butylbenzene	104.85	134.22	0.837
706	3,5-dimethyl-1-ethylbenzene	105.00	134.22	0.837
708	1,2-diethylbenzene	105.26	134.22	0.837
710	C ₁₁ -paraffin	105.39	156.32	0.886
712	C ₁₀ -aromatic	105.49	134.22	0.837
714	C ₁₀ -aromatic	105.64	134.22	0.837
716	C ₁₀ -aromatic	105.75	134.22	0.837
718	1-methyl-2-propylbenzenes	105.85	134.22	0.837
720	C ₁₀ -aromatic	105.95	134.22	0.837
722	5-methyldecane	106.11	156.32	0.886
724	4-methyldecane	106.26	156.32	0.886
726	2-methyldecane	106.39	156.32	0.886
728	C ₁₁ -paraffin	106.55	156.32	0.886
730	1,4-dimethyl-2-ethylbenzene	106.76	134.22	0.837
732	1,3-dimethyl-4-ethylbenzene	106.93	134.22	0.837
734	C ₁₁ -paraffin	107.06	156.32	0.886
736	3-methyldecane	107.27	156.32	0.886
738	C ₁ - indane	107.35	132.00	0.837
740	1,2-dimethyl-4-ethylbenzene	107.46	134.22	0.837
742	C ₁₁ -paraffin	107.76	156.32	0.886
744	1,3-dimethyl-2-ethylbenzene	108.01	134.22	0.837
746	C ₁₁ -paraffin	108.58	156.32	0.886
748	C ₁₁ -paraffin	108.75	156.32	0.886
750	1-methyl-4-tert-butylbenzene	108.98	148.25	0.840
752	1,2-dimethyl-3-ethylbenzene	109.17	134.22	0.837
754	1-ethyl-2-isopropylbenzene	109.50	148.25	0.840
756	N-undecane	109.62	156.32	0.886
758	1-ethyl-4-isopropylbenzene	109.80	148.25	0.840
760	C ₁₂ -paraffin	109.96	170.34	0.885
762	1,2,4,5-tetramethylbenzene	110.15	134.22	0.837
764	2-methylbutylbenzene	110.55	148.25	0.840
766	1,2,3,5-tetramethylbenzene	110.43	134.22	0.837
768	3 methylbutylbenzene	110.64	148.25	0.840
770	C ₁₁ -aromatic	110.74	148.25	0.840
772	C ₁₂ -paraffin	110.84	170.34	0.885
774	C ₁₁ -aromatic	110.94	148.25	0.840
776	C ₁₁ -aromatic	111.05	148.25	0.840
778	C ₁₁ -aromatic	111.12	148.25	0.840
780	1-tert-butyl-2-methylbenzene	111.56	148.25	0.840
782	C ₁₁ -aromatic	111.65	148.25	0.840
784	1-ethyl-2-propylbenzene	111.76	148.25	0.840
786	C ₁₁ -aromatic	112.00	148.25	0.840
788	C ₁₁ -aromatic	112.22	148.25	0.840
790	C ₁₁ -aromatic	112.34	148.25	0.840
792	1-methyl-3-butylbenzene	112.52	148.25	0.840
794	C ₁₁ -aromatic	112.63	148.25	0.840
796	1,2,3,4-tetramethylbenzene	112.79	148.25	0.840
798	Pentylbenzene	113.17	148.25	0.840
800	Trans-1-methyl-2-(4methylpentyl)-cyclopentane	113.44	168.33	0.874
802	C ₁₁ -aromatic	113.74	148.25	0.840
804	C ₁₁ -aromatic	113.85	148.25	0.840
806	C ₁₁ -aromatic	114.02	148.25	0.840
808	C ₁₂ -paraffin	114.12	170.34	0.885
810	1,2,3,4-tetrahydronaphthalene	114.17	132.09	0.824
812	1-tert-butyl-3,5-dimethylbenzene	114.32	162.30	0.843
814	Naphthalene	114.65	128.06	0.799
816	1,1-dimethylindane	114.94	146.10	0.829
818	1,2-dimethylindane	115.19	146.10	0.829
820	1,6-dimethylindane	115.33	146.10	0.829

TABLE A1.1 *Continued*

Peak No.	New Name	Retention Time	Molecular Mass, MWt	Theoretical Mass, Rf, (C1)
822	C ₁₁ -aromatic	115.55	148.25	0.885
824	1- ethylindane	115.65	146.10	0.829
826	2- ethylindane	115.88	146.10	0.829
828	Ethyl -1,3,5-trimethylbenzene	116.00	148.25	0.884
830	1,3-dipropylbenzene	116.21	162.34	0.843
832	<i>n</i> -dodecane	116.55	170.34	0.885
834	Ethyl -1,2,4-trimethylbenzene	116.69	148.25	0.840
836	C ₁₁ -aromatic	117.07	148.25	0.840
838	C ₁₁ -aromatic	117.19	148.25	0.840
840	C ₁₂ -aromatic+C ₂ -indane	117.55	162.30	0.843
842	2,4-dimethylindane	117.99	146.10	0.829
844	4-ethylindane	118.13	146.10	0.829
846	1-tert-butyl-4-ethylbenzene	118.59	162.30	0.843
848	1,3-dimethylindane	119.07	146.10	0.829
850	1-methyl-4-pentylbenzene	119.60	162.30	0.843
852	4,7-dimethylindane	119.65	146.10	0.829
854	5,6-dimethylindane	119.70	146.10	0.829
856	C ₁₂ -aromatic	119.77	162.30	0.843
858	Hexylbenzene	119.87	162.30	0.843
860	C ₆ -benzene	119.93	162.30	0.843
862	C ₆ -benzene	119.98	162.30	0.843
864	C ₆ -benzene	120.20	162.30	0.843
866	4,5-dimethylindane	120.30	146.10	0.829
868	C ₆ -benzene	120.80	163.30	0.843
870	2-methylnaphthalene	121.42	142.08	0.806
872	C ₆ -benzene	121.65	162.30	0.843
874	C ₆ -benzene	121.85	162.30	0.843
876	<i>n</i> -tridecane	122.06	184.22	0.884
878	1-methylnaphthalene	122.28	142.08	0.806
880	C ₆ -benzene	122.40	162.30	0.843
882	C ₂ -tetralin	122.80	160.20	0.843
884	C ₆ -benzene	123.20	162.30	0.843
886	C ₆ -benzene	124.00	162.30	0.843
888	C ₁₃ -paraffin	125.60	184.22	0.883
890	Trans-7-decene	126.34	140.20	0.874
892	<i>n</i> -tetradecane	126.60	198.34	0.883
895	2,6-dimethylnaphthalene	126.84	156.30	0.812
900	2,7-dimethylnaphthalene	126.97	156.30	0.812
905	<i>n</i> -tetradecane	127.10	198.34	0.883
910	1,3-dimethylnaphthalene	127.52	156.30	0.812
915	1,6-dimethylnaphthalene	127.69	156.30	0.812
920	1,5-dimethylnaphthalene	128.44	156.30	0.812
925	1,4-dimethylnaphthalene	128.31	156.30	0.812
930	Acenaphthalene	129.05	156.30	0.801
940	1,2-dimethylnaphthalene	129.92	156.30	0.812
950	<i>n</i> -pentadecane	131.10	212.34	0.883

TABLE A1.2 Repeatability and Reproducibility of IHA Determinations

NOTE 1—Brief explanation of header information:

- r_{\min} —lower 95 % confidence limit of r_{est} ,
- r_{est} —repeatability estimate in percent of concentration,
- r_{\max} —upper 95 % confidence limit of r_{est} ,
- R_{\min} , R_{est} , R_{\max} —same as above except for reproducibility,
- C_{\min} —lower concentration limit that r_{est} , R_{est} is applicable, and
- C_{\max} —upper concentration limit that r_{est} , R_{est} is applicable.

NOTE 2—Short analyte names were used for the above table and by using the number beside this name, it will correspond to a full name in [Annex A1](#).

NOTE 3—C₂benzene refer to grouping ethylbenzene, M, P, and O-xylene as a group.

NOTE 4—The numbers in the second column were used for the statistical analysis for the round robin for 1996. The numbers beside the names are the new numbers being used in the new presentation of the IHA Method.

GC/MS	No. for RR	IHA No.	IHA/ abbreviated Name	r_{\min}	r_{est}	r_{\max}	R_{\min}	R_{est}	R_{\max}	C_{\min}	C_{\max}
A	6	6	iC4	9.8	1.4	17.7	24.9	30.7	37.3	0.04	2.86
A	9	9	1C4=	10.4	16.7	25.1	28	36	45.4	0.01	0.14
A	11	12	nC4	10	12	14.2	27.1	31.7	36.6	0.92	8.51
A	12	14	t2C4=	12.1	15.7	19.8	28.2	36.8	47.1	0.03	0.31
A	14	18	c2C4=	14.2	15.4	16.7	25.2	31.1	37.9	0.03	0.29
A	20	24	3M1C4=	7.3	9.6	12.3	17.2	19.9	22.7	0.02	0.11
A	22	26	iC5	4.6	5.4	6.3	13.4	15.5	17.9	2.39	12.09

TABLE A1.2 *Continued*

GC/MS	No. for RR	IHA No.	IHA/ abbreviated Name	r _{min}	r _{est}	r _{max}	R _{min}	R _{est}	R _{max}	C _{min}	C _{max}
A	26	32	1C5=	5.9	7.5	9.4	17	20.6	24.7	0.06	0.4
A	28	36	2M1C4=	4.4	6.3	8.6	14.5	17.5	20.9	0.14	0.8
A	30	38	nC5	4.2	6.2	8.7	13.9	16.1	18.5	1	5.18
A	34	42	t2C5=	4.1	6.3	9.1	13	17.3	22.6	0.27	1.08
A	38	46	c2C5=	5.2	7.7	11	14.4	18.3	22.9	0.15	0.59
A	40	50	2M2C4=	3.9	6.2	9.2	15.2	18.1	21.4	0.44	1.78
A	42	52	t13C5=,=	4.5	10.2	19.6	22.1	31.1	42.2	0.01	0.05
A	52	62	222DMC4	2.9	3.7	4.7	9.8	12.9	16.6	0.07	2.16
A	54	64	cyC5=	4.6	9	15.5	15.6	20.3	25.9	0.07	0.25
A	56	66	4M1C5=	11.2	14.8	19	22.6	31.8	43.2	0.02	0.1
A	58	68	3M1C5=	8.3	12.1	17	37.1	50.5	66.8	0.04	0.12
A	62	72	cyC5	2.5	4.7	7.7	11.8	13.4	15.1	0.07	0.69
A	64	74	23DMC4	1.7	2.7	3.9	8.6	9.8	11.1	0.53	1.91
A	66	76	MTBE	1.9	3.2	5	9.1	12.3	16.2	0.12	15.73
A	70	80	c4M2C5=	5.1	7.1	9.7	27.4	43.7	65.4	0.02	0.09
A	74	82	2MC5	2.2	2.9	3.8	9.3	11	12.9	1.03	5.62
A	76	84	t4M2C5=	4.9	6.3	7.9	16.9	20.2	23.9	0.05	0.26
A	80	88	3MC5	2	2.7	3.5	7.7	9.1	10.7	0.58	3.25
A	84	92	2M1C5=	3.6	5.1	7	9.6	12.5	16.1	0.09	0.45
A	86	94	1C6=	3.9	6.4	9.9	15.1	19.9	25.7	0.04	0.26
A	96	104	nC6	2.5	4.6	7.7	11	13.3	15.8	0.25	3.23
A	98	106	c3C6=	4.4	6.5	9.1	12.5	16.3	20.9	0.08	0.48
A	102	110	t3C6=+C6=,=	2.9	5.2	8.4	9.4	12.4	15.9	0.17	0.93
A	103	112	2M2C5=	2.7	4.7	7.4	9.9	12	14.4	0.15	0.77
A	104	114	3McyC5=	7.8	11.3	15.9	22.7	25.2	28	0.02	0.11
A	105	116	t3M2C5=	4.3	6.9	10.2	10.1	12.5	15.4	0.1	0.48
A	106	118	c2C6=	4.1	6.7	10.2	14.3	17.4	21	0.07	0.4
A	109	122	c3M2C5=	3.1	4.5	6.4	9.1	10.5	12.1	0.14	0.75
A	112	128	McyC5	2.4	3.3	4.4	9.1	10.1	11.1	0.36	2.34
A	116	138	24DMC5	1.8	2.7	3.9	8	10.1	12.4	0.2	1.93
A	118	142	223TMC4	0.5	4.1	14.3	20.9	35.2	54.8	0.01	0.06
A	124	150	C7=,=	0	3.1	16.6	11.3	19.1	29.9	0.01	0.04
A	128	154	methyl- enecyC5	5.5	9.1	14.1	14.9	20.3	26.8	0.01	0.03
A	130	156	Benzene	2.6	4.7	7.8	11.5	13.8	16.5	0.15	1.86
A	131	158	1McyC5=	4.3	6.3	8.9	18.5	24.1	30.7	0.17	0.92
A	133	162	c2M3C6=	0	1.2	6.8	17	29.1	45.9	0.01	0.06
A	134	164	33DMC5+5M1C6=	2.3	3.9	6.2	8.5	14.8	23.6	0.02	0.22
A	136	166	cyC6	3.3	4.4	5.7	11.3	12.8	14.5	0.04	0.87
A	138	168	t2M3C6=	4.2	8.4	14.7	84.2	103.2	124.8	0.02	0.32
A	146	176	t2e3m1C4=	3.2	5.7	9.1	20.8	29.6	40.8	0.02	0.19
A	148	178	4M1C6=	0.1	2.4	11.5	16.8	29.3	46.6	0.01	0.05
A	154	184	4M2C6=	3	4.5	6.4	15.9	18.7	21.8	0.03	0.29
A	156	186	2MC6+C7=	1.4	2.1	3	6.2	7.7	9.5	1.09	3.54
A	160	190	cyC6=	3.9	7.2	12.1	30.1	45.4	65.2	0.02	0.13
A	166	198	3MC6	1.3	2	2.8	8.5	9.9	11.5	0.36	2.38
A	172	204	t13DMcyC5	1.7	2.4	3.3	10.5	11.3	12.2	0.12	0.6
A	174	206	c13DMcyC5	1.9	2.7	3.6	9.8	10.7	11.6	0.09	0.49
A	176	208	t12DMcyC5	2.2	3.2	4.3	7.6	9.1	10.8	0.05	0.46
A	180	210	3EC5	2.8	4.8	7.6	10	13.4	17.6	0.02	0.21
A	184	212	5M-1-C6=	1.8	5	10.6	24.1	35.2	49.1	0.03	0.19
A	186	214	224TMC5	2.3	3.4	4.9	7.6	13.2	21.1	0.09	23.25
A	188	218	1C7=	4.3	6.8	10.1	15.8	20.9	26.9	0.02	0.13
A	189	220	C7=	5.2	7.8	11.1	15.1	18.3	22	0.02	0.13
A	194	226	C7=	3.3	4.8	6.8	16.6	20.7	25.2	0.02	0.16
A	196	228	C7=,=	3.7	5	6.5	12.5	17.2	22.8	0.04	0.31
A	197	230	C7=,=	5.6	7.3	9.3	19.5	23	26.9	0.04	0.26
A	198	232	C7=	3.8	4.7	5.7	42.9	60.4	82.1	0.05	0.45
A	200	234	nC7	1.5	2.2	3.2	7.4	8.9	10.7	0.13	1.55
A	202	236	c3C7=	2.1	3	4.2	14.2	18.2	23	0.04	0.36
A	204	238	2M2C6=	2.1	3	4.3	14.4	16.5	18.7	0.05	0.43
A	206	240	c3M3C6=	3.3	4.5	6.1	21	24.9	29.3	0.03	0.29
A	208	242	t3C7=	1.8	2.7	4	12.9	15.2	17.8	0.04	0.35
A	210	244	3E2C5=	0.1	1.2	5.4	13.4	16.6	20.4	0.02	0.13
A	212	246	1,5DMcyC5=	3	5	7.8	10.3	16.2	24	0.03	0.27
A	214	248	t2M3C6=	2.8	3.6	4.7	13.8	17.9	22.9	0.04	0.33
A	218	252	23DM2C5=	3.1	4	5	9.1	13	17.8	0.04	0.56
A	222	256	McyC6	1.9	2.6	3.6	8.5	9.9	11.5	0.16	1.44
A	224	260	113TMCyC5	1.7	5.1	11.5	10.8	14.4	18.7	0.01	0.09
A	226	262	22DMC6	4.7	9.2	15.9	12.9	23.2	38.1	0.01	0.07
A	234	270	EcyC5	2.5	3.6	5	9.6	13.5	18.4	0.04	0.3
A	240	276	223TMC5	2.2	4.9	9.3	14.1	27.3	46.7	0.02	0.54
A	245	278	25DMC6+C8=	1.5	2.8	4.7	6.3	8.1	10.3	0.17	1.58

