



Standard Test Method for Determination of Individual Components in Spark Ignition Engine Fuels by 100–Metre Capillary (with Precolumn) High- Resolution Gas Chromatography¹

This standard is issued under the fixed designation D 6730; 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 from 0.01 to approximately 30 mass %. The test method may be applicable to higher and lower concentrations for the individual components; however, the user must verify the accuracy if the test method is used for components with concentrations outside the specified ranges.

1.3 This test method also determines methanol, ethanol, *t*-butanol, methyl *t*-butyl ether (MTBE), ethyl *t*-butyl ether (ETBE), and *t*-amyl methyl ether (TAME) in spark ignition engine fuels in the concentration range from 1 to 30 mass %. However, the cooperative study data provided insufficient statistical data for obtaining a precision statement for these compounds.

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 naphthenic (for example, virgin naphthas) constituents above *n*-octane may reflect significant errors in PONA-type groupings. Based on the gasoline samples in the interlaboratory cooperative study, this test method is applicable to samples containing less than 25 mass % of olefins. However, some interfering co-elution 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. Annex A1 of this test method compares results of the test method 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 the specific test methods listed in the reference section.

1.4.1 Total olefins in the samples may be obtained or confirmed, or both, if necessary, by Test Method D 1319 (volume %) or other test methods, such as those based on multidimensional PONA-type of instruments.

1.5 If water is or is suspected of being present, its concentration may be determined, if desired, by the use of Test Method D 1744 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 D 4815 and D 5599 for oxygenates, and Test Method D 5623 for sulfur compounds, or equivalent.

1.6 *This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.*

2. Referenced Documents

2.1 ASTM Standards:

- D 1319 Test Method for Hydrocarbon Types in Liquid Petroleum Products by Fluorescent Indicator Adsorption²
- D 1744 Test Method for Determination of Water in Liquid Petroleum Products by Karl Fisher Reagent³
- D 3700 Practice for Containing Hydrocarbon Fluid Samples Using a Floating Piston Cylinder⁴
- D 4057 Practice for Manual Sampling of Petroleum and

¹ This test method is under the jurisdiction of ASTM Committee D02 on Petroleum Products and Lubricants and is the direct responsibility of Subcommittee D02.04.01 on Gas Chromatographic Methods.

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² *Annual Book of Standards*, Vol 05.01.

³ Discontinued; see 1999 *Annual Book of ASTM Standards*, Vol 05.01.

⁴ *Annual Book of Standards*, Vol 05.02.

- Petroleum Products⁴
- D 4177 Practice for Automatic Sampling of Petroleum and Petroleum Products⁴
- D 4307 Practice for Preparation of Liquid Blends for Use as Analytical Standards⁴
- D 4626 Practice for Calculation of Gas Chromatographic Response Factors⁴
- D 4815 Test Method for Determination of MTBE, ETBE, TAME, DIPE, *tertiary*-Amyl Alcohol and C₁ to C₄ Alcohols in Gasoline by Gas Chromatography⁴
- D 5580 Test Method for Determination of Benzene, Toluene, Ethylbenzene, *p/m*-Xylene, *o*-Xylene, C₉ and Heavier Aromatics and Total Aromatics in Finished Gasoline by Gas Chromatography⁵
- D 5599 Test Method for Determination of Oxygenates in Gasoline by Gas Chromatography and Oxygen Selective Flame Ionization Detection⁵
- D 5623 Test Method for Sulfur Compounds in Light Petroleum Liquids by Gas Chromatography and Sulfur Selective Detection⁵
- E 355 Practice for Gas Chromatography Terms and Relationships⁶
- E 594 Practice for Testing Flame Ionization Detectors Used in Gas or Supercritical Fluid Chromatography⁶
- E 1510 Practice for Installing Fused Silica Open Tubular Capillary Columns in Gas Chromatography⁶

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 E 355.

4. Summary of Test Method

4.1 A representative sample of the petroleum liquid is introduced into a gas chromatograph equipped with an open tubular (capillary) column coated with a methyl silicone liquid phase, modified with a capillary precolumn. 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 presented on a strip chart recorder or digitally, or both, 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 % is determined by normalization of the peak areas after correction with detector response factors. Unknown components are reported as a total unknown mass %.

5. Significance and Use

5.1 Knowledge of the 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 can be determined through the use of this test method.

5.2 This test method is adopted from earlier development and enhancement.^{7,8,9,10} The chromatographic operating conditions and column tuning process, included in this test method, were developed to provide and enhance the separation and subsequent determination of many individual components not obtained with previous single-column analyses. The column temperature program profile is selected to afford the maximum resolution of possible co-eluting components, especially where these are of two different compound types (for example, a paraffin and a naphthene).

5.3 Although a majority of the individual hydrocarbons present in petroleum distillates are determined, some co-elution of compounds is encountered. If this test method is utilized to determine 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, or both, constituents above octane may reflect significant errors in PONA-type groupings.

5.4 If water is or is suspected of being present, its concentration is determined by the use of Test Method D 1744. Other compounds containing oxygen, sulfur, nitrogen, and so forth may also be present, and may co-elute with the hydrocarbons. When known co-elution exists, these are noted in the test method data tables. If determination of these specific compounds is required, it is recommended that test methods for these specific materials be used, such as Test Method D 4815 and D 5599 for oxygenates, Test Method D 5580 for aromatics, and Test Method D 5623 for sulfur compounds.

6. Apparatus

6.1 *Gas Chromatograph*—Instrumentation capable of column oven temperature programming, from subambient (5°C) to at least 200°C, in 0.1°C/min or less rate increments, is required. Multi-step column oven temperature programming is required, consisting of an initial hold time, an initial temperature program followed by an isothermal temperature hold and another programmed temperature rise. A heated flash vaporizing injector designed to provide a linear sample split injection (that is, 200:1) is required for proper sample introduction. The associated carrier gas controls must be of sufficient precision to provide reproducible column flows and split ratios in order to maintain analytical integrity. A hydrogen flame ionization

⁷ Johansen, N.G., and Ettre, L.S., "Retention Index Values of Hydrocarbons on Open Tubular Columns Coated with Methyl Silicone Liquid Phases," *Chromatographia*, Vol 5, No. 10, October 1982.

⁸ Johansen, N.G., Ettre, L.S., and Miller, R.L., "Quantitative Analysis of Hydrocarbons by Structural Group Type in Gasolines and Distillates. Part 1," *Journal of Chromatography*, 256, 1983, pp. 393-417.

⁹ Kopp, V.R., Bones, C.J., Doerr, D.G., Ho, S.P., and Schubert, A.J., "Heavy Hydrocarbon/Volatility Study: Fuel Blending and Analysis for the Auto/Oil Air Quality Improvement Research Program," SAE Paper No. 930143, March 1993.

¹⁰ Schubert, A.J. and Johansen, N.J., "Cooperative Study to Evaluate a Standard Test Method for the Speciation of Gasolines by Capillary Gas Chromatography," SAE Paper No. 930144, March 1993.

⁵ *Annual Book of Standards*, Vol 05.03.

⁶ *Annual Book of Standards*, Vol 14.02.

detector, with associated gas controls and electronics, designed for optimum response with open tubular columns, shall conform to the specifications as described in Practice E 594, as well as having an operating temperature range of up to at least 250°C.

6.2 Sample Introduction—Manual or automatic liquid sample injection to the splitting injector may be employed. Automated injections are highly recommended. Micro-syringes, auto-syringe samplers, or valves capable of 0.1 to 0.5 µL injections are suitable. It should be noted that some syringes and improper injection techniques as well as inadequate splitter design could result in sample fractionation. This must be determined in accordance with Section 10.

6.3 Electronic Integrator—Any electronic integration device used for quantitating these analyses shall meet or exceed these minimum requirements:

6.3.1 Capacity to handle 400 or more peaks per analysis.

6.3.2 Normalized area percent calculation with response factors.

6.3.3 Noise and spike rejection.

6.3.4 Accurate area determination of fast (1 to 2 s) peaks (10 Hz or greater sampling rate).

6.3.5 Maintain peak detection sensitivity for narrow and broad peaks.

6.3.6 Positive and negative sloping baseline correction.

6.3.7 Perpendicular drop and tangent skimming as needed.

6.3.8 Display of baseline used to ensure correct peak area determination.

6.4 Open Tubular Column—The column used for this test method consists of a primary (100 m) analytical column and a precolumn. The ability to provide the required component separations is dependent on the precise control of the column selectivity, which is typically slightly more than that exhibited by current commercially available columns. Some older columns, and columns that have a sample residue from repeated use without conditioning, may exhibit the required polarity. Until adequate columns are commercially available, the currently used methyl silicone columns can be modified or *tuned* to meet the method column specifications. See Section 11 for a description of the column performance specifications and Annex A1 for a description of the column modification procedure.

6.4.1 The primary gas chromatographic column used for this test method will meet the following specifications.

Material	fused silica
Length	100 m
Internal diameter	0.25 mm
Liquid phase	methyl silicone
Film thickness	0.50 µm
Theoretical plates, n, pentane at 35°C	~ 400 000 to 500 000
Retention factor, k, pentane at 35°C	0.45 to 0.50
Resolution, R, <i>t</i> -butanol and 2-methylbutene-2 at 35°C	3.25 to 5.25
Peak symmetry, <i>t</i> -butanol at 35°C	> 1.0 to < 5.0

6.4.2 Precolumn—A variable length (1 to 4 m) of 5 % phenyl/95 % dimethylpolysiloxane fused silica open tubular column (0.25 mm inside diameter) is added to the front (injector) end of the 100 m column, as described in Annex A1.

7. Reagents and Materials

7.1 *Carrier Gas*—Helium, 99.999 % pure. (**Warning**—Helium, air, nitrogen, compressed gas under pressure.)

7.2 *Oxidant*—Air, 99.999 % pure. (**Warning**—see 7.1.)

7.3 *Detector Makeup Gas*—Nitrogen, 99.999 % pure. (**Warning**—see 7.1.)

7.4 *Fuel Gas*—Hydrogen, 99.999 % pure. (**Warning**—Hydrogen, flammable gas under high pressure.)

7.5 *Reference Standards*:

7.5.1 *Purity of Reagents*—Reagent grade chemicals shall be used in all tests. Unless otherwise indicated, it is intended that all reagents conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society¹¹ where such specifications are available. Other grades may be used, provided it is first ascertained that the reagent is of sufficiently high purity to permit its use without lessening the accuracy of the determination.

7.5.2 *Methanol*—(**Warning**—These materials are flammable and may be harmful or fatal, if ingested or inhaled.)

7.5.3 *Ethanol*—Only absolute ethanol of 99.5 minimum percent meets the requirements of this test method. (**Warning**—see 7.5.2.)

7.5.4 *Hydrocarbon and Other Component References*—Individual and mixed component reference materials are commercially available and may be used to establish qualitative and quantitative calibration. (**Warning**—see 7.5.2.)

7.5.5 *System and Column Evaluation Mixture*—A quantitatively prepared mixture, complying with Practice D 4307, of individual hydrocarbons and oxygenates of interest is used for system and column evaluation (see Table 1). (**Warning**—see 7.5.2.) Fig. 1 is a chromatogram of the recommended mixture in Table 1.

8. Sampling

8.1 Hydrocarbon liquids with Reid vapor pressures of 110 kPa (16 psi) or less may be sampled either into a floating piston cylinder or into an open container (Practices D 4057 and D 4177). If the sample as received does not meet the upper boiling range requirements of 1.1, it may be necessary to extend the analysis time and raise the upper column temperature of this test method to ensure complete elution of higher boiling range sample material from the column.

8.1.1 *Piston Cylinder Sampling*—Refer to Practice D 3700 for instructions on transferring a representative sample of a hydrocarbon fluid from a source into a floating piston cylinder. Add inert gas to the ballast side of the floating piston cylinder to achieve a pressure of 350 kPa (45 psi) above the vapor pressure of the sample.

8.1.2 *Open Container Sampling*—Refer to Practice D 4057 for instructions on manual sampling from bulk storage into open containers. Stopper the container immediately after taking a sample.

¹¹ *Reagent Chemicals, American Chemical Society Specifications*, American Chemical Society, Washington, DC. For suggestions on the testing of reagents not listed by the American Chemical Society, see *Analar Standards for Laboratory Chemicals*, BDH Ltd., Poole, Dorset, U.K., and the United States Pharmacopeia and National Formulary, U.S. Pharmacopeial Convention, Inc. (USPC), Rockville, MD.

TABLE 1 System and Column Evaluation Mixture

	%
Ethanol	8.00
<i>n</i> -pentane	2.00
<i>t</i> -butanol	0.50
2-methylbutene-2	2.50
2,3-dimethylbutane	0.50
Methyl- <i>t</i> -butyl ether	10.00
<i>n</i> -hexane	2.00
1-methylcyclopentene	0.50
Benzene	1.00
Cyclohexane	28.90
3-ethylpentane	0.20
1,2- <i>t</i> -dimethylcyclopentane	0.50
<i>n</i> -heptane	2.00
2,3,3-trimethylpentane	0.50
Toluene	7.00
<i>n</i> -octane	2.00
Ethylbenzene	25.00
<i>p</i> -xylene	1.00
2,3-dimethylheptane	0.20
<i>n</i> -nonane	2.00
5-methylnonane	0.20
1-methyl-2-ethylbenzene	0.50
<i>n</i> -decane	1.00
<i>n</i> -undecane	0.50
1,2,3,5-tetramethylbenzene	0.25
Naphthalene	0.50
<i>n</i> -dodecane	0.25
1-methylnaphthalene	0.25
<i>n</i> -tridecane	0.25

8.2 Preserve the sample by cooling to approximately 4°C and maintaining that temperature prior to analysis.

8.3 Transfer an aliquot of the cooled sample to a precooled septum vial and seal immediately.

8.4 Obtain the test specimen for analysis directly from the sealed septum vial, for either manual or automatic injection.

9. Preparation of Apparatus

9.1 Install the 100-m column and, if required, a precolumn according to the manufacturer's or supplier's instructions and Annex A1. See Practice E 1510 for recommended installation procedures.

9.2 Determine the required length of the precolumn in accordance with Annex A1. Adjust the operating conditions of the gas chromatograph to those listed in Table 2 or as determined by Section 12 and Annex A1.

9.3 During setup and, when not performing analyses, it is advisable to turn off the cryogenic operation and set the column oven temperature at 35°C. Attach the column outlet to the flame ionization detector inlet and check for leaks throughout the system. If leaks are found, tighten or replace fittings before proceeding.

9.4 Confirm or adjust, or both, the column carrier gas flow rate by making injections of methane or natural gas. *The methane retention time shall be 7.00 ± 0.02 min with the column oven temperature at 35°C, which results in an average linear velocity of 24 cm/s, as determined using Eq 1. This will result in a methane retention time of 6.53 min at 5°C. Raising or lowering the carrier gas pressure to the injector makes flow rate adjustment. A starting point of 277-kPa (40-psig) helium pressure is recommended, although columns requiring as high as 332-kPa (48-psig) helium have been encountered.*

$$\text{average linear gas velocity: } u_{\text{ave}} (\text{cm/s}) = \text{column length (cm)} / t_{\text{M(s)}} \quad (1)$$

9.5 After final adjustment of the carrier gas flow rate, note the carrier gas inlet pressure. Measure and, if necessary, readjust the injector split flow rate to give the specified or desired split ratio. Calculate the column outlet flow rate using 9.5.1 and the split ratio using 9.5.2.

9.5.1 *Column Carrier Gas Flow Rate (at outlet):*

9.5.1.1 $P = (\text{head pressure (psig)} + \text{ambient pressure}) / \text{ambient pressure}.$

9.5.1.2 $j = \text{compressibility factor} = 3/2((P^2-1)/(P^3-1)).$

9.5.1.3 $u_o = u_{\text{ave}}/j = \text{column outlet velocity}.$

9.5.1.4 $A_c = \pi(r)^2 = \text{column cross-sectional area (cm}^2\text{)},$ where $r = \text{column internal radius (cm)}.$

9.5.1.5 $\text{Flow rate (cm}^3\text{/min)} = u_o \times A_c \times 60.$

9.5.2 *Injection Split Ratio*—(Split flow rate + column flow rate)/column flow rate.

9.5.3 *Example*—Using a 100-m × 0.25-mm capillary column:

9.5.3.1 $U_{\text{ave}} = 100 \times 100 / 6.98 \times 60 = 23.88 \text{ cm/s}.$

9.5.3.2 $P = 40 \text{ psig} + 12.0 / 12.0 = 4.33.$

9.5.3.3 $j = 3/2((18.778-1)/(81.370-1)) = 0.33$

9.5.3.4 $u_o = 23.88 / 0.33 = 71.96 \text{ cm/s}.$

9.5.3.5 $A_c = \pi(0.025/2)^2 = 4.9 \times 10^{-4} \text{ cm}^2.$

9.5.3.6 $\text{Flow rate} = 71.96 \times 4.9 \times 10^{-4} \times 60 = 2.12 \text{ cm}^3\text{/min}.$

9.5.3.7 $\text{Split Ratio} = (192 + 2.12) / 2.12 = 91.6:1.$

9.6 Make a blank analysis (no sample injection) run to ensure proper instrument operation and further condition the column and instrumentation. If stray peaks or a rising baseline signal is observed, the column oven shall be kept at the upper temperature until the baseline becomes steady and returns to within approximately 5 % of the starting temperature detector signal.

9.7 After any extended conditioning period, or if the instrument has been shut down, it is advisable to repeat 9.4, 9.5, and 9.6 to ensure proper carrier gas flows are being used and the column is clean.

10. Split Injection Linearity

10.1 Splitting injector linearity must be established to determine proper quantitative parameters and limits. The split ratio used is dependent upon the split linearity characteristics of the particular injector and the sample retention factor of the column. The retention factor of a particular column for a sample component is proportional to the amount of liquid phase (loading or film thickness) and the ratio of the column temperature to the component boiling point (vapor pressure). Overloading of the column may cause loss of resolution for some components and, since overloaded peaks are skewed, variance in retention times. This can lead to erroneous component identification. During column evaluations and split linearity studies, be aware of any peaks that may appear *front skewed*, indicating column overload. Note the component size and avoid conditions leading to this problem during actual analyses.

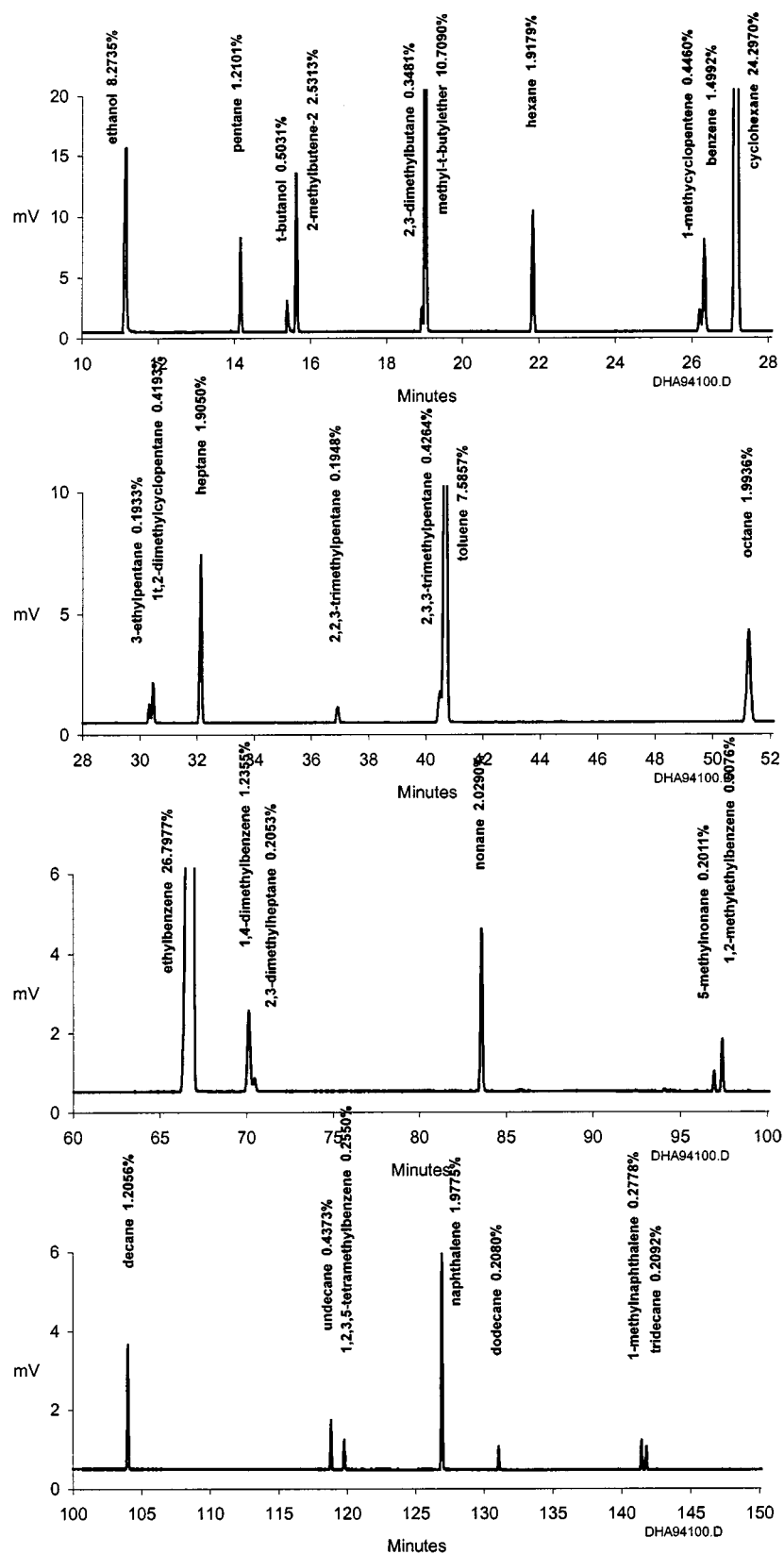


FIG. 1 DHA Speciation Analysis—System and Column Evaluation Mixture (7.5.5)

TABLE 2 GC Operating Conditions

Column Temperature Program	
Initial temperature	5°C
Initial time	10 min.
First program rate	5.0°/min
First hold temperature	50°C
First hold time	to the elution of ethylbenzene (~50 min)
Second program rate	1.5°/min
Final temperature	200°C
Final hold time	5 min
Injector	
Temperature	250°C
Split ratio	150:1
Sample size	0.1 - 0.2 µL
Detector	
Type	flame ionization
Temperature	250°C
Use manufacturers recommended detector gas flows or:	
Fuel gas	hydrogen at 30 mL/min
Oxidant	air at 300 mL/min
Make-up gas, where required	nitrogen at 20 mL/min
Carrier Gas	
Type	helium
Pressure	~ 277 kPa (40 psig)
Average linear velocity	24 cm/s at 35°C

10.2 Set the injector temperature and split ratio to the following values and, for each set of conditions inject the listed quantities of the system and column evaluation mixture (7.5.5), using the operating conditions listed in Table 2 or as determined in Section 12.

injector temperature: 250°C < $\begin{matrix} \text{split: 100:1} \\ \text{split: 200:1} \end{matrix}$ > sample: 0.2 µL, 0.5 µL, 1.0 µL

injector temperature: 300°C < $\begin{matrix} \text{split: 100:1} \\ \text{split: 200:1} \end{matrix}$ > sample: 0.2 µL, 0.5 µL, 1.0 µL

10.3 Compare the calculated concentrations to the known standard concentrations after calculating the corrected area normalization using the response factors from 13.2 and Table A1.1.

$$\begin{aligned} \% \text{ relative error} = \\ 100 \times (\text{concentration determined} \\ - \text{concentration known}) / \text{concentration known} \end{aligned} \quad (2)$$

10.4 Report and use only those combinations of conditions from 10.2 that result in 3 % or less relative error. This is the splitter linearity range.

11. Column Evaluation

11.1 In order to establish that a column will perform as required, the following specifications shall be determined for new column acceptability and are useful for periodic evaluation of column deterioration. These specification determinations can be made with or without a precolumn, since the precolumn will have little effect on their values. See Annex A1, Fig. A1.1, for examples of these determinations. After performing the steps in Sections 9 and 10, analyze the column performance mixture (7.5.5) at 35°C isothermal, at least through heptane. The remainder of the analysis may be ignored, but the remaining components must be eluted from the column prior to performing another analysis. Setting the column temperature to 220°C for an additional 20 min will be sufficient.

11.2 Calculate the retention factor (k) for pentane at 35°C:

$$k = (t_R - t_M) / t_M \quad (3)$$

where:

t_M = gas holdup time (methane), and
 t_R = retention time for pentane, min.

11.2.1 The retention factor must be between 0.45 and 0.50 for proper application of this test method.

11.3 Calculate the column efficiency using the pentane peak:

$$n = 5.545 (t_R / w_{1/2h})^2 \quad (4)$$

where:

n = column efficiency (theoretical plates),
 t_R = retention time of pentane, and
 $w_{1/2h}$ = peak width at half height.

11.3.1 The column efficiency must be at least 400 000 plates for proper application of this test method.

11.4 The selectivity of apparently identical columns toward hydrocarbons may vary regarding oxygenated compounds; either due to extraneous materials in the liquid phase, or due to activity of the column wall surface. The addition of a precolumn has little if any affect on the selectivity toward oxygenates (see Annex A1, Fig. A1.4). The relative resolution of oxygenates is inherent to the quality of the primary 100-m column, and is specified by the resolution of *t*-butanol from 2-methylbutene-2 at 35°C. Calculate the resolution:

$$R = 2(t_{R2-M-Butene-2} - t_{RTBA}) / 1.699(w_{1/2h2-M-Butene-2} + w_{1/2hTBA}) \quad (5)$$

11.4.1 The resolution for this pair at 35°C must be between 3.25 and 5.25.

11.5 Extraneous column effects, or instrumental effects such as an active injector liner, may cause adsorption of oxygenated compounds, commonly seen and referred to as *tailing*, and may increase their retention. If this effect is caused by instrumental activity, the problem should be corrected. If the column is inherently active, a new column should be obtained. A measure of the tailing can be made and specified by applying a *skewness* calculation, which determines a ratio of the distances from the peak apex perpendicular to the front and back of the peak at 5 % of the peak height. See Annex A1, Fig. A1.3 for an example of this calculation.

$$\text{skewness} = B/A \quad (6)$$

11.5.1 This test method shall be made using the *t*-butanol peak (0.5 %) in the analysis of the column performance mixture (7.5.5) at 35°C isothermal. The skewness ratio must be greater than 1.0 and not more than 5.0.

12. Optimization of Instrument Operating Conditions

12.1 The column temperature programming profile is dependent upon the individual column characteristics. Table 2 lists the programming profile determined for a 100-m methyl silicone column with a precolumn as determined in Annex A1. The profile is determined by establishing satisfactory separations for the sets of sample components listed in 12.3. It is not practical to expect complete separation of all components, so

the optimum for each column may contain some compromises, also dependent upon any particular other separations deemed important.

12.2 The use of retention indices to numerically express the relative location of components among themselves and to surrounding normal paraffins is a convenient convention. The indices are also useful in providing a system of component identification with complex analyses such as this. There are several schemes for calculating retention indices, the first of which is the Kovats method, developed to express the logarithmic relationship of retention times of a homologous series of compounds when chromatographed isothermally. While this test method is not an isothermal column temperature procedure, it does contain isothermal steps and the longer temperature program step is a slow rate. The use of the Kovats indices provides a closer relationship to previous work in this field than using the linear index format.

12.2.1 The formula for the calculation of Kovats retention indices is:

$$RI_i = 100 \times (n + (\log(t_i) - \log(t_n)) / (\log(t_{n+1}) - \log(t_n))) \quad (7)$$

where:

- RI = retention index,
- n = carbon number of n -paraffin,
- t_i = retention time of component,
- t_n = retention time of preceding n -paraffin, and
- t_{n+1} = retention time of next n -paraffin.

12.3 The following examples show the key or critical separations required for this analysis. Typical retention indices are given, and a description of the effect of instrumental conditions on the separation is provided.

12.3.1 *i*-butane/methanol and ethanol/3-methylbutene-1—The initial starting temperature of 5°C is dictated by these separations. A lower starting temperature is not necessary and a higher temperature would effect the next set. The retention indices should be about 380 for methanol and 456.5 for ethanol (Fig. 2).

12.3.2 *i*-propanol/2-methylbutene-1 and *t*-butanol/2-methylbutene-2—*i*-propanol will appear resolved between pentene-1 and 2-methyl-butene-1, *t*-butanol will appear resolved between *c*-pentene-2 and 2-methylbutene-2.

12.3.2.1 Higher temperatures will move the alcohols into the peaks ahead of them. At 35°C the alcohols will be located ahead of the pentene-1 and *c*-pentene-2, respectively (Fig. 3).

12.3.3 2,3-dimethylbutane/methyl-*t*-butylether—This separation is critical and the 5°C hold for 10 min determines its

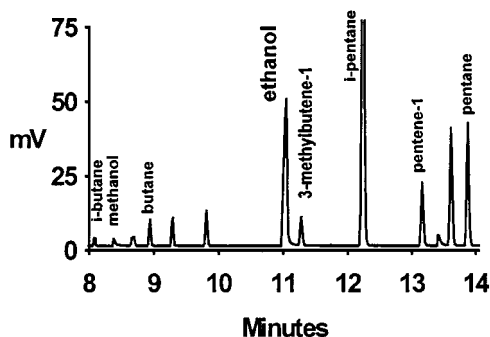


FIG. 2 *i*-butane/methanol and ethanol/3-methyl-butene-1

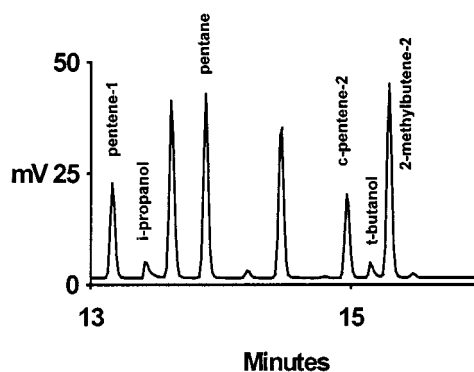


FIG. 3 *i* propanol/2-methylbutene-1 and *t* butanol/2-methylbutene-2

success. The retention indices should be about 569.5, 571.5, and 574.0 for 2,3-dimethylbutane, MTBE, and 2-methylpentane, respectively. If the MTBE is too close to the 2,3-DMC₄, use a 9 min initial hold. If too close to the 2-MC₅, use an 11 min hold (Fig. 4).

12.3.4 1-methylcyclopentene/benzene—This is a key separation that is used to specify the column selectivity. Changing column temperature produces only slight differences in this resolution (Fig. 5).

12.3.4.1 The 50°C column temperature is held isothermal until the elution of ethylbenzene. This is variable due to slight differences in the column retention factor.

12.3.5 2,3,3-trimethylpentane/toluene—This is a key separation that is used to specify the column selectivity. Column temperature has very little effect on this resolution, which is controlled by the column selectivity for aromatics (Fig. 6).

12.3.6 *p*-xylene/2,3-dimethylheptane—This is a key separation which limits the maximum length of the precolumn. If the column selectivity is too great the aromatics are retained and this separation is not achieved. If this resolution is excessive and the separation in 12.3.5 is insufficient, the precolumn should be lengthened slightly. Lowering the 50°C hold temperature to 48°C will increase this separation (Fig. 7).

12.3.7 117 (Unknown)/1,2-methylethylbenzene—The unknown isoparaffin (117) appears to be a component of alkylate

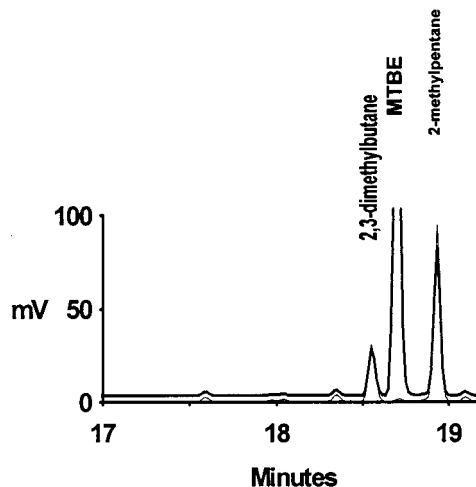


FIG. 4 2,3-dimethylbutane/methyl-*t* butylether

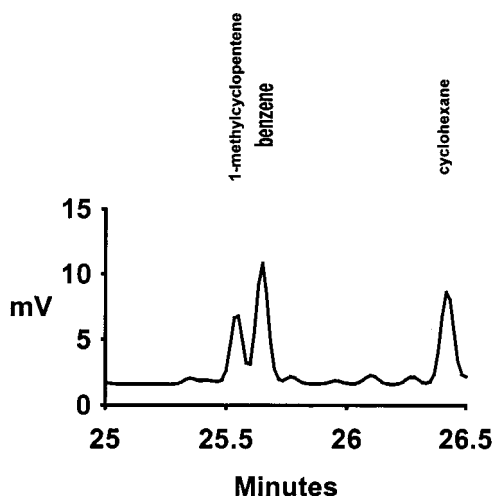


FIG. 5 1-methylcyclopentene/benzene

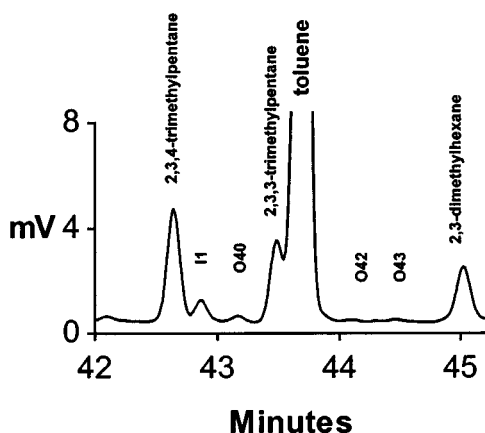


FIG. 6 2,3,3-trimethylpentane/toluene

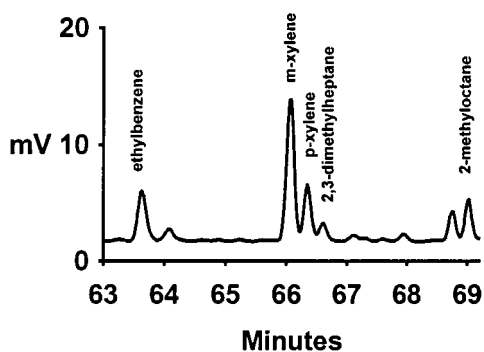


FIG. 7 *p*-xylene/2,3-dimethylheptane

and must be resolved from the aromatic. If the resolution is incomplete the final column temperature program rate of 1.5°/min. is adjusted to provide sufficient separation. Increase the rate in 0.1°/min increments to increase the resolution. This rate is also dictated by the separation requirements in 12.3.8. The proper rate will provide for both separations (Fig. 8).

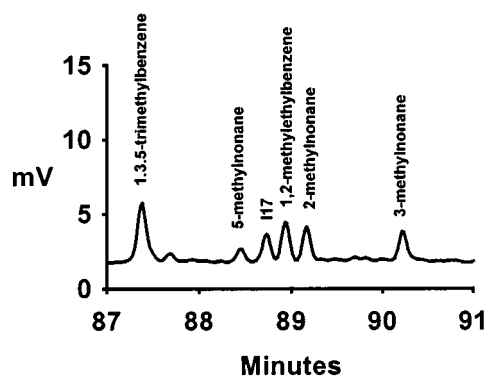


FIG. 8 I17 (unknown)/1,2-methylethylbenzene

12.3.8 *1-methylnaphthalene/tridecane*—The recommended final column temperature program rate of 1.5°/min. should also provide this separation. If the 1-MeNaph/*n*-C₁₃ resolution is incomplete this rate may be adjusted to provide sufficient separation. Lower the rate in 0.1°/min. increments to increase the resolution (Fig. 9).

13. Calibration

13.1 *Qualitative*—Determine the retention times of components by analyzing known reference mixtures or samples under identical conditions. Calculate retention indices from these data using 12.2. Table A1.1 provides a listing of typical values for this test method.

13.2 *Quantitative, Hydrocarbons*—Use theoretical response factors for correction of the detector response of hydrocarbons determined by this test method, unless response factors have been determined experimentally. The response of an FID to hydrocarbons is determined by the ratio of the molecular weight of the carbon in the analyte to the total molecular weight of the analyte. If experimentally determined response factors are to be used, they must be determined using known purity individual standards and calculated using Practice D 4626. The response factors, as listed in Table 3, are relative to that calculated for heptane. Calculations are based on the following equation:

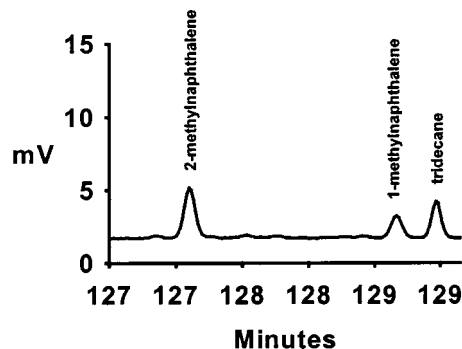


FIG. 9 1-methylnaphthalene/tridecane

TABLE 3 Theoretical FID Relative Response Factors

Carbon No.	Saturated Paraffins	Unsaturated Paraffins	Saturated Naphthenes	Unsaturated Naphthenes	Aromatics
1	1.1207	-	-	-	-
2	1.0503	-	-	-	-
3	1.0268	0.9799	-	-	-
4	1.0151	0.9799	-	-	-
5	1.0080	0.9799	0.9799	0.9517	-
6	1.0034	0.9799	0.9799	0.9564	0.9095
7	1.0000	0.9799	0.9799	0.9598	0.9195
8	0.9975	0.9799	0.9799	0.9623	0.9271
9	0.9955	0.9799	0.9799	0.9642	0.9329
10	0.9940	0.9799	0.9799	0.9658	0.9376
11	0.9927	0.9799	0.9799	0.9671	0.9415
12	0.9916	0.9799	0.9799	0.9681	0.9447
13	0.9907	0.9799	0.9799	0.9690	0.9474
14	0.9899	0.9799	0.9799	0.9698	0.9497
15	0.9893	0.9799	0.9799	0.9705	0.9517

$$F_i = (((C_{aw} \times C_n) + (H_{aw} \times H_n)/C_n) \times 0.83905)/C_{aw} \quad (8)$$

where:

F_i = relative response factor for a hydrocarbon type group of a particular carbon number.

C_{aw} = atomic weight of carbon 12.011,

C_n = number of carbon molecules in the group,

H_{aw} = atomic weight of hydrogen, 1.008,

H_n = number of hydrogen molecules in the group,
0.83905 is the correction factor with heptane as unity (1.0000), and
0.7487 is used with methane as unity.

13.3 Quantitative, Oxygenates—Determine response factors for methanol, ethanol, and other oxygenated compounds experimentally. The principles in Practice D 4626 should be applied when determining these response factors. The response of the flame ionization detector for oxygenated compounds is not directly (theoretically) related to mass concentration. A study has indicated that the FID response is linear for the conditions of this test method (see Figs. 10 and 11). Each individual apparatus must be calibrated using gravimetrically prepared standards, covering the sample concentration ranges expected and the scope of this test method. Standards used must comply with the requirements in Section 7. Figs. 10 and 11 present calibration data for six oxygenates as determined in a preliminary cooperative study report for calibration of this test method. Precision data will be prepared when more data becomes available.

14. Sample Analysis Procedure

14.1 Adjust the instrument operating variables to the values specified in Table 1 or as determined in Section 12.

14.2 Set the recorder or integration device, or both, for accurate presentation and collection of the data.

14.3 Inject an appropriate size sample (as determined in Section 10) into the injection port and start the analysis. Obtain a chromatogram and a peak integration report.

15. Calculation

15.1 Identify each peak by matching retention indices (or retention times) with those for known reference standards or sample components. If a computing integrator is used, examine the chromatographic data for proper peak integration. Examine the report to ensure peaks are properly identified.

15.1.1 Proper component identification using retention indices requires the use of *windows* surrounding each RI value in order to account for the analysis to analysis variations. The following windows have been found to provide satisfactory identification for this test method.

Indices	Window
100 – 300	±15
300 – 400	± 2.6
400 – 500	± 1.5
500 – 885	± 0.6
885 – 900	± 0.5
> 900	± 0.6

15.2 Obtain the area for each peak. Multiply each peak area by its appropriate response factor, taken from Table 2 or determined separately with standards, to obtain corrected peak areas. Use a response factor of 1.000 for unknown peaks.

15.3 If required, determine the concentration of water in the sample using Test Method D 1744, or an equivalent method. The total concentration of any other materials not determined by this test method should also be obtained.

15.4 The corrected peak areas are normalized to 100 % or to 100 % minus the concentrations determined in 15.3.

$$\text{component \% (m/m)} = \frac{\text{corrected peak area}}{\times (100 - \% \text{ undetected})/\text{total corrected peak area}} \quad (9)$$

16. Report

16.1 Report the concentration of each component as mass %, % (m/m), to the nearest 0.001 % (m/m).

16.2 These individual component data 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 calculation of other properties of petroleum liquids. See the caution in 5.3.

17. Precision and Bias ¹²

17.1 *Repeatability*—The difference in two test results obtained by the same operator with the same apparatus in a given laboratory under constant operating conditions on test samples

¹² Supporting data is available from ASTM International Headquarters in the form of a research report. Request RR: D02-1518.



Oxygenates Relative Response Factors

	Lab1	Lab 2	Lab 3	Lab4	Ave.	Std. Dev	%SD	Auto/Oil RRF
Methanol	3.0760	3.0477	2.9779	2.9230	3.0062	0.0691	2.30	3.0965
Ethanol	2.1888	2.0797	2.1755	2.0640	2.1270	0.0642	3.02	2.0953
t-Butanol	1.2975	1.3189	1.3312	1.2989	1.3116	0.0163	1.24	1.3368
MTBE	1.5279	1.5590	1.4860	1.5024	1.5188	0.0318	2.09	1.5016
ETBE	1.3848	1.3720	1.3804	1.3720	1.3773	0.0064	0.46	1.4032
TAME	1.3383	1.2993	1.3598	1.3340	1.3329	0.0250	1.88	1.3775

DHA Method Oxygenate Linearity Cooperative Study - peak area
Laboratory 4

Spl							Ave. Rf	RRF
MeOH	0.0100	1.0100	5.0500	10.0200	20.0100	29.8300		
	0.4037	34.7643	174.8862	340.9069	717.4781	1046.1427		
	0.3599	33.8017	179.9043	353.4087	717.1507	980.1566		
ave.	0.3818	34.2830	177.3953	347.1578	717.3144	1013.1496		
RF	0.0262	0.0295	0.0285	0.0289	0.0279	0.0294	0.0288	2.9230
EtOH	0.0100	1.0000	5.0000	10.1000	20.1500	30.1800		
	0.2883	50.5190	237.7223	496.9717	967.7888	1526.2755		
	0.4095	48.7438	242.3003	500.4514	1007.0434	1537.3776		
ave.	0.3489	48.6314	240.0113	498.2116	987.4161	1531.8265		
RF	0.0287	0.0206	0.0208	0.0203	0.0204	0.0197	0.0204	2.0640
TBA	0.0099	0.9640	4.9692	9.9583	19.8768	29.7953		
	1.0363	77.5423	408.5969	757.2307	1546.4197	2241.0530		
	1.1869	72.7672	392.8649	775.5192	1550.7498	2346.4085		
ave.	1.1116	75.1548	400.7309	766.3749	1548.5847	2293.7307		
RF	0.0089	0.0128	0.0124	0.0130	0.0128	0.0130	0.0128	1.2989
MTBE	0.0100	0.9992	5.0362	9.9724	20.0248	30.0471		
	0.7645	66.0865	345.4606	713.3773	1332.2069	2041.1591		
	0.5890	65.8994	325.8215	679.7792	1348.4042	2052.4822		
ave.	0.6767	65.9929	335.6411	696.5783	1340.3055	2048.8206		
RF	0.0148	0.0151	0.0150	0.0143	0.0149	0.0147	0.0148	1.5024
ETBE	0.0099	0.9851	4.9255	9.8707	19.6724	29.5727		
	0.4527	69.3251	374.3939	732.8740	1537.9746	2144.9023		
	0.6242	72.7316	374.7065	695.3345	1462.4055	2173.4412		
ave.	0.5384	71.0283	374.5502	714.1042	1500.1901	2159.1718		
RF	0.0183	0.0139	0.0132	0.0138	0.0131	0.0137	0.0135	1.3720
TAME	0.0100	0.9997	4.9788	9.8883	19.1530	29.7144		
	0.3702	75.3456	363.7452	762.9970	1488.8626	2346.1907		
	0.0072	75.1503	380.0280	763.6254	1420.3514	2230.3657		
ave.	0.1887	75.2480	371.8866	763.3112	1454.6070	2288.2782		
RF	0.0530	0.0133	0.0134	0.0130	0.0132	0.0130	0.0132	1.3340
C6	8.5050	8.4750	8.4400	8.4525	8.4525	8.6950		
	890.3467	843.5383	836.5459	803.1739	843.6532	847.7344		
	847.7681	854.2333	840.8679	834.8488	841.6083	802.3011		
ave.	869.0574	848.8858	838.7069	819.0113	842.6307	825.0177		
RF	0.0098	0.0100	0.0101	0.0103	0.0100	0.0105	0.0101	1.0262
C7	8.5050	8.4750	8.4400	8.4525	8.4525	8.6950		
	893.5123	847.7426	868.0640	834.6944	890.9965	869.6032		
	846.4708	858.0901	862.0443	871.7571	882.1653	834.0419		
ave.	869.9918	852.9164	865.0541	853.2258	881.5809	851.8225		
RF	0.0098	0.0099	0.0098	0.0099	0.0096	0.0102	0.0099	1.0000
C8	8.5050	8.4750	8.4400	8.4525	8.4525	8.6950		
	889.7205	846.6591	877.1065	838.8929	890.2631	873.8851		
	839.2188	855.2006	862.7846	884.2601	895.4804	854.5747		
ave.	864.4697	850.9298	869.9455	861.5765	892.8718	864.2299		
RF	0.0098	0.0100	0.0097	0.0098	0.0095	0.0101	0.0098	0.9944
C9	8.5050	8.4750	8.4400	8.4525	8.4525	8.6950		
	883.5337	843.1968	870.7139	832.1808	883.3178	868.7531		
	829.0626	849.0969	854.1742	881.9661	889.9074	860.0512		
ave.	856.2982	846.1469	862.4440	857.0734	886.6126	864.4021		
RF	0.0099	0.0100	0.0098	0.0099	0.0095	0.0101	0.0099	1.0003

FIG. 10 Determination of Oxygenate Response - DHA Speciation Analysis

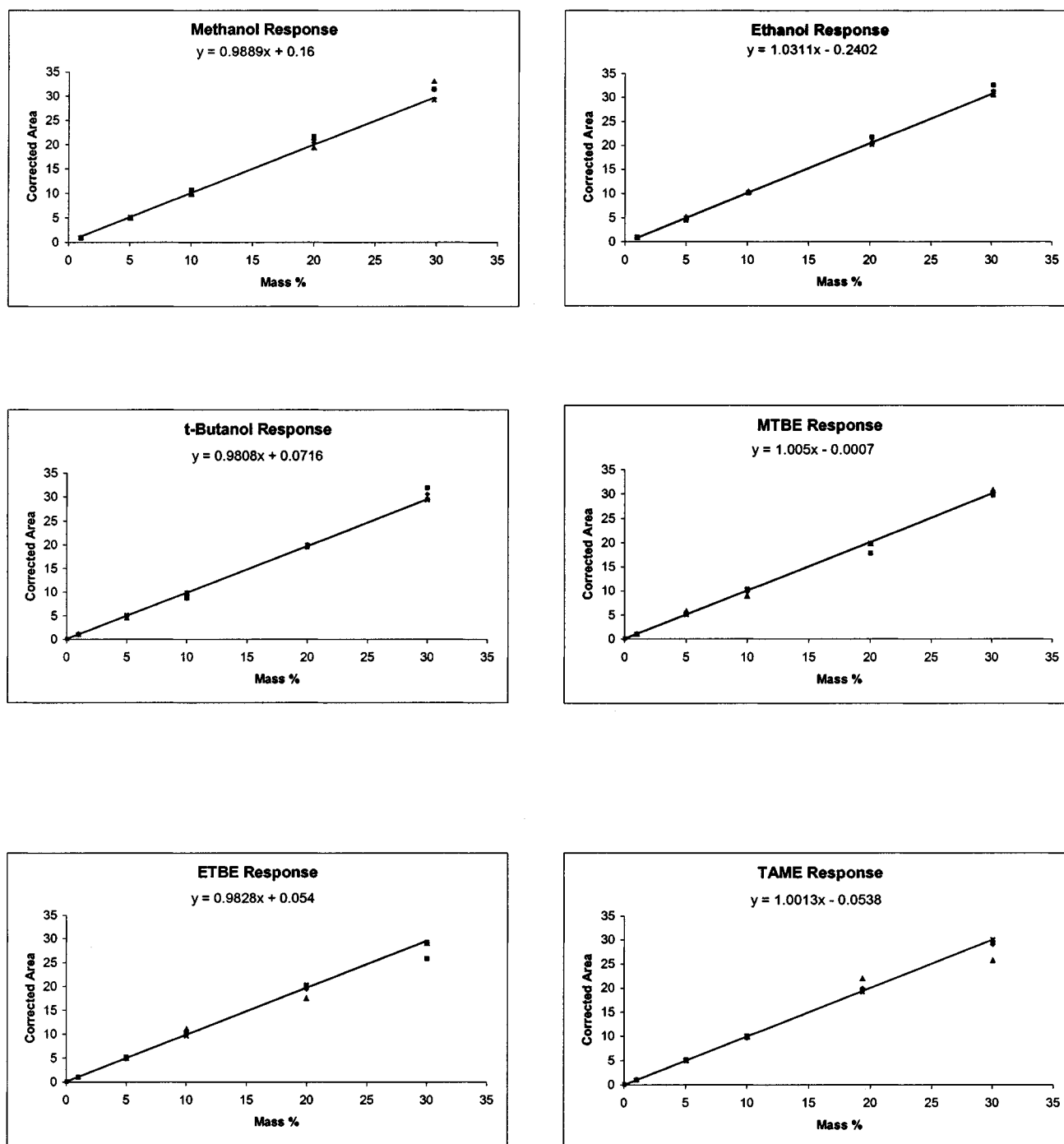


FIG. 11 Graphical Representation Determination of Oxygenate Response - DHA Speciation Analysis

taken from the same laboratory sample should, in the long run, in the normal and correct operation of the test method not exceed the values given in Table 4 and Table A1.3 for the gasoline components.

17.2 Reproducibility—The difference between two single and independent measurements on test samples taken from the same bulk sample should, in the long run, in the normal and correct operation of the test method, not exceed the values given in Table 4 and Table A1.3 for the gasoline components.

17.3 Bias—No information can be presented on the bias of the procedure in this test method for measuring hydrocarbon concentrations because no material having an accepted reference value is available.

18. Keywords

18.1 detailed hydrocarbon analysis; DHA; gas chromatography; hydrocarbons; open tubular column; oxygenates; PIONA; PONA

TABLE 4 Repeatability and Reproducibility of DHA Determinations

NOTE 1—The following is a partial list of precision data that has been prepared by statisticians of CS94 in accordance with RR: D2-1007, and represents their best estimate of the cooperative study data. The complete precision data set appears in Annex A1., Table A1.3.

NOTE 2—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. The (repeatability standard deviation)/mean value for each analyte/sample combination must be less than or equal to 0.1, as per LOQ requirements which, while not a standard, is what CS94 is recommending.

NOTE 3—

Legend:

$r_{\min}$	= 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} , $R_{\max}$	= 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.

Component	Average RI	$r_{\min}$	r_{est}	$r_{\max}$	$R_{\min}$	R_{est}	$R_{\max}$	$C_{\min}$	$C_{\max}$
n-butane	400.00	6.8	9.9	13.9	15.3	32.4	59.1	1.02	3.75
i-pentane	477.45	5.9	7.2	8.7	8.5	14.8	23.8	2.48	13.38
Pentene-1	490.83	5.2	7.5	10.5	9.7	13.8	19	0.06	0.43
n-pentane	500.00	5.2	6.5	8.1	7.1	10.4	14.8	1.06	3.49
Cyclopentane	566.84	3.8	4.9	6.2	7	10.1	14	0.07	0.59
2,3-dimethylbutane	569.24	2.9	3.2	3.5	5.1	8.5	13.1	0.7	1.91
n-hexane	600.00	2	2.4	2.9	3.6	5.1	6.9	0.33	2.52
Methylcyclopentane	625.86	2.2	2.6	3.1	4.5	6.4	8.7	0.37	2.35
1-methylcyclopentene	648.71	1.9	2.7	3.7	7.9	8.7	9.6	0.17	0.82
Benzene	649.92	2.6	3.6	4.8	5.5	9	13.7	0.17	1.58
Cyclohexane	657.81	2.7	3.7	4.9	8.2	14.8	24.3	0.07	0.9
2-methylhexane	667.61	1.6	2.2	2.9	5.1	6.1	7.2	0.39	1.09
2,2,4-trimethylpentane	688.48	2.4	3.2	4.1	7.4	11.4	16.7	0.1	11.26
n-heptane	700.00	2.5	3.4	4.5	7.7	10.8	14.7	0.21	1.06
Methylcyclohexane	717.89	2.8	3.4	4	4.1	5.9	8.2	0.11	1.2
2,3,4-trimethylpentane	746.83	2.3	3.8	6	5.8	7.8	10.3	0.08	4.26
Toluene	751.77	1.9	2.7	3.8	10.8	13.5	16.5	1.99	10.34
2-methylheptane	764.14	3.5	4.9	6.6	4.8	6.1	7.5	0.15	0.63
n-octane	800.00	2.2	3.6	5.5	6.5	15.7	30.9	0.14	0.75
Ethylbenzene	854.65	2.2	3.2	4.4	7.2	10.6	14.9	0.62	2.62
1,3-dimethylbenzene	864.22	2.6	3.3	4.2	9.7	12.5	15.7	1.55	6.66
3-methyloctane	880.24	5.1	8.5	13	8.7	15.5	24.9	0.07	0.29
n-nonane	900.20	3.9	6.4	9.8	8.6	10.3	12.2	0.06	0.34
n-propylbenzene	946.33	2.8	5	8.1	7.6	11.9	17.7	0.21	0.77
1,4-methylethylbenzene	956.22	3.5	5.3	7.7	5.1	7.7	11.1	0.32	1.19
1,3,5-trimethylbenzene	961.92	3.7	5.5	7.7	5.4	8.3	12.1	0.39	1.21
2-methylnonane	971.77	6.5	10.6	16.2	17.5	25.9	36.6	0.03	0.19
1,2,4-trimethylbenzene	983.40	4.2	5.7	7.5	7.8	10.6	13.9	1.19	4.32
n-decane	1000.20	7.5	9.2	11.1	12.1	17.9	25.3	0.03	0.25
1,2,3-trimethylbenzene	1006.88	3.8	5.8	8.5	7.2	8.5	10	0.28	0.96
n-undecane	1100.00	8.6	13.9	21	24.4	40	61.2	0.03	0.18
1,2,3,5-tetramethylbenzene	1108.79	6.4	7.8	9.3	10.2	13.9	18.3	0.21	0.51
Naphthalene	1168.01	6.1	8.5	11.3	12.9	16.9	21.5	0.13	0.4
n-dodecane	1200.00	12.2	16.7	22.1	20.2	32.9	50	0.01	0.11
2-methylnaphthalene	1282.57	7.6	11.1	15.4	17.5	22.3	28	0.05	0.5



(Mandatory Information)

A1. PROCEDURE FOR ADJUSTING THE SELECTIVITY OF A DHA METHYL SILICONE OPEN TUBULAR COLUMN

A1.1 The successful application of this test method is highly dependent upon the selectivity of the column used. New 100 m $\times$ 0.25 mm 0.5 μ m methyl silicone open tubular fused silica columns will likely not have sufficient selectivity for aromatics to function properly. Critical to the successful analysis of *reformulated* and oxygenated spark engine motor fuels is column inertness and component selectivity. Inertness of the primary 100-m column affects the retention and adsorption of the oxygenates such as alcohols and ethers, while selectivity for the aromatic compounds is controlled by the liquid phase. Until adequate commercial columns are available, it will be necessary to slightly increase the column selectivity, which is accomplished by the addition of a short precolumn containing a moderately selective liquid phase.

A1.2 Prior to making any precolumn additions to the 100-m methyl silicone capillary column, determine that the main column meets the column specifications outlined in 6.4.1 and determined in Section 9. Section 9 describes the preliminary evaluation of the 100-m methyl silicone capillary column, using a 35°C isothermal analysis to determine the basic column characteristics of efficiency, retention factor, inertness, and selectivity. Figs. A1.1-A1.3 provide examples of the column quality specification determinations. These determinations may also be made with a precolumn attached, since the precolumn has little if any affect on the results. Fig. A1.4 illustrates that the addition of different lengths of precolumn has negligible influence on the retention characteristics of oxygenated compounds. Poor peak shape and resolution of these oxygenates cannot be corrected by the addition of the precolumn. Tailing peaks may also be the result of an active injector liner or packing material, or both, in the injector liner. An increase in retention of the oxygenates is likely due to column activity. The relative position of the oxygenates to the hydrocarbons is dependent upon column temperature, thus a faulty column oven temperature control could also result in shifted peaks.

A1.3 When necessary, a precolumn is added to the primary 100-m column to adjust the column selectivity for aromatic compounds. Precolumns that have been used successfully are variable lengths of 0.25 mm internal diameter fused silica open tubular column containing a 1.0 μ m film thickness of 5 % phenyl methyl silicone. The film thickness is likely not critical, only the total amount of phase. Lengths ranging from 1 to more than 3 m have been necessary to provide sufficient selectivity, depending on the initial selectivity of the methyl silicone column used. One metre of 1.0 μ m precolumn is equivalent to a 100-m column with 0.5 μ m of 0.1 % phenyl methyl silicone liquid phase.

A1.4 Figs. A1.5-A1.8 illustrate the resolution of the methylcyclopentene-1 and benzene pair with a new column and one, two, and three metres of precolumn. The key segment of the chromatogram is expanded to better illustrate the resolution of this component pair.

A1.5 The preliminary evaluation of the 100-m column will provide the user with information regarding the initial length of precolumn with which to start the tuning process. Dependent upon the methylcyclopentene-1 and benzene resolution, an initial precolumn of between 1 and 4 m is selected; which ever provides a resolution greater than 1.2.

A1.6 The final tuning will consist of reducing the precolumn length, probably in increments of 0.25 m, until the proper resolution is achieved between 2,3,3-trimethylpentane and toluene, and 1,4-dimethylbenzene and 2,3-dimethylheptane; using the actual analysis temperature conditions.

A1.7 Fig. A1.9 illustrates graphically the effect of different lengths of precolumn, attached to the same 100-m column. The key component separations are shown. These analyses were made using the conditions given in Table 2. In this case, the use

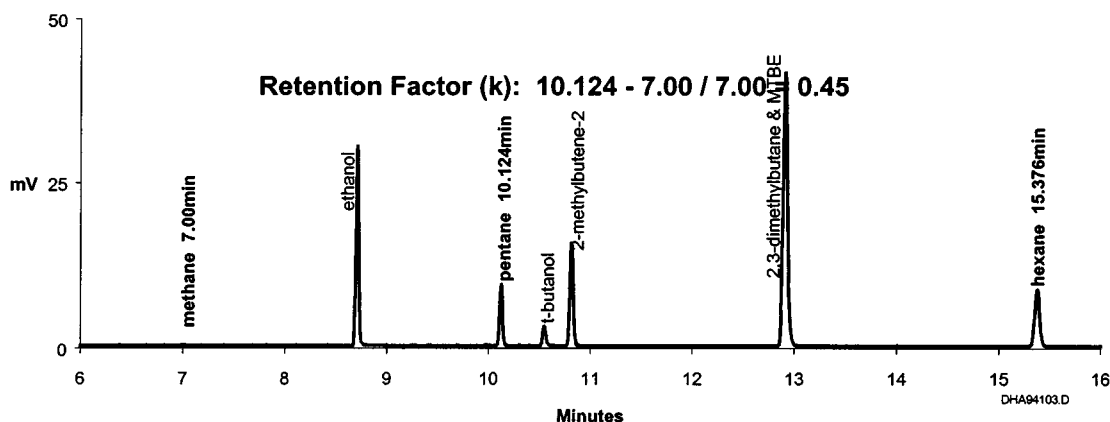


FIG. A1.1 Column Retention Factor Calculation (9.1)

Efficiency (n): $5.545 (10.124 / 0.032)^2 = 555,016$
 Resolution (R): $2(10.817 - 10.547) / 1.699(0.034 + 0.034) = 4.673$

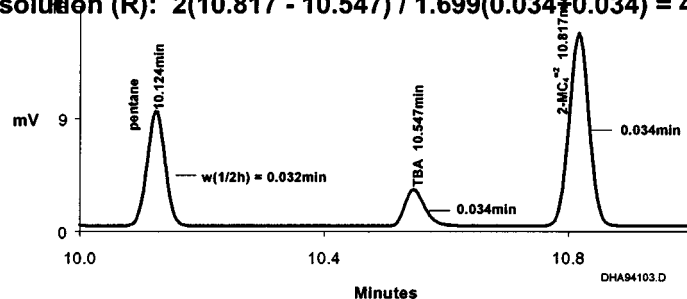


FIG. A1.2 Column Efficiency and Resolution Calculations (9.2 and 9.3)

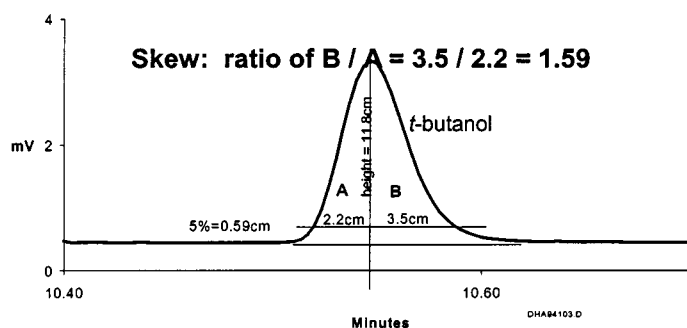


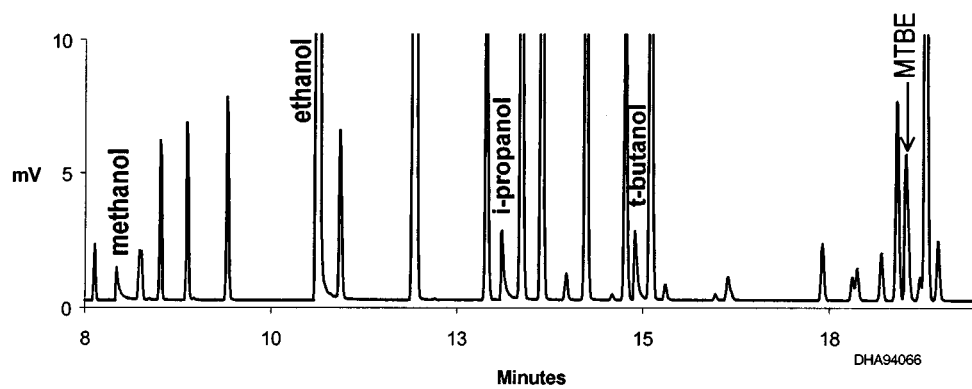
FIG. A1.3 Column Inertness - Peak Skewness Calculation (11.5)

of the 1.25-m precolumn provides the best compromise for the three key separations.