TABLE A1.2 *Continued*

GC/MS	No. for RR	IHA No.	IHA/ abbreviated Name	r _{min}	r _{est}	r _{max}	R _{min}	R _{est}	R _{max}	C _{min}	C _{max}
A	250	280	24DMC6	1.8	2.9	4.5	6.1	8.1	10.4	0.25	2.19
A	260	284	tc124TMcyC5	2.4	3.7	5.4	10.8	15.1	20.5	0.03	0.16
A	265	286	3,3DMC6+C8=	1.3	5.4	14.1	8.7	14.8	23.2	0.01	0.07
A	278	292	tc123TMcyC5	6.1	11.5	19.5	40.9	70	110.3	0.03	0.09
A	290	298	C8=S	0.3	3.2	11.8	15.5	20.3	26.1	0.02	0.23
A	292	300	234TMC5	1.9	3.2	5	8.7	12	16	0.09	9.14
A	294	302	C7=,=	2.9	4.2	5.8	19.2	41.1	75.2	0.06	0.51
A	300	304	Toluene	1.7	3.1	5.3	8.7	16.6	28.2	2.52	13.14
A	312	316	C8=	3.9	6	8.7	26	35.7	47.6	0.02	0.2
A	314	320	23DMC6	2.2	3.5	5.2	16.1	30.6	51.9	0.18	2.06
A	316	322	2M3EC5	2.3	4.5	7.9	21.3	40	67.2	0.03	0.31
A	318	324	112TMcyC5+C7=,=,=	3.3	3.3	11.8	26.6	33.7	42	0.02	0.23
A	326	332	2MC7	3.3	4.4	5.9	8.4	11.2	14.5	0.14	0.93
A	328	334	4MC7	3.5	5.6	8.3	12.5	24.4	42.4	0.15	0.5
A	334	340	c13DMcyC6	3.7	4.8	6.2	18.7	32.6	52.1	0.04	0.25
A	336	344	3MC7	2.3	3.3	4.5	17.8	21.9	26.5	0.15	1.04
A	338	346	3EC6	4.1	6.4	9.4	34.8	53	76.7	0.04	0.21
A	352	360	c1E3McyC5	3.1	4.3	5.7	8.6	23.2	48.7	0.09	2.32
A	356	364	t1E3McyC5	3.8	5.1	6.7	24.4	35.5	49.7	0.03	0.21
A	360	366	t1E2McyC5	4.5	7.7	12.3	32.3	54.1	84.1	0.02	0.11
A	362	368	1M1EcyC5	0.2	3.1	12.5	24.1	33.3	44.7	0.01	0.08
A	366	372	C8=	7.2	9.9	13.3	27.1	37	49	0.01	0.08
A	368	374	t12DMcyC6	2.2	4.8	9	63.9	97.3	140.6	0.02	0.15
A	372	378	C8=S	3.4	5.3	7.9	109.3	124.4	141	0.02	0.26
A	374	380	t-3-C8=	0	1.5	9.4	50.8	67.2	86.9	0.02	0.12
A	380	382	C8=	3.6	5.4	7.9	21.1	38.9	64.7	0.03	0.33
A	385	384	t13DMcyC6	3.1	5.4	8.4	34.1	48.5	66.5	0.04	0.31
A	400	388	nC8	3	3.7	4.5	8.8	11.9	15.6	0.1	0.89
A	406	394	t2C8=	3	6.5	12.2	45.6	72.5	108.4	0.02	0.28
A	408	396	iPrcyC5	5.8	7.4	9.3	31.7	50.8	76.5	0.03	0.36
A	416	404	C9=	0.3	2.9	9.9	46.9	63.8	84.4	0.02	0.14
A	422	410	C9=S	4.8	8	12.4	30.5	43.2	58.9	0.02	0.17
A	432	420	c12DMcyC6	3.4	4.9	6.8	22.1	39.3	63.8	0.04	0.39
A	434	422	24DMC7	5.6	9.9	15.9	54.5	105.5	181.2	0.02	0.09
A	436	424	C9=	1.9	6	13.7	34.7	47.5	63.1	0.01	0.07
A	438	426	C9=	4.1	6.6	10	19	27.7	38.7	0.02	0.11
A	440	428	EcyC6	2.7	5	8.2	14.1	22	32.5	0.03	0.28
A	444	432	2M4EC6	7.7	11.1	15.3	20.2	27.4	36	0.01	0.03
A	446	434	26DMC7	5.9	7.3	8.9	21.9	27.7	34.4	0.03	0.14
A	450	438	114TMcyC6	5.9	8.2	11	28	42.1	60.3	0.03	0.21
A	458	446	25&35DMc7	3.7	5.9	8.7	10.5	14.9	20.5	0.07	0.25
A	460	448	C9=S	3.3	8.4	17.1	40.1	56.4	76.6	0.01	0.07
A	462	450	33DMC7	0.1	3.3	15.7	25	44	70.9	0.01	0.05
A	475	460	EBenzene	2.8	3.9	5.4	7.2	8.9	10.9	0.66	3.12
A	480	464	t124TMcyC6	6.9	10.9	16.3	84.7	109.3	138.2	0.02	0.15
A	500	474	M-Xylene	2.7	3.7	5	7.5	9.2	11	1.67	7.93
A	502	476	P-Xylene	3.1	4.4	5.9	8.8	11.6	14.8	0.63	3.26
A	503	478	23DMC7	5.1	7.6	10.9	45.3	73.5	111.5	0.03	0.16
A	504	480	35DMC7	7.2	9.8	13	44.1	82.8	139.2	0.02	0.07
A	506	482	34DMC7	6.5	10.1	15	42.5	67.7	101.4	0.02	0.07
A	510	486	3M3EC6	6.3	10	15	38	61	92	0.02	0.14
A	518	492	4MC8+C9=	4.1	5.9	8.1	12.4	14.3	16.3	0.05	0.3
A	520	494	2MC8	4.4	5.9	7.7	12.4	15.9	20.1	0.07	0.38
A	522	496	C9=	6.8	10.6	15.7	22.3	33.3	47.4	0.01	0.1
A	528	502	3EC7	4.5	6.8	9.8	24.7	34.4	46.3	0.02	0.11
A	530	504	3MC8	5	8	12	12.4	17.9	24.9	0.08	0.45
A	550	512	O-Xylene	2.1	3	4.1	7.7	9.8	12.3	0.92	4.18
A	564	518	C9P	3.1	6.6	12	31.1	50.4	76.3	0.01	0.37
A	568	522	t1E4McyC6	6.5	9.7	13.8	26.3	46.1	74.1	0.02	0.13
A	570	524	c1E4McyC6	4.7	7.4	10.8	22.1	35.8	54.2	0.02	0.15
A	572	526	C9P	4.5	7.2	10.7	28.7	55.7	95.9	0.03	0.6
A	582	532	C9P	7.5	11.1	15.6	16.9	23.1	30.8	0.02	0.24
A	586	534	t3C9=	4.6	9.1	16	27.3	38.8	53.2	0.01	0.16
A	590	536	c3C9=	7.1	11.1	16.4	23.5	36.1	52.7	0.01	0.17
A	600	540	nC9	5.8	7.2	8.7	18.3	30	45.8	0.1	0.51
A	606	546	1M1EcyC6	0.4	3.1	10.7	46.2	75.8	116	0.02	0.11
A	608	548	1M2PrcyC5	0.2	3	12.2	19.2	30.1	44.5	0.01	0.1
A	616	556	iPrbenz	4.3	6.9	10.4	11.2	18.9	29.6	0.04	0.41
A	626	566	iPrcyC6	4.4	7.7	12.4	21.8	40.2	66.9	0.01	0.35
A	636	576	sBucyC5	0.5	4.5	16.1	22.9	36.7	55.1	0.01	0.06
A	638	578	PrcyC6	4	7.3	12	77.9	96.8	118.6	0.02	0.12
A	644	584	26DMC8	4.6	8.6	14.4	41.2	68.2	105	0.03	0.23
A	646	586	36DMC8	4.5	7.7	12.3	31.6	40.4	50.8	0.03	0.11

TABLE A1.2 *Continued*

GC/MS	No. for RR	IHA No.	IHA/ abbreviated Name	r _{min}	r _{est}	r _{max}	R _{min}	R _{est}	R _{max}	C _{min}	C _{max}
A	651	592	nPrbenz	3.5	5.8	9	11.6	17.3	24.6	0.21	0.83
A	655	600	1E3Mbenz	2.8	4.5	6.9	6.5	8.3	10.3	0.85	2.8
A	656	602	1E4Mbenz	3.1	4.5	6.3	7.8	9.7	11.9	0.36	1.26
A	658	606	135TMbenz	3.4	5.8	9.1	8.5	12.5	17.7	0.46	1.53
A	660	610	5MC9	10.9	12.9	15.1	76.7	104.7	138.8	0.02	0.13
A	661	612	4MC9	7.1	10.2	14	29.7	44.5	63.5	0.02	0.13
A	662	614	2MC9	4.4	7.1	10.9	14.9	24.2	36.6	0.1	2.07
A	663	616	1E2Mbenz	3.6	5.5	8.1	10.3	15.9	23.2	0.3	1.1
A	668	622	3MC9	7.2	12.9	21	41.8	59	80.3	0.04	0.19
A	671	626	C10-P	0.5	5.4	19.5	30.3	52.1	82.6	0.01	0.47
A	673	630	124TMbenz	2.8	4.7	7.4	9.3	12.5	16.4	1.29	4.65
A	674	632	C10-P	7.1	12.9	21.4	35.6	81.2	155.1	0.01	0.32
A	675	634	C10P	2.6	6.2	12.3	25.2	55.1	102.4	0.01	0.34
A	684	648	C10A	5.2	9.3	15.1	22.8	38.2	59.4	0.01	0.3
A	688	652	naphthene	4.8	7.5	11	40.2	63.2	93.7	0.03	0.27
A	700	662	nC10	7.3	8.9	10.7	14.3	29.5	52.8	0.07	0.29
A	705	668	123TMbenz	4	6.3	9.2	18.2	23.2	29.1	0.28	1.15
A	708	674	1M4iPrbenz	3	6.6	12.1	22	34.2	50.1	0.01	0.08
A	709	676	C11P	5.1	8.9	14.1	34.9	68.2	118.1	0.02	0.12
A	712	680	indan	4	6.6	10.1	15.7	23.6	33.8	0.15	0.4
A	714	682	sBucyC6	8.7	12.7	17.6	46.7	70.2	100.5	0.01	0.06
A	718	686	1M2iPrbenz	4.6	8.4	13.7	48	88.1	146	0.02	0.33
A	723	694	C11P	5	7.8	11.4	29.6	60.7	108.3	0.02	0.19
A	724	696	13DEbenz	4.6	6.1	8	11.1	19.5	31.5	0.07	0.22
A	725	698	1M3Prbenz	3.5	5.2	7.3	8.5	13	18.8	0.18	0.71
A	727	702	1M4Prbenz	4.8	7.8	11.7	16.7	22.8	30.2	0.1	0.35
A	728	704	Bubenz	7.2	11	16.1	15.8	21.8	29.3	0.04	0.14
A	729	706	35DM1EBenz	3.5	6.4	10.5	9.1	14	20.3	0.18	0.56
A	730	708	12DEbenz	6.4	9.7	14	38.6	57.4	81.4	0.02	0.09
A	740	718	1M2PrBenz	6.8	10.7	15.8	27.3	41.7	60.4	0.06	0.21
A	746	722	5MC10	7.1	11.5	17.5	30.8	44.5	61.8	0.02	0.08
A	748	724	4MC10	4.2	6.9	10.4	15.3	32.1	57.9	0.01	0.68
A	750	726	2MC10	6.5	9.5	13.3	52.7	68.9	88.2	0.02	0.15
A	756	730	14DM2Ebenz	4.1	6.1	8.7	17.4	26.3	37.9	0.12	0.42
A	758	732	13DM4Ebenz	4.5	6.2	8.3	18.5	22.9	27.8	0.12	0.54
A	762	736	3MC10	10.9	15.7	21.7	35.8	54.5	78.8	0.02	0.17
A	764	740	12DM4Ebenz+ C1indane	3.1	5.3	8.5	8.2	12.5	18.2	0.27	0.75
A	768	744	13DM2Ebenz	6.2	9.6	14	37.9	68.9	113.3	0.03	0.35
A	780	750	1M4tBubenz	6.1	10.3	16.1	45.8	83.5	137.7	0.03	0.11
A	785	752	12DM3Ebenz	4.1	7.3	11.7	28.2	45.3	68.2	0.09	0.2
A	800	756	nC11	8.7	11.1	13.9	31.2	40.2	50.6	0.04	0.21
A	806	762	1245tertM- benz	5.4	6.8	8.6	12.3	16.8	22.2	0.12	0.39
A	810	766	1235tertM- benz	4.7	7.7	11.6	12.7	19.9	29.3	0.16	0.56
A	824	782	C11A	8.7	11.3	14.2	32.9	55.6	86.9	0.02	0.07
A	826	784	1E2Prbenz?	5	7.5	10.7	14.2	25.2	40.8	0.09	0.44
A	828	786	C11A	8.5	11.8	15.7	23.4	35.1	50.3	0.02	0.1
A	830	788	C11A	8.8	12.3	16.7	35.7	49.9	67.5	0.02	0.1
A	832	790	C11A	9.7	13.4	17.8	22.9	39.6	63	0.02	0.1
A	834	792	1M3Bubenz	5.6	7.9	10.9	11.1	14.8	19.2	0.08	0.35
A	836	796	1234tetraM- benz +C11A	6.8	9.3	12.5	24.4	36.5	52.1	0.1	0.28
A	840	800	t1M2(4MC5)cyC5	10.2	15.5	22.3	41	56.7	75.8	0.02	0.11
A	844	804	C11A	9.1	13.5	19	34	54.7	82.5	0.02	0.07
A	846	806	C11A	9.6	13.6	18.5	65.6	96.4	135.5	0.02	0.08
A	854	812	1tBu35DMbenz	11.2	15.5	20.7	36.6	62.3	97.7	0.02	0.1
A	858	814	naphthalene	4.9	6.7	8.9	15.3	25.8	40.3	0.12	0.52
A	862	817	C11A	9.7	14.4	20.5	46.5	66.5	91.4	0.02	0.16
A	870	820	16DMIND- ANE	9	12.3	16.3	25.7	42.6	65.8	0.02	0.17
A	875	822	C11A	15.6	19.4	23.8	43.8	68.4	100.9	0.02	0.09
A	884	824	2ETHYL IN- DANE	5.8	9.8	15.4	18.4	29	42.9	0.03	0.19
A	888	826	2ETHYL135TMBZ	7.9	12.8	19.5	39.4	59.9	86.6	0.01	0.07
A	895	832	nC12	13.4	16.7	20.6	53.4	73.9	99.1	0.02	0.15
A	915	842	24DMIND- ANE	10.3	16.3	24.2	27.2	40	56.2	0.02	0.05
A	925	846	1tBu4Ebenz	7.7	13.1	20.7	60.2	101.8	159.2	0.04	0.16
A	930	848	13DM IN- DANE	5.3	10.3	17.9	31.3	43	57.3	0.01	0.18

TABLE A1.2 *Continued*

GC/MS	No. for RR	IHA No.	IHA/ abbreviated Name	r_{\min}	r_{est}	r_{\max}	R_{\min}	R_{est}	R_{\max}	C_{\min}	C_{\max}
^A	940	858	HEXYLbenz	9.8	15.1	21.9	61.2	96	141.8	0.01	0.13
^A	942	870	2Mnaphtha- lene	6.4	8.9	12.1	17	21.6	27	0.04	0.64
^A	947	879	1Mnaphtha- lene	7.5	11.6	16.9	25	29.8	35.2	0.02	0.27

^A The components that have been checked by GCMS by one of the participants on one of the samples that was used in the 1996 ASTM round robin.