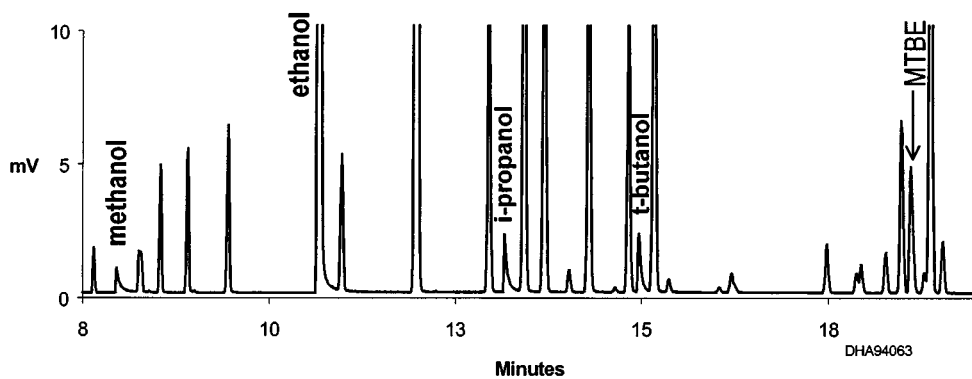
A1.8 Fig. A1.10 illustrates the use of different lengths of precolumn to achieve the specified selectivity for three differ-

ent 100-m columns. The final precolumn length will provide adequate resolution of all three of the key separations.

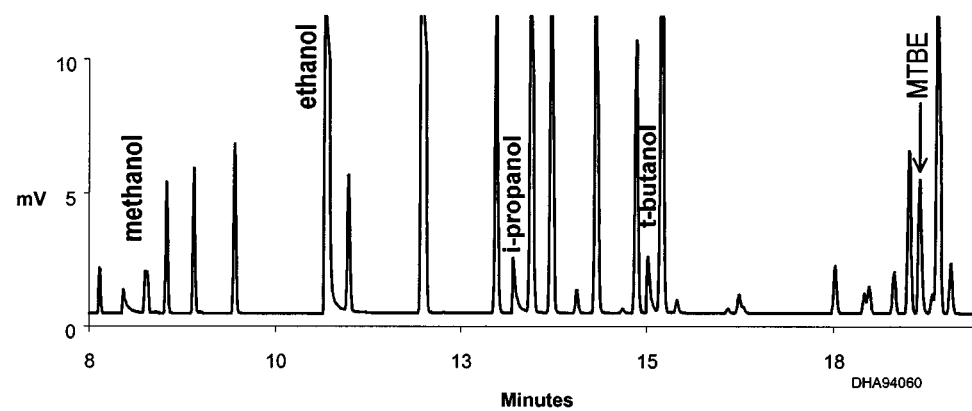
A1.9 Figs. A1.11-A1.17 illustrate DHA analyses.



0.5 μm_{df} methylsilicone column + 1.25m 1 μm_{df} precolumn



0.5 μm_{df} methylsilicone column + 1.5m 1 μm_{df} precolumn



0.5 μm_{df} methylsilicone column + 2.0m 1 μm_{df} precolumn

FIG. A1.4 Oxygenates Separations - Effect of Different Precolumn Lengths

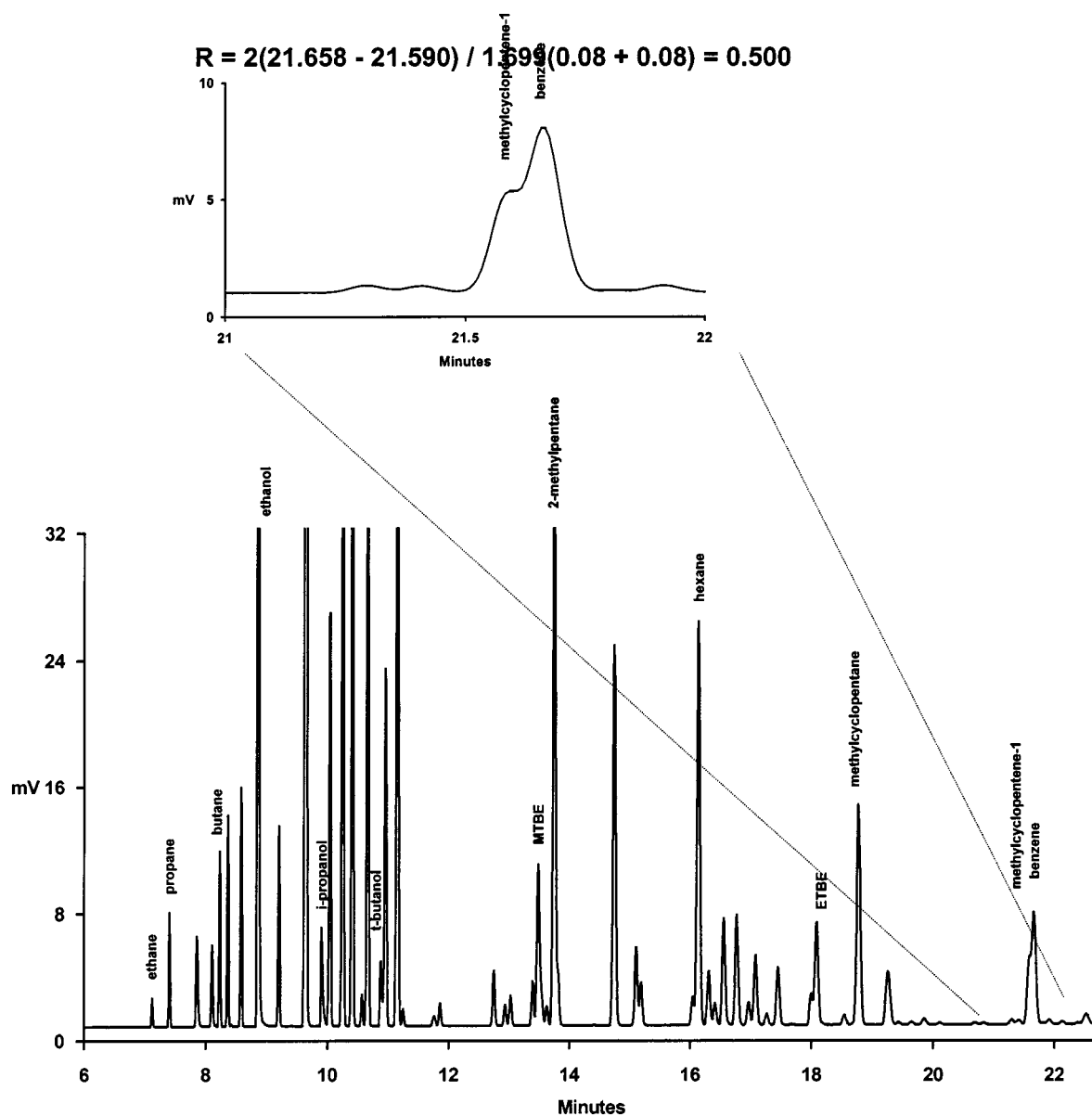


FIG. A1.5 PONA-V Standard—Analysis through Benzene New DHA Column—Analyzed at 35°C Isothermal (Conditions in Accordance with Table 2)

$$R = 2(21.025 - 20.925) / 1.699(0.076 + 0.081) = 0.750$$

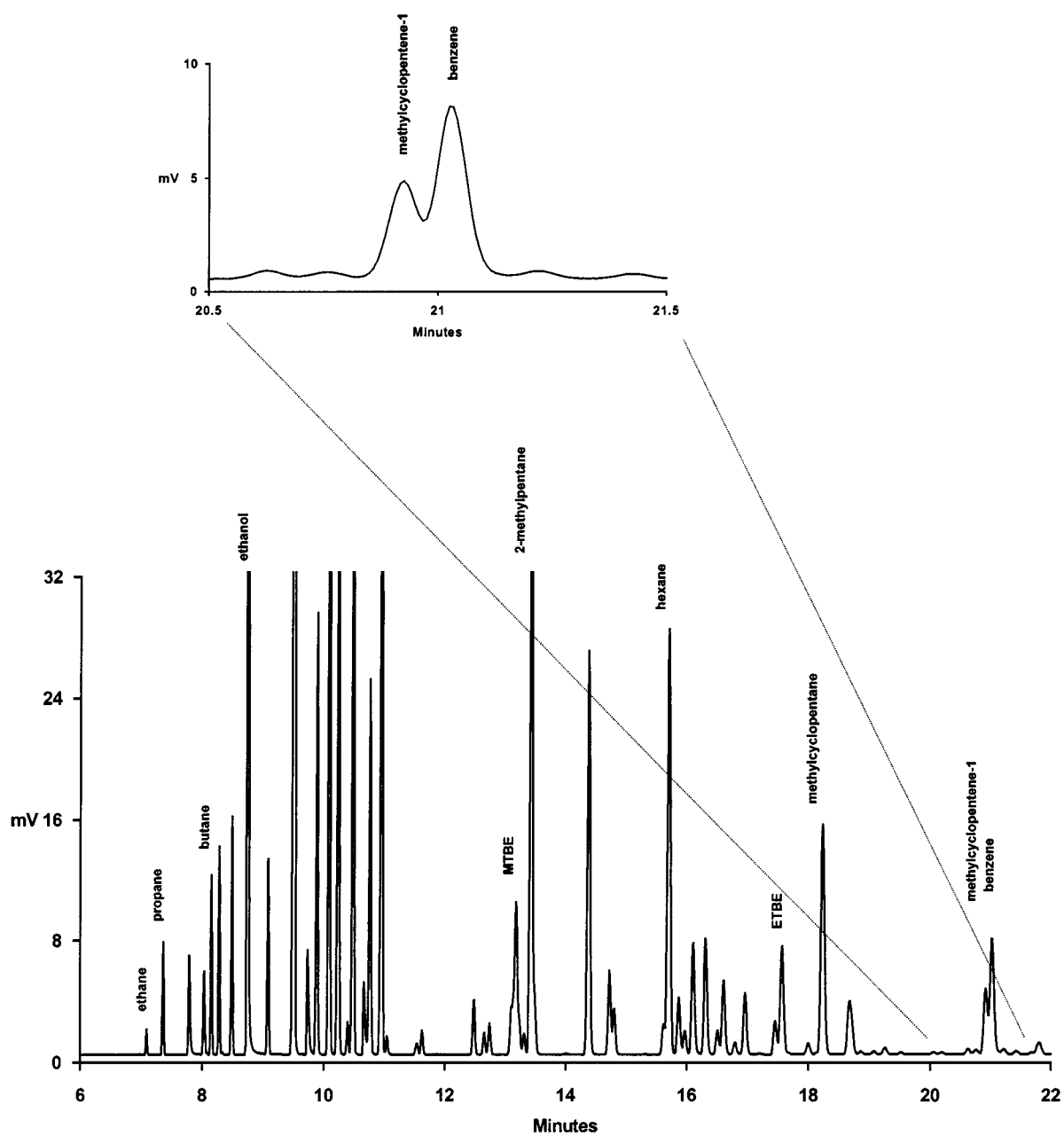


FIG. A1.6 PONA-V Standard - Analysis through Benzene New DHA Column plus 1 m × 0.25 mm 1 μm Precolumn Analyzed at 35 °C Isothermal

$$R = 2(21.258 - 21.125) / 1.699(0.079 + 0.081) = 0.978$$

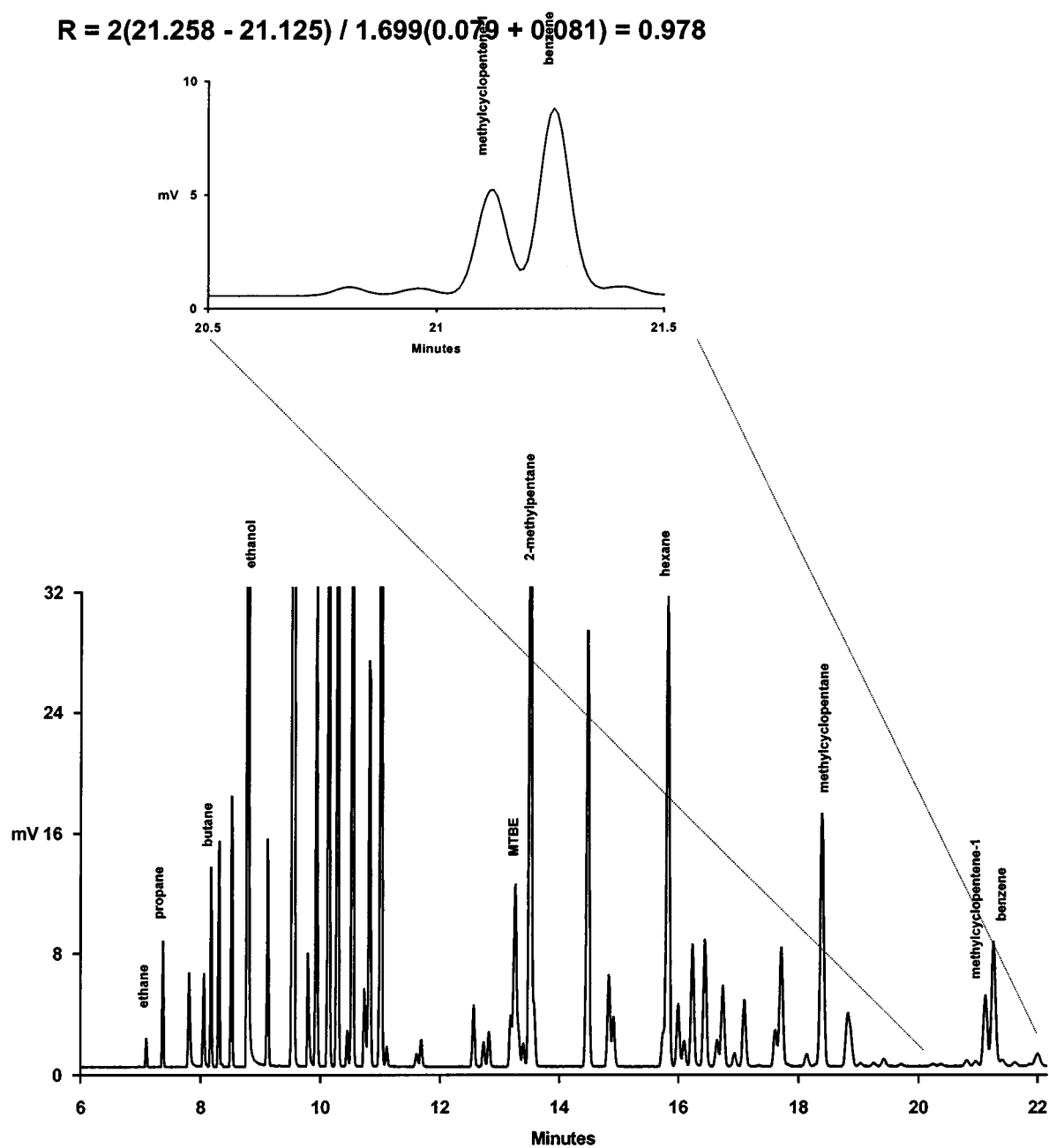


FIG. A1.7 PONA-V Standard - Analysis through Benzene New DHA Column plus 2 m × 0.25 mm 1 μm Precolumn Analyzed at 35 °C Isothermal



$$R = 2(22.642 - 22.458) / 1.699(0.089 + 0.093) = 1.190$$

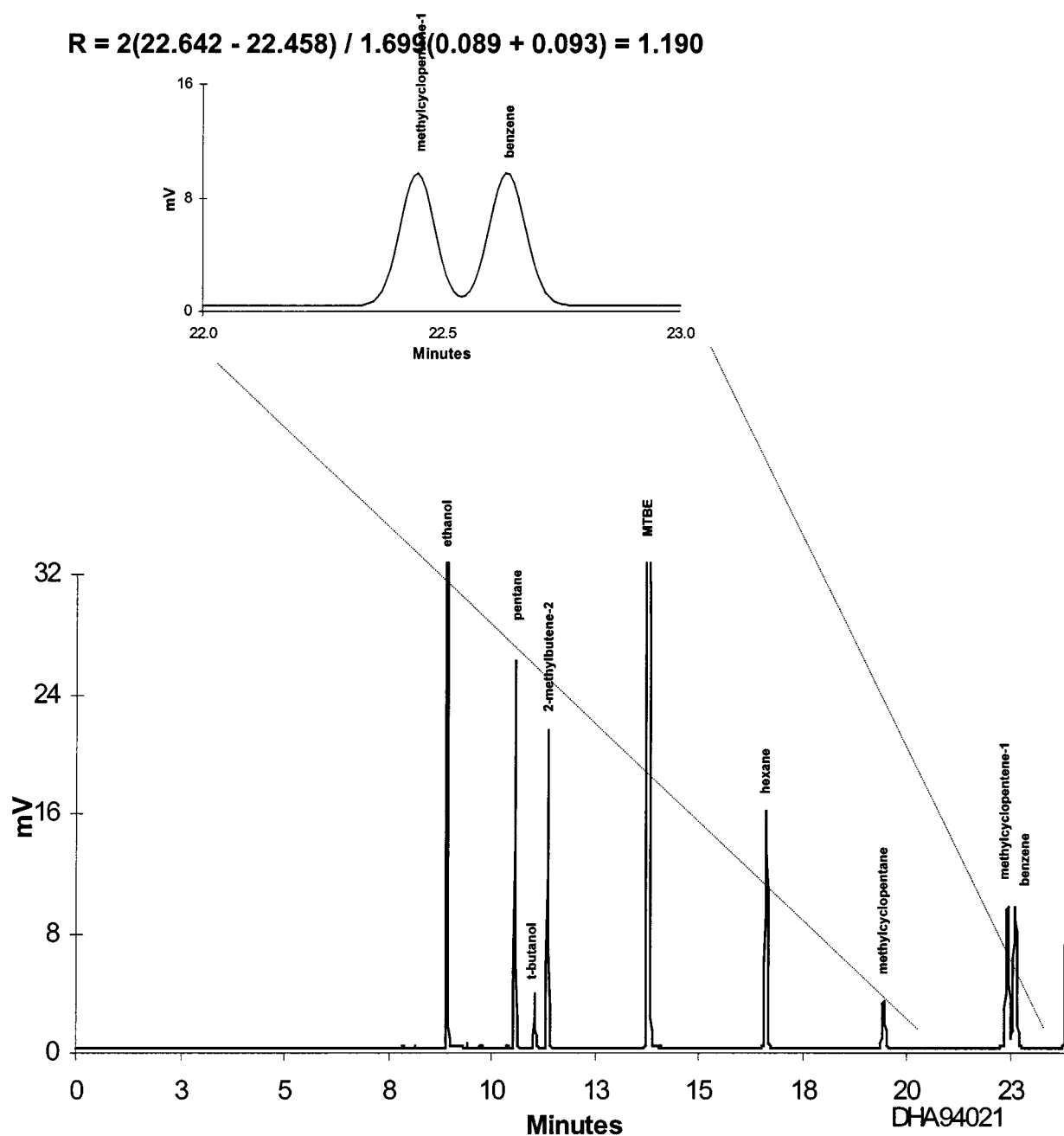


FIG. A1.8 DHA Calibration Standard - Analysis through Benzene New DHA Column plus 3 m × 0.25 mm 1 μm Precolumn Analyzed at 35 °C Isothermal

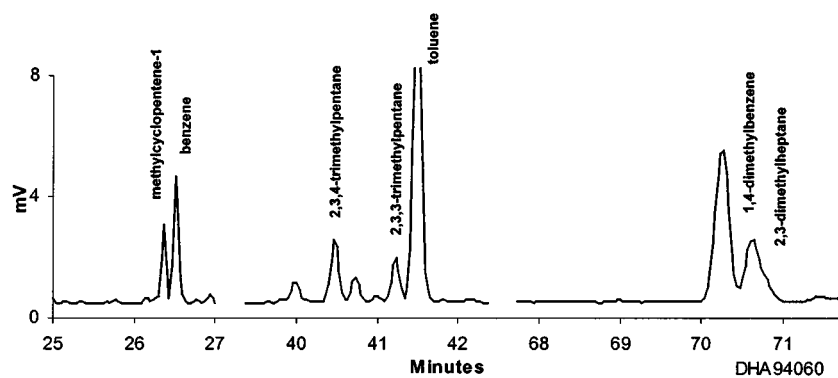
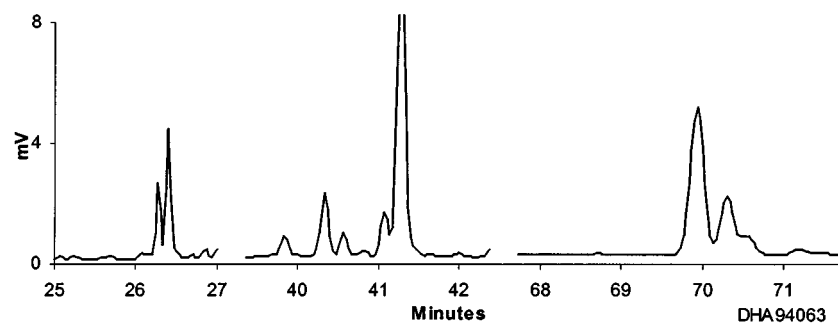
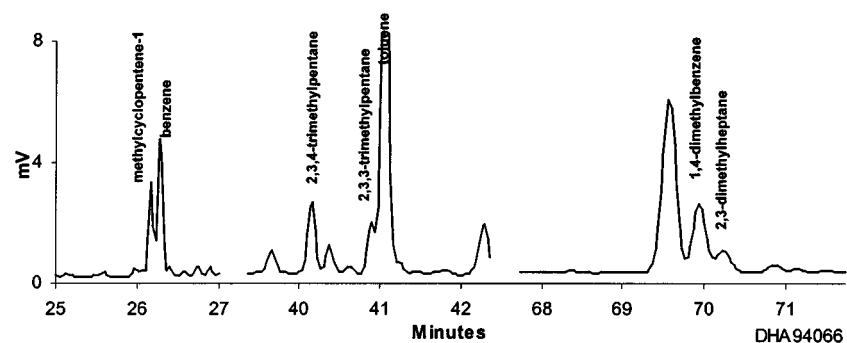
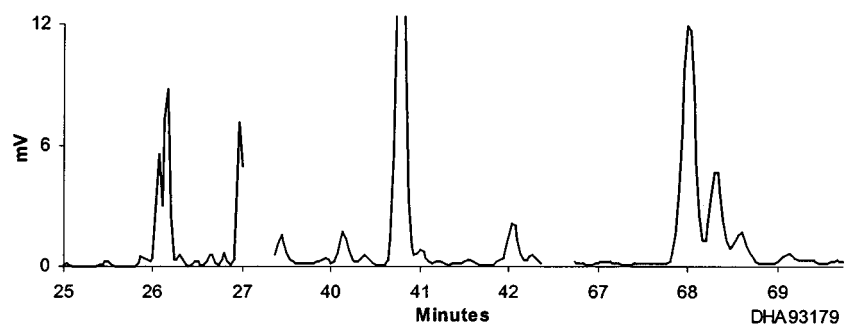
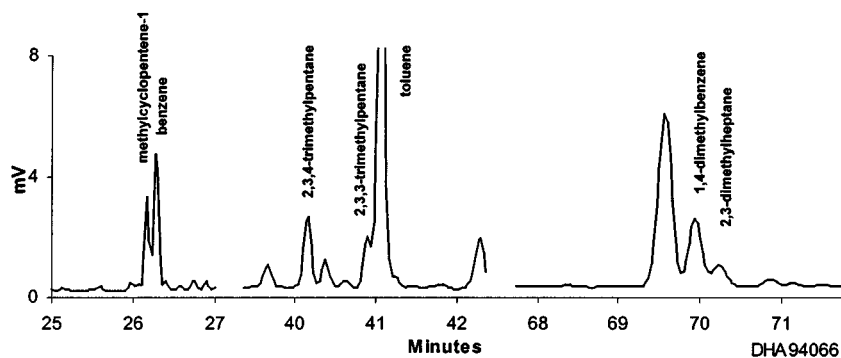
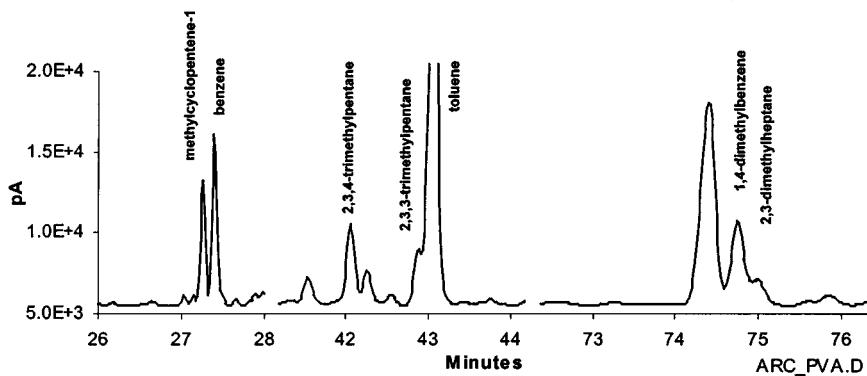


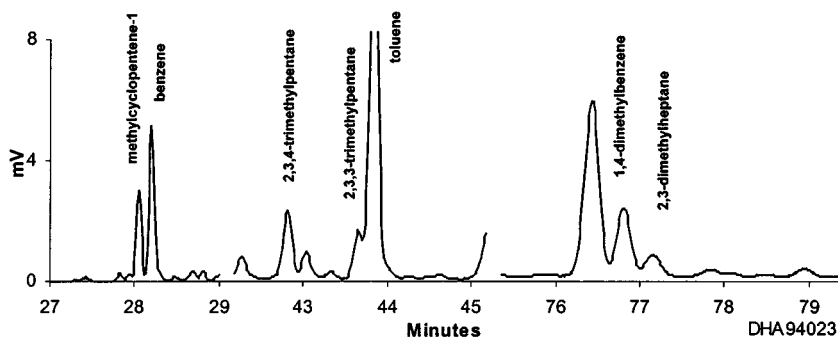
FIG. A1.9 Key Separations - Effect of Different Precolumn Lengths Same Primary Column. Conditions in Accordance with Table 2 Top to Bottom - 1.00 m, 1.25 m, 1.50 m, and 2.00 m Precolumn



(Old) 0.5 μ m_{df} methylsilicone column + 1.25m 1 μ m_{df} precolumn

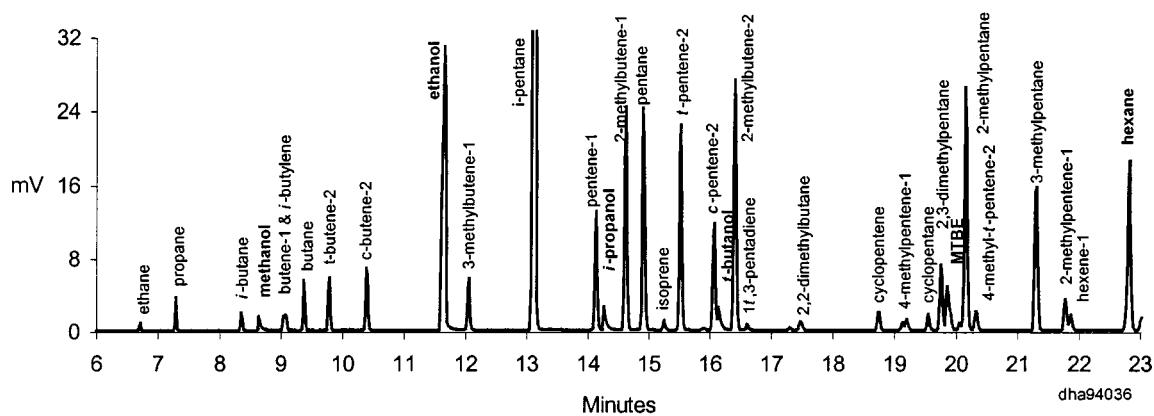


0.5 μ m_{df} methylsilicone column + 2.4m 1 μ m_{df} precolumn

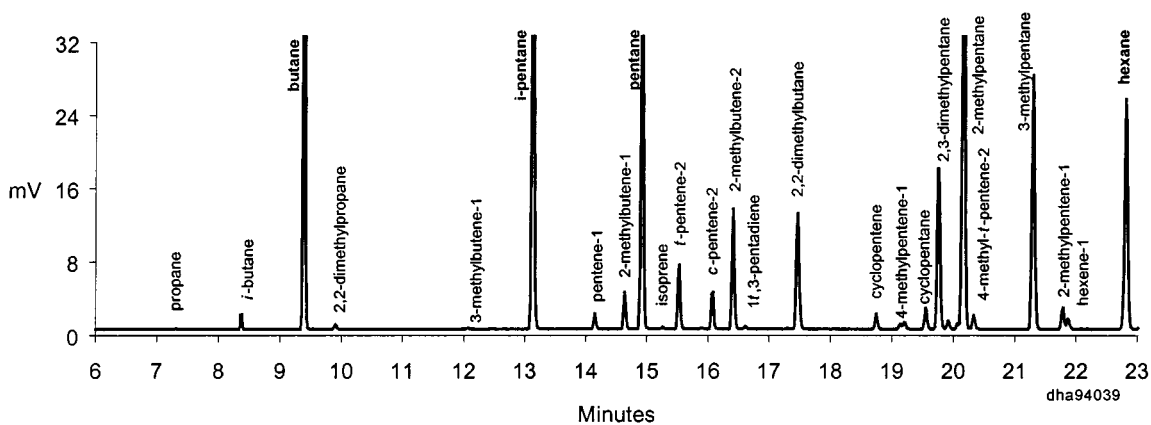


100m x 0.25mm HP 0.5 μ m_{df} methylsilicone column + 3.0m 1 μ m_{df} precolumn

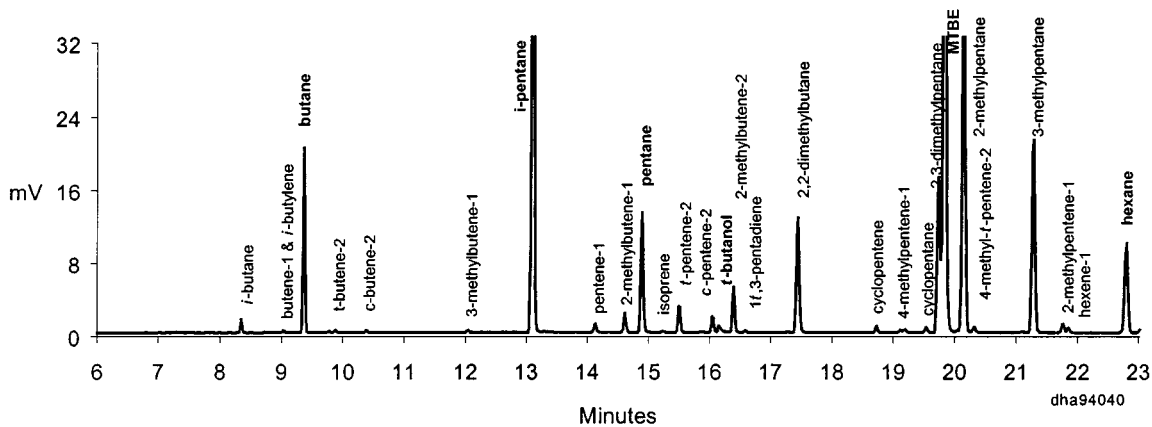
FIG. A1.10 Key Separations - Tuning of Different Columns Conditions in Accordance with Table 2