TABLE A1.3 Group Summaries for the Gasolines Run in the 1996 ASTM Interlaboratory Cooperative Study

Name	r_{\min}	r_{est}	r_{\max}	R_{\min}	R_{est}	R_{\max}	C_{\min}	C_{\max}
Paraffin	0.0562	0.0646	0.08	0.125	0.186	0.373	1	20
Isoparaffin	0.0209	0.024	0.03	0.047	0.065	0.102	20	65
C ₂ Benzene	0.0334	0.0384	0.05	0.057	0.073	0.102	3	20
Oxygenates	0.0418	0.0491	0.06	0.104	0.141	0.221	3	20

A2. OXYGENATE LINEARITY STUDY

A2.1 This information is presented in **Tables A2.1-A2.7** and **Figs. A2.1-A2.13**.

A2.2 **Tables A2.2-A2.7** show comparisons between this test method and other methods for several compound types. Multidimensional PIONA is included since it tends to give reasonable peak compound type groupings for total olefins, total paraffins, and total naphthenes. The differences for benzene

and toluene among the indicated methods are well within the reproducibilities of the methods. The sample numbers refer to the interlaboratory cooperative study samples. It should be noted that the interlaboratory cooperative study samples included only spark ignition fuels and different results may be obtained with pure blending components.

TABLE A2.1 Oxygenates Relative Response Factors

NOTE 1—All RRF relative to $N-C_7 = 1.000$; this also applies to the cooperative study.

	Laboratory No. 1	Laboratory No. 2	Laboratory No. 3	Laboratory No. 4	Laboratory No. 5	Laboratory No. 6	Laboratory No. 7	Average RRF	Standard Deviation	% Standard Deviation
Methanol	2.921	2.957	2.903	2.795	3.085	3.391	2.923	2.996	.194	6.465
Ethanol	1.997	2.043	2.003	2.057	2.138	2.354	2.014	2.087	.127	6.1
t-butanol	1.274	1.282	1.329	1.305	1.297	1.429	1.2	1.302	.069	5.281
MTBE	1.508	1.523	1.552	1.791	1.508	1.658	1.498	1.577	.109	6.932
ETBE	1.352	1.349	1.406	1.543	1.369	1.509	1.319	1.407	.086	6.108
TAME	1.308	1.323	1.342	1.451	1.336	1.471	1.264	1.356	.076	5.593

TABLE A2.2 Benzene

Sample	Benzene (mass percent)	
	D5580	D6729
2	1.52	1.61
6	1.05	1.12
8	1.10	1.16
10	1.13	1.18
13	0.14	0.16
14	0.62	0.70
Average	0.93	0.99

TABLE A2.3 Toluene

Sample	Toluene (mass percent)	
	D5580	D6729
2	4.3	4.6
6	2.1	1.9
8	10.1	11.4
10	5.0	6.1
13	3.3	2.9
14	4.4	5.3
Average	4.9	5.4

TABLE A2.4 Total Aromatics

Sample	Total Aromatics (mass percent)		
	D5580	PIONA ^A	D6729
2	30.3	28.2	32.6
6	18.9	18.7	20.0
8	49.1	49.0	51.0
10	23.9	24.5	25.4
13	19.7	19.8	22.4
14	23.8	24.6	27.5
Average	27.6	27.5	29.8

^A Multidimensional PIONA.

TABLE A2.5 Total Olefins

Sample	Total Olefins (mass percent)	
	PIONA ^A	D6729
2	7.1	4.4
6	9.8	9.4
8	6.6	6.2
10	15.1	13.7
13	11.1	11.1
14	24.6	22.2
Average	12.4	11.2

^A Multidimensional PIONA.

TABLE A2.6 Total Oxygenates

Sample	Total Oxygenates (mass percent)	
	PIONA ^A	Procedure B
2 ^B	15.3	16.1
6 ^B	7.0	8.1
8 ^B	4.2	4.5
10 ^C	>8	10.0
13 ^B	20.5	19.9
14 ^B	2.8	3.2
Average	N/A	10.3

^A Multidimensional PIONA.

^B Major oxygenate = MTBE.

^C Major oxygenate = Ethanol.

TABLE A2.7 Total Paraffins and Total Naphthenes

Sample	Total Paraffins (mass percent)		Total Naphthenes (mass percent)	
	PIONA ^A	D6729	PIONA ^A	D6729
8	35.6	35.0	2.2	2.8
10	41.1	42.3	5.6	6.7
13	42.6	43.0	1.3	3.5
14	34.1	37.9	5.9	7.6
Average	38.4	39.6	3.8	5.2

^A Multidimensional PIONA.

Spl(mass%)						Avg RF	RRF
MEOH	1.01	5.05	10.02	20.01	29.83		
	37792	201545	406795	816960	1208524		
	38002	200204	409233	820596	1225686		
avg	37897	200874.5	408014	818778	1217105		
RF	2.67E-05	2.51E-05	2.46E-05	2.44E-05	2.45E-05	2.51E-05	2.920678
Spl(mass%)						Avg RF	RRF
ETOH	1	5	10.1	20.15	30.18		
	56107	288820	604107	1214248	1807248		
	52935	285869	597366	1223531	1830666		
avg	54521	287344.5	600736.5	1218890	1818957		
RF	1.83E-05	1.74E-05	1.68E-05	1.65E-05	1.66E-05	1.71E-05	1.997164
Spl(mass%)						Avg RF	RRF
TBA	0.964	4.9692	9.9583	19.8768	29.7953		
	89751	443262	899170	1830312	2742339		
	92269	441843	893544	1820174	2765568		
avg	91010	442552.5	896357	1825243	2753954		
RF	1.06E-05	1.12E-05	1.11E-05	1.09E-05	1.08E-05	1.09E-05	1.273649
Spl(mass%)						Avg RF	RRF
MTBE	0.9992	5.0362	9.9724	20.0248	30.0471		
	76166	391956	765248	1537935	2332931		
	77640	399654	761273	1535598	2332734		
avg	76903	395805	763260.5	1536767	2332833		
RF	1.3E-05	1.27E-05	1.31E-05	1.3E-05	1.29E-05	1.29E-05	1.507996
Spl(mass%)						Avg RF	RRF
ETBE	0.9851	4.9255	9.8707	19.6724	29.5727		
	86770	420851	852468	1689595	2515456		
	85993	420221	867050	1690395	2506966		
avg	86381.5	420536	859759	1689995	2511211		
RF	1.14E-05	1.17E-05	1.15E-05	1.16E-05	1.18E-05	1.16E-05	1.352309
Spl(mass%)						Avg RF	RRF
TAME	0.9997	4.9788	9.8883	19.153	29.7144		
	90368	443934	876234	1740744	2576420		
	88502	444981	874999	1762466	2584069		
avg	89435	444457.5	875616.5	1751605	2580245		
RF	1.12E-05	1.12E-05	1.13E-05	1.09E-05	1.15E-05	1.12E-05	1.308241
Spl(mass%)						Avg RF	RRF
Nc7	8.475	8.44	8.4525	8.4525	8.695		
	994302	951197	991971	982424	1006023		
	997469	983612	990664	1002009	1006083		
avg	995885.5	967404.5	991317.5	992216.5	1006053		
RF	8.51E-06	8.72E-06	8.53E-06	8.52E-06	8.64E-06	8.58E-06	1

FIG. A2.1 IHA Method Oxygenates Linearity Cooperative Study–Laboratory 1

Spl(mass%)						Avg RF	RRF
MEOH	1.01	5.05	10.02	20.01	29.83		
	44097	236256	478801	985095	1454605		
	44051	237455	480020	992190	1465533		
avg	44074	236855.5	479410.5	988642.5	1460069		
RF	2.29E-05	2.13E-05	2.09E-05	2.02E-05	2.04E-05	2.12E-05	2.956773
Spl(mass%)						Avg RF	RRF
ETOH	1	5	10.1	20.15	30.18		
	63749	332568	698238	1430974	2178293		
	62784	332799	701430	1431363	2204197		
avg	63266.5	332683.5	699834	1431169	2191245		
RF	1.58E-05	1.5E-05	1.44E-05	1.41E-05	1.38E-05	1.46E-05	2.04331
Spl(mass%)						Avg RF	RRF
TBA	0.964	4.9692	9.9583	19.8768	29.7953		
	108001	526541	1055347	2147710	3316200		
	110407	524386	1061356	2163089	3322481		
avg	109204	525463.5	1058352	2155400	3319341		
RF	8.83E-06	9.46E-06	9.41E-06	9.22E-06	8.98E-06	9.18E-06	1.282428
Spl(mass%)						Avg RF	RRF
MTBE	0.9992	5.0362	9.9724	20.0248	30.0471		
	90887	473216	910349	1794640	2777855		
	91715	476896	904173	1794196	2780266		
avg	91301	475056	907261	1794418	2779061		
RF	1.09E-05	1.06E-05	1.1E-05	1.12E-05	1.08E-05	1.09E-05	1.523223
Spl(mass%)						Avg RF	RRF
ETBE	0.9851	4.9255	9.8707	19.6724	29.5727		
	103792	516002	1020170	2007710	2980345		
	104863	518258	1035091	2007448	2983391		
avg	104327.5	517130	1027631	2007579	2981868		
RF	9.44E-06	9.52E-06	9.61E-06	9.8E-06	9.92E-06	9.66E-06	1.349418
Spl(mass%)						Avg RF	RRF
TAME	0.9997	4.9788	9.8883	19.153	29.7144		
	103829	523120	1050222	2077446	3083066		
	104085	517930	1057409	2115710	3084788		
avg	103957	520525	1053816	2096578	3083927		
RF	9.62E-06	9.56E-06	9.38E-06	9.14E-06	9.64E-06	9.47E-06	1.322771
Spl(mass%)						Avg RF	RRF
Nc7	8.475	8.44	8.4525	8.4525	8.695		
	1198960	1190806	1178498	1177607	1195493		
	1198844	1190899	1178015	1176611	1212114		
avg	1198902	1190853	1178257	1177109	1203804		
RF	7.07E-06	7.09E-06	7.17E-06	7.18E-06	7.22E-06	7.15E-06	1

FIG. A2.2 IHA Method Oxygenates Linearity Cooperative Study–Laboratory 2

Spl(mass%)						Avg RF	RRF
MEOH	1.01	5.05	10.02	20.01	29.83		
	151533	864732	1741799	3589766	5293556		
	164863	854798	1759435	3746174	5368227		
avg	158198	859765	1750617	3667970	5330892		
RF	6.38E-06	5.87E-06	5.72E-06	5.46E-06	5.6E-06	5.81E-06	2.903282
Spl(mass%)						Avg RF	RRF
ETOH	1	5	10.1	20.15	30.18		
	245820	1078429	2521533	5099484	7899031		
	257618	1197628	2511218	5200823	8259533		
avg	251719	1138029	2516376	5150154	8079282		
RF	3.97E-06	4.39E-06	4.01E-06	3.91E-06	3.74E-06	4.01E-06	2.002794
Spl(mass%)						Avg RF	RRF
TBA	0.964	4.9692	9.9583	19.8768	29.7953		
	399808	1793750	3184446	7393280	11429736		
	409171	1908282	3579163	7370104	11664000		
avg	404489.5	1851016	3381805	7381692	11546868		
RF	2.38E-06	2.68E-06	2.94E-06	2.69E-06	2.58E-06	2.66E-06	1.32856
Spl(mass%)						Avg RF	RRF
MTBE	0.9992	5.0362	9.9724	20.0248	30.0471		
	353648	1719976	3016380	5400167	9756443		
	365624	1734192	3207775	6049396	9486117		
avg	359636	1727084	3112078	5724782	9621280		
RF	2.78E-06	2.92E-06	3.2E-06	3.5E-06	3.12E-06	3.1E-06	1.55197
Spl(mass%)						Avg RF	RRF
ETBE	0.9851	4.9255	9.8707	19.6724	29.5727		
	368857	1916504	3651460	6366342	8631784		
	370528	1990928	3698002	6858897	9781590		
avg	369692.5	1953716	3674731	6612620	9206687		
RF	2.66E-06	2.52E-06	2.69E-06	2.97E-06	3.21E-06	2.81E-06	1.405891
Spl(mass%)						Avg RF	RRF
TAME	0.9997	4.9788	9.8883	19.153	29.7144		
	373564	1867693	3846963	7398715	9605677		
	364642	1876735	4016568	7511412	10394700		
avg	369103	1872214	3931766	7455064	10000189		
RF	2.71E-06	2.66E-06	2.51E-06	2.57E-06	2.97E-06	2.68E-06	1.342326
Spl(mass%)						Avg RF	RRF
Nc7	8.475	8.44	8.4525	8.4525	8.695		
	3E+06	4E+06	312404	4E+06	4E+06		
	4E+06	4E+06	4E+06	5E+06	4E+06		
avg	3691763	4064455	2253742	4516374	4371883		
RF	2.3E-06	2.08E-06	3.75E-06	1.87E-06	1.99E-06	2E-06	1

FIG. A2.3 IHA Method Oxygenates Linearity Cooperative Study–Laboratory 3

Spl(mass%)						Avg RF	RRF
MEOH	1.01	5.05	10.02	20.01	29.83		
	658639	3389850	6670376	13542502	18749414		
	601443	3019715	6368637	13051539	17165160		
avg	630041	3204783	6519507	13297021	17957287		
RF	1.6E-06	1.58E-06	1.54E-06	1.5E-06	1.66E-06	1.58E-06	2.794957
Spl(mass%)						Avg RF	RRF
ETOH	1	5	10.1	20.15	30.18		
	826854	4450557	9154374	18060524	28066595		
	734856	4082467	8580584	17505672	28072314		
avg	780855	4266512	8867479	17783098	28069455		
RF	1.28E-06	1.17E-06	1.14E-06	1.13E-06	1.08E-06	1.16E-06	2.056683
Spl(mass%)						Avg RF	RRF
TBA	0.964	4.9692	9.9583	19.8768	29.7953		
	1578407	4266396	14460028	29135138	43225116		
	1435170	6337881	13565261	27794630	42612348		
avg	1506789	5302139	14012645	28464884	42918732		
RF	6.4E-07	9.37E-07	7.11E-07	6.98E-07	6.94E-07	7.36E-07	1.305022
Spl(mass%)						Avg RF	RRF
MTBE	0.9992	5.0362	9.9724	20.0248	30.0471		
	1252485	5941164	10848222	17786018	23089928		
	1255790	6142349	10162313	17011562	22404206		
avg	1254138	6041757	10505268	17398790	22747067		
RF	7.97E-07	8.34E-07	9.49E-07	1.15E-06	1.32E-06	1.01E-06	1.791283
Spl(mass%)						Avg RF	RRF
ETBE	0.9851	4.9255	9.8707	19.6724	29.5727		
	1310455	6926229	12417871	20398546	27031106		
	1306372	7052557	12595757	19329114	26122426		
avg	1308414	6989393	12506814	19863830	26576766		
RF	7.53E-07	7.05E-07	7.89E-07	9.9E-07	1.11E-06	8.7E-07	1.542526
Spl(mass%)						Avg RF	RRF
TAME	0.9997	4.9788	9.8883	19.153	29.7144		
	1400316	6820054	13673677	22152636	28646506		
	1357511	6857019	13936737	22286660	27439076		
avg	1378914	6838537	13805207	22219648	28042791		
RF	7.25E-07	7.28E-07	7.16E-07	8.62E-07	1.06E-06	8.18E-07	1.450677
Spl(mass%)						Avg RF	RRF
Nc7	8.475	8.44	8.4525	8.4525	8.695		
	15260819	15252480	14899327	15397626	14345822		
	14816484	14876828	14956987	15670374	15233576		
avg	15038652	15064654	14928157	15534000	14789699		
RF	5.64E-07	5.6E-07	5.66E-07	5.44E-07	5.88E-07	5.64E-07	1

FIG. A2.4 IHA Method Oxygenate Linearity Cooperative Study–Laboratory 4

Spl(mass%)						Avg RF	RRF
MEOH	1.01	5.05	10.02	20.01	29.83		
avg	130.85	729.625	1474.483	3103.843	4600.484		
RF	0.007719	0.006921	0.006796	0.006447	0.006484	0.006873	3.08498
Spl(mass%)						Avg RF	RRF
ETOH	1	5	10.1	20.15	30.18		
avg	195.402	1054.59	2115.254	4301.374	6707.759		
RF	0.005118	0.004741	0.004775	0.004685	0.004499	0.004763	2.138015
Spl(mass%)						Avg RF	RRF
TBA	0.964	4.9692	9.9583	19.8768	29.7953		
avg	347.107	1725.706	3442.236	6695.103	10183.1		
RF	0.002777	0.00288	0.002893	0.002969	0.002926	0.002889	1.296638
Spl(mass%)						Avg RF	RRF
MTBE	0.9992	5.0362	9.9724	20.0248	30.0471		
avg	290.368	1518.529	3008.79	6043.303	8800.898		
RF	0.003441	0.003316	0.003314	0.003314	0.003414	0.00336	1.508054
Spl(mass%)						Avg RF	RRF
ETBE	0.9851	4.9255	9.8707	19.6724	29.5727		
avg	308.613	1630.908	3253.559	6580.098	9806.89		
RF	0.003192	0.00302	0.003034	0.00299	0.003016	0.00305	1.369041
Spl(mass%)						Avg RF	RRF
TAME	0.9997	4.9788	9.8883	19.153	29.7144		
avg	322.928	1631.466	3351.751	6693.316	10161.7		
RF	0.003096	0.003052	0.00295	0.002862	0.002924	0.002977	1.336026
Spl(mass%)						Avg RF	RRF
Nc7	8.475	8.44	8.4525	8.4525	8.695		
avg	3915.73	3733.39	3714.828	3835.85	3889.013		
RF	0.002164	0.002261	0.002275	0.002204	0.002236	0.002228	1
note: average area counts are the average of two runs							

FIG. A2.5 IHA Method Oxygenate Linearity Cooperative Study–Laboratory 5

Spl(mass%)						Avg RF	RRF
MEOH	1.01	5.05	10.02	20.01	29.83		
avg	128.825	795.291	1607.186	3383.189	5800.591		
RF	0.00784	0.00635	0.006234	0.005915	0.005143	0.006296	3.390586
Spl(mass%)						Avg RF	RRF
ETOH	1	5	10.1	20.15	30.18		
avg	212.988	1149.503	2305.626	4688.498	7300.836		
RF	0.004695	0.00435	0.004381	0.004298	0.004134	0.004371	2.354003
Spl(mass%)						Avg RF	RRF
TBA	0.964	4.9692	9.9583	19.8768	29.7953		
avg	378.347	1881.019	3752.037	7297.662	11045.72		
RF	0.002548	0.002642	0.002654	0.002724	0.002697	0.002653	1.428645
Spl(mass%)						Avg RF	RRF
MTBE	0.9992	5.0362	9.9724	20.0248	30.0471		
avg	316.501	1655.196	3279.581	6587.2	9660.288		
RF	0.003157	0.003043	0.003041	0.00304	0.00311	0.003078	1.657594
Spl(mass%)						Avg RF	RRF
ETBE	0.9851	4.9255	9.8707	19.6724	29.5727		
avg	336.388	1777.69	3546.379	7172.307	10609.51		
RF	0.002928	0.002771	0.002783	0.002743	0.002787	0.002803	1.509178
Spl(mass%)						Avg RF	RRF
TAME	0.9997	4.9788	9.8883	19.153	29.7144		
avg	351.991	1778.298	3653.409	7295.715	11076.25		
RF	0.00284	0.0028	0.002707	0.002625	0.002683	0.002731	1.47059
Spl(mass%)						Avg RF	RRF
Nc7	8.475	8.44	8.4525	8.4525	8.695		
avg	4698.033	4477.402	4454.942	4601.379	4665.706		
RF	0.001804	0.001885	0.001897	0.001837	0.001864	0.001857	1
note: average area counts are an average of three runs							

FIG. A2.6 IHA Method Oxygenate Linearity Cooperative Study–Laboratory 6

Spl(mass%)						Avg RF	RRF
MEOH	1.01	5.05	10.02	20.01	29.83		
	35419	207968	408281	807253	1208115		
	36040	195967	408281	874729	1301947		
avg	35729.5	201967.5	408281	840991	1255031		
RF	2.83E-05	2.5E-05	2.45E-05	2.38E-05	2.38E-05	2.51E-05	2.922508
Spl(mass%)						Avg RF	RRF
ETOH	1	5	10.1	20.15	30.18		
	45510	292874	642031	1234541	1824287		
	50885	281463	594198	1259869	2005196		
avg	48197.5	287168.5	618114.5	1247205	1914742		
RF	2.07E-05	1.74E-05	1.63E-05	1.62E-05	1.58E-05	1.73E-05	2.014392
Spl(mass%)						Avg RF	RRF
TBA	0.964	4.9692	9.9583	19.8768	29.7953		
	93315	475528	979360	2031219	2865032		
	102421	476914	888766	1840517	2928378		
avg	97868	476221	934063	1935868	2896705		
RF	9.85E-06	1.04E-05	1.07E-05	1.03E-05	1.03E-05	1.03E-05	1.200454
Spl(mass%)						Avg RF	RRF
MTBE	0.9992	5.0362	9.9724	20.0248	30.0471		
	75952	405208	705631	1548681	2380261		
	77415	417553	757750	1580147	2408423		
avg	76683.5	411380.5	731690.5	1564414	2394342		
RF	1.3E-05	1.22E-05	1.36E-05	1.28E-05	1.25E-05	1.29E-05	1.497693
Spl(mass%)						Avg RF	RRF
ETBE	0.9851	4.9255	9.8707	19.6724	29.5727		
	83107	436772	890514	1713524	2609194		
	85993	442601	917344	1720724	2604325		
avg	84550	439686.5	903929	1717124	2606760		
RF	1.17E-05	1.12E-05	1.09E-05	1.15E-05	1.13E-05	1.13E-05	1.31875
Spl(mass%)						Avg RF	RRF
TAME	0.9997	4.9788	9.8883	19.153	29.7144		
	89539	455171	900734	1836776	2713677		
	90145	461944	915196	1883508	2658665		
avg	89842	458557.5	907965	1860142	2686171		
RF	1.11E-05	1.09E-05	1.09E-05	1.03E-05	1.11E-05	1.08E-05	1.264195
Spl(mass%)						Avg RF	RRF
Nc7	20%	5%	1%	10%	30%		
	8.475	8.44	8.4525	8.4525	8.695		
	1034198	1392371	989383	983168	1077830		
	889948	935398	1051329	1067382	1010624		
avg	962073	1163885	1020356	1025275	1044227		
RF	8.81E-06	7.25E-06	8.28E-06	8.24E-06	8.33E-06	8.18E-06	1

FIG. A2.7 IHA Method Oxygenate Linearity Cooperative Study–Laboratory 7

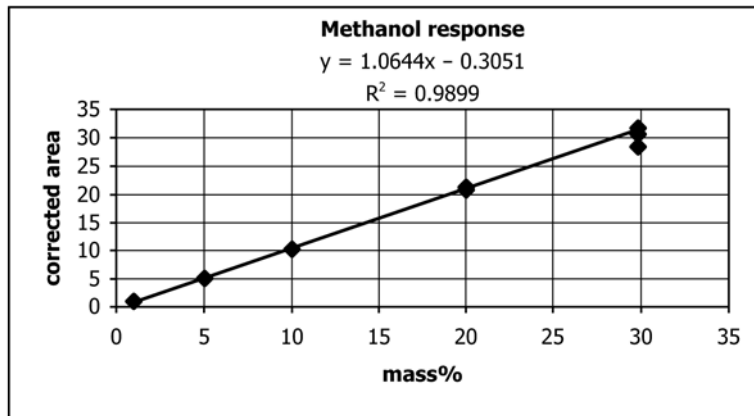


FIG. A2.8 Determination of Calculated Oxygenate Response from IHA Method—Methanol

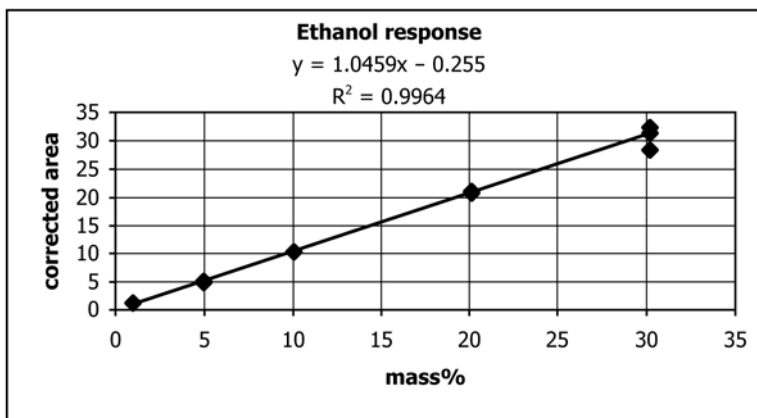


FIG. A2.9 Determination of Calculated Oxygenate Response from IHA Method—Ethanol

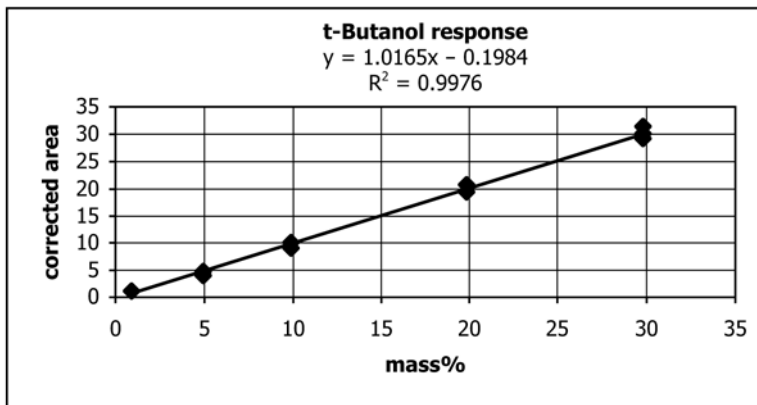


FIG. A2.10 Determination of Calculated Oxygenate Response from IHA Method—t-Butanol

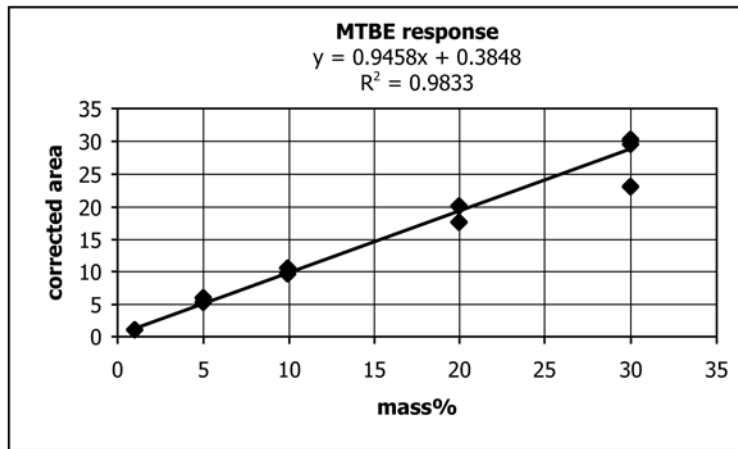


FIG. A2.11 Determination of Calculated Oxygenate Response from IHA Method–MTBE

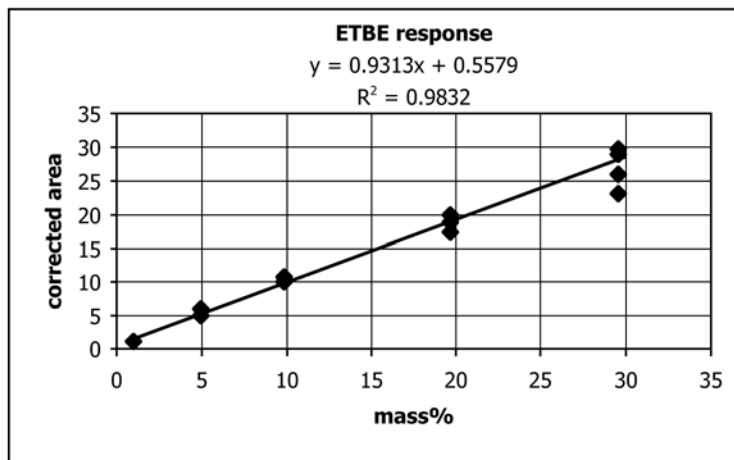


FIG. A2.12 Determination of Calculated Oxygenate Response from IHA Method–ETBE

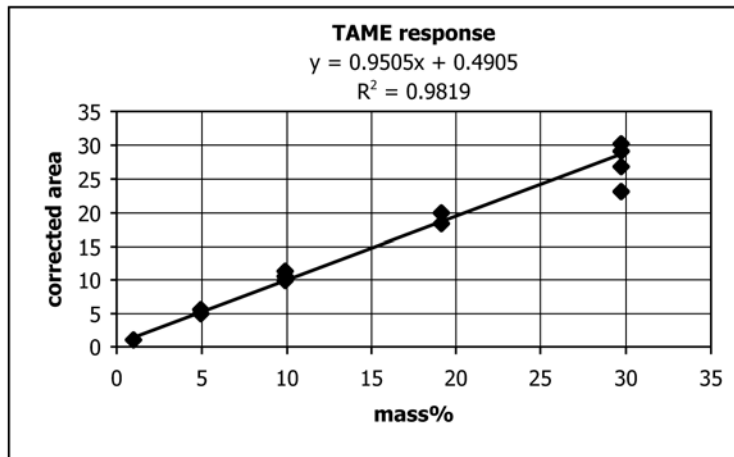


FIG. A2.13 Determination of Calculated Oxygenate Response from IHA Method–TAME

APPENDIXES

(Nonmandatory Information)

X1. BIBLIOGRAPHY

X1.1 The following publications on DHA analyses may be useful as background and are recommended to the user of these test procedures:

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X1.1.5 Di Sanzo, F. P. and Giarrocco, V. G., “Analysis of Pressurized Gasoline-Range Liquid Hydrocarbon Samples by Capillary Column and PIONA Analyzer Gas Chromatography,” *Journal of Chromatographic Science*, Vol 26, June 1988, pp 258-266

X1.1.6 Durand, J. P., Beboluene, J. J. and Ducrozet, A., “Detailed Characterization of Petroleum Products with Capillary GC Analyzers,” *Analisis*, 23, 1995, pp 481-483

X1.1.7 Canadian General Standards Board: CAN/CGSB-3.0, No.14.3-94, “Test Method for Individual Hydrocarbon Component Analysis (IHA) in Spark Ignition Engine Fuels by Gas Chromatography”

X1.1.8 French Standard NF N07-086, December 1995, “Determination of Hydrocarbon Type Contents in Motor Gasolines from Detailed Analysis Capillary Gas Chromatography”

X2. HYDROCARBON DATA USING HYDROGEN CARRIER

X2.1 This appendix presents a modification of the test method that decreases the total analysis time from 150 minutes to 82 minutes. The primary change is the use of hydrogen rather than helium as the carrier gas. The oven temperature ramp and column flow rates are also modified in order to ensure optimum resolution and linear velocity. All remaining operating parameters of the GC remain as stated in the method. Column performance tests are included in order to evaluate method performance. This appendix includes a chromatogram with components identified, as well as the component list analyzed with the test conditions specified in [Table X2.1](#).