PONA-Va Mixture



RFA Gasoline

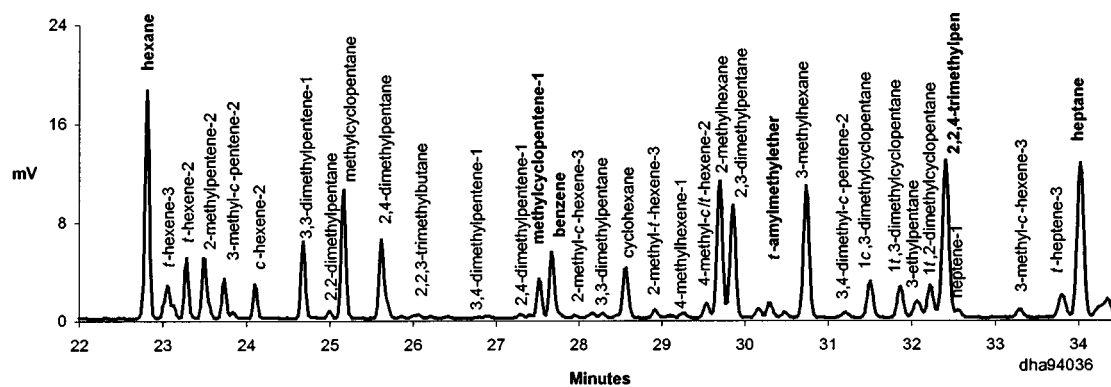


CCF Gasoline

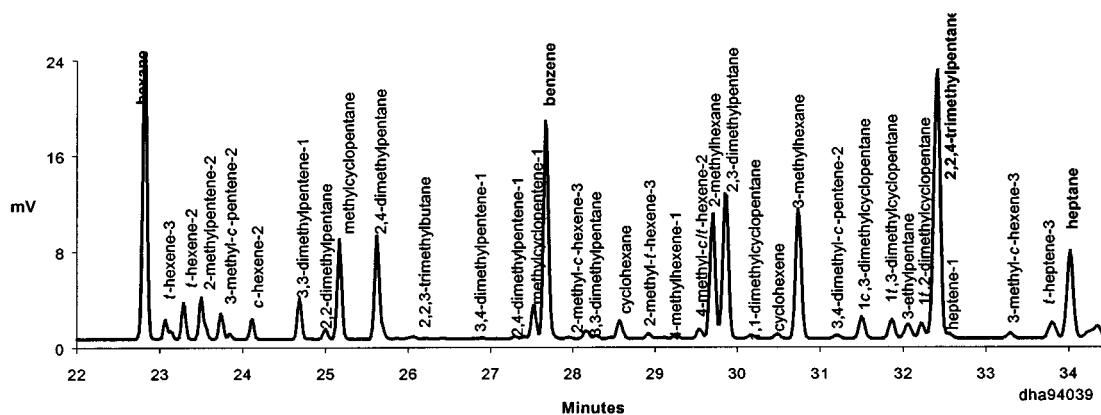
FIG. A1.11 DHA Analyses - Methane through Hexane



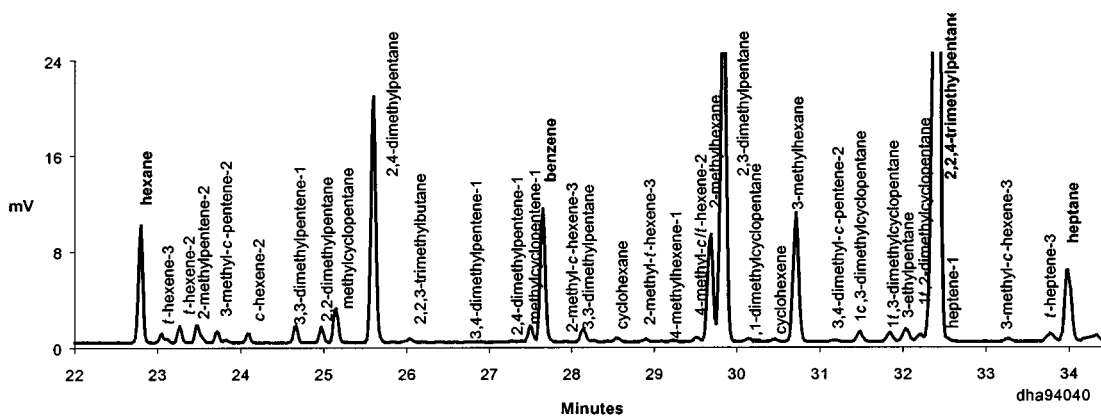
D 6730 - 01



PONA-Va Mixture



RFA Gasoline

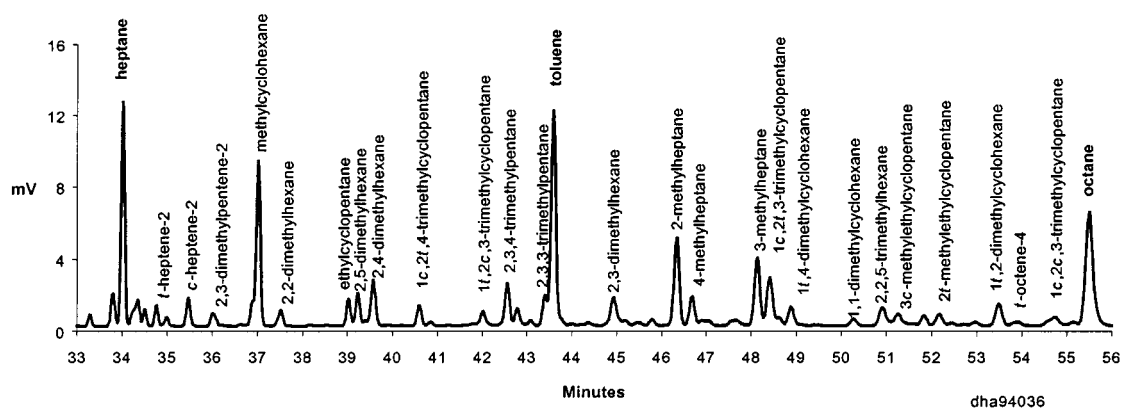


CCF Gasoline

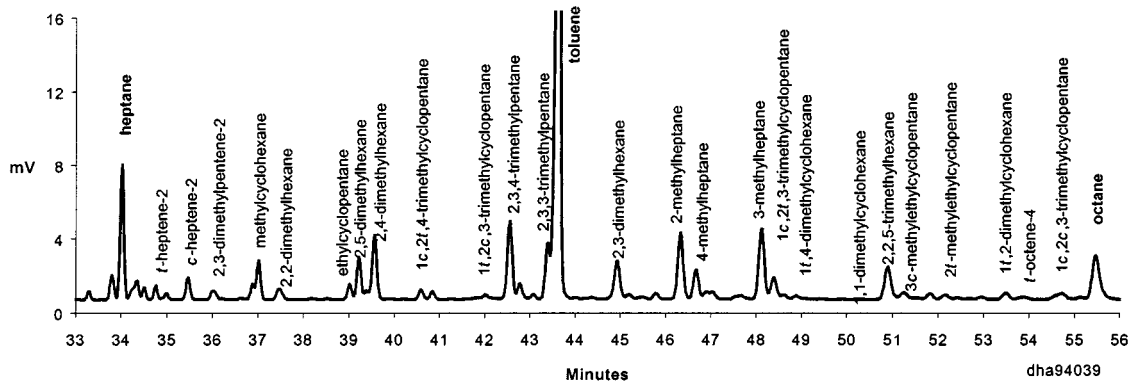
FIG. A1.12 DHA Analyses - Hexane through Heptane



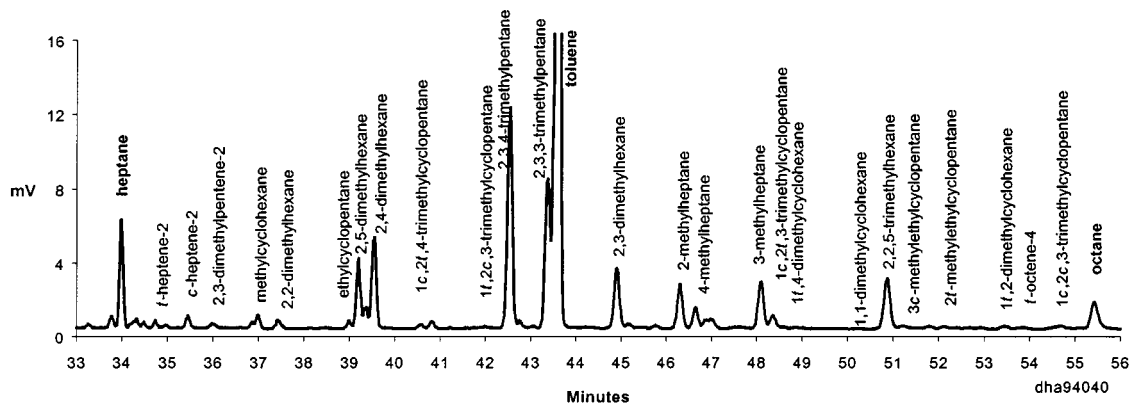
D 6730 - 01



PONA-Va Mixture



RFA Gasoline

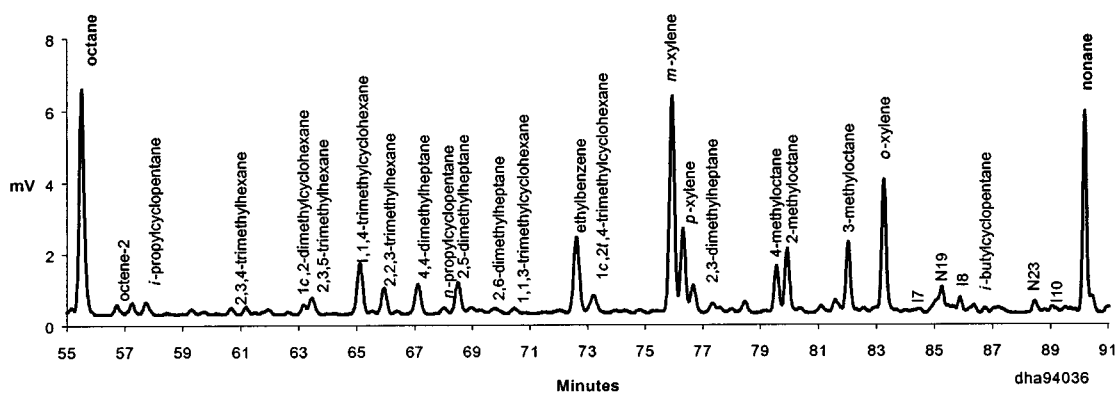


CCF Gasoline

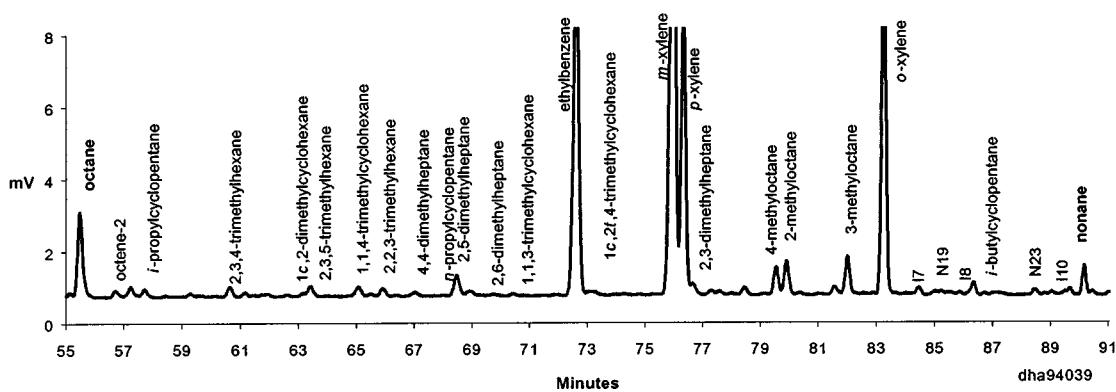
FIG. A1.13 DHA Analyses - Heptane through Octane



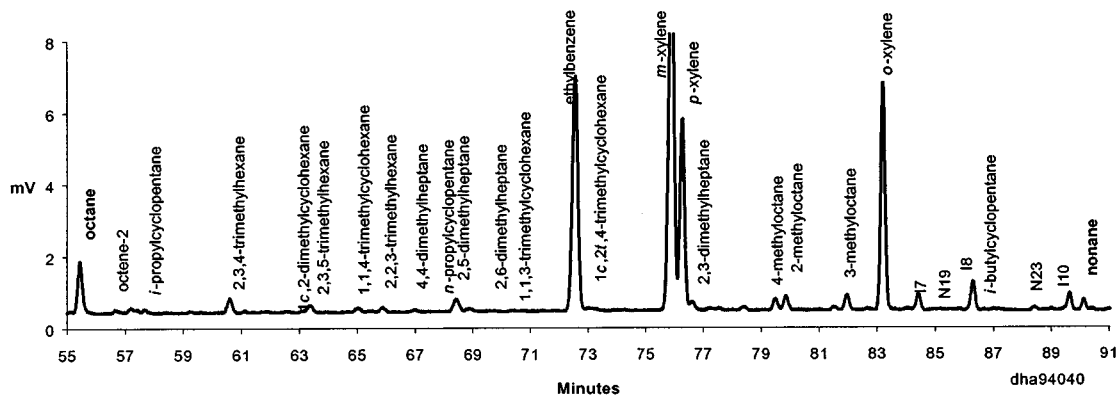
D 6730 - 01



PONA-Va Mixture

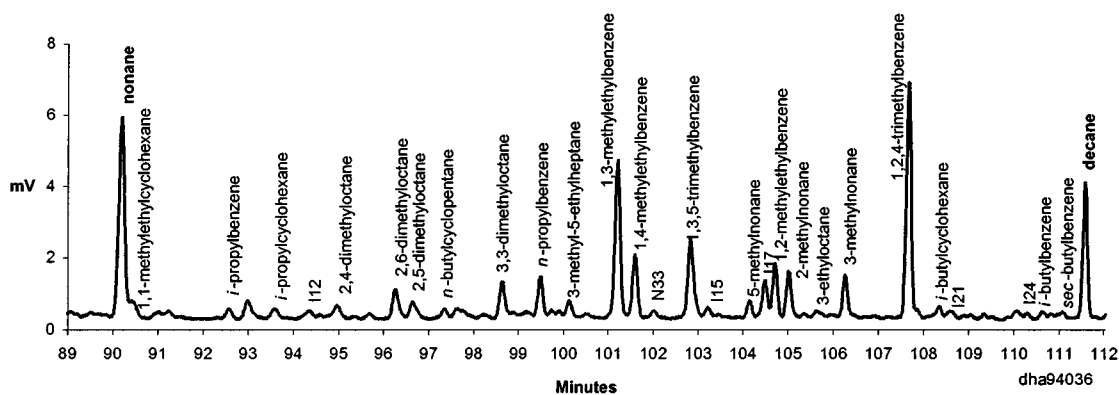


RFA Gasoline

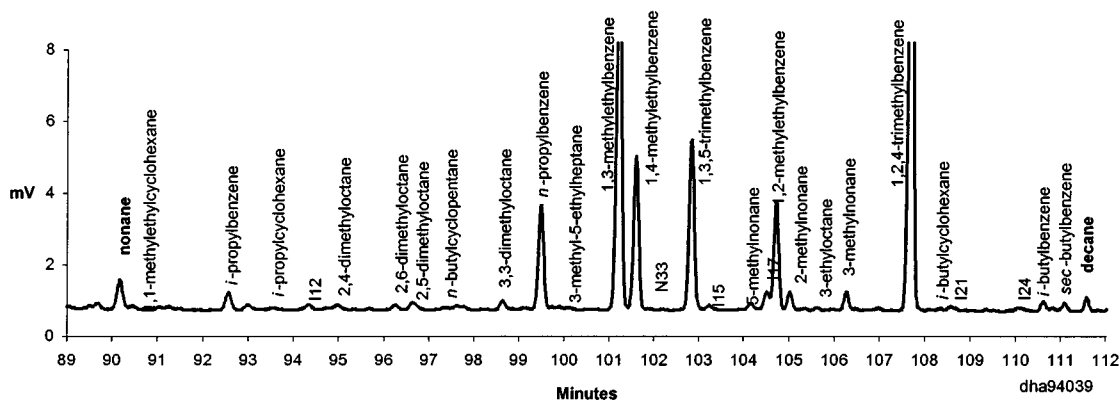


CCF Gasoline

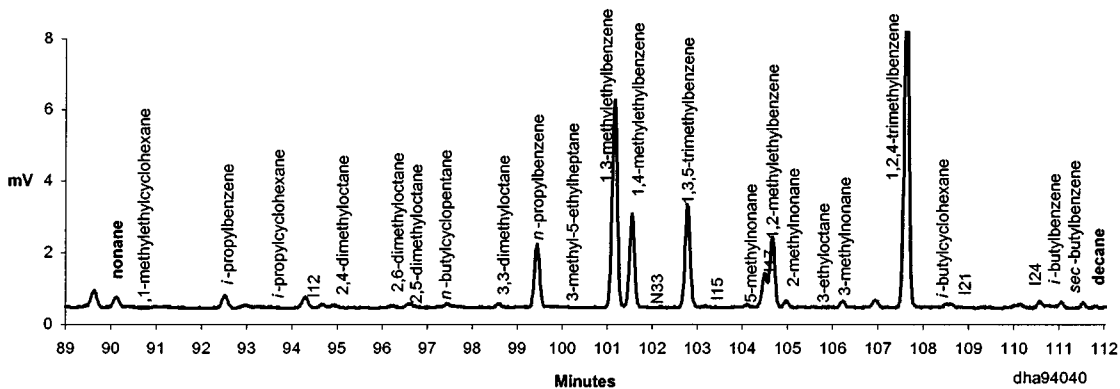
FIG. A1.14 DHA Analyses - Octane through Nonane



PONA-Va Mixture



RFA Gasoline

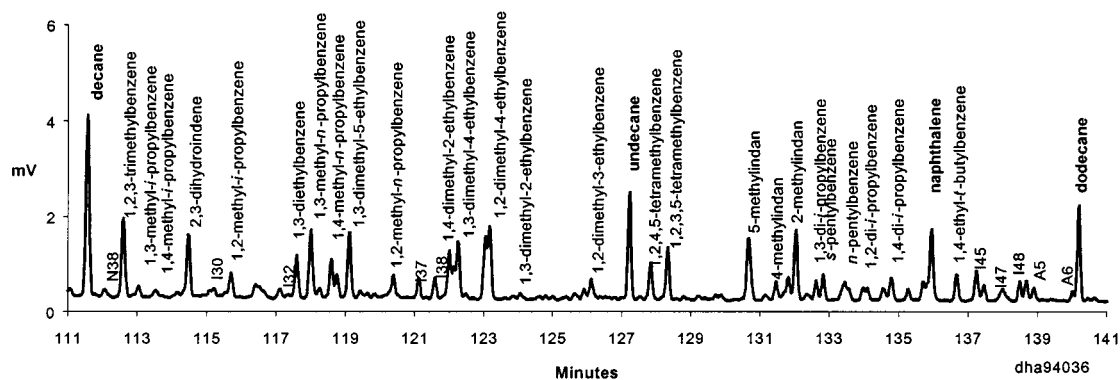


CCF Gasoline

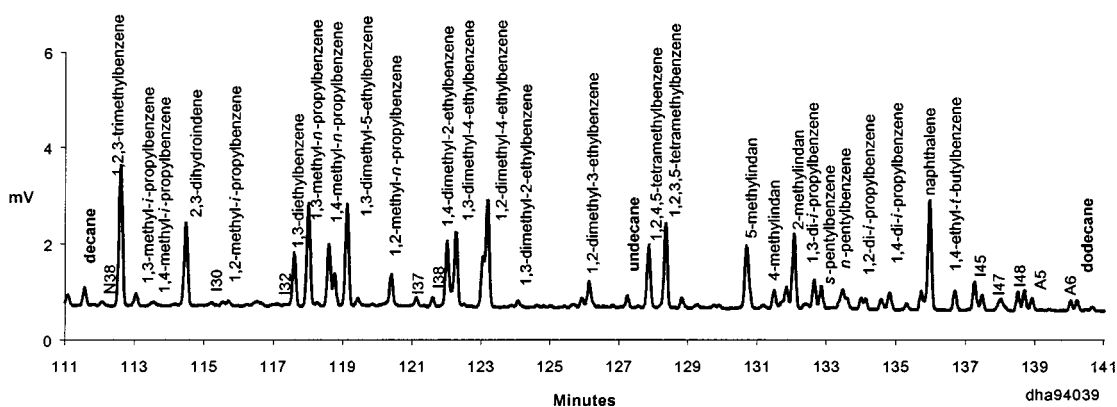
FIG. A1.15 DHA Analyses - Nonane through Decane



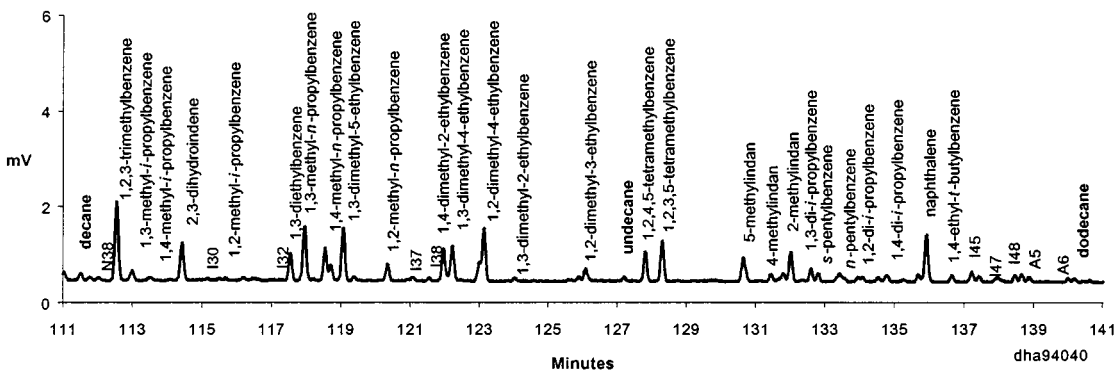
D 6730 - 01



PONA-Va Mixture

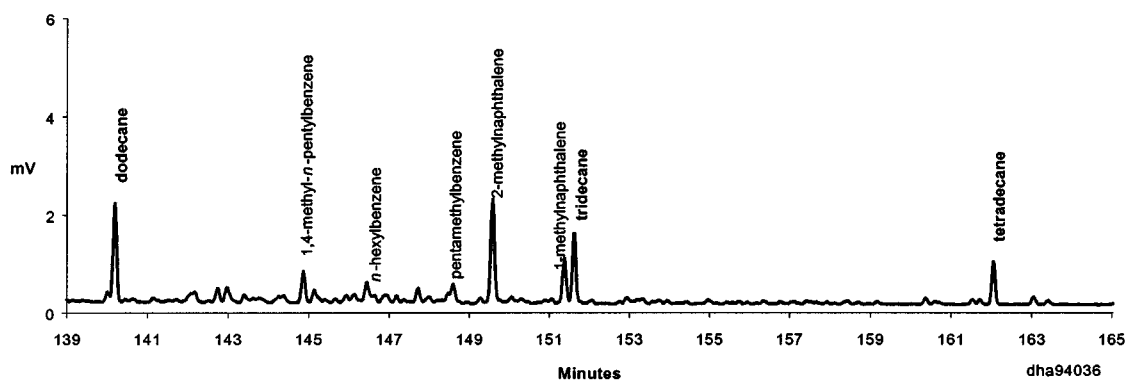


RFA Gasoline

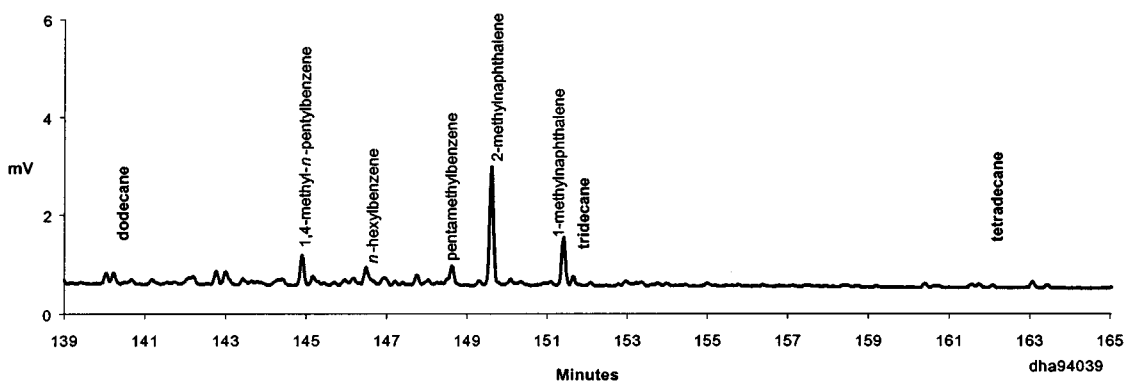


CCF Gasoline

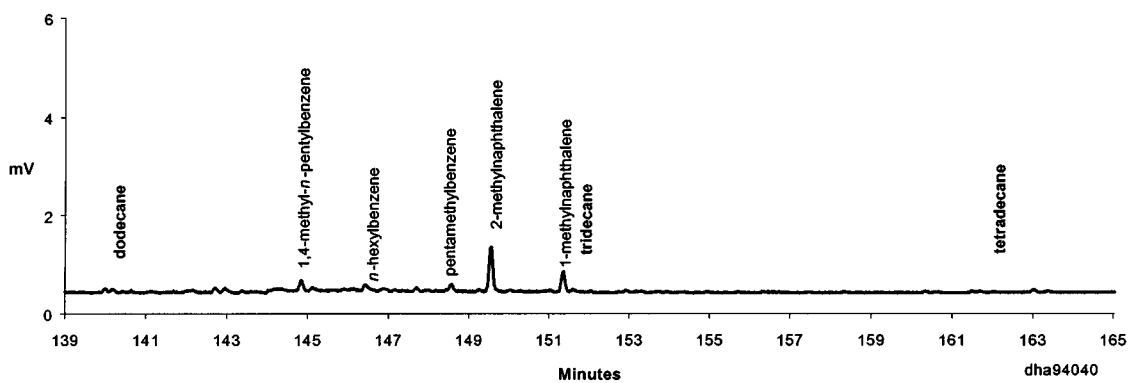
FIG. A1.16 DHA Analyses - Decane through Dodecane



PONA-Va Mixture



RFA Gasoline



CCF Gasoline

FIG. A1.17 DHA Analyses - Dodecane through Tetradecane

TABLE A1.1 DHA Component Data

NOTE 1—These data consist of the current physical constants used in the cooperative study. The average RI are those accumulated in a ruggedness test of the *tuning* process. The RFA and CCF gasoline data are the averages determined in the cooperative study. RFA is an industry average regular gasoline and CCF is a California Certification Fuel (reformulated gasoline).

Component	Average RI	RRF	MW	Relative Density	RFA Gasoline			CCF Gasoline		
					Min.	Index	Mass %	Min.	Index	Mass %
Methane	100.00	1.121	16.043	0.2600						
Ethylene	178.10	1.050	28.054	0.3000						
Ethane	200.00	1.051	30.070	0.3399						
Propylene	284.00	1.030	42.081	0.5053				7.173	293.43	0.000
Propane	300.00	1.027	44.097	0.5005	7.270	300.32	0.003	7.266	299.79	0.003
i-Butane	366.15	1.015	58.124	0.5572	8.266	365.46	0.088	8.262	365.29	0.078
Methanol	378.82	3.148	32.040	0.7914				8.506		0.021
Butene-1	390.72	0.980	56.108	0.5951				8.893	390.31	0.019
Isobutylene	391.51	0.980	56.108	0.5951						
1,3-Butadiene	394.93	0.980	54.092	0.6211						
n-Butane	400.00	1.015	58.124	0.5788	9.195	400.00	4.637	9.193	400.00	1.201
Vinyl acetylene	409.00	1.100	54.090	0.6500						
t-Butene-2	412.09	0.980	56.108	0.6042	9.441	411.72	0.002	9.567	412.10	0.013
2,2-Dimethylpropane	415.10	1.008	72.151	0.5910	9.670	415.09	0.036	9.666	415.05	0.017
c-Butene-2	427.74	0.980	56.108	0.6213	9.983	427.70	0.004	10.128	427.73	0.018
1,2-Butadiene	450.00	0.945	54.092	0.6520						
Ethanol	455.33	2.193	46.070	0.7890	11.063	452.52	0.006			
3-Methylbutene-1	460.84	0.980	70.135	0.6272	11.670	460.81	0.010	11.665	460.76	0.020
O ₁	469.00	0.980	70.135	0.6300	12.030	470.03	0.005			
O ₂	474.00	0.980	70.135	0.6300						
i-Pentane	477.45	1.008	72.151	0.6196	12.653	477.15	4.773	12.650	477.13	7.163
Acetone ^A	477.55	1.850	58.080	0.7899				12.649		0.134
1,4-Pentadiene	481.18	0.952	68.119	0.6607				13.464	482.77	0.005
?	483.00				13.122	486.32	0.014	12.675	482.80	0.003
Butyne-2	488.00	0.945	54.092	0.6910						
Pentene-1	490.83	0.980	70.135	0.6405	13.616	490.86	0.152	13.613	490.85	0.091
i-Propanol	493.38	1.400	60.110	0.8000						
2-Methylbutene-1	496.66	0.980	70.135	0.6504	14.074	496.73	0.334	14.071	496.72	0.185
n-Pentane	500.00	1.008	72.151	0.6262	14.341	500.00	3.627	14.339	500.00	1.094
Isoprene	506.02	0.952	68.119	0.6809	14.666	506.00	0.013	14.664	505.98	0.009
?	508.00				14.644	508.07	0.003			
t-Pentene-2	510.56	0.980	70.135	0.6482	14.917	510.41	0.653	14.916	510.41	0.285
3,3-dimethylbutene-1	516.79	1.050	70.135	0.6500	15.277	516.60	0.011	15.141	516.58	0.004
c-Pentene-2	519.53	0.980	70.135	0.6556	15.439	519.25	0.378	15.438	519.28	0.160
t-Butanol	521.64	1.154	74.120	0.7887				15.468	522.58	0.065
?	522.40							16.198	522.66	0.038
2-Methylbutene-2	524.92	0.980	70.135	0.6623	15.765	524.49	1.100	15.763	524.51	0.461
1t,3-Pentadiene	527.97	0.952	68.119	0.6760	15.960	527.59	0.022	15.956	527.56	0.015
3-Methylbutadiene-1,2	535.00	0.952	68.120	0.6500						
Cyclopentadiene	538.05	0.938	67.100	0.6500	16.478	537.58	0.004	16.475	537.57	0.003
2,2-Dimethylbutane	540.54	1.004	86.178	0.6491	16.779	539.78	1.102	16.776	539.75	1.106
1c,3-Pentadiene	541.90	0.952	68.119	0.6910						
?	543.00				16.895	543.44	0.006			
O ₅	547.70	1.020	70.135	0.6500						
O ₆	549.70	1.020	70.135	0.6500						
Cyclopentene	557.21	0.952	68.119	0.7720	18.026	556.65	0.160	18.025	556.67	0.070
n-Propanol	560.00	1.400	60.110	0.8035						
4-Methylpentene-1	562.02	0.980	84.162	0.6673	18.411	561.26	0.050	18.402	561.42	0.021
3-Methylpentene-1	562.81	0.980	84.162	0.6637	18.468	562.21	0.083	18.469	562.26	0.032
Cyclopentane	566.84	0.980	70.135	0.7454	18.811	566.40	0.216	18.813	566.45	0.052
2,3-Dimethylbutane	569.24	1.004	86.178	0.6616	19.003	568.67	1.723	19.001	568.69	1.655
Methyl-t-butylether	570.65	1.417	88.150	0.7405				19.110	570.03	11.282
4-Methyl-c-pentene-2	571.00	0.980	84.162	0.6741	19.154	570.47	0.113			
2,3-Dimethylbutene-1	572.67	0.980	84.162	0.6830	19.306	572.01	0.048	19.520	572.52	0.028
2-Methylpentane	573.70	1.004	86.178	0.6531	19.388	573.19	5.145	19.389	573.23	3.967
4-Methyl-t-pentene-2	575.47	0.980	84.162	0.6736	19.542	574.94	0.167	19.546	575.03	0.083
O ₈	578.00	0.980	84.162	0.6736	20.042		0.002	19.893		0.002
2-Methyl-1,4-pentadiene	579.00	0.980	82.146	0.6940	20.078		0.002			
?	581.00							20.002		0.002
1,5-Hexadiene	581.90	0.980	82.146	0.6923	20.123	581.47	0.002	20.210		0.011
?	583.90				20.250	583.99	0.002			
3-Methylpentane	585.52	1.004	86.178	0.6643	20.477	585.25	2.589	20.476	585.25	2.189
2-Methylpentene-1	590.19	0.980	84.162	0.6848	20.933	590.01	0.241	20.934	590.05	0.103
Hexene-1	591.06	0.980	84.162	0.6780	21.021	590.91	0.127	21.021	590.94	0.059

TABLE A1.1 *Continued*

Component	Average RI	RRF	MW	Relative Density	RFA Gasoline			CCF Gasoline		
					Min.	Index	Mass %	Min.	Index	Mass %
O ₁₁	592.00	0.980	84.162	0.6780						
i-Butanol	593.50	1.337	74.120	0.8030						
?	596.00				21.474	596.35	0.007			
1c/t,4-Hexadiene	597.14	0.980	84.146	0.7000	21.571	597.22	0.004			
2-Ethylbutene-1	598.95	0.980	84.162	0.6944						
n-Hexane	600.00	1.004	86.178	0.6594	21.937	600.00	2.598	21.935	600.00	1.057
Diisopropylether	601.90	1.100	102.180	0.7241						
t-Hexene-3	602.83	0.980	84.162	0.6821	22.169	602.83	0.191	22.170	602.86	0.080
c-Hexene-3	603.56	0.980	84.162	0.6847	22.258	603.60	0.063	22.234	603.65	0.028
t-Hexene-2	605.44	0.980	84.162	0.6827	22.382	605.40	0.347	22.383	605.43	0.157
2-Methylpentene-2	607.86	0.980	84.162	0.6912	22.583	607.77	0.462	22.584	607.80	0.193
4-Methylcyclopentene	609.00				21.685	608.90	0.113	22.589	608.65	0.047
3-Methyl-c-pentene-2	610.54	0.980	84.162	0.6980	22.816	610.51	0.240	22.817	610.54	0.102
3-Methylcyclopentene	611.61	0.980	82.146	0.7622	22.920	611.74	0.055	22.921	611.77	0.025
O ₁₃	613.08	0.980	84.162	0.6920						
c-Hexene-2	614.67	0.980	84.162	0.6920	23.171	614.60	0.194	23.172	614.63	0.088
O ₁₄	617.06	0.980	84.162	0.6920	23.007	617.10	0.004	23.088	617.08	0.002
Ethyl-t-butylether	619.00	1.342	102.180	0.7519						
3,3-Dimethylpentene-1	620.91	0.980	98.189	0.7019	23.722	620.77	0.371	23.723	620.80	0.158
3-Methyl-t-pentene-2	622.11	0.980	84.162	0.7023	23.603	622.17	0.006	23.480	622.19	0.003
2-Butanol	622.40	0.980	74.120	0.8080						
4-4-Dimethyl-t-pentene-2	623.10	0.980	98.189	0.6936						
2,2-Dimethylpentane	624.17	1.000	100.205	0.6738	24.025	624.11	0.084	24.024	624.12	0.128
Methylcyclopentane	625.86	0.980	84.162	0.7486	24.189	625.88	0.963	24.190	625.91	0.355
Cyclic diolefin or triolefin	627.00	0.957	82.140	0.7092						
2,4-Dimethylpentane	630.60	1.000	100.205	0.6727	24.622	630.47	1.036	24.623	630.48	2.437
2,3,3-Tirmethylbutene-1	631.00	0.980	98.189	0.7092						
Cyclic diolefin or triolefin	632.90	0.957	82.140	0.7092	24.846	632.82	0.010	24.844	632.80	0.008
?	634.20				25.835	634.14	0.009			
2,2,3-Trimethylbutane	634.86	1.000	100.205	0.6901	25.049	634.91	0.031	25.050	634.93	0.038
?	636.30				25.186	636.30	0.007	25.091	636.33	0.004
Cyclic diolefin or triolefin	638.30	0.957	82.140	0.7092	25.380	639.26	0.008	25.378	638.27	0.005
O ₁₇	641.97	0.980	84.160	0.7039	25.745	641.92	0.005	25.622	642.14	0.002
3,4-Dimethylpentene-1	642.87	0.980	98.189	0.7022	25.846	642.92	0.014	25.845	642.92	0.008
4,4-Dimethyl-c-pentene-2	646.65	0.980	98.189	0.7039	26.223	646.57	0.024	26.224	646.61	0.011
2,4-Dimethylpentene-1	647.67	0.980	98.189	0.6988	26.332	647.63	0.021	26.334	647.65	0.011
Diolefin	647.70	0.957	82.140	0.6988						
1-Methylcyclopentene	648.71	0.957	82.146	0.7795	26.443	648.69	0.374	26.444	648.72	0.180
Benzene	649.92	0.910	78.114	0.8789	26.580	649.98	1.969	26.579	649.99	1.242
3-Ethylpentene-1	650.00	0.980	98.189	0.7005						
n-Butanol ⁴	650.02	1.295	74.120	0.8000						
3-Methylhexene-1	650.95	0.980	98.189	0.6959	26.420	651.56	0.029	26.434	651.55	0.015
2-Methyl-c-hexene-3	652.60	0.980	98.189	0.6980	27.081	652.56	0.018	27.059	652.59	0.009
3,3-dimethylpentane	654.43	1.000	100.205	0.6932	27.057	654.47	0.094	27.055	654.46	0.139
5-Methylhexene-1	655.56	0.980	98.189	0.6965	27.198	655.83	0.031	26.985	656.10	0.016
?	656.93				28.233	656.74	0.014	27.752	656.78	0.007
Cyclohexane	657.81	0.980	84.162	0.7785	27.440	657.97	0.225	27.445	658.05	0.050
2-Methyl-t-hexene-3	661.03	0.980	98.189	0.6941	27.763	660.87	0.057	27.766	660.91	0.027
Diolefin (hexadiene)	661.30	0.980	98.189	0.6941	27.946	661.74	0.007	27.794		0.003
2-Ethyl-3-methylbutene-1	662.60	0.980	98.189	0.7135	27.941	662.47	0.018	27.944	662.51	0.009
4-Methylhexene-1	663.81	0.980	98.189	0.7030	28.087	663.77	0.040	28.089	663.80	0.019
4-Methyl-t/c-hexene-2	666.23	0.980	98.189	0.7040	28.357	666.13	0.107	28.361	666.18	0.051
2-methylhexane	667.61	1.000	100.205	0.6786	28.510	667.45	1.342	28.518	667.54	1.236
2,3-Dimethylpentane	668.84	1.000	100.205	0.6951	28.663	668.79	1.635	28.673	668.88	4.375
5-Methyl-t-hexene-2	669.80	0.980	98.189	0.6971						
1,1-Dimethylcyclopentane	671.25	0.980	98.189	0.7545	28.958	671.32	0.045	28.961	671.35	0.042
t-Amylmethylether	672.48	1.318	102.180	0.7517				28.973		0.002
Cyclohexene	673.69	0.980	82.146	0.8110	29.254	673.82	0.058	29.255	673.84	0.032
3-Methylhexane	675.89	1.000	100.205	0.6871	29.496	675.82	1.449	29.497	675.83	1.450
1,6-Heptadiene	677.40	0.980	98.190	0.7500						
3,4-Dimethyl-c-pentene-2	679.46	0.980	98.189	0.7180	29.935	679.42	0.043	29.938	679.46	0.021
5-Methyl-c-hexene-2	680.00	0.980	98.189	0.7060						
1c,3-Dimethylcyclopentane	681.68	0.980	98.189	0.7448	30.225	681.78	0.261	30.228	681.82	0.114
1t,3-Dimethylcyclopentane	684.37	0.980	98.189	0.7488	30.567	684.51	0.228	30.572	684.54	0.104
3-Ethylpentane	685.98	1.000	100.205	0.6981	30.751	685.96	0.202	30.757	686.01	0.169
1t,2-Dimethylcyclopentane	687.07	0.980	98.189	0.7514	30.911	687.21	0.185	30.921	687.30	0.085
2,2,4-Trimethylpentane	688.48	0.998	114.232	0.6919	31.086	688.57	3.273	31.115	688.81	9.481
Heptene-1	688.60	0.980	98.189	0.6970						
2-Ethylpentene-1	689.58	0.980	98.189	0.6970	31.231	689.58	0.059			
1,5-Heptadiene	691.60	0.980	93.168	0.7500						
O ₂₅	692.89	0.980	98.189	0.6900	31.502	692.93	0.007	31.422	692.95	0.004
3-Methyl-c-hexene-3	694.82	0.980	98.189	0.7181	31.912	694.87	0.070	31.913	694.88	0.034

TABLE A1.1 *Continued*

Component	Average RI	RRF	MW	Relative Density	RFA Gasoline			CCF Gasoline		
					Min.	Index	Mass %	Min.	Index	Mass %
?	696.80									
t-Heptene-3	698.39	0.980	98.189	0.7026	32.392	698.43	0.250	31.826	696.87	0.004
n-Heptane	700.00	1.000	100.205	0.6837	32.605	700.00	1.164	32.393	698.44	0.126
c-Heptene-3	701.00	0.980	98.189	0.7028				32.604	700.00	0.996
2-Methyl-2-hexene	701.30	0.980	98.189	0.7126						
3-Methyl-c-hexene-2	702.30	0.980	98.189	0.7126	32.916	702.07	0.283	32.917	702.08	0.139
3-Methyl-t-hexene-3	702.99	0.980	98.189	0.6941	33.064	703.05	0.103	33.065	703.06	0.049
t-Heptene-2	704.58	0.980	98.189	0.7057	33.306	704.63	0.124	33.307	704.64	0.063
3-Ethylpentene-2	705.96	0.980	98.189	0.7249	33.526	706.06	0.065	33.527	706.07	0.030
c-Heptene-2	708.82	0.980	98.189	0.7116	33.974	708.92	0.221	33.990	709.04	0.129
3-Methyl-t-hexene-2	709.50	0.980	98.189	0.7188						
O ₂₈	710.53	0.980	98.189	0.7188						
2,3-Dimethylpentene-2	712.07	0.980	98.189	0.7322	34.488	712.18	0.123	34.488	712.17	0.059
3-Ethylcyclopentene	713.22	0.980	96.173	0.7830	34.789	713.45	0.010	34.562	713.41	0.004
O ₂₉	715.67	0.980	98.189	0.7190	35.083	715.85	0.018	35.084	715.86	0.009
1c,2-Dimethylcyclopentane	717.13	0.980	98.189	0.7322	35.329	717.35	0.138	35.331	717.36	0.060
Methylcyclohexane	717.89	0.980	98.189	0.7694	35.451	718.09	0.380	35.452	718.09	0.139
O ₃₀	719.00	0.980	98.189	0.7322	35.359	720.75	0.079			
2,2-Dimethylhexane	720.70	0.998	114.232	0.6953	35.884	720.70	0.117	35.854	720.51	0.099
1,1,3-Trimethylcyclopentane ^A	720.72	0.980	112.216	0.7482						
O ₃₂	721.00	0.980	98.189	0.7322						
O ₃₃	722.00	0.980	112.216	0.7322						
O ₃₄	723.00	0.980	112.216	0.7322						
O ₃₅	724.35	0.980	98.189	0.7322	36.531	724.50	0.013	36.398	724.50	0.005
O ₃₆	726.26	0.980	98.189	0.7322	36.872	726.47	0.010	36.652	725.97	0.010
?	727.00							36.637	726.09	0.003
Ethylcyclopentane	728.90	0.980	98.189	0.7664	37.342	729.16	0.146	37.381	729.09	0.072
2,5-Dimethylhexane	730.05	0.998	114.232	0.6935	37.522	730.17	0.422	37.523	730.17	0.693
2,2,3-Trimethylpentane	730.90	0.998	114.232	0.7160	37.775	730.96	0.064	37.682	731.06	0.188
2,4-Dimethylhexane	731.84	0.998	114.232	0.7003	37.846	731.99	0.697	37.848	731.98	1.056
?	733.53				38.081	733.63	0.011	37.823		0.007
O ₃₇	735.18	0.980	98.189	0.7322	38.047	735.41	0.004	38.027	735.34	0.003
1c,2t,4-Trimethylcyclopentane	737.11	0.980	112.216	0.7634	38.824	737.36	0.102	38.825	737.35	0.046
3,3-Dimethylhexane	738.39	0.998	114.232	0.7100	39.052	738.59	0.077	39.053	738.59	0.075
O ₃₈	740.43	0.980	98.189	0.7322	39.432	740.61	0.005	38.887	740.62	0.004
?	742.18				39.825	742.27	0.007	39.387	742.53	0.003
?	743.20				40.364	743.28	0.024	40.227	743.50	0.015
?	743.80				39.701	744.00	0.033	39.868	743.90	0.014
1t,2c,3-Trimethylcyclopentane	744.21	0.980	112.216	0.7704	40.162	744.46	0.077	40.163	744.44	0.033
O ₃₉	745.34	0.980	98.189	0.7322				40.019	744.72	0.005
2,3,4-Trimethylpentane	746.83	0.998	114.232	0.7190	40.667	747.06	0.862	40.678	747.11	2.585
l1	747.91	0.998	114.232	0.7190	40.874	748.12	0.195	40.876	748.11	0.093
O ₄₀	749.37	0.980	98.189	0.7322	41.157	749.56	0.058	41.160	749.54	0.032
2,3,3-Trimethylpentane	750.84	0.998	114.232	0.7262	41.539	751.10	0.525	41.470	751.10	1.716
Toluene	751.77	0.920	92.143	0.8670	41.666	752.08	6.421	41.688	752.18	8.999
O ₄₁	752.20	0.980	112.220	0.7322						
O ₄₂	753.63	0.980	112.220	0.7322	42.037	753.73	0.030	42.362	753.75	0.014
?	754.63				42.054	754.65	0.009	42.183	754.77	0.020
O ₄₃	755.33	0.980	112.220	0.7322	42.351	755.48	0.049	42.334	755.36	0.031
2,3-Dimethylhexane	757.87	0.998	114.232	0.7121	42.890	758.08	0.508	42.898	758.11	0.812
2-Methyl-3-ethylpentane	759.04	0.998	114.232	0.7121	43.139	759.28	0.060	43.149	759.31	0.062
1,1,2-Trimethylcyclopentane	760.33	0.980	112.216	0.7725	43.380	760.44	0.045	43.379	760.42	0.025
O ₄₄	761.73	0.980	112.220	0.7322	43.709	761.99	0.076	43.712	761.97	0.041
O ₄₅	762.20	0.980	112.220	0.7322						
O ₄₆	763.00	0.980	112.220	0.7322						
2-Methylheptane	764.14	0.998	114.232	0.6979	44.199	764.29	0.831	44.198	764.26	0.571
2-ethylhexene-1 ^A	764.20	0.980	112.220	0.7650						
4-Methylheptane	765.62	0.998	114.232	0.7046	44.521	765.78	0.362	44.521	765.75	0.266
3-Methyl-3-ethylpentane	766.62	0.980	114.232	0.7121	44.753	766.83	0.084	44.750	766.80	0.104
3,4-Dimethylhexane	767.18	0.998	114.232	0.7192	44.865	767.35	0.086	44.867	767.33	0.114
1c,2c,4-Trimethylcyclopentane	768.95	0.980	112.216	0.7620	45.430	769.91	0.090	45.427	769.88	0.041
1c,3-Dimethylcyclohexane	769.80	0.980	112.216	0.7625						
3-Methylheptane	771.78	0.998	114.232	0.7058	45.880	771.92	0.911	45.877	771.88	0.651
1c,2t,3-Trimethylcyclopentane	772.98	0.980	112.216	0.7704	56.135	773.05	0.291	46.127	772.98	0.185
3-Ethylhexane	773.76	0.998	114.232	0.7136	46.360	774.03	0.055	46.362	774.01	0.022
1t,4-Dimethylcyclohexane	774.89	0.980	112.216	0.7625	46.689	775.16	0.059	46.621	775.15	0.024
?	775.65				46.439	775.62	0.009			
1,3-Octadiene	777.16	0.980	110.200	0.7650	47.117	777.33	0.011	46.298	777.29	0.006
O ₄₈	778.50	0.980	112.220	0.7322	46.756	779.08	0.004	46.542	779.02	0.003
1,1-Dimethylcyclohexane	780.48	0.980	112.216	0.7809	47.922	780.76	0.009	47.413	780.75	0.005
2,2,5-Trimethylhexane	782.93	0.996	128.259	0.7072	48.473	783.08	0.470	48.473	783.04	0.740
3c-Ethylmethylcyclopentane	784.35	0.980	112.216	0.7670	48.831	784.57	0.130	48.833	784.53	0.060

TABLE A1.1 *Continued*

Component	Average RI	RRF	MW	Relative Density	RFA Gasoline			CCF Gasoline		
					Min.	Index	Mass %	Min.	Index	Mass %
2,6-Dimethylheptene-1	785.55	0.980	126.240	0.7196	49.152	785.64	0.018	49.046	785.66	0.010
3t-Ethylmethylcyclopentane	786.55	0.980	112.216	0.7670	49.366	786.75	0.081	49.369	786.74	0.036
2t-Ethylmethylcyclopentane	787.86	0.980	112.216	0.7690	49.679	788.03	0.071	49.682	788.00	0.036
Octene-1 ^A	787.87	0.980	112.220	0.7650						
1,1-Methylethylcyclopentane	788.78	0.980	112.216	0.7809	49.894	788.90	0.028	49.896	788.89	0.013
?	789.88				49.436	789.88	0.013	50.166	789.96	0.013
2,2,4-Trimethylhexane	790.75	0.996	128.259	0.7392	50.384	790.88	0.046	50.386	790.86	0.020
1t,2-Dimethylcyclohexane	792.77	0.980	112.216	0.7760	50.911	792.96	0.096	50.915	792.94	0.045
t-Octene-4	794.21	0.980	112.216	0.7185	51.252	794.31	0.052	51.259	794.31	0.029
3,5,5-Trimethylhexene-1	795.00	0.980	126.240	0.7196	50.669		0.002			
t-Octene-3	796.00	0.980	112.216	0.7196						
1c,2c,3-Trimethylcyclopentane	797.25	0.980	112.216	0.7792	52.033	797.34	0.151	52.042	797.34	0.080
1t,3-Dimethylcyclohexane	798.80	0.980	112.216	0.7760	52.443	798.90	0.033	52.450	798.89	0.017
n-Octane	800.00	0.998	114.232	0.7025	52.733	800.00	0.811	52.740	800.00	0.502
1c,4-Dimethylcyclohexane	801.05	0.980	112.216	0.7828						
3,3-Dimethylheptene-1	802.50	0.980	126.240	0.7196						
Octene-2	804.40	0.980	112.216	0.7196	53.872	804.52	0.065	53.885	804.54	0.035
O ₅₀	805.50	0.980	112.216	0.7196						
I ₂	806.39	0.996	128.259	0.7300	54.389	806.51	0.112	54.406	806.56	0.052
?	807.00							54.600	807.29	0.020
i-Propylcyclopentane	808.06	0.980	112.216	0.7765	54.822	808.20	0.074	54.840	808.23	0.035
2,4,4-Trimethylhexane	808.50	0.996	128.259	0.7392						
O ₅₂	810.62	0.980	126.240	0.7196	55.508	810.81	0.011	55.545	810.84	0.006
O ₅₃	813.47	0.980	126.240	0.7196	56.268	813.67	0.035	56.295	813.72	0.016
N ₁	815.02	0.980	112.216	0.7800	56.655	815.09	0.012	55.395	814.95	0.007
?	815.60				56.673		0.006			
2,2,3,4-Tetramethylpentane	816.45	0.996	128.259	0.7389	57.221	816.59	0.009	56.554	816.50	0.005
2,3,4-Trimethylhexane	818.10	0.996	128.259	0.7392	57.536	818.33	0.081	57.567	818.38	0.127
N ₂	819.93	0.980	112.216	0.7800	58.170	820.20	0.034	58.093	820.25	0.014
?	820.85				58.049	821.28	0.014	57.612	821.13	0.007
?	821.10				59.376	820.70	0.014	59.939	821.10	0.009
N ₃	822.29	0.980	112.216	0.7800	58.668	822.41	0.039	58.725	822.55	0.022
2,3,3-Trimethylhexene-1	824.74	0.980	126.240	0.6826	59.404	824.98	0.017	59.582	825.08	0.010
?	825.00									
1c,2-Dimethylcyclohexane	826.48	0.980	112.216	0.7962	59.179	826.84	0.023			
1,3,5-Trimethylhexane	827.51	0.996	128.259	0.7219	60.088	827.38	0.122	60.141	827.50	0.084
?	828.95				60.649	828.50	0.006			
?	829.20							59.422		0.002
2,2-Dimethylheptane	829.76	0.996	128.259	0.7105	59.233	829.20	0.012	59.294	829.15	0.005
?	831.80							60.231		0.004
1,1,4-Trimethylcyclohexane	832.56	0.980	126.243	0.7722	61.689	832.81	0.104	61.744	832.94	0.049
N ₄	834.07	0.980	112.216	0.7800	62.028	833.97	0.021	62.098	834.13	0.014
?	834.40				60.862		0.013	60.851		0.008
2,2,3-Trimethylhexane	834.96	0.996	128.259	0.7153	62.399	835.22	0.074	62.472	835.37	0.043
2,4-Dimethylheptane	836.47	0.996	128.259	0.7153	62.878	836.79	0.011	61.246	836.60	0.005
4,4-Dimethylheptane	838.68	0.996	128.259	0.7153	63.454	838.68	0.048	63.530	838.83	0.024
Ethylcyclohexane	840.20	0.980	112.216	0.7839	62.675		0.004	62.624		0.003
n-Propylcyclopentane	841.38	0.980	112.216	0.7763	64.307	842.00	0.008	62.897	841.45	0.004
1c,3c,5-trimethylcyclohexane ^A	841.40	0.980	126.243	0.7697						
2,5-Dimethylheptane	842.63	0.996	128.259	0.7167	64.762	842.88	0.208	64.855	843.09	0.132
3,3-Dimethylheptane	843.96	0.996	128.259	0.7256	65.189	844.25	0.059	65.283	844.44	0.032
3,5-Dimethylheptane	845.02	0.996	128.259	0.7225	65.042	844.55	0.015	64.178		0.002
?	845.60							64.526		0.002
2,6-Dimethylheptane	846.47	0.996	128.259	0.7089	65.987	846.73	0.023	64.434	846.51	0.007
?	847.00							64.667		0.002
1,1,3-Trimethylcyclohexane	848.43	0.980	126.243	0.7870	66.612	848.67	0.024	66.718	848.90	0.011
2,4-Dimethylheptene-1	849.43	0.980	126.240	0.6826	66.462	848.89	0.006	65.309	849.56	0.005
N ₇	850.89	0.980	112.216	0.7800	66.993	850.52	0.008	65.931	851.33	0.005
N ₈	852.36	0.980	112.216	0.7800	68.017	852.57	0.017	67.247	852.64	0.010
N ₁₀	853.04	0.980	126.240	0.7800						
Ethylbenzene	854.65	0.927	106.168	0.8670	68.687	854.92	3.131	68.809	855.18	2.395
N ₁₁	854.70	0.980	126.240	0.7800	67.693		0.015			
1c,2t,4t-Trimethylcyclohexane	856.34	0.980	126.243	0.7800	69.261	856.61	0.060	69.377	857.03	0.037
I ₃	858.51	0.996	128.259	0.7300	70.138	858.77	0.016	68.290	858.74	0.011
?	859.00							68.711		0.002
2-Methyloctene-1	859.80	0.980	126.240	0.6826	70.172	859.99	0.021	70.169	859.87	0.015
?	860.50							69.131		0.002
I ₄	860.89	0.996	128.259	0.7300	70.240	860.78	0.011	69.125	861.19	0.008
?	861.40				69.536		0.005	69.853		0.002
2-Methyloctene-2	862.14	0.980	126.240	0.6826	71.310	862.00	0.014	69.565	862.44	0.009
N ₁₂	863.00	0.980	126.243	0.7800	72.374	863.22	0.017	70.111		0.005
N ₁₃	863.77	0.980	126.243	0.7800						

TABLE A1.1 *Continued*

Component	Average RI	RRF	MW	Relative Density	RFA Gasoline			CCF Gasoline		
					Min.	Index	Mass %	Min.	Index	Mass %
1,3-Dimethylbenzene	864.22	0.927	106.168	0.8642	72.171	864.48	5.181	72.190	864.82	3.649
1,4-Dimethylbenzene	865.20	0.927	106.168	0.8610	72.389	865.45	2.299	72.556	865.82	1.698
2,3-Dimethylheptane	866.02	0.996	128.259	0.7260	71.154	866.59	0.111	72.052	866.55	0.088
3,4-Dimethylheptane	867.94	0.996	128.259	0.7314	73.307	868.00	0.040	73.490	868.38	0.025
?	868.00							72.160		0.002
3,4-Dimethylheptane	868.78	0.996	128.259	0.7314	73.584	868.75	0.041	73.782	869.18	0.023
N ₁₄	869.70	0.980	126.243	0.7800	74.035	869.96	0.018	72.336	870.12	0.009
I ₅	870.95	0.996	128.259	0.7300	74.478	871.15	0.073	74.658	871.53	0.036
4-Ethylheptane	872.73	0.996	128.259	0.7202	75.147	872.96	0.016	75.560	873.30	0.010
4-Methyloctane	873.81	0.996	128.259	0.7202	75.557	874.05	0.237	75.727	874.35	0.104
2-Methyloctane	874.76	0.996	128.259	0.7134	75.918	874.99	0.298	76.089	875.30	0.128
?	875.00							75.116		0.002
N ₁₅	876.00	0.980	126.243	0.7800	76.402	876.26	0.023	74.595	876.35	0.010
1c,2t,3-Trimethylcyclohexane	877.98	0.980	126.243	0.7580	76.601	878.00	0.016	76.726	878.30	0.007
?	878.00							75.982		0.005
3-Ethylheptane	879.11	0.996	128.259	0.7265	77.373	879.55	0.075	77.742	879.58	0.031
3-Methyloctane	880.24	0.996	128.259	0.7205	78.034	880.47	0.336	78.185	880.70	0.144
?	881.04				80.286	881.32	0.016	76.824	881.25	0.004
1c,2t,4c-Trimethylcyclohexane	881.67	0.980	126.243	0.7722	78.628	881.99	0.019	78.869	882.43	0.020
1,1,2-Trimethylcyclohexane	882.78	0.980	126.243	0.8000	78.917	882.72	0.015			
1,2,-Dimethylbenzene	883.47	0.927	106.168	0.8802	79.328	883.76	2.652	79.453	883.89	1.784
?	884.87				77.949	884.74	0.015	77.734	884.66	0.004
I ₆	885.34	0.996	128.259	0.7300	80.135	885.49	0.022	80.257	885.89	0.023
?	885.88				79.109		0.008			
I ₇	886.38	0.996	128.259	0.7300	80.474	886.61	0.062	80.608	886.75	0.139
N ₁₈	887.87	0.980	126.243	0.7800	81.060	888.07	0.038	81.621	888.18	0.017
N ₁₉	888.36	0.980	126.243	0.7800	81.279	888.60	0.030	81.398	888.69	0.011
Nonene-1	889.00	0.980	126.240	0.7684						
?	889.40				81.509	889.16	0.040	80.759	889.21	0.016
I ₈	889.78	0.996	128.259	0.7300	81.888	890.09	0.024	81.476	889.94	0.011
N ₂₀	890.51	0.980	126.243	0.7800	85.070	890.70	0.111	86.286	890.45	0.010
I ₉	891.29	0.996	128.259	0.7300	82.375	891.28	0.102	82.454	891.24	0.235
i-Butylcyclopentane	892.11	0.980	126.243	0.7809	82.743	892.17	0.017	82.390	892.18	0.008
N ₂₁	892.96	0.980	126.243	0.7800	83.054	892.92	0.030	82.938	892.86	0.015
?	893.20				83.607	893.23	0.030	83.590	893.22	0.009
?	894.00				80.659	893.15	0.033	81.076	893.14	0.015
t-7-Methyloctene-3	895.10	0.980	126.241	0.6826						
N ₂₂	895.99	0.980	126.243	0.7800	84.408	896.11	0.057	84.519	896.14	0.029
?	896.76				84.742	896.88	0.014	84.855	896.93	0.007
N ₂₃ /c-nonene-2	897.24	0.980	126.243	0.7800	84.967	897.41	0.035	85.075	897.44	0.018
t-Nonene-3	897.94	0.980	126.241	0.6826						
?	898.44				84.255	898.49	0.034			
I ₁₀	898.70	0.996	128.259	0.7300	85.566	898.78	0.051	85.709	898.90	0.147
?	899.19				87.478	898.94	0.046			
n-Nonane	900.20	0.996	128.259	0.7176	86.082	900.00	0.214	86.186	900.01	0.086
1,1-Methylethylcyclohexane	901.39	0.980	126.243	0.8062	86.378	901.62	0.035	86.476	901.59	0.016
3,7-Dimethyloctene-1	903.40	0.980	140.270	0.7013						
?	904.38				86.929	904.59	0.023	87.029	904.57	0.012
N ₂₅	905.50	0.980	126.243	0.7900	87.138	905.71	0.026	87.239	905.70	0.010
t-2,2,5,5-Tetramethylhexene-3	906.68	0.980	140.270	0.7013	87.352	906.86	0.012	85.841	906.79	0.006
i-Propylbenzene	912.28	0.933	120.195	0.8618	88.419	912.54	0.112	88.510	912.51	0.082
N ₂₆	913.43	0.980	126.243	0.7900						
N ₂₇	914.45	0.980	126.243	0.7900	88.839	914.76	0.045	88.957	914.88	0.029
c-Nonene-3	915.00	0.980	126.240	0.6826	88.198		0.004			
I ₁₁	916.40	0.994	142.286	0.7300				89.216	916.24	0.010
i-Propylcyclohexane	917.51	0.980	126.243	0.8022	89.365	917.50	0.020	88.849	917.53	0.009
?	918.60				90.523	918.03	0.006			
I ₁₂	921.30	0.994	142.286	0.7300	90.138	921.53	0.042	90.227	921.50	0.082
2,2-Dimethyloctane	922.59	0.994	142.286	0.7245	89.707		0.018	90.591	923.41	0.025
2,4-Dimethyloctane	924.39	0.994	142.286	0.7264	90.743	924.65	0.061	90.829	924.62	0.026
N ₂₈	926.32	0.980	126.243	0.7900	89.078	926.37	0.005	90.227		0.003
N ₂₉	927.99	0.980	126.243	0.7900	91.453	928.31	0.008	88.997	928.03	0.005
2,6-Dimethyloctane	930.83	0.994	142.286	0.7276	91.999	931.09	0.038	92.075	931.02	0.018
2,5-Dimethyloctane	932.66	0.994	142.286	0.7302	92.361	932.90	0.065	92.438	932.86	0.034
?	934.00							92.042		0.002
?	934.50				92.053		0.004			
n-Butylcyclopentane	936.13	0.980	126.243	0.7846	93.070	936.48	0.022			
I ₁₃	937.41	0.994	142.286	0.7300	93.309	937.65	0.033	93.261	937.01	0.045
?	937.60							92.567		0.002
N ₃₀	938.04	0.980	140.270	0.8000	92.877	938.25	0.025	93.516	938.28	0.011
I ₁₄	940.39	0.994	142.286	0.7300	93.378	940.53	0.009	91.869	940.47	0.005
?	941.00				93.027		0.005			