X2.2 [Table X2.1](#) presents the chromatographic operating conditions, column requirements, and data acquisition requirements. These conditions will elute all components up to and including pentadecane ($n\text{-C}_{15}$).

X2.3 [Table X2.2](#) presents the column resolution performance requirements.

X2.4 [Figs. X2.1-X2.5](#) present chromatographic examples of each critical separation described in [Table X2.2](#).

X2.5 [Table X2.3](#) presents the column efficiency results based on a 35 °C isothermal analysis of pentane ($n\text{-C}_5$).

X2.6 [Figs. X2.6-X2.11](#) presents, in detail, the identified CGSB0496 gasoline standard analyzed under the conditions stated in [Table X2.1](#).

X2.7 [Table X2.4](#) presents the summary of coeluting compounds.

X2.8 [Table X2.5](#) presents the component retention times and properties of the compounds identified in the CGSB0496 standard.

X2.9 All sections of Test Method D6729 should be followed as written, except for the following:

X2.9.1 In [7.2](#), hydrogen gas is used rather than helium as the carrier gas.

X2.9.2 In [8.1](#), the Linear Gas Velocity is now set to 31.0 psi constant pressure that corresponds to an average linear velocity of 42 cm/s at 35 °C. This linear velocity value is optimum for hydrogen. Under these isothermal conditions, methane should elute at 3.9 min. Using the conditions described in [Table X2.1](#), methane should elute at 3.6 min.

X2.9.3 The [Table 1](#) chromatographic operating conditions are now modified as in [Table X2.1](#).

TABLE X2.1 Instrument Parameters

Chromatographic Conditions	Requirements
Carrier gas	Hydrogen
Injector settings	
Injector temperature, °C	270
Split ratio	200:1
Liner	deactivated glass
Injection volume	0.2 µL
Detector settings	
FID temperature, °C ^A	300
Hydrogen flow, mL/min ^B	40
Air flow, mL/min	450
Nitrogen make-up flow, mL/min	20
Column oven settings	
Initial temperature, °C	0
Initial hold time, min	9.5
1st ramp rate, °C/min	1.5
Final temperature, °C	50
Final hold time, min	0
2nd ramp rate, °C/min	3.14
Final temperature, °C	130
Final hold time, min	0
3rd ramp rate, °C/min	6.28
Final temperature, °C	270
Final hold time, min ^C	0
Column requirements	
Length, m	100
Inside diameter, mm	0.25
Liquid phase	100 % polydimethylsiloxane
Film thickness, µm	0.5
Pressure (Constant), psi	31.0
Flow, mL/min (range)	3.3 – 1.0
Linear gas velocity, cm/s (range)	46 – 29
Data acquisition, Hz	20
Total analysis time, min (including clean-up time)	90.6

^A Set to 25 °C to 50 °C above the highest column temperature.

^B Values to be set as recommended by instrument manufacturer.

^C Final hold time or temperature may be adjusted to ensure complete elution of the sample components.

TABLE X2.2 Column Resolution Performance Requirements

Component Pair	Minimum Resolution	Concentration of Each Component, w/w
74 2,3-Dimethylbutane	1.0	0.99 %
78 Methyl <i>t</i> -butyl ether		1.23 %
156 Benzene	1.0	0.83 %
158 1-methylcyclopentene		0.49 %
304 Toluene	0.4	7.65 %
306 2,3,3-trimethylpentane		0.65 %
474 <i>m</i> -Xylene	0.4	3.95 %
476 <i>p</i> -Xylene		1.58 %
876 <i>n</i> -Tridecane	1.0	0.01 %
878 1-methylnaphthalene		0.02 %

**TABLE X2.3 Column Efficiency
(Based on 35°C Isothermal Analysis)**

Component	RT (min)	Peak Width (½ h)	Theoretical Plates
38 <i>n</i> -Pentane	5.406	0.0178	511 462

TABLE X2.4 Coeluting Compounds^{A,B}

RT	Predominant Peak	Coeluting Compound
23.00	methylcyclopentane	2,2-dimethylpentane
27.13	3,3-dimethylpentane	5-methyl-1-hexene
29.36	2-methylhexane	2,3-dimethylpentane and C ₇ -olefin
38.76	2,5-dimethylhexane	C ₈ -olefin
41.32	toluene	2,3,3-trimethylpentane ^C
44.59	3-methylheptane	<i>t</i> -1,4-dimethylcyclohexane
54.27	4-methyloctane	C ₉ -olefin
55.27	<i>o</i> -xylene	1,1,2-trimethylcyclohexane
72.54	1,2,3,4-tetramethylbenzene	C ₁₁ -aromatic

^A Due to the possibility of coeluting peaks in other areas, the user is cautioned in the interpretation of the data.

^B Manual integration may be necessary in many areas of the chromatogram.

^C The 233-TMC5 is partially resolved as a shoulder of the toluene peak.

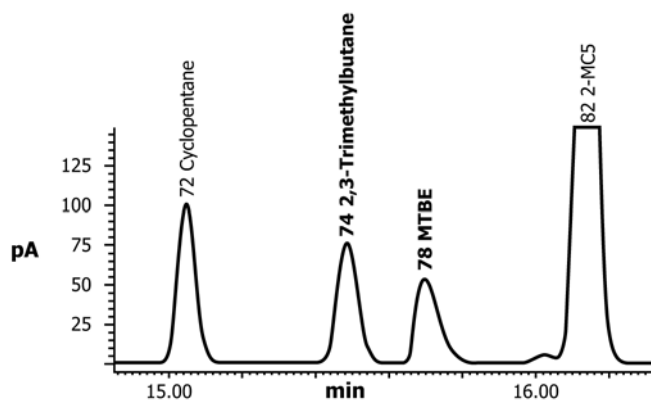


FIG. X2.1 2,3-dimethylbutane and methyl *t*-butyl ether
 $R = 2(15.698 - 15.487) / 1.699(0.0603 + 0.0633) = 2.01$

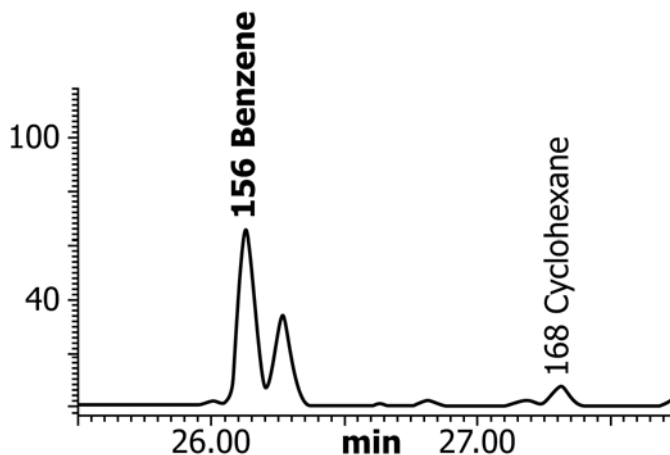


FIG. X2.2 Benzene and 1-methylcyclopentene
 $R = 2(26.269 - 26.131) / 1.699(0.0654 + 0.0684) = 1.21$

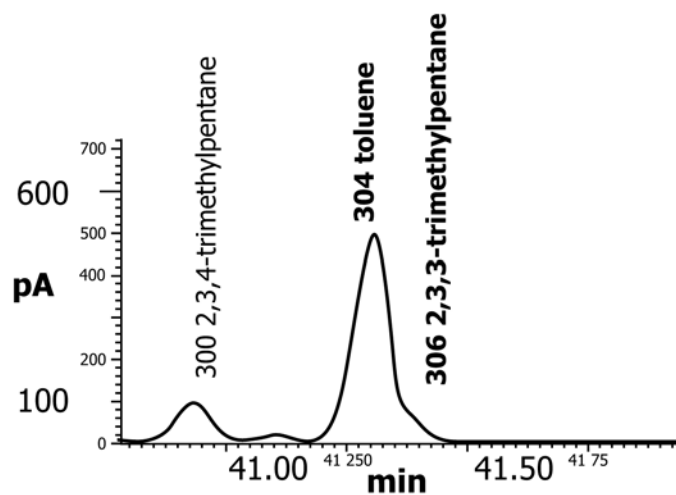


FIG. X2.3 Toluene and 2,3,3-trimethylpentane
 $R = 2(41.378 - 41.317) / 1.699(0.0832 + 0.0383) = 0.59$

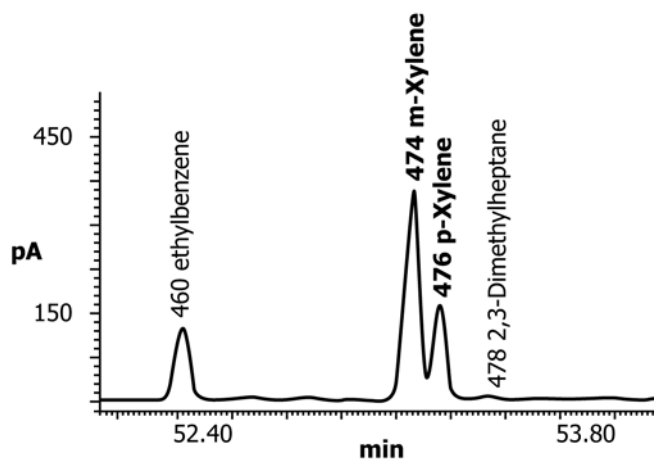


FIG. X2.4 *m*-xylene and *p*-xylene
 $R = 2(53.371 - 53.275) / 1.699(0.0558 + 0.0506) = 1.08$

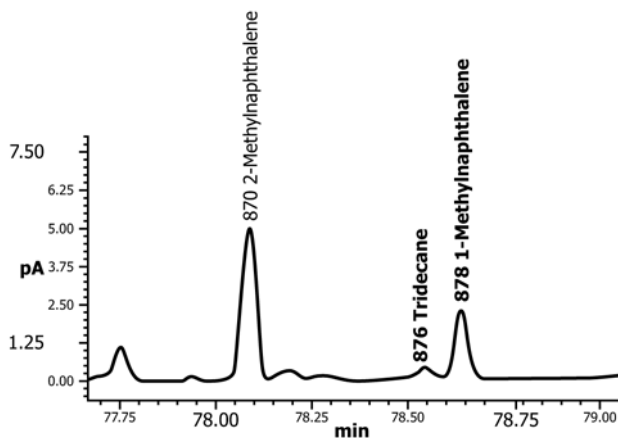


FIG. X2.5 Tridecane and 1-methylnaphthalene
 $R = 2(78.639 - 78.542) / 1.699(0.0486 + 0.0375) = 1.32$

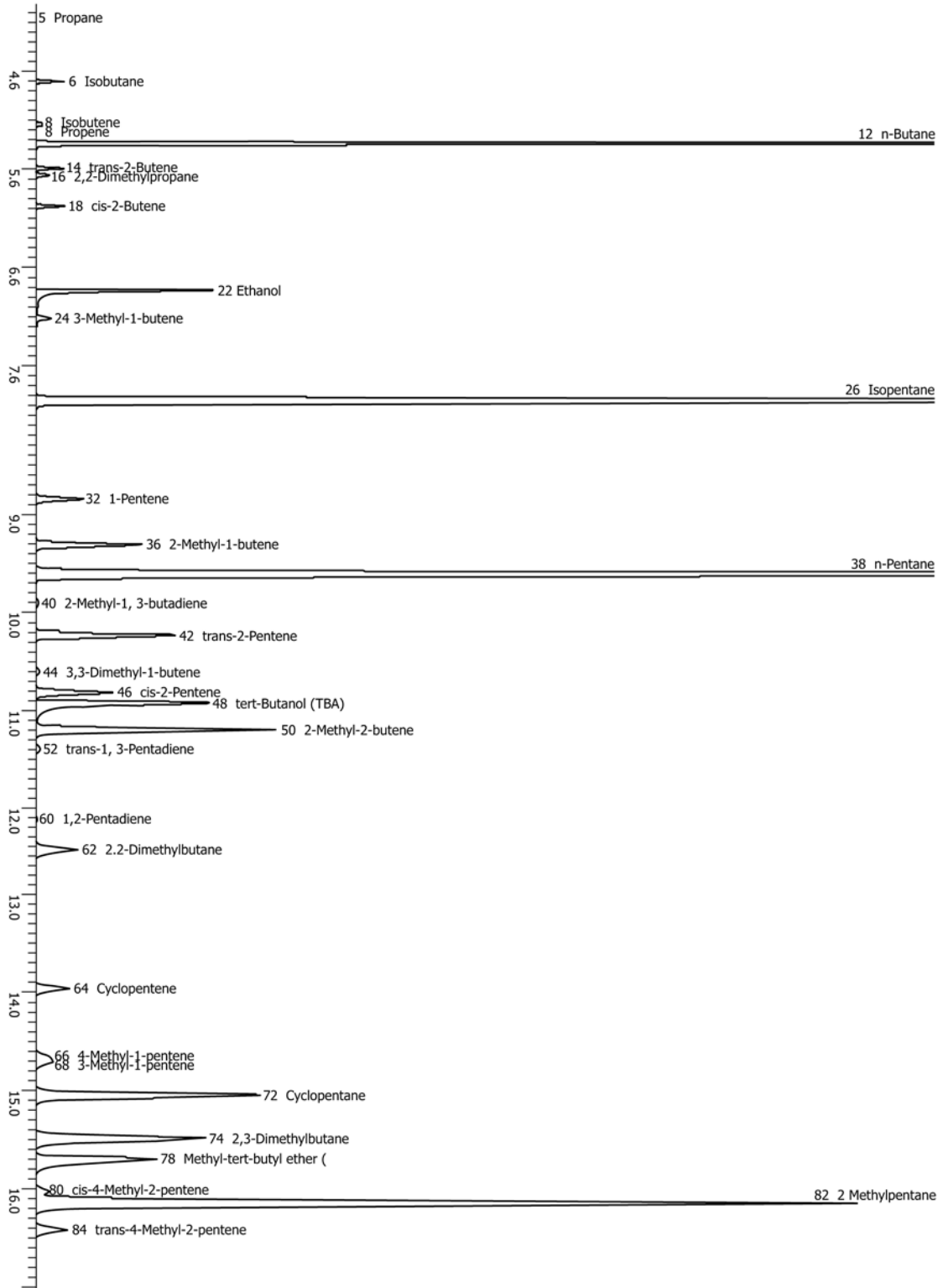


FIG. X2.6 Detailed Chromatogram for CGSB0496 Reference Gasoline (0 min to 17 min)