TABLE A1.1 *Continued*

Component	Average RI	RRF	MW	Relative Density	RFA Gasoline			CCF Gasoline		
					Min.	Index	Mass %	Min.	Index	Mass %
3,3-Dimethyloctane	942.30	0.994	142.286	0.7390	94.291	942.54	0.064	94.358	942.50	0.028
N ₃₁	943.42	0.980	140.270	0.8000	92.409	943.48	0.006	92.497	943.66	0.004
?	944.55				98.078	945.10	0.018	97.967	945.45	0.006
?	944.95				92.576	944.76	0.019	92.740	944.90	0.009
n-Propylbenzene	946.33	0.933	120.195	0.8620	95.116	946.61	0.627	95.182	946.55	0.401
?	947.54				93.127	947.53	0.020	93.521	947.53	0.008
3,6-Dimethyloctane	948.31	0.994	142.286	0.7363	95.496	948.44	0.023	95.564	948.39	0.011
3-Methyl-5-ethylheptane	949.41	0.994	142.286	0.7264	95.692	949.41	0.022	95.536	949.35	0.010
?	950.00				95.233		0.002	95.262		0.002
N ₃₂	951.22	0.980	140.270	0.8000	96.296	951.32	0.018	95.351		0.002
?	951.50				95.460		0.004	95.442		0.002
?	953.00				95.589		0.003	95.580		0.002
1,3-Methylethylbenzene	954.42	0.933	120.195	0.8645	96.789	954.72	2.027	96.840	954.63	1.276
1,4-Methylethylbenzene	956.22	0.933	120.195	0.8612	97.157	956.49	0.878	97.212	956.42	0.571
?	957.40				94.626	957.43	0.011	96.594		0.005
N ₃₃	958.16	0.980	140.270	0.800						
?	958.90				95.762	958.89	0.008	96.900		0.003
?	960.80				96.911		0.006			
1,3,5-Trimethylbenzene	961.92	0.933	120.195	0.8652	98.348	962.16	0.996	98.400	962.09	0.638
2,3-Dimethyloctane	961.99	0.994	142.286	0.7379						
I15	963.67	0.994	142.286	0.7400	98.713	963.86	0.043	98.765	963.81	0.019
N34	964.76	0.980	140.270	0.8000						
?	965.40				98.438		0.003			
I16	966.53	0.994	142.286	0.7400	97.299	966.78	0.011	97.660	966.93	0.007
?	967.10				103.953	967.90	0.018	98.644		0.004
5-Methylnonane	967.89	0.994	142.286	0.7326	99.600	968.02	0.051	99.652	967.98	0.023
I17	969.41	0.994	142.286	0.7400	99.944	969.61	0.116	100.383	969.77	0.216
1,2-Methylethylbenzene	970.33	0.933	120.195	0.8807	100.146	970.57	0.605	100.194	970.52	0.447
2-Methylnonane	971.77	0.994	142.286	0.7264	100.431	971.87	0.115	100.479	971.83	0.048
?	973.00				100.210	973.37	0.017	99.457	973.29	0.011
3-Ethyl-octane	974.47	0.994	142.286	0.7399	101.009	974.55	0.022	100.826	974.53	0.010
?	975.05							100.504		0.003
N35	975.89	0.980	140.270	0.8000	101.669	976.12	0.018	99.640	976.08	0.006
3-Methylnonane	977.26	0.994	142.286	0.7334	101.629	977.38	0.119	101.675	977.35	0.046
?	978.30				101.271	978.34	0.011	100.504	978.16	0.007
N36	979.33	0.980	140.270	0.8000	100.066	979.21	0.008	99.919	979.30	0.005
3-Ethyl-2-methylheptene-2	979.35	0.980	140.270	0.7013						
I18	980.12	0.994	142.286	0.7400	102.306	980.46	0.029	102.362	980.49	0.065
I19	981.56	0.994	142.286	0.7400	101.282	981.67	0.007	99.704	981.50	0.007
1,2,4-Trimethylbenzene	983.40	0.933	120.195	0.8758	103.003	983.63	2.813	103.032	983.55	1.829
t-butylbenzene ^A	983.42	0.933	120.200	0.8665						
I20	985.82	0.994	142.286	0.7400	103.376	985.29	0.014	102.881	985.32	0.011
i-Butylcyclohexane	986.27	0.980	140.270	0.7960	103.606	986.32	0.023	103.402	986.29	0.010
I21	987.40	0.994	142.286	0.7400	103.819	987.26	0.044	103.845	987.19	0.025
?	987.60							104.866	987.79	0.026
I22	988.00	0.994	142.286	0.7400	105.334	988.43	0.018	102.938	987.84	0.019
?	988.60				102.239	988.63	0.009	102.157	988.55	0.005
I23	989.12	0.994	142.286	0.7400	103.648	989.04	0.011	103.593		0.002
N ₃₇	990.53	0.980	140.270	0.8000	104.581	990.68	0.015	104.365	990.61	0.006
?	991.24				104.174	991.37	0.010	103.370	991.29	0.005
Decene-1	992.81	0.990	140.270	0.7408	103.762	992.78	0.009	103.952	992.90	0.004
1t-Methyl-2-n-propylcyclohexane	993.55	0.980	140.270	0.8000						
2,3-Dimethyloctene-2	993.56	0.990	140.270	0.7400						
I24	993.70	0.994	142.286	0.7400	105.255	993.65	0.053	105.375	993.99	0.041
?	994.20							107.662	994.18	0.029
i-Butylbenzene	995.95	0.938	134.222	0.8532	105.781	995.97	0.063	105.813	995.95	0.046
I25	996.84	0.994	142.286	0.7400	105.356	996.81	0.021	105.398	996.72	0.024
Sec-Butylbenzene	997.79	0.938	134.222	0.8620	106.237	997.97	0.054	106.270	997.95	0.040
?	998.70				105.784		0.005	105.732		0.002
?	999.30							105.862		0.004
n-Decane	1000.20	0.994	142.286	0.7300	106.708	999.99	0.080	106.737	999.97	0.038
I26	1001.71	0.993	156.313	0.7400	106.952	1001.70	0.011	106.990	1001.72	0.016
N ₃₈	1003.39	0.980	140.260	0.8000	107.189	1003.32	0.028	107.218	1003.28	0.017
1,2,3-Trimethylbenzene	1006.88	0.933	120.195	0.8944	107.705	1006.91	0.539	107.732	1006.87	0.323
1,3-Methyl-i-propylbenzene	1009.84	0.938	134.222	0.8610	108.117	1009.73	0.058	108.144	1009.70	0.048
N ₃₉	1011.33	0.980	154.290	0.8000						
1,4-Methyl-i-propylbenzene	1013.24	0.938	134.222	0.8573	108.602	1013.08	0.034	108.638	1013.12	0.025
I27	1014.33	0.993	156.313	0.7400	106.759	1014.42	0.005	105.253	1014.26	0.002
I28	1015.86	0.993	156.313	0.7400	112.126	1016.63	0.012	113.222	1016.31	0.010
I29	1017.87	0.993	156.313	0.7400	107.874	1017.86	0.012	106.471	1017.20	0.009
2-3-Dihydroindene	1019.44	0.918	118.179	0.9640	109.504	1019.21	0.341	109.532	1019.21	0.163
?	1022.40				109.027		0.006	109.056		0.002

TABLE A1.1 *Continued*

Component	Average RI	RRF	MW	Relative Density	RFA Gasoline			CCF Gasoline		
					Min.	Index	Mass %	Min.	Index	Mass %
Sec-butylcyclohexane	1023.07	0.980	140.270	0.8140	107.271	1022.40	0.008	108.837	1022.14	0.007
I30	1024.82	0.993	156.313	0.7400	110.213	1023.97	0.028	110.225	1023.86	0.013
?	1026.50				108.990	1026.17	0.022	107.795	1026.59	0.011
1,2-Methyl-i-propylbenzene	1027.73	0.938	134.222	0.8766	110.677	1027.07	0.031	110.675	1026.86	0.017
?	1028.40				110.218		0.002			
3-Ethylnonane	1029.40	0.993	156.313	0.7440	110.240		0.003	110.276		0.002
?	1031.13				111.214	1030.68	0.011	111.238	1030.69	0.015
N ₄₀	1032.29	0.980	154.290	0.8000						
I31	1033.20	0.993	156.313	0.7400	111.495	1032.55	0.048	111.515	1032.54	0.020
I32	1036.92	0.993	154.290	0.8000	112.055	1036.34	0.021	111.809	1036.22	0.008
?	1038.53				112.295	1037.88	0.012	112.295	1037.71	0.008
1,3-Diethylbenzene	1039.97	0.938	134.222	0.8639	112.538	1039.49	0.205	112.563	1039.49	0.112
1,3-Methyl-n-propylbenzene	1042.60	0.938	134.222	0.8609	112.944	1042.17	0.400	112.967	1042.18	0.232
I33	1044.35	0.993	156.313	0.7400	112.500	1044.04	0.020	112.163	1043.97	0.008
1,4-Diethylbenzene	1045.25	0.938	134.222	0.8620						
1,4-Methyl-n-propylbenzene	1046.40	0.938	134.222	0.8584	113.528	1046.00	0.237	113.550	1046.00	0.135
n-Butylbenzene	1047.48	0.938	134.222	0.8610	113.688	1047.04	0.121	113.710	1047.05	0.063
1,3-Dimethyl-5-ethylbenzene	1049.78	0.938	134.222	0.8800	114.044	1049.36	0.392	114.064	1049.35	0.218
1,2-Diethylbenzene	1051.72	0.938	134.222	0.8799	114.349	1051.35	0.043	114.372	1051.36	0.024
I34	1051.80	0.993	156.313	0.7400				113.736		0.003
t-Decahydronaphthalene	1053.12	0.980	154.290	0.8000	110.769	1052.71	0.004	110.626	1052.67	0.002
N41	1054.60	0.980	154.290	0.8000	114.769	1054.07	0.017	111.829	1054.53	0.008
?	1055.80							114.310		0.007
?	1056.50				115.938	1056.62	0.013	114.197	1056.55	0.007
1,2-Methyl-n-propylbenzene	1057.87	0.938	134.222	0.8736	115.304	1057.54	0.150	115.324	1057.56	0.078
I35	1058.87	0.993	156.313	0.7400	114.740		0.004	114.800		0.003
?	1059.00				114.838		0.002			
?	1059.50				114.918		0.003			
I36	1060.15	0.993	156.313	0.7400	115.058		0.008	115.622	1061.18	0.013
I37	1062.62	0.993	156.313	0.7400	116.030	1062.17	0.045	116.044	1062.16	0.020
?	1063.96				113.810	1063.86	0.006	115.398		0.005
I38	1065.53	0.993	156.313	0.7400	116.492	1065.12	0.038	116.508	1065.12	0.015
1,4-Dimethyl-2-ethylbenzene	1068.05	0.938	134.222	0.8772	116.905	1067.76	0.264	116.920	1067.77	0.140
A3	1068.90	0.938	134.222	0.8594						
1,3-Dimethyl-4-ethylbenzene	1069.53	0.938	134.222	0.8594	117.158	1069.36	0.307	117.173	1069.38	0.158
I39	1071.12	0.993	156.313	0.7400	114.335	1071.04	0.015	114.937	1071.08	0.005
?	1072.49				113.681	1072.50	0.003	116.744		0.006
?	1073.00				116.736		0.005			
I40	1074.39	0.993	156.313	0.7400	117.933	1074.24	0.178	118.598	1073.91	0.068
1,2-Dimethyl-4-ethylbenzene	1075.25	0.938	134.222	0.8745	118.068	1075.08	0.426	118.079	1075.09	0.250
?	1076.00				117.400		0.002			
?	1077.00							117.638		0.003
?	1078.00				115.592	1078.25	0.007	114.048	1078.31	0.005
I41	1079.65	0.993	156.313	0.7400	118.759	1079.41	0.012	117.600	1079.52	0.006
1,3-Dimethyl-2-ethylbenzene	1080.68	0.938	134.222	0.8904	118.945	1080.60	0.031	118.958	1080.62	0.017
I42	1081.60	0.993	156.313	0.7400	114.969	1081.44	0.003	114.988	1081.58	0.003
?	1083.35							118.280		0.002
?	1083.60							118.402		0.003
I43	1084.18	0.993	156.313	0.7400	119.495	1084.00	0.016	118.753	1084.00	0.008
?	1085.30				118.946	1085.34	0.009	116.463	1085.44	0.004
?	1086.54				119.122	1086.45	0.009	118.927		0.006
?	1087.50							117.379	1088.22	0.004
?	1088.80				120.278	1088.86	0.011	117.484	1088.96	0.004
Undecene-1	1090.45	0.980	154.300	0.7503	120.533	1090.42	0.022	120.544	1090.44	0.012
1,4-Methyl-t-butylbenzene	1092.00	0.942	148.240	0.8500	120.778	1091.92	0.038	120.788	1091.96	0.018
1,2-Dimethyl-3-ethylbenzene	1093.12	0.938	134.222	0.8921	120.985	1093.18	0.113	120.993	1093.22	0.058
?	1094.89				117.966	1094.87	0.008	117.921	1094.88	0.004
?	1095.78				118.105	1095.78	0.013	118.273	1095.78	0.007
1,2-Ethyl-i-propylbenzene	1097.22	0.942	148.240	0.8900	121.661	1097.33	0.011	119.139	1097.36	0.005
?	1098.54				118.119	1098.85	0.009	119.095	1098.88	0.003
?	1099.00				120.899		0.005	125.366	1099.10	0.004
n-Undecane	1100.00	0.993	156.313	0.7440	122.105	1100.03	0.053	122.106	1100.03	0.020
1,4-Ethyl-i-propylbenzene	1102.50	0.942	148.240	0.8900	122.417	1102.56	0.012	121.163	1102.56	0.006
1,2,4,5-Tetramethylbenzene	1104.83	0.938	134.222	0.8875	122.718	1104.99	0.234	122.720	1105.00	0.116
1,2-Methyl-n-butylbenzene	1107.30	0.942	148.240	0.8900						
1,2,3,5-Tetramethylbenzene	1108.79	0.938	134.222	0.8903	123.207	1108.93	0.319	123.208	1108.94	0.158
?	1110.82				119.539	1110.82	0.006	122.376		0.005
?	1112.39				122.367		0.002			
?	1113.53				123.033	1112.50	0.047	123.661	1112.56	0.007
?	1115.92				126.210	1113.03	0.045	122.689		0.003
?	1117.49				124.100	1116.08	0.023	123.509	1115.99	0.008
?	1119.00				120.791	1117.53	0.007	124.824	1116.58	0.011

TABLE A1.1 *Continued*

Component	Average RI	RRF	MW	Relative Density	RFA Gasoline			CCF Gasoline		
					Min.	Index	Mass %	Min.	Index	Mass %
?	1119.50							123.251		0.001
?	1120.13				124.606	1120.12	0.017	124.435	1120.07	0.006
?	1121.30				124.756	1121.30	0.015	123.419	1121.29	0.005
1,2-Methyl-t-butylbenzene	1122.80	0.942	148.240	0.8900						
?	1124.62				123.227	1124.59	0.006	124.027		0.006
?	1126.18				127.869	1126.31	0.007			
5-Methylindan	1127.35	0.938	132.200	0.8900	125.539	1127.50	0.300	125.538	1127.50	0.122
?	1127.30				124.651		0.002			
I43	1131.42	0.992	170.340	0.7530	126.032	1131.37	0.020	125.747	1131.30	0.008
4-Methylindan	1133.70	0.938	132.200	0.8900	126.337	1133.74	0.072	126.333	1133.72	0.031
?	1134.90				134.919	1134.95	0.032	135.048	1134.95	0.013
1,2-Ethyl-n-propylbenzene	1136.52	0.942	148.240	0.8900	126.698	1136.56	0.105	126.692	1136.53	0.043
2-Methylindan	1138.11	0.938	132.200	0.9034	126.908	1138.22	0.289	126.903	1138.20	0.121
1,3-Methyl-n-butylbenzene	1140.67	0.942	148.240	0.8900	127.238	1140.79	0.029	127.233	1140.74	0.012
1,3-Di-i-propylbenzene	1142.70	0.945	162.272	0.8900	127.496	1142.79	0.103	127.490	1142.77	0.050
s-Pentylbenzene	1144.27	0.942	148.240	0.8900	127.697	1144.35	0.084	127.692	1144.33	0.033
?	1146.90				129.097	1146.97	0.008	126.758		0.004
n-Pentylbenzene	1149.04	0.942	148.240	0.8900	128.303	1149.00	0.123	128.297	1148.99	0.053
?	1149.83				127.775	1149.87	0.039	129.737	1149.75	0.015
1t-M-2-(4-MP)cyclopentane	1151.80	0.980	168.320	0.8000						
1,2-Di-propylbenzene	1153.16	0.945	162.272	0.8900	128.847	1153.19	0.045	128.839	1153.17	0.019
?	1154.09				128.970	1154.13	0.046	128.961	1154.11	0.019
?	1155.98				127.861		0.005	127.862		0.004
?	1157.64				129.427	1157.63	0.051	128.886	1157.45	0.021
?	1158.00							130.655	1158.06	0.023
1,4-Di-i-propylbenzene	1159.52	0.945	162.272	0.8900	129.675	1159.53	0.079	129.666	1159.51	0.033
Tetrahydronaphthalene	1163.30	0.924	132.206	0.9695	130.166	1163.27	0.034	130.156	1163.24	0.013
?	1165.13				128.975		0.004	128.924		0.002
?	1166.34				129.516	1166.43	0.076	130.568	1166.34	0.030
Naphthalene	1168.01	0.896	128.174	1.0253	130.803	1168.08	0.438	130.792	1168.05	0.190
?	1168.50				129.780		0.004	129.802		0.006
1-t-Butyl-3,5-dimethylbenzene	1169.25	0.945	162.272	0.8900	140.612	1169.00	0.013			
1,4-Ethyl-t-butylbenzene	1173.72	0.945	162.272	0.8900	131.554	1173.75	0.083	131.544	1173.73	0.030
I45	1177.88	0.942	170.300	0.7530	132.112	1177.91	0.124	132.101	1177.89	0.047
I46	1179.46	0.942	170.300	0.7530	132.325	1179.51	0.062	132.313	1179.48	0.022
?	1181.20				131.043		0.003	131.107		0.003
I47	1183.44	0.942	170.300	0.7530	132.864	1183.52	0.083	132.850	1183.47	0.032
I48	1187.14	0.942	170.300	0.7530	133.358	1187.20	0.071	133.347	1187.17	0.027
1,3-Di-n-propylbenzene	1188.64	0.945	162.272	0.8900	133.560	1188.67	0.077	133.547	1188.66	0.032
A5	1190.24	0.945	162.272	0.8900	133.778	1190.29	0.052	133.765	1190.26	0.023
?	1191.00				132.431		0.009	132.415		0.005
Dodecene-1	1192.19	0.980	168.330	0.7584	129.949	1192.19	0.012	132.673		0.005
?	1193.83				130.211	1193.93	0.012	130.350	1193.95	0.004
?	1194.60				132.870	1194.59	0.011			
?	1196.00				129.167	1196.71	0.006	133.044		0.003
A6	1198.52	0.945	162.272	0.8900	134.904	1198.56	0.040	134.888	1198.53	0.016
n-Dodecane	1200.00	0.992	170.340	0.7530	135.106	1200.07	0.046	135.089	1200.03	0.016
?	1202.51				135.377	1202.51	0.013	135.454	1202.41	0.007
?	1204.12				132.955	1204.05	0.025	132.926	1204.09	0.009
?	1205.70				132.109	1205.60	0.002	134.038		0.004
?	1208.41				136.041	1208.47	0.029	136.025	1208.45	0.010
1,3,5-Triethylbenzene	1211.79	0.945	162.272	0.8897	136.427	1211.94	0.015	135.785	1211.76	0.005
?	1212.90				132.913	1213.33	0.015	132.196	1213.37	0.007
?	1213.71				135.006	1213.74	0.017	139.107	1214.01	0.006
?	1215.50				147.556	1215.90	0.028	141.394	1215.65	0.014
?	1216.27				137.053	1217.68	0.059	137.053	1217.63	0.020
?	1217.50				135.296		0.036			
?	1220.12				133.673	1220.92	0.007	135.640		0.005
?	1220.90				135.780	1220.94	0.009	131.628	1220.92	0.002
?	1222.36				137.009	1222.52	0.057	136.986	1222.46	0.021
?	1223.70				144.395	1223.59	0.051	144.409	1223.50	0.020
?	1225.08				137.897	1225.05	0.061	137.878	1224.99	0.023
?	1228.60				138.310	1228.69	0.031	138.291	1228.66	0.011
?	1230.00				136.676		0.020	136.675		0.008
1,2,4-Triethylbenzene	1230.83	0.945	162.272	0.8897	138.551	1230.60	0.019	137.335	1230.73	0.008
?	1232.23				139.172	1232.17	0.027	136.248	1232.12	0.009
?	1235.00				149.881	1235.00	0.008	137.282		0.013
?	1236.42				140.655	1236.40	0.030	141.732	1236.33	0.013
?	1237.42				136.485	1237.39	0.038	136.151	1237.46	0.014
?	1238.00				150.631	1238.00	0.009	137.580		0.002
1,4-Methyl-n-pentylbenzene	1241.71	0.945	162.272	0.8897	139.782	1241.65	0.111	139.761	1241.62	0.041
?	1242.50				151.463	1243.85	0.047			

TABLE A1.1 *Continued*

Component	Average RI	RRF	MW	Relative Density	RFA Gasoline			CCF Gasoline		
					Min.	Index	Mass %	Min.	Index	Mass %
?	1244.15				140.120	1244.60	0.040	140.045	1244.10	0.018
?	1245.00				142.998	1246.66	0.013	136.979	1245.15	0.005
?	1246.48				140.387	1246.86	0.016	140.890	1246.61	0.006
?	1248.73				140.651	1249.22	0.015	139.999	1248.99	0.006
?	1251.16				140.918	1251.54	0.033	140.895	1251.51	0.012
n-Hexylbenzene	1252.85	0.945	162.272	0.8897	141.130	1253.37	0.048	141.107	1253.34	0.019
?	1254.25				137.178	1254.09	0.006	139.194		0.003
?	1255.61				138.479	1255.85	0.108	136.643	1255.74	0.042
?	1257.39				144.513	1257.88	0.040	146.044	1257.31	0.030
?	1259.54				138.912	1259.82	0.053	138.882	1259.85	0.020
?	1262.15				142.153	1262.20	0.020	142.129	1262.17	0.007
?	1262.55				139.337	1263.68	0.017	137.487	1263.60	0.006
?	1264.03				144.425	1265.33	0.014	149.847	1265.65	0.008
?	1265.92				142.689	1266.78	0.043	142.663	1266.76	0.016
?	1266.71				155.002	1268.55	0.025	154.983	1268.25	0.010
?	1269.02				139.934	1269.13	0.031	139.907	1269.18	0.011
I49	1270.79	0.991	184.370	0.7560	143.152	1270.72	0.016	138.284	1270.93	0.005
?	1271.58				140.243	1271.91	0.011	138.379	1271.81	0.004
?	1273.13				143.553	1273.26	0.020	143.558	1273.31	0.009
1,2,3,4,5-Pentamethylbenzene	1274.04	0.942	148.240	1.0000	143.534	1273.98	0.094	143.510	1273.98	0.035
?	1276.70							141.702		0.004
?	1277.23				143.318	1277.27	0.007			
?	1279.96				144.231	1279.89	0.023	144.205	1279.88	0.008
2-Methylnaphthalene	1282.57	0.903	143.170	1.0200	144.522	1282.34	0.431	144.494	1282.33	0.171
?	1286.59				144.932	1285.80	0.034	142.460	1285.86	0.014
?	1287.50				157.493	1285.55	0.036	152.473	1286.23	0.013
?	1288.77				145.273	1288.67	0.032	144.965	1288.52	0.013
Tridecene-1	1290.10	0.980	182.350	0.7658				143.309		0.002
?	1292.41				150.934	1292.15	0.015	143.429		0.002
?	1293.81				145.906	1293.96	0.020	144.668	1293.93	0.007
?	1295.08				142.847	1295.31	0.019	142.817	1295.36	0.008
1-Methylnaphthalene	1297.72	0.903	143.170	1.0200	146.330	1297.50	0.175	146.303	1297.49	0.074
n-Tridecane	1300.00	0.991	184.370	0.7564	146.635	1300.11	0.040	146.604	1300.09	0.013
C ₁₄ ⁺	1300.50						0.361			0.127

^A Components known to co-elute. Unknown components whose group type are known are named with a letter (that is, O for olefin) and a consecutive number. If consecutive numbers are missing, they have been identified by name.

TABLE A1.2 Analyses of Naphtha, Reformate, Alkylate, and Other Refinery Streams

NOTE 1—These data consist of analyses of various refinery samples and the purpose for this table is to illustrate that the DHA test method is equally applicable to other refinery samples as well as to the analysis of spark-ignition engine gasolines. This table presents the retention times and retention indices of the components covering a range of different concentrations as well as the actual analyses.