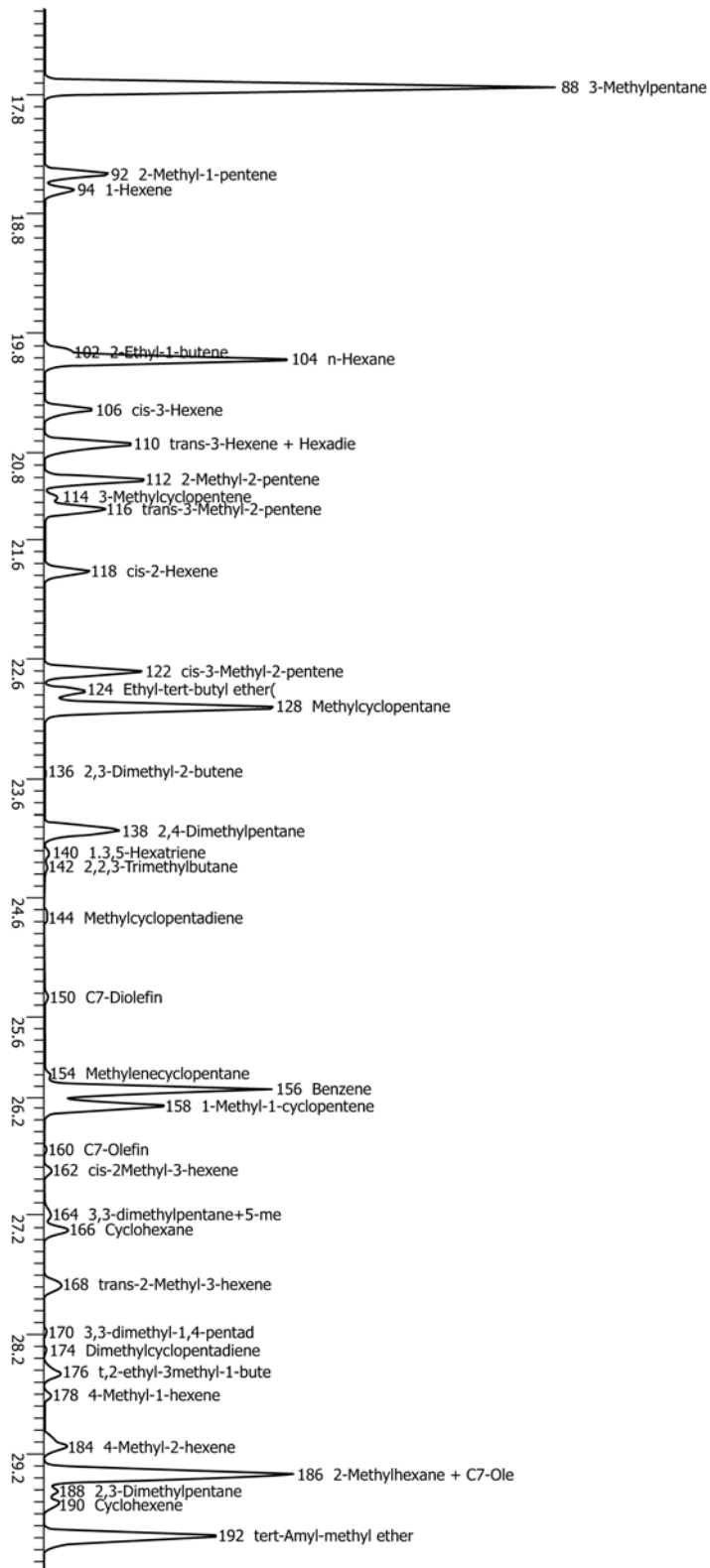


FIG. X2.7 Detailed Chromatogram for CGSB0496 Reference Gasoline (17 min to 30.5 min)

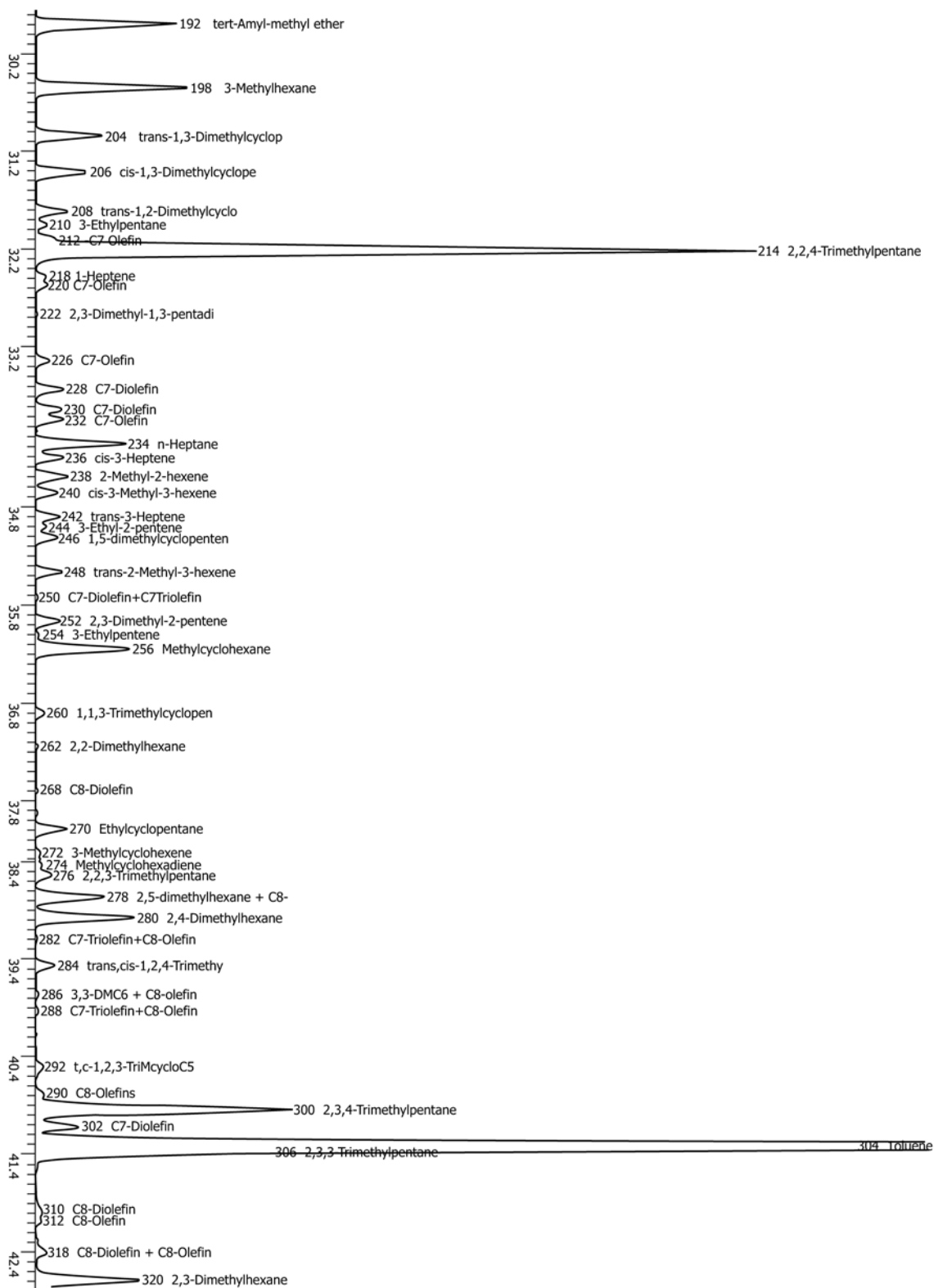


FIG. X2.8 Detailed Chromatogram for CGSB0496 Reference Gasoline (30.5 min to 42.5 min)

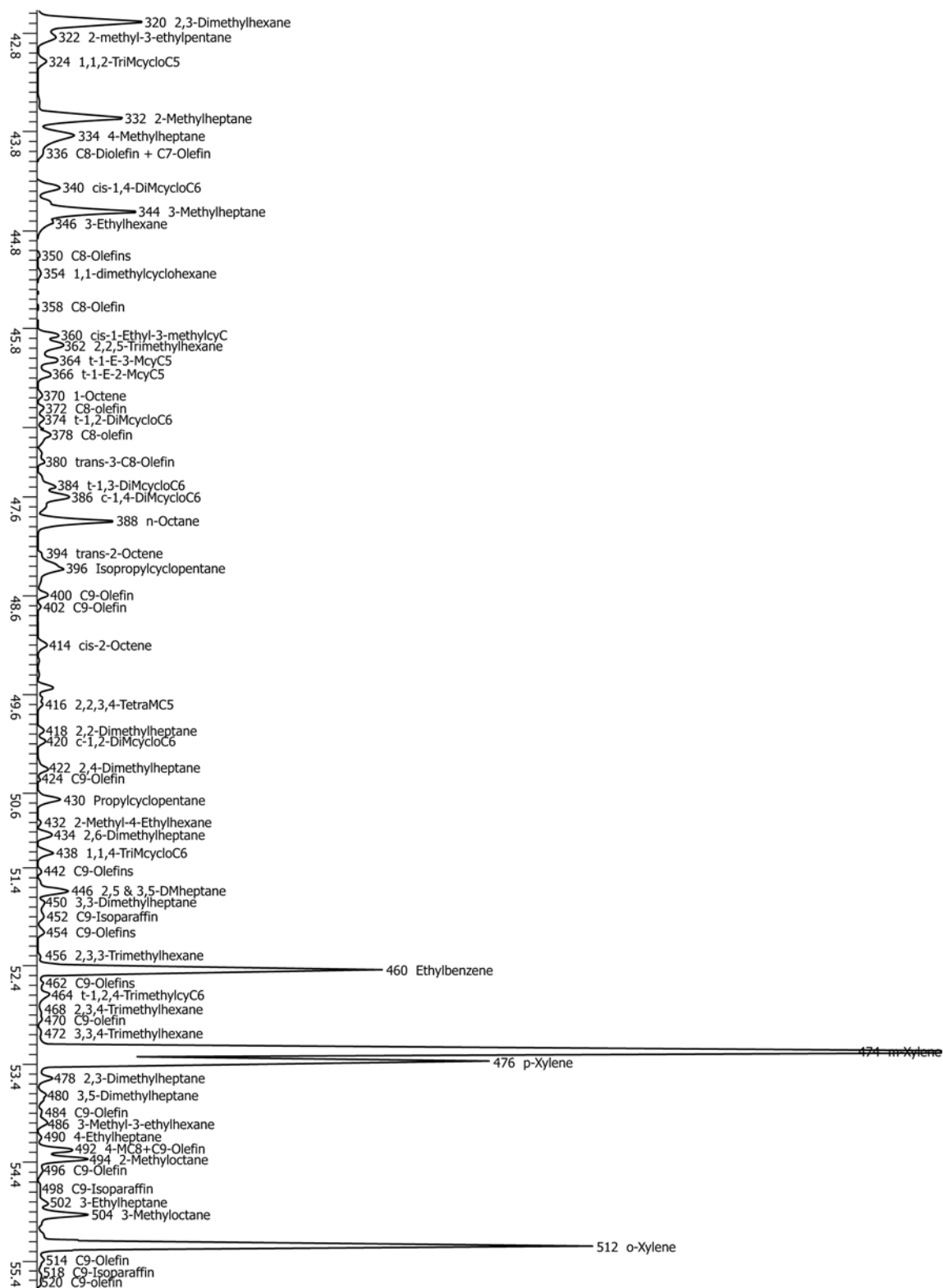


FIG. X2.9 Detailed Chromatogram for CGSB0496 Reference Gasoline (42.5 min to 55.5 min)

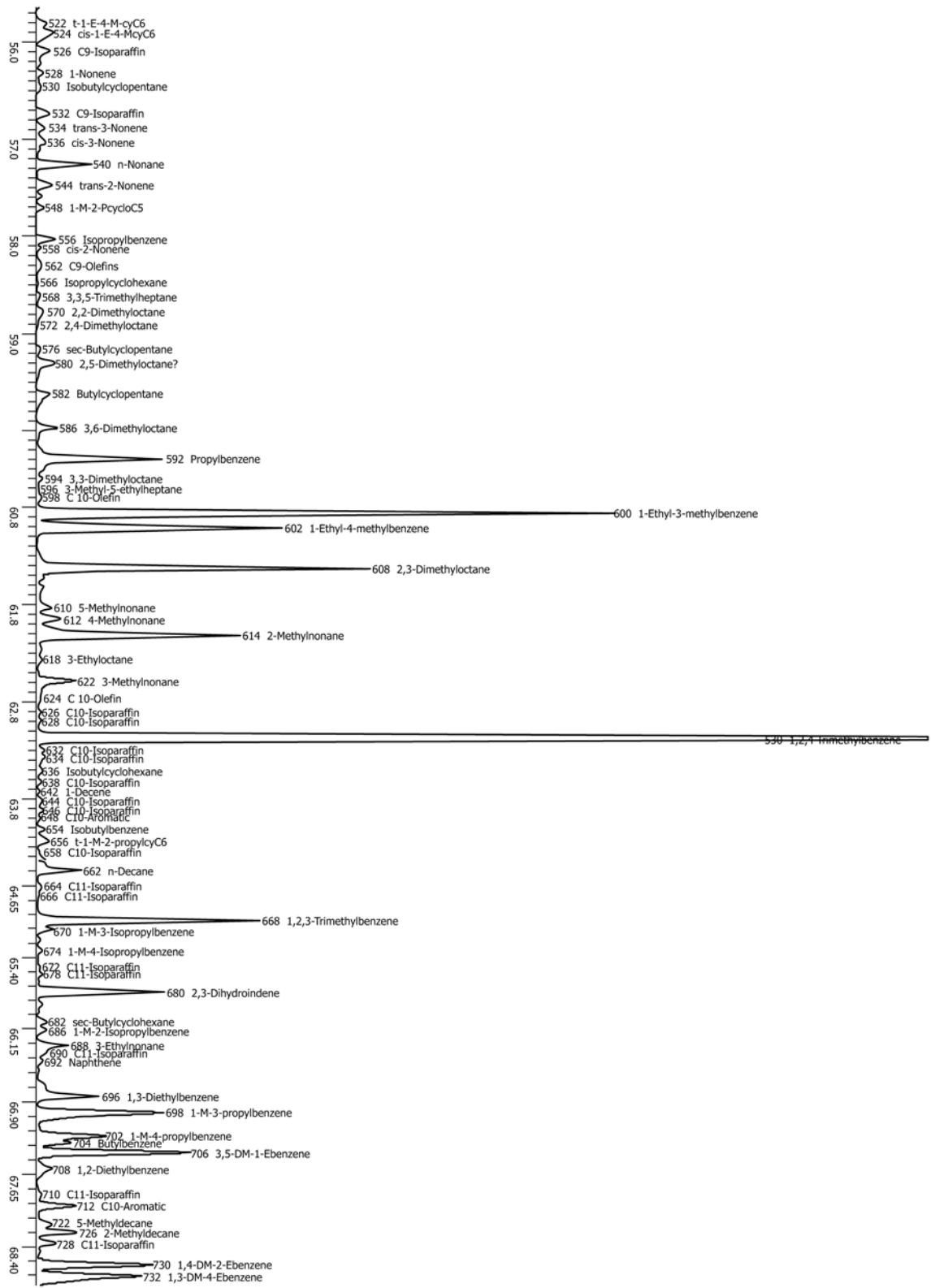


FIG. X2.10 Detailed Chromatogram for CGSB0496 Reference Gasoline (55.5 min to 68.5 min)

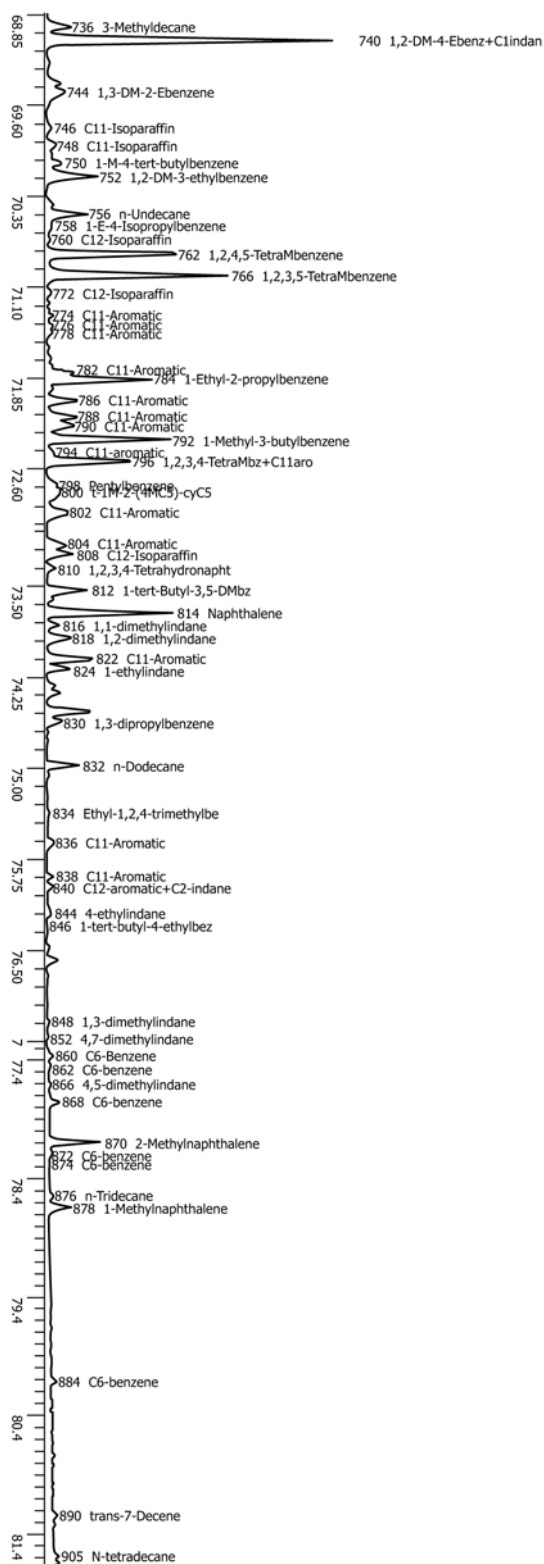


FIG. X2.11 Detailed Chromatogram for CGSB0496 Reference Gasoline (68.5 min to 82 min)

TABLE X2.5 Component Retention Times and Properties

Peak Number and Component Name	RT (min)	MW	RRF	
1	Methane	3.89	16.04	1.000
2	Ethene	3.93	28.03	0.874
3	Ethane	3.97	30.05	0.937
4	Propene	4.01	42.05	0.874
5	Propane	4.05	44.06	0.916
6	Isobutane	4.70	58.08	0.906
7	Methanol	4.92	32.03	2.672
8	Isobutene	5.13	56.06	0.874
9	1-Butene	5.15	56.06	0.874
10	1,3-Butadiene	5.24	54.09	0.843
12	n-Butane	5.33	58.08	0.906
14	trans-2-Butene	5.59	56.06	0.874
16	2,2-Dimethylpropane	5.65	72.09	0.899
18	cis-2-Butene	5.97	56.06	0.874
20	1,2-Butadiene	6.39	54.09	0.843
22	Ethanol	6.83	46.04	1.862
24	3-Methyl-1-butene	7.12	70.08	0.874
26	Isopentane	7.96	72.09	0.899
28	1,4-Pentadiene	8.25	68.06	0.849
30	2-Butyne (Dimethylacetylene)	8.55	54.05	0.843
32	1-Pentene	8.84	70.08	0.874
34	Isopropanol	9.07	60.06	1.950
36	2-Methyl-1-butene	9.31	70.08	0.874
38	n-Pentane	9.60	72.09	0.899
40	2-Methyl-1,3-butadiene	9.89	68.06	0.849
42	trans-2-Pentene	10.23	70.08	0.874
44	3,3-Dimethyl-1-butene	10.60	84.09	0.874
46	cis-2-Pentene	10.81	70.08	0.874
48	tert-Butanol (TBA)	10.92	74.12	1.161
50	2-Methyl-2-butene	11.19	70.08	0.874
52	trans-1,3-Pentadiene	11.39	68.06	0.849
54	3-Methyl-1,2-butadiene	11.39	68.06	0.849
56	Cyclopentadiene	11.63	67.10	0.824
58	cis-1,3-Pentadiene	11.87	68.06	0.849
60	1,2-Pentadiene	12.11	68.06	0.849
62	2,2-Dimethylbutane	12.42	86.11	0.895
64	Cyclopentene	13.96	68.06	0.849
66	4-Methyl-1-pentene	14.68	84.09	0.874
68	3-Methyl-1-pentene	14.70	84.09	0.874
70	n-Propanol	14.89	60.06	1.770
72	Cyclopentane	15.05	70.08	0.874
74	2,3-Dimethylbutane	15.49	86.11	0.895
76	2,3-Dimethyl-1-butene	15.62	84.09	0.874
78	Methyl-tert-butyl ether (MTBE)	15.70	88.09	1.407
80	cis-4-Methyl-2-pentene	16.03	84.09	0.874
82	2-Methylpentane	16.15	86.11	0.895
84	trans-4-Methyl-2-pentene	16.42	84.09	0.874
86	Methylethylketone (MEK)	17.08	72.06	1.570
88	3-Methylpentane	17.73	86.11	0.895
90	C6-Olefin	18.10	84.09	0.874
92	2-Methyl-1-pentene	18.46	84.09	0.874
94	1-Hexene	18.60	84.09	0.874
96	Methyl-sec-butyl ether	18.95	88.09	1.550
98	C6-Olefin	19.29	84.09	0.874
100	2-Butanol	19.63	74.07	1.600
102	2-Ethyl-1-butene	19.97	84.09	0.874
104	n-Hexane	20.03	86.11	0.895
106	cis-3-Hexene	20.45	84.09	0.874
108	Di-isopropylether (DIPE)	20.60	102.00	1.600
110	trans-3-Hexene + Hexadiene	20.74	84.09	0.874
112	2-Methyl-2-pentene	21.04	84.09	0.874
114	3-Methylcyclopentene	21.19	82.10	0.853
116	trans-3-Methyl-2-pentene	21.28	84.09	0.874
118	cis-2-Hexene	21.86	84.09	0.874
120	3,3-Dimethyl-1-pentene	22.29	98.19	0.874
122	cis-3-Methyl-2-pentene	22.70	84.09	0.874
124	Ethyl-tert-butyl ether (ETBE)	22.87	102.18	1.255
126	2,3-Dimethyl-1,3-butadiene	22.94	82.00	0.853
128	Methylcyclopentane	23.00	84.09	0.874
130	2,2-dimethylpentane	23.05	100.13	0.892
132	4,4-Dimethyl-1-pentene	23.19	98.19	0.874
134	Isobutanol	23.37	74.12	1.500
136	2,3-Dimethyl-2-butene	23.55	84.09	0.874
138	2,4-Dimethylpentane	24.03	100.13	0.892
140	1,3,5-Hexatriene	24.22	80.00	0.832

TABLE X2.5 *Continued*

	Peak Number and Component Name	RT (min)	MW	RRF
142	2,2,3-Trimethylbutane	24.33	100.13	0.892
144	Methylcyclopentadiene	24.77	80.00	0.832
146	C7-Olefin	24.83	98.19	0.874
148	C7-Olefin	25.13	98.19	0.874
150	C7-Diolefin	25.43	96.00	0.856
152	4-Methylcyclopentene	25.71	82.10	0.853
154	Methylenecyclopentane	26.01	82.10	0.853
156	Benzene	26.13	78.05	0.812
158	1-Methyl-1-cyclopentene	26.27	82.10	0.853
160	C7-Olefin	26.64	98.19	0.874
162	cis-2-Methyl-3-hexene	26.82	98.19	0.874
164	3,3-dimethylpentane+5-me	27.19	100.13	0.892
166	Cyclohexane	27.32	84.09	0.874
168	trans-2-Methyl-3-hexene	27.78	98.19	0.874
170	3,3-dimethyl-1,4-pentad	28.18	96.00	0.856
172	n-Butanol	28.29	74.07	1.500
174	Dimethylcyclopentadiene	28.33	94.16	0.838
176	t,2-ethyl-3methyl-1-bute	28.52	98.19	0.874
178	4-Methyl -1-hexene	28.71	98.19	0.874
180	C7-Olefin	28.86	98.19	0.874
182	3-Methyl-1-hexene	29.00	98.19	0.874
184	4-Methyl-2-hexene	29.14	98.19	0.874
186	2-Methylhexane + C7-Ole	29.37	100.13	0.892
188	2,3-Dimethylpentane	29.52	100.13	0.892
190	Cyclohexene	29.62	82.10	0.853
192	tert-Amyl-methyl ether	29.89	102.18	1.210
194	C7-Olefin	30.11	98.19	0.874
196	C7-Olefin	30.33	98.19	0.874
198	3-Methylhexane	30.55	100.13	0.892
200	C7-Olefin	30.71	98.19	0.874
202	C7-Olefin	30.88	98.19	0.874
204	trans-1,3-Dimethylcyclop	31.04	98.11	0.874
206	cis-1,3-Dimethylcyclope	31.42	98.11	0.874
208	trans-1,2-Dimethylcyclo	31.82	98.11	0.874
210	3-Ethylpentane	31.95	100.10	0.892
212	C7-Olefin	32.12	98.19	0.874
214	2,2,4-Trimethylpentane	32.22	114.14	0.890
216	C7-Olefin	32.47	98.19	0.874
218	1-Heptene	32.49	98.19	0.874
220	C7-Olefin	32.57	98.19	0.874
222	2,3-Dimethyl-1,3-pentadi	32.87	96.18	0.874
224	C7-Diolefin	33.11	96.00	0.856
226	C7-Olefin	33.35	98.19	0.874
228	C7-Diolefin	33.64	96.00	0.856
230	C7-Diolefin	33.85	96.00	0.856
232	C7-Olefin	33.95	98.19	0.874
234	n-Heptane	34.15	100.13	0.892
236	cis-3-Heptene	34.29	98.19	0.874
238	2-Methyl-2-hexene	34.49	98.19	0.874
240	cis-3-Methyl-3-hexene	34.65	98.19	0.874
242	trans-3-Heptene	34.89	98.19	0.874
244	3-Ethyl-2-pentene	35.00	96.17	0.856
246	1,5-dimethylcyclopenten	35.11	96.17	0.856
248	trans-2-Methyl-3-hexene	35.46	98.19	0.874
250	C7-Diolefin+C7Triolefin	35.72	96.00	0.856
252	2,3-Dimethyl-2-pentene	35.96	98.19	0.874
254	3-Ethylpentene	36.12	98.19	0.874
256	Methylcyclohexane	36.25	98.11	0.874
258	C7-Olefin	36.58	98.19	0.874
260	1,1,3-Trimethylcyclopen	36.90	112.10	0.874
262	2,2-Dimethylhexane	37.24	114.10	0.890
264	2,3,4-TriM-1,4-C5diene	37.60	110.00	0.859
266	3,3-dimethyl-1,5-hexadie	37.63	110.21	0.859
268	C8-Diolefin	37.69	98.19	0.874
270	C7-Olefin	37.93	98.19	0.874
272	Ethylcyclopentane	38.10	98.11	0.874
274	3-Methylcyclohexene	38.34	96.17	0.856
276	Methylcyclohexadiene	38.43	94.17	0.838
278	2,2,3-Trimethylpentane	38.53	114.10	0.890
280	2,5-dimethylhexane + C8-	38.76	114.14	0.890
282	2,4-Dimethylhexane	38.97	114.14	0.890
284	C7-Triolefin+C8-Olefin	39.19	112.24	0.856
286	trans,cis-1,2,4-Trimethy	39.46	112.10	0.874
288	3,3-DMC6 + C8-olefin	39.75	114.14	0.890
290	C7-Triolefin+C8-Olefin	39.92	112.24	0.856

TABLE X2.5 *Continued*

	Peak Number and Component Name	RT (min)	MW	RRF
292	t,c-1,2,3-TriMcyCloC5	40.50	112.10	0.874
294	C8-Olefins	40.57	112.13	0.874
296	C8-Olefins	40.63	112.13	0.874
298	C8-Olefins	40.69	112.13	0.874
290	C8-Olefins	40.79	112.13	0.874
300	2,3,4-Trimethylpentane	40.94	114.14	0.890
302	C7-Diolefin	41.11	96.00	0.856
304	Toluene	41.32	92.06	0.821
306	2,3,3-Trimethylpentane	41.38	114.23	0.890
308	C8-Olefin	41.54	112.13	0.874
310	C8-Diolefin	41.99	110.00	0.859
312	C8-Olefin	42.08	112.13	0.874
314	C8-Olefin	42.26	112.13	0.874
316	C8-Olefin	42.35	112.13	0.874
318	C8-Diolefin + C8-Olefin	42.41	110.00	0.859
320	2,3-Dimethylhexane	42.68	114.14	0.890
322	2-methyl-3-ethylpentane	42.84	114.14	0.890
324	1,1,2-TriMcyCloC5	43.09	112.10	0.874
326	C8DiolefinC8-paraffin	43.23	114.23	0.859
328	C8-Olefins	43.37	112.13	0.874
330	C8-Olefins	43.52	112.13	0.874
332	2-Methylheptane	43.66	114.14	0.890
334	4-Methylheptane	43.83	114.14	0.890
336	C8-Diolefin + C7-Olefin	44.01	112.10	0.856
338	C8-Olefins	44.11	112.13	0.874
340	cis-1,4-DiMcyCloC6	44.37	112.10	0.874
342	trans-1,4-DiMcyCloC6	44.51	112.10	0.874
344	3-Methylheptane	44.61	114.14	0.890
346	3-Ethylhexane	44.71	114.14	0.890
348	C8-Diolefin	44.89	110.00	0.874
350	C8-Olefins	45.06	112.13	0.874
352	C8-Olefin	45.14	112.13	0.874
354	1,1-dimethylcyclohexane	45.24	112.13	0.874
356	C8-Olefin	45.43	112.13	0.874
358	C8-Olefin	45.58	112.13	0.874
360	cis-1-Ethyl-3-methylcyC	45.87	112.10	0.874
362	2,2,5-Trimethylhexane	45.96	128.20	0.888
364	t-1-E-3-McyC5	46.11	112.10	0.874
366	t-1-E-2-McyC5	46.26	112.10	0.874
368	1-methyl-1-ethylcyclope	46.34	112.22	0.874
370	1-Octene	46.48	112.13	0.874
372	C8-Olefin	46.61	112.13	0.874
374	t-1,2-DiMcyCloC6	46.71	112.10	0.874
376	C8-Olefins	46.79	112.13	0.874
378	C8-olefin	46.96	112.22	0.874
382	C8-olefins	47.12	112.22	0.874
380	trans-3-C8-Olefin	47.23	112.11	0.874
384	t-1,3-DiMcyCloC6	47.49	112.13	0.874
386	c-1,4-DiMcyCloC6	47.59	112.10	0.874
388	n-Octane	47.84	114.14	0.890
390	C8-Olefin	47.92	112.13	0.874
392	C8-Olefin	48.03	112.13	0.874
394	trans-2-Octene	48.19	112.13	0.874
396	Isopropylcyclopentane	48.32	112.10	0.874
398	C9-Olefin	48.35	126.14	0.874
400	C9-Olefin	48.59	126.14	0.874
402	C9-Olefin	48.71	126.14	0.874
404	C9-Olefin	48.76	126.14	0.874
406	2,2,4-Trimethylhexane	48.88	128.16	0.888
408	2,4,4-Trimethylhexane	48.89	128.16	0.888
410	C9-Olefins	48.92	126.14	0.874
412	2,3,5-Trimethylhexane	49.03	128.16	0.888
414	cis-2-Octene	49.10	112.13	0.874
416	2,2,3,4-TetraMC5	49.70	128.16	0.888
418	2,2-Dimethylheptane	49.96	128.16	0.888
420	c-1,2-DiMcyCloC6	50.08	112.10	0.874
422	2,4-Dimethylheptane	50.36	128.16	0.888
424	C9-Olefin	50.46	126.14	0.874
426	C9-Olefin	50.54	126.14	0.874
428	Ethylcyclohexane	50.59	112.10	0.874
430	Propylcyclopentane	50.67	112.10	0.874
432	2-Methyl-4-Ethylhexane	50.90	128.20	0.888
434	2,6-Dimethylheptane	51.02	128.20	0.888
436	C9-Olefin	51.14	126.14	0.874
438	1,1,4-TriMcyCloC6	51.21	126.14	0.874