Component	Coopera- tive Study Average RI	File: DHA94098.D Sample: PONA-Va *NJ* Reference Mixture			File: DHA94099.D Sample: HFLA C-Matrix Auto/Oil C-Matrix Alkylate			File: DHA94101.D Sample: Platformate Auto/Oil C-Matrix Reformate			File: DHA94105.D Sample: No. 1 Ref Naphtha ASTM No. 1 Reference Sample			File: DHA94108.D Sample: ASTM Indolene Standard		
		Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass%	Min.	Index	Mass %
Methane	100.00															
Ethylene	178.10															
Ethane	200.00	6.70	201.5	0.033												
Propylene	284.00															
Propane	300.00	7.20	300.0	0.166												
i-Butane	366.15	8.13	366.0	0.099				8.15	366.1	0.036				8.15	366.0	0.350
Methanol	378.82	8.42	378.6	0.362												
Butene-1	390.72	8.74	390.6	0.087										8.75	390.6	0.007
Isobutylene	391.51	8.76	391.5	0.090												
1,3-Butadiene	394.93	8.87	395.1	0.005												
n-Butane	400.00	9.02	400.0	0.315	9.02	400.0	0.155	9.04	400.0	0.596				9.04	400.0	1.844
Vinyl acetylene	409.00															
t-Butene-2	412.09	9.38	412.0	0.353				9.40	412.0	0.003				9.40	412.1	0.005
2,2-Dimethylpropane	415.10	9.46	414.6	0.005										9.49	414.7	0.024
c-Butene-2	427.74	9.92	427.5	0.438				9.94	427.6	0.004				9.95	427.6	0.012
1,2-Butadiene	450.00															
Ethanol	455.33	11.13	454.8	6.914												
3-Methylbutene-1	460.84	11.43	460.5	0.418				11.46	460.6	0.005				11.47	460.6	0.021
O ₁	469.00															
O ₂	474.00															
i-Pentane	477.45	12.43	477.2	9.757	12.43	477.2	11.716	12.45	477.1	3.248	12.45	477.1	0.158	12.46	477.1	5.163
*Acetone	477.55															
1,4-Pentadiene	481.18	12.70	481.1	0.004												
Butyne-2	488.00															
Pentene-1	490.83	13.40	490.8	0.951				13.43	490.9	0.007				13.44	490.9	0.041
i-Propanol	493.38	13.58	493.1	0.310												
2-Methylbutene-1	496.66	13.87	496.7	1.811				13.90	496.8	0.029				13.91	496.8	0.099
n-Pentane	500.00	14.14	500.0	1.850	14.14	500.0	0.599	14.17	500.0	2.693	14.17	500.0	0.428	14.18	500.0	2.875
Isoprene	506.02	14.47	506.0	0.072												
t-Pentene-2	510.56	14.74	510.7	1.641				14.77	510.9	0.022				14.78	510.8	0.090
3,3-Dimethylbutene-1	516.79	15.09	516.8	0.016												
c-Pentene-2	519.53	15.27	519.7	0.888				15.30	519.9	0.012				15.31	519.8	0.055
t-Butanol	521.64	15.37	521.4	0.250												
2-Methylbutene-2	524.92	15.61	525.2	1.980	15.61	525.3	0.005	15.64	525.3	0.060				15.65	525.2	0.165
1t,3-Pentadiene	527.97	15.81	528.3	0.042												
3-Methylbutadiene-1,2	535.00															
Cyclopentadiene	538.05	16.48	538.3	0.015												
2,2-Dimethylbutane	540.54	16.64	540.7	0.087	16.64	540.6	0.005	16.67	540.6	0.185	16.67	540.6	0.067	16.68	540.6	0.330
1c,3-Pentadiene	541.90															
O ₅	547.70															
O ₆	549.70															
Cyclopentene	557.21	17.91	557.5	0.157										17.94	557.5	0.020
n-Propanol	560.00															
4-Methylpentene-1	562.02	18.31	562.5	0.062										18.35	562.5	0.011
3-Methylpentene-1	562.81	18.37	563.2	0.097										18.41	563.2	0.013
Cyclopentane	566.84	18.70	567.1	0.140	18.70	567.1	0.007	18.72	567.1	0.094	18.72	567.0	0.190	18.73	567.1	0.232
2,3-Dimethylbutane	569.24	18.91	569.6	0.611	18.90	569.5	2.601	18.93	569.6	0.346	18.94	569.6	0.228	18.95	569.6	1.522
Methyl-t-butylether	570.65	19.01	570.8	0.731												



TABLE A1.2 Continued

Component	Cooperative Study Average RI	File: DHA94098.D Sample: PONA-Va			File: DHA94099.D Sample: HFLA C-Matrix			File: DHA94101.D Sample: Platformate			File: DHA94105.D Sample: No. 1 Ref Naphtha			File: DHA94108.D Sample: ASTM Indolene Standard		
		NJ Reference Mixture			Auto/Oil C-Matrix Alkylate			Auto/Oil C-Matrix Reformate			ASTM No. 1 Reference Sample			ASTM D02 Reference Standard		
		Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %
4-Methyl-c-pentene-2	571.00	19.21	573.1	0.061	19.29	573.9	0.754	19.09	571.4	0.008	19.33	574.0	1.857	19.10	571.4	0.018
2,3-Dimethylbutene-1	572.67	19.30	574.1	2.213				19.32	574.1	1.878				19.25	573.0	0.009
2-Methylpentane	573.70	19.46	575.9	0.175				19.49	575.9	0.012				19.34	574.0	2.536
4-Methyl-t-pentene-2	575.47													19.50	575.9	0.027
O ₈	578.00															
2-Methyl-1,4-pentadiene	579.00															
1,5-Hexadiene	581.90															
3-Methylpentane	585.52	20.38	585.7	1.450	20.37	585.6	0.356	20.40	585.7	1.441	20.41	585.7	1.332	20.42	585.7	1.831
2-Methylpentene-1	590.19	29.83	590.4	0.288				20.86	590.4	0.015				20.88	590.4	0.041
Hexene-1	591.06	20.92	591.2	0.157				20.95	591.3	0.005				20.97	591.2	0.020
O ₁₁	592.00															
i-Butanol	593.50															
1-c/4-Hexadiene	597.14															
2-Ethylbutene-1	598.95															
n-Hexane	600.00	21.83	600.0	1.754				21.85	600.0	1.735	21.86	600.0	3.609	21.87	600.0	2.573
Diisopropylether	601.90															
t-Hexene-3	602.83	22.06	602.9	0.323				22.09	603.0	0.010				22.11	603.0	0.035
c-Hexene-3	603.56	22.11	603.6	0.076				22.15	603.8	0.002						
r-Hexene-2	605.44	22.27	605.5	0.436				22.30	605.6	0.014				22.32	605.5	0.049
2-Methylpentene-2	607.86	22.47	608.0	0.547				22.50	608.0	0.028				22.52	608.0	0.073
4-Methylcyclopentene																
3-Methyl-c-pentene-2	610.54	22.68	610.6	0.308				22.71	610.7	0.018				22.73	610.6	0.041
3-Methylcyclopentene	611.61	22.76	611.5	0.040										22.81	611.5	0.007
O ₁₃	613.08	22.90	613.2	0.003												
c-Hexene-2	614.67	23.04	614.8	0.250				23.07	614.8	0.008				23.08	614.8	0.028
O ₁₄	617.06	23.23	617.1	0.005												
Ethyl-t-butylether	619.00							23.60	621.1	0.026				23.63	621.1	0.053
3,3-Dimethylpentene-1	620.91	23.58	621.1	0.602												
3-Methyl-t-pentene-2	622.11	23.70	622.4	0.009												
2-Butanol	622.40															
4,4-Dimethyl-t-pentene-2	623.10															
2,2-Dimethylpentane	624.17	23.86	624.3	0.053				23.88	624.2	0.118	23.89	624.2	0.104	23.91	624.2	0.174
Methylcyclopentane	625.86	23.99	625.7	1.057	23.86	624.0	0.006	24.02	625.7	0.310	24.03	625.7	2.226	24.04	625.7	0.982
Cyclic Diolfin or triolefin	627.00															
2,4-Dimethylpentane	630.60	24.45	630.8	0.768	24.47	630.8	8.169	24.47	630.7	0.243	24.48	630.7	0.192	24.50	630.7	1.039
2,3,3-Trimethylbutene-1	631.00															
Cyclic Diolfin or triolefin	632.90	24.67	633.1	0.013												
2,2,3-Trimethylbutane	634.86	24.83	634.9	0.046	24.83	634.7	0.057	24.86	634.9	0.010	24.87	634.8	0.027	24.88	634.8	0.091
Cyclic diolfin or triolefin	636.30	25.00	636.6	0.010												
O ₁₇	638.30	25.16	638.4	0.011												
3,4-Dimethylpentene-1	641.97	25.52	642.0	0.006												
4,4-Dimethyl-c-pentene-2	642.87	25.60	642.9	0.017												
2,4-Dimethylpentene-1	646.65	25.99	646.8	0.029				26.00	646.6	0.005				26.04	646.8	0.006
Diolfin	647.67	26.06	647.6	0.021												
1-Methylcyclopentene	647.70															
	648.71	26.18	648.7	0.328										26.24	648.7	0.042



TABLE A1.2 Continued

Component	Cooperative Study Average RI	File: DHA94098.D Sample: PONA-Va *NJT Reference Mixture			File: DHA94099.D Sample: HFLA C-Matrix Auto/Oil C-Matrix Alkylate			File: DHA94101.D Sample: Platformate Auto/Oil C-Matrix Reformate			File: DHA94105.D Sample: No. 1 Ref Naphtha ASTM No. 1 Reference Sample			File: DHA94108.D Sample: ASTM Indolene Standard		
		Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %
Benzene	649.92	26.29	649.8	0.541				26.32	649.8	1.786	26.33	649.7	0.445	26.35	649.8	1.506
3-Ethylpentene-1	650.00															
*n-Butanol	650.02															
3-Methylhexene-1	650.95	26.42	651.0	0.038				26.61	652.7	0.003						
2-Methyl-c-hexene-3	652.60	26.60	652.8	0.020				26.80	654.5	0.109	26.81	654.4	0.073	26.82	654.4	0.161
3,3-Dimethylpentane	654.43	26.77	654.4	0.049	26.77	654.2	0.009	26.95	656.0	0.007				26.97	655.8	0.007
5-Methylhexene-1	655.56	26.91	655.8	0.038				27.13	657.7	0.040	27.14	657.5	1.886	27.16	657.6	0.717
Cyclohexane	657.81	27.10	657.6	0.513				27.51	661.2	0.006				27.54	661.2	0.011
2-Methyl-t-hexene-3	661.03	27.49	661.2	0.078												
Diolefin (hexadiene)	661.30															
2-Ethyl-3-methylbutene-1	662.60	27.65	662.7	0.014												
4-Methylhexene-1	663.81	27.77	663.9	0.049										27.84	663.9	0.008
4-Methyl-t-c-hexene-2	666.23	28.05	666.4	0.138				28.08	666.4	0.012				28.09	666.2	0.020
2-Methylhexane	667.61	28.19	667.7	1.337	28.23	667.8	0.310	28.22	667.7	1.382	28.23	667.6	1.464	28.25	667.7	1.452
2,3-Dimethylpentane	668.84	28.32	668.8	1.179	28.37	669.0	13.267	28.35	668.8	0.484	28.37	668.8	0.539	28.38	668.8	1.229
5-Methyl-t-hexene-2	669.80															
1,1-Dimethylcyclopentane	671.25	28.58	671.1	0.085				28.61	671.1	0.048	28.62	671.0	0.508	28.64	671.1	0.130
t-Amylmethylether	672.48	28.71	672.3	0.200												
Cyclohexene	673.69	28.85	673.5	0.050										28.91	673.4	0.009
3-Methylhexane	675.89	29.13	675.9	1.351	29.13	675.7	0.310	29.17	676.0	1.743	29.17	675.8	1.775	29.20	675.9	1.771
1,6-Heptadiene	677.40															
3,4-Dimethyl-c-pentene-2	679.46	29.55	679.5	0.049				29.59	679.6	0.005				29.61	679.5	0.012
5-Methyl-c-hexene-2	680.00															
1c,3-Dimethylcyclopentane	681.68	29.79	681.5	0.376				29.83	681.6	0.076	29.84	681.4	1.014	29.86	681.6	0.279
1t,3-Dimethylcyclopentane	684.37	30.12	684.2	0.331				30.15	684.3	0.081	30.16	684.1	0.950	30.18	684.2	0.255
3-Ethylpentane	685.98	30.34	686.0	0.189	30.36	686.0	0.009	30.34	685.8	0.175	30.36	685.7	0.169	30.38	685.9	0.183
1t,2-Dimethylcyclopentane	687.07	30.44	686.9	0.359				30.48	686.9	0.097	30.49	686.8	1.785	30.51	686.9	0.446
2,2,4-Trimethylpentane	688.48	30.63	688.4	1.852	30.75	689.1	30.506				30.66	688.2	0.006	30.71	688.6	5.212
Heptene-1	688.60															
2-Ethylpentene-1	689.58	30.78	689.6	0.071										31.05	691.3	0.003
1,5-Heptadiene	691.60	30.97	691.2	0.002												
O ₂₅	692.89															
3-Methyl-c-hexene-3	694.82	31.45	694.9	0.087				31.48	694.9	0.009				31.51	694.9	0.013
t-Heptene-3	698.39	31.90	698.4	0.289				31.94	698.5	0.017				31.97	698.5	0.035
n-Heptane	700.00	32.11	700.0	1.810				32.14	700.0	1.304	32.17	700.0	4.808	32.18	700.0	1.914
c-Heptene-3	701.00															
2-Methyl-2-hexene	701.30															
3-Methyl-c-hexene-2	702.30	32.41	702.1	0.355				32.45	702.2	0.032				32.48	702.1	0.043
3-Methyl-t-hexene-3	702.99	32.55	703.1	0.130				32.59	703.1	0.012				32.62	703.1	0.016
t-Heptene-2	704.58	32.78	704.7	0.170				32.82	704.7	0.009				32.85	704.7	0.017
3-Ethylpentene-2	705.96	32.99	706.1	0.074				33.03	706.1	0.005				33.06	706.1	0.010
c-Heptene-2	708.82	33.40	708.8	0.252				33.44	708.9	0.020				33.48	708.9	0.041
3-Methyl-t-hexene-2	709.50															
O ₂₈	710.53															
2,3-Dimethylpentene-2	712.07	33.90	742.1	0.160				33.98	712.5	0.014				33.97	712.1	0.014
3-Ethylcyclopentene	713.22															
O ₂₉	715.67	34.43	715.5	0.015												

TABLE A1.2 *Continued*

Component	Cooperative Study Average RI	File: DHA94098.D Sample: PONA-Va *NJT Reference Mixture			File: DHA94099.D Sample: HFLA C-Matrix Auto/Oil C-Matrix Alkylate			File: DHA94101.D Sample: Platformate Auto/Oil C-Matrix Reformate			File: DHA94105.D Sample: No. 1 Ref Naphtha ASTM No. 1 Reference Sample			File: DHA94108.D Sample: ASTM Indolene Standard		
		Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %
1c,2-Dimethylcyclopentane	717.13	34.75	717.6	1.667				34.68	717.0	0.028						
Methylcyclohexane	717.89				34.74	717.4	0.005	34.78	717.6	0.094	34.82	717.6	5.093	34.82	717.6	1.113
O ₃₀	719.00															
2,2-Dimethylhexane	720.70	35.21	720.5	0.174	35.19	720.1	0.008	35.23	720.4	0.126	35.27	720.4	0.748	35.28	720.5	0.199
*1,1,3-Trimethylcyclopentane	720.72															
O ₃₂	721.00															
O ₃₃	722.00															
O ₃₄	723.00															
O ₃₅	724.35	35.82	724.3	0.019												
O ₃₆	726.26	36.14	726.2	0.014												
Ethylcyclopentane	728.90	36.56	728.7	0.254				36.59	728.7	0.066	36.61	728.6	0.373	36.64	728.7	0.112
2,5-Dimethylhexane	730.05	36.78	730.0	0.386	36.80	729.9	2.941	36.82	730.1	0.211	36.84	729.9	0.258	36.87	730.1	1.326
2,2,3-Trimethylpentane	730.90															
2,4-Dimethylhexane	731.84	37.08	731.8	0.480	37.10	731.7	3.098	37.12	731.9	0.387	37.14	731.7	0.295	37.17	731.9	0.849
O ₃₇	733.53	37.37	733.5	0.015												
1c,2i,4-Trimethylcyclopentane	735.18															
O ₃₉	737.11	37.97	737.0	0.208				38.01	737.0	0.052	38.03	736.8	0.703	38.06	737.0	0.137
3,3-Dimethylhexane	738.39	38.20	738.3	0.045												
O ₃₈	740.43	38.60	740.5	0.008				38.24	738.3	0.084	38.26	738.1	0.098	38.29	738.3	0.088
	742.18	38.91	742.2	0.015												
	743.20	39.07	743.1	0.019												
1,1,2c,3-Trimethylcyclopentane	744.21	39.24	744.0	0.196				39.28	744.1	0.035	39.30	743.9	0.871	39.33	744.1	0.156
O ₃₉	745.34															
2,3,4-Trimethylpentane	746.83	39.73	746.7	0.497	39.79	746.8	8.132									
I ₁	747.91	39.95	747.9	0.198				39.77	746.8	0.016	39.79	746.6	0.052	39.84	746.8	2.781
O ₄₀	749.37	40.21	749.3	0.060				40.01	748.0	0.009	40.02	747.8	0.004			
2,3,3-Trimethylpentane	750.84	40.47	750.6	0.302	40.52	750.7	6.021									
Toluene	751.77	40.64	751.5	2.306				40.78	752.1	12.895	40.70	751.4	1.930	40.58	750.8	2.818
O ₄₁	752.20													40.77	751.8	7.664
O ₄₂	753.63	41.03	753.6	0.031												
O ₄₃	755.33	41.35	755.3	0.060												
2,3-Dimethylhexane	757.87	41.86	757.8	0.417	41.86	757.6	2.284	41.92	758.0	0.263	41.94	757.8	0.472	41.96	757.9	0.856
2-Methyl-3-ethylpentane	759.04	42.08	759.0	0.065	42.09	758.8	0.099	42.14	759.1	0.045	42.15	758.8	0.136	42.19	759.1	0.062
1,1,2-Trimethylcyclopentane	760.33	42.36	760.4	0.055												
O ₄₄	761.73	42.59	761.5	0.074												
O ₄₅	762.20															
O ₄₆	763.00															
2-Methylheptane	764.14	43.12	764.1	1.092	43.10	763.8	0.075	43.17	764.2	0.836	43.20	764.0	2.228	43.22	764.1	0.785
*2-Ethylhexene-1	764.20															
4-Methylheptane	765.62	43.42	765.6	0.359	43.41	765.3	0.022	43.47	765.7	0.433	43.50	765.5	0.524	43.52	765.6	0.305
3-Methyl-3-ethylpentane	766.62	43.64	766.7	0.056	43.60	766.2	0.229	43.67	766.6	0.070	43.69	766.4	0.064	43.71	766.6	0.100
3,4-Dimethylhexane	767.18	43.71	767.0	0.061	43.72	766.8	0.267	43.78	767.2	0.073	43.80	766.9	0.096	43.83	767.1	0.107
1c,2c,4-Trimethylcyclopentane	768.95							44.23	769.3	0.024	44.27	769.2	0.100	44.28	769.3	0.028
1c,3-Dimethylcyclohexane	769.80	44.27	769.7	0.128												



TABLE A1.2 Continued

Component	Cooperative Study Average RI	File: DHA94098.D Sample: PONA-Va			File: DHA94099.D Sample: HFLA C-Matrix			File: DHA94101.D Sample: Platformate			File: DHA94105.D Sample: No. 1 Ref Naphtha			File: DHA94108.D Sample: ASTM Indolene Standard		
		*NJT Reference Mixture			Auto/Oil C-Matrix Alkylate			Auto/Oil C-Matrix Reformate			ASTM No. 1 Reference Sample			ASTM D02 Reference Standard		
		Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %
3-Methylheptane	771.78	44.71	771.7	0.873	44.70	771.5	0.064	44.76	771.8	1.057	44.79	771.6	1.166	44.81	771.8	0.718
1c,2i,3-Trimethylcyclopentane	772.98	44.92	772.7	0.734	44.92	772.5	0.009	44.98	772.9	0.299	45.00	772.6	1.845	45.02	772.8	0.412
3-Ethylhexane	773.76															
1i,4-Dimethylcyclohexane	774.89	45.33	774.6	0.241				45.39	774.7	0.013	45.41	774.5	0.662	45.44	774.7	0.107
1,3-Octadecene	777.16	45.89	777.2	0.016												
O ₄₈	778.50										46.16	777.9	0.002			
1,1-Dimethylcyclohexane	780.48	46.55	780.2	0.099							46.62	780.0	0.291	46.67	780.3	0.047
2,2,5-Trimethylhexane	782.93	47.16	782.9	0.274	47.19	782.8	3.951	47.22	783.0	0.010	47.25	782.8	0.030	47.28	783.0	1.175
3c-Ethylmethylcyclopentane	784.35	47.44	784.1	0.199				47.50	784.2	0.037	47.52	784.0	0.216	47.56	784.2	0.054
2,6-Dimethylheptene-1	785.55	47.64	785.0	0.034				48.01	786.4	0.034	48.03	786.2	0.192	48.07	786.4	0.049
3i-Ethylmethylcyclopentane	786.55	47.95	786.4	0.154												
2i-Ethylmethylcyclopentane	787.86	48.26	787.7	0.172				48.32	787.8	0.037	48.34	787.5	0.484	48.38	787.7	0.094
*Octene-1	787.87															
1,1-Methylethylcyclopentane	788.78	48.54	788.9	0.039	48.70	789.3	0.050	48.79	789.8	0.012	48.82	789.5	0.065	48.83	789.7	0.026
2,2,4-Trimethylhexane	789.88	48.73	789.7	0.023												
1i,2-Dimethylcyclohexane	790.75	49.00	790.8	0.053												
t-Octene-4	792.77	49.41	792.6	0.314				49.47	792.7	0.021	49.49	792.4	0.902	49.52	792.6	0.141
3,5,5-Trimethylhexene-1	794.21	49.83	794.3	0.075												
t-Octene-3	795.00															
1c,2c,3-Trimethylcyclopentane	796.00							50.61	797.3	0.018				50.68	797.3	0.014
1i,3-Dimethylcyclohexane	797.25	50.57	797.3	0.200												
n-Octane	798.80	50.91	798.7	0.051							50.94	798.3	0.031	50.97	798.5	0.006
1c,4-Dimethylcyclohexane	800.00	51.23	800.0	1.978				51.27	800.0	0.837	51.36	800.0	5.227	51.34	800.0	1.159
3,3-Dimethylheptene-1	801.05															
Octene-2	802.50															
O ₅₀	804.40	52.33	804.5	0.082												
i2	805.50															
i-Propylcyclopentane	806.39	52.79	806.3	0.109	52.92	806.6	0.119							53.06	806.9	0.041
2,4,4-Trimethylhexane	808.06	53.21	808.0	0.123				53.28	808.1	0.019	53.36	808.0	0.125	53.25	807.7	0.028
O ₅₂	808.50															
	810.62	53.85	810.5	0.013												
	812.50										54.39	812.1	0.013			
O ₅₃	813.47	54.60	813.4	0.044												
N1	815.02	54.93	814.6	0.031							55.04	814.5	0.073	55.08	814.8	0.013
2,2,3,4-Tetramethylpentane	816.45	55.35	816.2	0.014												
2,3,4-Trimethylhexane	818.10	55.77	817.8	0.057	55.75	817.5	0.585									
N ₂	819.93	56.21	819.4	0.063				55.83	817.8	0.022	55.87	817.7	0.049	55.90	817.9	0.207
	820.85	56.45	820.3	0.016				56.25	819.4	0.014	56.30	819.3	0.067	56.33	819.4	0.017
N ₃	822.29	56.89	821.9	0.069				56.96	822.0	0.028	57.00	821.8	0.107	57.02	822.0	0.038



TABLE A1.2 Continued

Component	Cooperative Study Average RI	File: DHA94098.D Sample: PONA-Va *N)* Reference Mixture			File: DHA94099.D Sample: HFLA C-Matrix Auto/Oil C-Matrix Alkylate			File: DHA94101.D Sample: Platformate Auto/Oil C-Matrix Reformate			File: DHA94105.D Sample: No. 1 Ref Naphtha ASTM No. 1 Reference Sample			File: DHA94108.D Sample: ASTM Indolene Standard		
		Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %
2,3,3-Trimethylhexene-1	824.74	57.55	824.3	0.024							57.58	824.0	0.003			
1c,2-Dimethylcyclohexane	826.48	57.95	825.7	0.084							58.04	825.6	0.174	58.10	825.9	0.030
2,3,5-Trimethylhexane	827.51	58.23	826.8	0.172	58.20	826.4	0.116	58.28	826.8	0.094	58.35	826.7	0.304	58.37	826.8	0.095
2,2-Dimethylheptane	829.76										59.00	829.0	0.059			
1,1,4-Trimethylcyclohexane	832.56	59.66	831.8	0.530	59.60	831.4	0.006	59.71	831.8	0.030	59.79	831.7	1.549	59.81	831.9	0.264
N ₄	834.07	60.06	833.1	0.031	60.05	832.9	0.012	60.12	833.2	0.022	60.17	833.1	0.033	60.19	833.2	0.020
2,2,3-Trimethylhexane	834.96	60.42	834.4	0.263	60.40	834.1	0.056	60.49	834.5	0.059	60.55	834.3	0.862	60.57	834.5	0.183
2,4-Dimethylheptane	836.47	60.80	835.6	0.030							60.91	835.6	0.061			
4,4-Dimethylheptane	838.68	61.43	837.8	0.323	61.41	837.5	0.020				61.55	837.7	1.205	61.58	837.9	0.175
Ethylcyclohexane	840.20										62.33	840.3	0.156	62.35	840.5	0.027
n-Propylcyclopentane	841.38	62.20	840.4	0.076	62.20	840.2	0.011									
*1c,3c,5-Trimethylcyclohexane	841.40															
2,5-Dimethylheptane	842.63	62.68	841.9	0.326	62.66	841.7	0.192	62.74	842.0	0.179	62.80	841.9	0.421	62.84	842.1	0.212
3,3-Dimethylheptane	843.96	63.09	843.3	0.086	63.04	842.9	0.020	63.16	843.4	0.062	63.23	843.3	0.101	63.27	843.5	0.069
3,5-Dimethylheptane	845.02	63.30	843.9	0.038							63.46	844.0	0.100			
2,6-Dimethylheptane	846.47	63.79	845.5	0.083							63.93	845.5	0.188	63.95	845.7	0.033
1,1,3-Trimethylcyclohexane	848.43	64.40	847.5	0.070							64.51	847.4	0.095	64.55	847.6	0.022
2,4-Dimethylheptene-1	849.43															
N ₇	850.89										65.98	852.0	0.103	66.03	852.2	0.020
N ₈	852.36	65.77	851.8	0.046												
N ₁₀	853.04										66.54	853.7	0.797	66.61	854.0	2.653
Ethylbenzene	854.65	66.41	853.7	0.728	66.43	853.7	0.032	66.64	854.3	7.364						
N ₁₁	854.70	66.91	855.3	0.222	66.90	855.1	0.049				67.04	855.3	0.580	67.08	855.5	0.106
1c,2,4i-Trimethylcyclohexane	856.34															
I ₃	858.51	67.63	857.5	0.032							67.75	857.4	0.042			
2-Methyloctene-1	859.80	68.01	858.6	0.030												
I ₄	860.89	68.51	860.1	0.033							68.61	859.9	0.030			
2-Methyloctene-2	862.14	69.05	861.7	0.021												
N ₁₂	863.00	69.25	862.3	0.016												
N ₁₃	863.77															
1,3-Dimethylbenzene	864.22	69.69	836.6	1.828				69.98	864.3	10.485	69.81	863.5	1.833	69.94	863.9	5.660
1,4-Dimethylbenzene	865.20	70.06	864.6	0.693				70.31	865.2	4.479	70.18	864.6	0.558	70.28	864.9	2.475
2,3-Dimethylheptane	866.02	70.36	865.5	0.261	70.32	865.3	0.127				70.47	865.4	0.560			
3,4-Dimethylheptane	867.94	70.99	867.3	0.102	70.93	867.0	0.019	71.09	867.5	0.029	71.11	867.2	0.207	71.15	867.4	0.049
3,4-Dimethylheptane	868.78	71.30	868.2	0.042	71.23	867.9	0.021	71.38	868.3	0.029	71.36	868.0	0.065	71.48	868.4	0.032
N ₁₄	869.70	71.62	869.1	0.033							71.74	869.0	0.048	71.82	869.3	0.012
I ₅	870.95	72.11	870.5	0.106				72.23	870.7	0.053	72.21	870.3	0.131	72.28	870.6	0.051
4-Ethylheptane	872.73	72.83	872.5	0.025												
4-Methyloctane	873.81	73.19	873.5	0.393				73.27	873.6	0.217	73.30	873.4	0.557	73.34	873.6	0.187
2-Methyloctane	874.76	73.55	874.4	0.560	73.53	874.3	0.006	73.63	874.6	0.215	73.66	874.4	0.822	73.69	874.5	0.222
N ₁₅	876.00	73.95	875.5	0.045							74.05	875.4	0.072	74.10	875.7	0.018
1c,2,3-Trimethylcyclohexane	877.98	74.66	877.5	0.073							74.76	877.4	0.208	74.86	877.7	0.033
3-Ethylheptane	879.11	75.19	878.9	0.135				75.24	878.9	0.063	75.29	878.8	0.206	75.32	878.9	0.066
3-Methyloctane	880.24	75.62	880.0	0.577				75.68	880.1	0.288	75.72	879.9	1.004	75.75	880.1	0.280
1c,2,4c-Trimethylcyclohexane	881.67	76.05	881.2	0.061	76.23	881.6	0.042				76.17	881.1	0.090	76.40	881.8	0.124

TABLE A1.2 Continued

Component	Coopera- tive Study Average RI	File: DHA94098.D Sample: PONA-Va *NJ* Reference Mixture				File: DHA94099.D Sample: HFLA C-Matrix Auto/Oil C-Matrix Alkylate				File: DHA94101.D Sample: Platformate Auto/Oil C-Matrix Reformate				File: DHA94105.D Sample: No. 1 Ref Naphtha ASTM No. 1 Reference Sample				File: DHA94108.D Sample: ASTM Indolene Standard			
		Min.	Index	Mass %		Min.	Index	Mass %		Min.	Index	Mass %		Min.	Index	Mass %		Min.	Index	Mass %	
1,1,2-Trimethyl- cyclohexane	882.78	76.49	882.3	0.065	76.49	882.3	0.007			76.62	887.8	0.011		76.60	882.3	0.158		76.64	882.5	0.014	
1,2-Dimethylbenzene	883.47	76.83	883.2	1.011	76.86	883.2	0.062			76.98	883.5	5.364		76.92	883.1	0.970		76.99	883.4	3.212	
I6	884.87	77.43	884.8	0.022										77.56	884.8	0.021		77.48	884.6	0.003	
I7	885.34				77.60	885.2	0.044							77.91	885.7	0.020		77.76	885.4	0.082	
N18	886.38	77.99	886.2	0.054	77.97	886.1	0.296							78.08	886.1	0.022		78.13	886.3	0.141	
N19	887.87																				
Nonene-1	888.36	78.70	888.1	0.323						78.62	887.8	0.011		78.80	888.0	0.990		78.83	888.1	0.172	
I8	889.00	78.91	888.6	0.036										79.02	888.5	0.059					
N20	889.40	79.08	889.0	0.051										79.15	888.9	0.109					
I9	889.78	79.32	889.6	0.112										79.41	889.5	0.332		79.45	889.7	0.047	
N21	890.51	79.81	890.9	0.083	79.80	890.8	0.442							79.90	890.8	0.030		79.95	890.9	0.141	
i-Butylcyclopentane	891.29	80.20	891.8	0.033										80.26	891.6	0.057		80.30	891.8	0.011	
N22	892.11	80.54	892.7	0.091										80.70	892.7	0.116		80.69	892.8	0.024	
t-7-Methyloctene-3	892.96																	81.25	894.2	0.003	
I10	894.00																				
N23/c-nonene-2	895.10	81.89	896.0	0.113										81.81	895.5	0.068					
t-Nonene-3	895.99	82.22	896.8	0.019										82.22	896.5	0.010					
I11	896.76													82.57	897.3	0.083		82.61	897.5	0.016	
N24	897.24	82.46	897.4	0.070																	
N25	897.94																				
1,1-Methylethylcyclo- hexane	898.44																				
3,7-Dimethyloctene-1	898.70	82.93	898.5	0.081	83.01	898.7	0.242							83.04	898.5	0.029		83.15	898.8	0.110	
N26	899.19	83.17	899.1	0.026	83.33	899.4	0.007							83.26	899.0	0.140					
N27	900.20	83.55	900.0	1.495						83.57	900.0	0.224		83.68	900.0	4.771		83.64	900.0	0.691	
2,2,2,5,5-Tetramethylhexene-3	901.39																				
I12	903.40																				
N28	904.38	84.17	903.4	0.033																	
N29	905.50	84.43	904.9	0.108										84.27	903.2	0.078		84.58	905.1	0.013	
N30	906.68													85.54	904.7	0.056					
i-Propylbenzene	911.02	85.55	910.9	0.008																	
N31	912.28	85.80	912.3	0.061						85.83	912.3	0.295		85.87	911.9	0.132		85.90	912.3	0.207	
N32	913.43																				
N33	914.45	86.15	914.1	0.145	86.27	914.7	0.020							86.22	913.8	0.493		86.25	914.1	0.068	
c-Nonene-3	915.00																				
I13	916.40																				
i-Propylcyclohexane	917.51	86.75	917.3	0.086	86.50	915.9	0.015														
N34	918.60	87.06	919.0	0.011						86.79	917.4	0.009		86.82	917.0	0.234		86.85	917.4	0.036	
I14	921.30	87.53	921.4	0.098	87.51	921.2	0.130							87.07	918.3	0.025					
N35	922.59				87.86	923.1	0.037			87.57	921.4	0.013		87.60	921.1	0.228		87.64	921.5	0.093	
2,2-Dimethyloctane	924.39	88.13	924.6	0.135	88.11	924.4	0.017			88.16	924.5	0.029		88.20	924.2	0.274		88.21	924.5	0.042	
N36	926.32	88.49	926.4	0.018										88.56	926.1	0.051					
N37	927.99	88.75	927.8	0.037										88.82	927.5	0.101		88.85	927.8	0.014	
2,6-Dimethyloctane	930.83	89.33	930.7	0.221	89.36	930.8	0.006			89.39	930.8	0.014		89.40	930.5	0.800		89.43	930.8	0.107	
N38	932.66	89.75	932.9	0.144	89.72	932.7	0.021			89.78	932.8	0.038		89.81	932.6	0.245		89.84	932.9	0.050	
2,5-Dimethyloctane	934.50				90.51	936.7	0.053														
n-Butylcyclopentane	936.13	90.38	936.1	0.078										90.44	935.8	0.212		90.48	936.2	0.028	
I15	937.41	90.68	937.7	0.078						90.71	937.6	0.015		90.76	937.4	0.179		90.66	937.1	0.031	
N39	938.04	90.83	938.4	0.059						90.84	938.2	0.016		90.93	938.3	0.125		90.78	937.7	0.028	