TABLE X2.5 *Continued*

	Peak Number and Component Name	RT (min)	MW	RRF
440	C9-Olefins	51.29	126.14	0.874
442	C9-Olefins	51.44	126.14	0.874
444	1,1,3-trimethylcyclohex	51.45	126.24	0.874
446	2,5 & 3,5-DMheptane	51.64	128.16	0.888
448	C9-Olefins	51.69	126.14	0.874
450	3,3-Dimethylheptane	51.76	128.20	0.888
452	C9-Isoparaffin	51.90	128.26	0.888
454	C9-Olefins	52.05	126.14	0.874
456	2,3,3-Trimethylhexane	52.30	128.16	0.888
458	C9-Olefins	52.38	126.14	0.874
460	Ethylbenzene	52.44	106.08	0.827
462	C9-Olefins	52.58	126.14	0.874
464	t-1,2,4-TrimethylcyC6	52.69	126.14	0.874
466	C9-Olefins	52.76	126.14	0.874
468	2,3,4-Trimethylhexane	52.88	128.20	0.888
470	C9-olefin	52.94	126.24	0.874
472	3,3,4-Trimethylhexane	53.07	128.16	0.888
474	m-Xylene	53.28	106.08	0.827
476	p-Xylene	53.37	106.08	0.827
478	2,3-Dimethylheptane	53.54	128.16	0.888
480	3,5-Dimethylheptane	53.71	128.20	0.888
482	3,4-Dimethylheptane	53.76	128.16	0.888
484	C9-Olefin	53.89	126.14	0.874
486	3-Methyl-3-ethylhexane	53.99	128.16	0.888
488	C9-Olefin	54.09	126.14	0.874
490	4-Ethylheptane	54.14	128.16	0.888
492	4-MC8+C9-Olefin	54.27	128.20	0.888
494	2-Methyloctane	54.36	128.20	0.888
496	C9-Olefin	54.47	126.14	0.874
498	C9-Isoparaffin	54.67	128.20	0.888
500	C9-Olefin	54.74	126.14	0.874
502	3-Ethylheptane	54.81	128.20	0.888
504	3-Methyloctane	54.93	128.20	0.888
506	C9-Isoparaffin	55.02	126.14	0.874
508	c-1,2,4-TriMcyC6	55.08	126.14	0.874
510	1,1,2-TriMcyC6	55.10	126.14	0.874
512	o-Xylene	55.24	106.08	0.827
514	C9-Olefin	55.38	126.14	0.874
516	C9-Isoparaffin	55.47	128.26	0.888
518	C9-Isoparaffin	55.56	128.26	0.888
520	C9-olefin	55.63	128.20	0.874
522	t-1-E-4-M-cyC6	55.81	126.14	0.874
524	cis-1-E-4-McyC6	55.90	126.14	0.874
526	C9-Isoparaffin	56.09	128.20	0.888
528	1-Nonene	56.32	126.14	0.874
530	Isobutylcyclopentane	56.45	126.14	0.874
532	C9-Isoparaffin	56.74	128.20	0.888
534	trans-3-Nonene	56.89	126.14	0.874
536	cis-3-Nonene	57.03	126.14	0.874
538	C9-Isoparaffin	57.16	128.20	0.888
540	n-Nonane	57.26	128.16	0.888
542	C10-Olefin	57.40	140.16	0.874
544	trans-2-Nonene	57.48	126.14	0.874
546	1-M-1-Ecyclohexane	57.63	126.14	0.874
548	1-M-2-PcycloC5	57.70	126.14	0.874
550	C10-Olefin	57.79	140.16	0.874
552	C10-Isoparaffin	57.87	142.17	0.887
554	C10-Isoparaffin	57.96	142.17	0.887
556	Isopropylbenzene	58.03	118.08	0.832
558	cis-2-Nonene	58.12	126.14	0.874
560	tert-Butylcyclopentane	58.15	126.14	0.874
562	C9-Olefins	58.30	126.14	0.874
564	nonene	58.42	126.24	0.874
566	Isopropylcyclohexane	58.47	126.14	0.874
568	3,3,5-Trimethylheptane	58.61	142.17	0.887
570	2,2-Dimethyloctane	58.78	142.17	0.887
572	2,4-Dimethyloctane	58.90	142.17	0.887
574	1-M-4-isopropylcyC6	59.00	140.16	0.874
576	sec-Butylcyclopentane	59.16	126.14	0.874
578	Propylcyclohexane	59.19	126.14	0.874
580	2,5-Dimethyloctane	59.31	142.17	0.887
582	Butylcyclopentane	59.62	126.14	0.874
584	2,6-Dimethyloctane	59.63	142.17	0.887
586	3,6-Dimethyloctane	59.97	142.17	0.887
588	1-M-2-EcycloC6	60.05	126.14	0.874

TABLE X2.5 *Continued*

Peak Number	Component Name	RT (min)	MW	RRF
590	C10-Olefin	60.21	140.16	0.874
592	Propylbenzene	60.30	120.09	0.832
594	3,3-Dimethyloctane	60.51	142.17	0.887
596	3-Methyl-5-ethylheptane	60.61	142.17	0.887
598	C10-Olefin	60.69	140.16	0.874
600	1-Ethyl-3-methylbenzene	60.87	120.09	0.832
602	1-Ethyl-4-methylbenzene	61.01	120.09	0.832
604	Naphthene	61.26	140.27	0.874
606	1,3,5-Trimethylbenzene	61.30	120.09	0.832
608	2,3-Dimethyloctane	61.43	142.17	0.887
610	5-Methylnonane	61.83	142.17	0.887
612	4-Methylnonane	61.95	142.17	0.887
614	2-Methylnonane	62.11	142.17	0.887
616	1-Ethyl-2-methylbenzene	62.21	120.09	0.832
618	3-Ethylheptane	62.37	142.17	0.887
620	Naphthene	62.45	140.16	0.874
622	3-Methylnonane	62.58	142.17	0.887
624	C10-Olefin	62.77	140.16	0.874
626	C10-Isoparaffin	62.92	142.17	0.887
628	C10-Isoparaffin	62.99	142.17	0.887
630	1,2,4-Trimethylbenzene	63.18	120.09	0.832
632	C10-Isoparaffin	63.30	142.17	0.887
634	C10-Isoparaffin	63.37	142.17	0.887
636	Isobutylcyclohexane	63.51	140.27	0.874
638	C10-Isoparaffin	63.62	142.28	0.887
640	C10-Isoparaffin	63.67	142.37	0.887
642	1-Decene	63.72	140.27	0.874
644	C10-Isoparaffin	63.81	142.28	0.887
646	C10-Isoparaffin	63.92	142.17	0.887
648	C10-Aromatic	63.99	134.11	0.837
654	Isobutylbenzene	64.11	134.11	0.837
656	t-1-M-2-propylcycC6	64.24	140.16	0.874
652	Naphthene	64.26	140.16	0.874
650	C10-Isoparaffin	64.29	142.17	0.887
658	C10-Isoparaffin	64.36	142.17	0.887
660	sec-Butylbenzene	64.38	134.11	0.837
662	n-Decane	64.49	142.17	0.887
664	C11-Isoparaffin	64.65	156.19	0.886
666	C11-Isoparaffin	64.74	156.19	0.886
668	1,2,3-Trimethylbenzene	65.01	120.09	0.832
670	1-M-3-isopropylbenzene	65.10	134.11	0.837
674	1-M-4-isopropylbenzene	65.32	134.11	0.837
676	C11-Isoparaffin	65.47	156.19	0.886
672	C11-Isoparaffin	65.50	156.19	0.886
678	C11-Isoparaffin	65.57	156.19	0.886
680	2,3-Dihydroindene	65.75	118.08	0.819
682	sec-Butylcyclohexane	66.07	140.16	0.874
684	C11-IsoParaffin	66.12	156.19	0.886
686	1-M-2-isopropylbenzene	66.15	134.11	0.837
688	3-Ethylnonane	66.31	156.19	0.886
690	C11-Isoparaffin	66.38	156.19	0.886
692	Naphthene	66.48	140.16	0.874
694	C11-Isoparaffin	66.63	126.19	0.886
696	1,3-Diethylbenzene	66.84	134.11	0.837
698	1-M-3-propylbenzene	67.01	134.11	0.837
700	1,4-Diethylbenzene	67.10	134.11	0.837
702	1-M-4-propylbenzene	67.25	134.11	0.837
704	Butylbenzene	67.32	134.11	0.837
706	3,5-DM-1-Ebenzene	67.42	134.11	0.837
708	1,2-Diethylbenzene	67.59	134.11	0.837
710	C11-Isoparaffin	67.86	156.19	0.886
712	C10-Aromatic	67.98	134.11	0.837
714	C10-Aromatic	68.01	134.11	0.837
716	C10-Aromatic	68.09	134.11	0.837
718	1-M-2-propyl benzene	68.14	134.11	0.837
720	C10-Aromatic	68.14	134.22	0.837
722	5-Methyldecane	68.17	156.19	0.886
724	4-Methyldecane	68.20	156.19	0.886
726	2-Methyldecane	68.25	156.32	0.886
728	C11-Isoparaffin	68.36	156.19	0.886
730	1,4-DM-2-Ebenzene	68.59	134.11	0.837
732	1,3-DM-4-Ebenzene	68.71	134.11	0.837
734	C11-Isoparaffin	68.77	156.19	0.886
736	3-Methyldecane	68.95	156.19	0.886
738	C1-indane	68.99	132.21	0.837

TABLE X2.5 *Continued*

Peak Number and Component Name	RT (min)	MW	RRF	
740	1,2-DM-4-Ebenz+C1indan	69.05	134.11	0.837
742	C11-Isoparaffin	69.12	156.19	0.886
744	1,3-DM-2-Ebenzene	69.48	134.11	0.837
746	C11-Isoparaffin	69.78	156.19	0.886
748	C11-Isoparaffin	69.92	156.19	0.886
750	1-M-4-tert-butylbenzene	70.06	148.13	0.840
752	1,2-DM-3-ethylbenzene	70.18	134.11	0.837
754	1-E-2-isopropylbenzene	70.35	148.13	0.840
756	n-Undecane	70.49	156.19	0.886
758	1-E-4-isopropylbenzene	70.57	148.13	0.840
760	C12-Isoparaffin	70.70	170.20	0.885
762	1,2,4,5-TetraMbenzene	70.82	134.11	0.837
764	2-Methylbutylbenzene	70.90	148.13	0.840
766	1,2,3,5-TetraMbenzene	71.00	134.11	0.837
768	3-methylbutylbenzene	71.03	148.25	0.840
770	C11-Aromatic	71.04	148.13	0.840
772	C12-Isoparaffin	71.14	170.20	0.885
774	C11-Aromatic	71.33	148.13	0.840
776	C11-Aromatic	71.41	148.13	0.840
778	C11-Aromatic	71.47	148.13	0.840
780	1-tert-B-2-methylbenzen	71.69	148.13	0.840
782	C11-Aromatic	71.80	148.13	0.840
784	1-Ethyl-2-propylbenzene	71.86	148.13	0.840
786	C11-Aromatic	72.03	148.13	0.840
788	C11-Aromatic	72.17	148.13	0.840
790	C11-Aromatic	72.24	148.13	0.840
792	1-Methyl-3-butylbenzene	72.35	148.13	0.840
794	C11-aromatic	72.47	148.25	0.840
796	1,2,3,4-TetraMbz+C11aro	72.54	148.25	0.840
798	Pentylbenzene	72.73	148.13	0.840
800	t-1M-2-(4MC5)-cyC5	72.80	168.33	0.874
802	C11-Aromatic	72.95	148.13	0.840
804	C11-Aromatic	73.16	148.13	0.840
806	C11-Aromatic	73.19	148.13	0.840
808	C12-Isoparaffin	73.23	170.20	0.885
810	1,2,3,4-Tetrahydronapht	73.35	132.09	0.824
812	1-tert-Butyl-3,5-DMbz	73.53	162.30	0.843
814	Naphthalene	73.72	128.06	0.799
816	1,1-dimethylindane	73.82	146.10	0.829
818	1,2-dimethylindane	73.92	146.10	0.829
820	1,6-dimethylindane	74.00	146.10	0.829
822	C11-Aromatic	74.10	148.25	0.840
824	1-ethylindane	74.18	146.10	0.829
826	2-ethylindane	74.33	146.10	0.829
828	Ethyl-1,3,5-trimethylbe	74.44	148.25	0.840
830	1,3-dipropylbenzene	74.61	162.34	0.843
832	n-Dodecane	74.98	170.20	0.885
834	Ethyl-1,2,4-trimethylbe	75.38	148.25	0.840
836	C11-Aromatic	75.61	148.10	0.840
838	C11-Aromatic	75.90	148.13	0.840
840	C12-aromatic+C2-indane	75.98	162.30	0.843
842	2,4-dimethylindane	76.07	146.10	0.829
844	4-ethylindane	76.20	146.10	0.829
846	1-tert-butyl-4-ethylbez	76.30	162.30	0.843
848	1,3-dimethylindane	77.10	146.10	0.829
850	1-methyl-4-pentylbenzene	77.20	162.30	0.843
852	4,7-dimethylindane	77.23	146.10	0.829
854	5,6-dimethylindane	77.26	146.10	0.829
856	C12-aromatic	77.30	162.30	0.843
858	Hexylbenzene	77.33	162.30	0.843
860	C6-Benzene	77.36	162.30	0.843
862	C6-benzene	77.45	162.30	0.843
864	C6-benzene	77.50	162.30	0.843
866	4,5-dimethylindane	77.60	146.10	0.829
868	C6-benzene	77.75	162.30	0.843
870	2-Methylnaphthalene	78.09	142.08	0.806
872	C6-benzene	78.19	162.30	0.843
874	C6-benzene	78.28	162.30	0.843
876	n-Tridecane	78.54	184.22	0.884
878	1-Methylnaphthalene	78.64	142.08	0.806
880	C6-benzene	79.15	162.30	0.843
882	C2-tetralin	79.66	160.20	0.843
884	C6-benzene	80.11	162.30	0.843
886	C6-benzene	80.49	162.30	0.843
888	C13-Isoparaffin	80.87	184.22	0.883

TABLE X2.5 *Continued*

	Peak Number and Component Name	RT (min)	MW	RRF
890	trans-7-Decene	81.24	140.20	0.874
895	2,6-Dimethylnaphthalene	81.38	156.30	0.812
900	2,7-Dimethylnaphthalene	81.50	156.30	0.812
905	N-tetradecane	81.59	198.34	0.883
910	1,3-Dimethylnaphthalene	82.43	156.30	0.812
940	1,2-Dimethylnaphthalene	82.55	156.30	0.812
950	n-Pentadecane	82.57	212.34	0.883
915	1,6-Dimethylnaphthalene	82.59	156.30	0.812
925	1,4-Dimethylnaphthalene	82.86	156.30	0.812
920	1,5-Dimethylnaphthalene	83.01	156.30	0.812
930	Acenaphthalene	83.26	156.30	0.801

SUMMARY OF CHANGES

Subcommittee D02.04 has identified the location of selected changes to this standard since the last issue (D6729 – 04 (2009)) that may impact the use of this standard. (Approved Oct. 1, 2014.)

(1) Revised Scope and Referenced Documents sections.

(2) Revised **Table A1.1** and **Table X2.5**.

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