TABLE A1.2 Continued

File: DHA94108.D																		
Component	Coopera- tive Study	File: DHA94098.D				File: DHA94101.D				File: DHA94105.D					Sample: ASTM Indolene Standard			
		Sample: PONA-Va		Sample: HFLA C-Matrix		Sample: Platformate		Sample: No. 1 Ref Naphtha		Sample: No. 1 Reference		Sample: No. 1 Reference		Sample: No. 1 Reference		Sample: No. 1 Reference		
		Average RI	*NJ* Reference	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %
114	940.39	91.21	940.3	0.042	91.62	942.3	0.011	91.68	942.4	0.039	91.26	939.9	0.121	91.75	942.6	0.158		
3,3-Dimethyloctane	942.30	91.66	942.5	0.295							91.73	942.3	1.121					
N ₃₁	943.42										92.15	944.4	0.241					
	944.55	92.07	944.6	0.077										92.18	944.7	0.015		
n-Propylbenzene	944.95													92.27	945.2	0.011		
	946.33	92.43	946.4	0.242				92.46	946.3	1.417	92.49	946.1	0.203	92.52	946.4	0.863		
	947.54	92.70	947.7	0.056				92.73	947.6	0.009	92.76	947.4	0.190	92.80	947.8	0.024		
3,6-Dimethyloctane	948.31	92.85	948.5	0.051				92.89	948.4	0.007	92.92	948.2	0.069					
3-Methyl-5-ethylheptane	949.41	93.08	949.6	0.125							93.15	949.4	0.669	93.18	949.6	0.063		
	951.22	93.39	951.1	0.058							93.48	951.0	0.158					
N ₃₂	954.42	94.07	954.4	0.925				94.16	954.6	4.421	94.13	954.1	0.725	94.19	954.6	2.428		
Methylethylbenzene																		
1,4-Methylethylbenzene	956.22	94.44	956.2	0.359				94.51	956.2	1.972	94.50	955.9	0.300	94.55	956.3	1.063		
	958.16	94.79	957.9	0.066														
N ₃₃	961.92	95.63	961.9	0.544				95.69	961.9	2.101	94.86	957.7	0.383	94.89	957.9	0.037		
1,3,5-Trimethylbenzene	961.99				95.72	962.2	0.019				95.70	961.7	0.985	95.73	962.0	1.370		
2,3-Dimethyloctane	963.67	96.05	963.9	0.081				96.08	963.7	0.025	96.11	963.7	0.144	96.14	963.9	0.025		
N ₃₄	964.76	96.20	964.6	0.024							96.27	964.4	0.158					
16	966.53																	
5-Methylnonane	967.89	96.93	968.1	0.159				96.96	967.9	0.023	97.00	967.9	0.262	97.03	968.1	0.033		
117	969.41	97.25	969.6	0.219	97.26	969.5	0.417				97.32	969.4	0.604					
1,2-Methylethylbenzene	970.33	97.39	970.2	0.306				97.43	970.1	1.356	97.45	970.0	0.218	97.49	970.3	1.308		
	971.77	97.75	971.9	0.294				97.78	971.7	0.051	97.82	971.7	0.700	97.84	971.9	0.082		
2-Methylnonane	973.33	98.04	973.3	0.049							98.07	972.9	0.100					
3-Ethyl-2-methylheptene-2	974.47	98.33	974.6	0.080				98.35	974.4	0.013	98.40	974.4	0.255	98.43	974.7	0.027		
N ₃₅	975.89	98.57	975.7	0.059							98.65	975.6	0.120					
3-Methylnonane	977.26	98.93	977.4	0.260				98.96	977.2	0.060	99.00	977.2	0.622	99.03	977.4	0.077		
	978.30	99.14	978.3	0.030														
N ₃₆	979.33										99.55	979.8	0.028					
3-Ethyl-2-methylheptene-2	979.35																	
	980.12	99.51	980.0	0.048				99.57	980.2	0.116				99.67	980.4	0.111		
118	981.56				99.79	981.2	0.005			0.010	99.78	980.8	0.069					
119					100.12	982.6	0.006				99.99	981.8	0.049					
					100.33	983.6	0.042											
1,2,4-Trimethylbenzene	983.40	100.22	983.2	1.286				100.32	983.3	5.960	100.28	983.1	1.477	100.34	983.4	3.921		
*t-Butylbenzene	983.42																	
984.20											100.45	983.9	0.266					
120	985.82	100.72	985.5	0.035				100.63	985.0	0.007								
n-Butylcyclohexane	986.27	100.87	986.2	0.081							100.93	986.0	0.445	100.96	986.2	0.026		
121	987.40	101.12	987.3	0.082				101.06	986.9	0.029	101.18	987.2	0.113	101.18	987.2	0.050		
122	988.60	101.38	988.5	0.059				101.19	987.8	0.038								
123	989.12										101.54	988.8	0.181					
N ₃₇	990.53	101.80	990.4	0.035							101.86	990.2	0.091					
	991.24	102.02	991.4	0.019							102.13	991.4	0.022					
Decene-1	992.81																	
11-Methyl-2-n-propylcyclohexane	993.55	102.53	993.6	0.078							102.51	993.1	0.122					
2,3-Dimethyloctene-2	993.56																	
124	994.53	102.70	994.3	0.052							102.75	994.2	0.173	102.72	994.1	0.098		
n-Butylbenzene	995.95	103.00	995.7	0.027	102.60	993.7	0.051	103.00	995.3	0.127	103.05	995.5	0.065	103.07	995.6	0.049		



TABLE A1.2 Continued

Component	Cooperative Study Average RI	File: DHA94098.D Sample: PONA-Va *NJT Reference Mixture				File: DHA94099.D Sample: HFLA C-Matrix Auto/Oil C-Matrix Alkylate				File: DHA94101.D Sample: Platformate Auto/Oil C-Matrix Reformate				File: DHA94105.D Sample: No. 1 Ref Naphtha				File: DHA94108.D Sample: ASTM Indolene Standard			
		Min.		Index		Mass %		Min.		Index		Mass %		Min.		Index		Min.		Index	
I25	996.20 996.84	103.10 103.29	996.1 997.0	0.030 0.056	996.2 996.9	0.023 0.017	103.17 103.31	996.2 996.9	0.023 0.017	996.2 996.9	0.023 0.017	103.17 103.31	996.2 996.9	0.023 0.017	103.17 103.31	996.2 996.9	0.023 0.017	103.17 103.31	996.2 996.9	0.023 0.017	103.17 103.31
Sec-butylbenzene	997.79 998.70	103.44 103.99	997.6 1000.0	0.051 0.739	997.6 1000.0	0.051 0.739	103.44 103.99	997.6 1000.0	0.051 0.739	997.6 1000.0	0.051 0.739	103.44 103.99	997.6 1000.0	0.051 0.739	103.44 103.99	997.6 1000.0	0.051 0.739	103.44 103.99	997.6 1000.0	0.051 0.739	103.44 103.99
n-Decane	1001.71 1003.39	104.46 104.87	1003.4 1006.4	0.049 0.311	1001.1 1001.1	0.028 0.028	104.18 104.18	999.0 1001.1	0.008 0.028	999.6 1001.1	0.048 0.028	104.00 104.18	999.6 1001.1	0.048 0.028	104.00 104.18	999.6 1001.1	0.048 0.028	104.00 104.18	999.6 1001.1	0.048 0.028	104.00 104.18
I26	1008.88 1008.70	105.16 105.33	1008.5 1009.7	0.018 0.057	1008.2 1008.2	0.007 0.007	105.16 105.33	1008.2 1008.2	0.007 0.007	1005.9 1009.1	1.075 0.132	104.90 105.34	1005.9 1009.1	1.075 0.132	104.90 105.34	1005.9 1009.1	1.075 0.132	104.90 105.34	1005.9 1009.1	1.075 0.132	104.90 105.34
1,2,3-Trimethylbenzene	1009.84 1011.33	106.62 106.62	1018.9 1013.1	0.257 0.050	1019.4 1013.7	0.011 0.006	106.73 105.94	1019.4 1013.7	0.011 0.006	1018.4 1013.7	0.463 0.006	106.64 105.94	1018.4 1013.7	0.463 0.006	106.64 105.94	1018.4 1013.7	0.463 0.006	106.64 105.94	1018.4 1013.7	0.463 0.006	106.64 105.94
I27	1015.86 1017.87	106.19 106.45	1015.8 1017.7	0.013 0.028	1016.5 1016.5	0.013 0.013	106.33 106.33	1016.5 1016.5	0.013 0.013	1016.5 1016.5	0.013 0.013	106.33 106.33	1016.5 1016.5	0.013 0.013	106.33 106.33	1016.5 1016.5	0.013 0.013	106.33 106.33	1016.5 1016.5	0.013 0.013	106.33 106.33
I28	1019.44 1022.40	107.23 107.45	1023.1 1024.7	0.056 0.051	1023.7 1023.7	0.009 0.009	107.35 107.35	1023.7 1023.7	0.009 0.009	1025.7 1025.7	0.029 0.029	107.68 107.68	1025.7 1025.7	0.029 0.029	107.68 107.68	1025.7 1025.7	0.029 0.029	107.68 107.68	1025.7 1025.7	0.029 0.029	107.68 107.68
2-3-Dihydroindene	1024.50 1024.82	107.68 107.68	1026.3 1026.3	0.034 0.034	1026.3 1026.3	0.034 0.034	107.68 107.68	1026.3 1026.3	0.034 0.034	1026.3 1026.3	0.034 0.034	107.68 107.68	1026.3 1026.3	0.034 0.034	107.68 107.68	1026.3 1026.3	0.034 0.034	107.68 107.68	1026.3 1026.3	0.034 0.034	107.68 107.68
Sec-butylcyclohexane	1025.70 1026.50	107.90 107.90	1027.8 1027.8	0.109 0.109	1027.8 1027.8	0.109 0.109	107.90 107.90	1027.8 1027.8	0.109 0.109	1027.8 1027.8	0.109 0.109	107.90 107.90	1027.8 1027.8	0.109 0.109	107.90 107.90	1027.8 1027.8	0.109 0.109	107.90 107.90	1027.8 1027.8	0.109 0.109	107.90 107.90
I30	1029.40 1031.13	108.48 108.48	1031.8 1031.8	0.063 0.063	1030.9 1030.9	0.044 0.044	108.38 108.38	1030.9 1030.9	0.044 0.044	1030.9 1030.9	0.044 0.044	108.38 108.38	1030.9 1030.9	0.044 0.044	108.38 108.38	1030.9 1030.9	0.044 0.044	108.38 108.38	1030.9 1030.9	0.044 0.044	108.38 108.38
1,2-Methyl-i-propylbenzene	1032.29 1032.60	108.67 108.67	1033.2 1033.2	0.089 0.089	1033.2 1033.2	0.089 0.089	108.67 108.67	1033.2 1033.2	0.089 0.089	1033.2 1033.2	0.089 0.089	108.67 108.67	1033.2 1033.2	0.089 0.089	108.67 108.67	1033.2 1033.2	0.089 0.089	108.67 108.67	1033.2 1033.2	0.089 0.089	108.67 108.67
3-Ethylnonane	1035.50 1035.92	109.17 109.45	1036.6 1038.5	0.053 0.020	1037.7 1037.7	0.014 0.014	109.37 109.37	1037.7 1037.7	0.014 0.014	1039.4 1039.4	0.477 0.477	109.64 109.64	1039.4 1039.4	0.477 0.477	109.64 109.64	1039.4 1039.4	0.477 0.477	109.64 109.64	1039.4 1039.4	0.477 0.477	109.64 109.64
I31	1038.53 1039.97	109.63 109.63	1039.7 1039.7	0.173 0.173	1041.5 1041.5	0.030 0.030	109.92 109.92	1041.5 1041.5	0.030 0.030	1042.1 1042.1	0.942 0.942	110.04 110.04	1042.1 1042.1	0.942 0.942	110.04 110.04	1042.1 1042.1	0.942 0.942	110.04 110.04	1042.1 1042.1	0.942 0.942	110.04 110.04
I32	1040.50 1042.60	110.01 110.01	1042.4 1042.4	0.257 0.257	1044.35 1044.35	0.064 0.064	110.60 110.60	1044.35 1044.35	0.064 0.064	1045.9 1045.9	0.596 0.596	110.59 110.59	1045.9 1045.9	0.596 0.596	110.59 110.59	1045.9 1045.9	0.596 0.596	110.59 110.59	1045.9 1045.9	0.596 0.596	110.59 110.59
1,3-Diethylbenzene	1044.35 1045.25	110.31 110.31	1044.4 1044.4	0.064 0.064	1046.1 1046.1	0.008 0.008	110.60 110.60	1046.1 1046.1	0.008 0.008	1045.9 1045.9	0.596 0.596	110.59 110.59	1045.9 1045.9	0.596 0.596	110.59 110.59	1045.9 1045.9	0.596 0.596	110.59 110.59	1045.9 1045.9	0.596 0.596	110.59 110.59
1,4-Diethylbenzene	1046.40 1047.48	110.57 110.72	1046.2 1047.2	0.152 0.097	1049.6 1049.6	0.249 0.249	110.07 110.07	1049.6 1049.6	0.249 0.249	1051.1 1051.1	0.070 0.070	111.36 111.36	1051.1 1051.1	0.070 0.070	111.36 111.36	1051.1 1051.1	0.070 0.070	111.36 111.36	1051.1 1051.1	0.070 0.070	111.36 111.36
1,4-Methyl-n-propylbenzene	1049.78 1051.72	111.07 111.32	1049.6 1051.2	0.249 0.038	1055.5 1055.5	0.013 0.013	111.95 111.95	1055.5 1055.5	0.013 0.013	1055.5 1055.5	0.013 0.013	111.95 111.95	1055.5 1055.5	0.013 0.013	111.95 111.95	1055.5 1055.5	0.013 0.013	111.95 111.95	1055.5 1055.5	0.013 0.013	111.95 111.95
n-Butylbenzene	1051.80 1053.12	111.49 111.83	1052.4 1054.7	0.040 0.032	1052.4 1052.4	0.040 0.040	111.49 111.49	1052.4 1052.4	0.040 0.040	1052.4 1052.4	0.040 0.040	111.49 111.49	1052.4 1052.4	0.040 0.040	111.49 111.49	1052.4 1052.4	0.040 0.040	111.49 111.49	1052.4 1052.4	0.040 0.040	111.49 111.49
1,3-Dimethyl-5-ethylbenzene	1053.80 1054.60	111.83 111.83	1054.7 1054.7	0.032 0.032	1055.7 1055.7	0.062 0.062	112.05 112.05	1055.7 1055.7	0.062 0.062	1055.7 1055.7	0.062 0.062	112.05 112.05	1055.7 1055.7	0.062 0.062	112.05 112.05	1055.7 1055.7	0.062 0.062	112.05 112.05	1055.7 1055.7	0.062 0.062	112.05 112.05
I33	1055.80 1056.80	111.95 111.95	1055.5 1055.5	0.013 0.013	1055.5 1055.5	0.013 0.013	111.95 111.95	1055.5 1055.5	0.013 0.013	1055.5 1055.5	0.013 0.013	111.95 111.95	1055.5 1055.5	0.013 0.013	111.95 111.95	1055.5 1055.5	0.013 0.013	111.95 111.95	1055.5 1055.5	0.013 0.013	111.95 111.95
I34	1056.80 1057.80	112.05 112.05	1056.7 1056.7	0.013 0.013	1056.7 1056.7	0.013 0.013	112.05 112.05	1056.7 1056.7	0.013 0.013	1056.7 1056.7	0.013 0.013	112.05 112.05	1056.7 1056.7	0.013 0.013	112.05 112.05	1056.7 1056.7	0.013 0.013	112.05 112.05	1056.7 1056.7	0.013 0.013	112.05 112.05
t-Decalhydronaphthalene	1057.80 1058.80	112.05 112.05	1057.7 1057.7	0.013 0.013	1057.7 1057.7	0.013 0.013	112.05 112.05	1057.7 1057.7	0.013 0.013	1057.7 1057.7	0.013 0.013	112.05 112.05	1057.7 1057.7	0.013 0.013	112.05 112.05	1057.7 1057.7	0.013 0.013	112.05 112.05	1057.7 1057.7	0.013 0.013	112.05 112.05
I41	1058.80 1059.80	112.05 112.05	1058.7 1058.7	0.013 0.013	1058.7 1058.7	0.013 0.013	112.05 112.05	1058.7 1058.7	0.013 0.013	1058.7 1058.7	0.013 0.013	112.05 112.05	1058.7 1058.7	0.013 0.013	112.05 112.05	1058.7 1058.7	0.013 0.013	112.05 112.05	1058.7 1058.7	0.013 0.013	112.05 112.05



TABLE A1.2 Continued

Component	Cooperative Study Average RI	File: DHA94098.D Sample: PONA-Va *NJ* Reference Mixture			File: DHA94099.D Sample: HFLA C-Matrix Auto/Oil C-Matrix Alkylate			File: DHA94101.D Sample: Platformate Auto/Oil C-Matrix Reformate			File: DHA94105.D Sample: No. 1 Ref Naphtha ASTM No. 1 Reference Sample			File: DHA94108.D Sample: ASTM Indolene Standard		
		Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %
1,2-Methyl-n-propylbenzene	1057.87	112.26	1057.5	0.123				112.28	1057.3	0.277	112.31	1057.4	0.176	112.34	1057.6	0.139
I35	1058.87	112.46	1058.9	0.018	112.83	1061.1	0.038				112.50	1058.7	0.047			
I36	1060.15	112.59	1059.8	0.014							112.66	1059.8	0.036	112.93	1061.5	0.400
I37	1062.62	113.03	1062.7	0.091	113.10	1062.9	0.007	113.05	1062.5	0.009	113.08	1062.6	0.199	113.20	1063.3	0.059
I38	1063.96													113.57	1065.8	0.037
	1065.53	113.47	1065.6	0.090	113.47	1065.4	0.006	113.49	1065.4	0.009	113.52	1065.5	0.231			
											113.67	1066.5	0.052			
1,4-Dimethyl-2-ethylbenzene	1068.05	113.80	1067.7	0.185				113.82	1067.6	0.502	113.85	1067.7	0.108	113.87	1067.8	0.229
A3	1068.90															
1,3-Dimethyl-4-ethylbenzene	1069.53	114.01	1069.2	0.316				114.05	1069.2	0.530	114.05	1069.0	0.380	114.11	1069.3	0.230
I39	1071.12	114.32	1071.2	0.033							114.36	1071.0	0.061	114.27	1070.4	0.202
I40	1072.49				114.17	1070.0	0.025	114.74	1073.7	0.050				114.80	1073.8	0.022
	1074.39	114.72	1073.8	0.147				114.92	1074.9	0.886	114.93	1074.8	0.388	114.98	1075.0	0.419
1,2-Dimethyl-4-ethylbenzene	1075.25	114.89	1074.9	0.360												
	1076.00				115.06	1075.8	0.005									
I41	1077.91	115.28	1077.5	0.017							115.30	1077.2	0.080			
	1079.65										116.65	1079.4	0.064			
1,3-Dimethyl-2-ethylbenzene	1080.68	115.72	1080.3	0.053				115.74	1080.2	0.045	115.77	1080.2	0.036	115.82	1080.5	0.040
I42	1081.60	115.94	1081.7	0.012	115.90	1081.3	0.015				116.07	1082.2	0.035	116.00	1081.7	0.168
I43	1084.18	116.32	1084.1	0.017							116.35	1084.0	0.044			
	1085.30	116.44	1084.9	0.016							116.49	1084.9	0.055			
	1086.54										116.71	1086.3	0.072	116.70	1086.2	0.013
	1088.20										117.08	1088.7	0.059	117.00	1088.1	0.024
	1088.88	117.04	1088.8	0.027												
Undecene-1	1090.45	117.30	1090.4	0.035							117.28	1090.0	0.041			
1,4-Methyl-t-butylbenzene	1092.00	117.51	1091.8	0.032				117.50	1091.5	0.051	117.55	1091.7	0.063	117.40	1090.7	0.029
1,2-Dimethyl-3-ethylbenzene	1093.12	117.66	1092.7	0.115				117.68	1092.7	0.226	117.71	1092.7	0.118	117.74	1092.8	0.147
	1094.89	118.05	1095.2	0.016												
	1095.78	118.15	1095.8	0.020							118.17	1095.6	0.052			
1,2-Ethyl-i-propylbenzene	1097.22	118.34	1097.0	0.014				118.34	1097.0	0.015	118.37	1096.9	0.021	118.42	1097.1	0.025
n-Undecane	1098.54	118.53	1098.3	0.018				118.61	1098.7	0.013	118.61	1098.4	0.034			
	1100.00	118.81	1100.0	0.381				118.82	1100.0	0.009	118.86	1100.0	0.753	118.86	1099.9	0.058
	1101.00													119.07	1101.6	0.021
1,4-Ethyl-i-propylbenzene	1102.50	119.13	1102.7	0.018												
1,2,4,5-Tetramethylbenzene	1104.83	119.32	1104.4	0.147				119.33	1104.4	0.473	119.36	1104.3	0.073	119.39	1104.4	0.283
1,2-Methyl-n-butylbenzene	1107.30															
1,2,3,5-Tetramethylbenzene	1108.79	119.77	1108.3	0.200				119.79	1108.3	0.621	119.82	1108.3	0.073	119.85	1108.3	0.393
	1110.82													120.14	1110.8	0.013
	1112.39	120.16	1111.5	0.028												



TABLE A1.2 Continued

Component	Cooperative Study Average RI	File: DHA94098.D Sample: PONA-Va *NJ* Reference Mixture			File: DHA94099.D Sample: HFLA C-Matrix Auto/Oil C-Matrix Alkylate			File: DHA94101.D Sample: Platformate Auto/Oil C-Matrix Reformate			File: DHA94105.D Sample: No. 1 Ref Naptha ASTM No. 1 Reference Sample			File: DHA94108.D Sample: ASTM Indolene Standard		
		Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %
1,2-Methyl-t-butylbenzene	1115.92	120.65	1115.8	0.023				120.75	1116.4	0.007				120.62	1114.9	0.017
	1120.13	121.18	1120.2	0.023										121.25	1120.2	0.012
	1121.30	121.30	1121.2	0.025							121.36	1121.4	0.015			
	1122.80															
5-Methylindan	1124.62													121.83	1125.1	0.013
	1127.35	121.95	1126.7	0.283				121.97	1126.7	0.372				122.02	1126.7	0.083
	1129.83				122.42	1130.4	0.008				122.09	1127.5	0.019			
	1131.42	122.52	1131.5	0.026							122.57	1131.5	0.012	122.52	1130.9	0.041
4-Methylindan	1133.70	122.75	1133.4	0.072				122.76	1133.4	0.124				122.82	1133.4	0.022
	1134.90	122.90	1134.6	0.031							122.79	1133.4	0.011	123.01	1135.0	0.011
	1136.52	123.09	1136.2	0.078				123.11	1136.2	0.161				123.16	1136.3	0.021
											123.13	1136.2	0.022			
1,2-Ethyl-n-propylbenzene	1138.11	123.23	1137.4	0.266				123.25	1137.4	0.353				123.31	1137.5	0.084
	1140.67	123.60	1140.4	0.036				123.56	1140.0	0.009						
2-Methylindan	1142.70	123.81	1142.1	0.071				123.82	1142.2	0.172				123.88	1142.2	0.113
	1144.27	124.03	1144.0	0.094							123.85	1142.2	0.019	124.10	1144.0	0.013
	1148.00	124.52	1148.0	0.047				124.54	1148.0	0.053				124.59	1148.0	0.017
	1149.04	124.62	1148.8	0.049												
n-Pentylbenzene	1149.83	124.71	1149.5	0.047				124.73	1149.6	0.034						
1-M-2-(4-MP)cyclopentane	1151.80															
1,2-Di-i-propylbenzene	1153.16	125.11	1152.8	0.042				125.13	1152.8	0.071				125.18	1152.9	0.015
	1154.09	125.23	1153.8	0.056				125.24	1153.8	0.066				125.29	1153.7	0.012
	1157.64	125.65	1157.2	0.045				125.67	1157.3	0.068				125.72	1157.3	0.021
	1159.52	125.91	1159.4	0.110				125.90	1159.2	0.085				125.95	1159.1	0.017
1,4-Di-i-propylbenzene	1161.30													126.29	1161.8	0.013
Tetrahydronaphthalene	1163.30	126.41	1163.3	0.051				126.37	1162.9	0.007				126.61	1164.5	0.021
	1165.13													126.80	1166.0	0.021
	1166.34	126.73	1166.0	0.060				126.74	1165.9	0.117				126.96	1167.3	0.148
	1168.01	126.89	1167.3	0.285				126.91	1167.3	0.891						
Naphthalene	1169.25	127.15	1169.3	0.015												
1-t-Butyl-3,5-dimethylbenzene	1173.72	127.68	1173.6	0.099				127.66	1173.3	0.010				127.81	1174.1	0.057
	1177.88	128.15	1177.3	0.114				128.18	1177.4	0.036				128.15	1176.8	0.043
	1179.46	128.34	1178.8	0.063				128.29	1178.3	0.010						
	1181.20							128.35	1178.8	0.010						
1,4-Ethyl-t-butylbenzene	1183.44	128.85	1182.9	0.081				128.87	1182.9	0.031						
	1187.14	129.31	1186.5	0.068				129.33	1186.6	0.011						
	1188.64	129.53	1188.2	0.075				129.55	1188.3	0.077				129.60	1188.3	0.026
	1190.24	129.73	1189.8	0.058				129.75	1189.8	0.050				129.80	1189.9	0.022
1,3-Di-n-propylbenzene	1192.19															
Dodecene-1	1198.52	130.79	1198.1	0.037				130.80	1198.1	0.045				130.86	1198.1	0.017
	1200.00	131.03	1200.0	0.332										131.10	1200.0	0.014
	1202.51	131.30	1202.6	0.025												
	1208.41	131.86	1208.0	0.027				131.84	1207.7	0.009						
n-Dodecane	1211.79															
1,3,5-Triethylbenzene	1216.27	132.68	1215.9	0.030				132.68	1215.8	0.009				132.91	1217.6	0.009
	1217.50	132.83	1217.3	0.061				132.86	1217.4	0.024						
	1222.36	133.32	1222.0	0.060				133.33	1221.9	0.023						
	1223.70	133.52	1223.9	0.055				133.53	1223.9	0.022				133.72	1225.3	0.006
	1225.08	133.66	1225.2	0.025				133.65	1225.0	0.014						



TABLE A1.2 Continued

Component	Cooperative Study Average RI	File: DHA94098.D Sample: PONA-Va *NJ* Reference Mixture			File: DHA94099.D Sample: HFLA C-Matrix Auto/Oil C-Matrix Alkylate			File: DHA94101.D Sample: Platformate Auto/Oil C-Matrix Reformate			File: DHA94105.D Sample: No. 1 Ref Naphtha ASTM No. 1 Reference Sample			File: DHA94108.D Sample: ASTM Indolene Standard ASTM D02 Reference Standard		
		Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %	Min.	Index	Mass %
1,2,4-Triethylbenzene	1228.64	133.97	1228.2	0.040				134.03	1228.6	0.010				134.15	1229.8	0.004
	1230.00															
	1230.83															
	1232.23	134.40	1232.2	0.052				134.87	1236.5	0.002						
	1236.42	134.84	1236.4	0.058												
1,4-Methyl-n-pentylbenzene	1237.42	134.98	1237.7	0.012												
	1241.71	135.34	1241.1	0.110				135.35	1241.0	0.083				135.39	1241.1	0.015
	1244.15	135.63	1243.8	0.068												
	1246.48	135.86	1246.0	0.013				135.89	1246.1	0.007						
n-Hexylbenzene	1248.73	136.12	1248.4	0.013				136.12	1248.3	0.007						
	1251.16	136.38	1250.8	0.029												
	1252.85	136.57	1252.5	0.052												
	1255.61	136.84	1255.1	0.101				136.84	1254.9	0.055						
	1257.39	137.11	1257.6	0.029				137.25	1258.7	0.021						
1,2,3,4,5-Pentamethylbenzene	1259.54	137.25	1258.9	0.050												
	1262.15	137.61	1262.2	0.041												
	1266.71	138.08	1266.5	0.062												
	1269.02	138.31	1268.6	0.030										138.30	1268.3	0.015
	1270.79	138.43	1269.8	0.015										138.59	1270.9	0.007
2-Methylnaphthalene	1274.04	138.82	1273.3	0.107				138.85	1273.5	0.067				138.92	1273.9	0.027
	1277.23													139.25	1277.0	0.010
	1279.96	139.49	1279.4	0.022												
	1282.57	139.75	1281.7	0.353				139.77	1281.8	0.786				139.82	1282.1	0.059
Tridecene-1	1286.59	140.11	1285.1	0.010												
	1287.50	140.22	1286.0	0.034										140.40	1287.4	0.019
	1288.77	140.46	1288.2	0.029												
	1290.10															
	1292.41													140.84	1291.5	0.011
1-Methylnaphthalene	1295.08	141.16	1294.6	0.042												
	1297.72	141.42	1296.9	0.152				141.44	1296.8	0.377				141.49	1297.3	0.035
	1300.00	141.77	1300.0	0.234										141.85	1300.6	0.007
C14+	1300.50															

TABLE A1.3 Repeatability and Reproducibility of DHA Determinations

NOTE 1—The following data has been prepared by statisticians of CS94 in accordance with RR: D02-1007, and represents their best estimate of the cooperative study data. Not all of the data qualified for this evaluation since:

(a) 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.

(b) 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.

NOTE 2—

Legend:

$r_{\min}$	=	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} , $R_{\max}$	=	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.

Component	Average RI	$r_{\min}$	r_{est}	$r_{\max}$	$R_{\min}$	R_{est}	$R_{\max}$	$C_{\min}$	$C_{\max}$
i-Butane	366.15	5.6	10.1	16.4	22.8	46.1	81.5	0.06	0.38
Butene-1	390.72	6.4	11.1	17.6	31.2	64.5	115.8	0.01	0.14
n-Butane	400.00	6.8	9.9	13.9	15.3	32.4	59.1	1.02	3.75
2,2-Dimethylpropane	415.10	3.3	8.8	18.6	32.1	50.1	73.7	0.01	0.02
i-Pentane	477.45	5.9	7.2	8.7	8.5	14.8	23.8	2.48	13.38
Pentene-1	490.83	5.2	7.5	10.5	9.7	13.8	19.0	0.06	0.43
2-Methylbutene-1	496.66	4.9	6.9	9.4	8.3	12.9	19.0	0.14	0.86
n-Pentane	500.00	5.2	6.5	8.1	7.1	10.4	14.8	1.06	3.49
t-Pentene-2	510.56	4.5	6.5	9.0	7.2	10.3	14.4	0.28	1.16
c-Pentene-2	519.53	4.7	6.3	8.1	7.6	13.2	20.9	0.16	0.63
2-Methylbutene-2	524.92	4.3	6.0	8.1	7.8	11.4	15.9	0.50	1.85
1t,3-Pentadiene	527.97	6.7	14.0	25.3	17.0	25.3	35.9	0.01	0.06
2,2-Dimethylbutane	540.54	3.1	4.7	6.8	6.4	9.9	14.6	0.08	2.18
Cyclopentene	557.21	4.0	5.8	8.1	7.6	10.5	13.9	0.07	0.27
4-Methylpentene-1	562.02	2.7	4.2	6.1	8.9	11.3	14.0	0.02	0.09
3-Methylpentene-1	562.81	3.5	5.0	6.9	6.1	8.8	12.0	0.03	0.12
Cyclopentane	566.84	3.8	4.9	6.2	7.0	10.1	14.0	0.07	0.59
2,3-Dimethylbutane	569.24	2.9	3.2	3.5	5.1	8.5	13.1	0.70	1.91
2-Methylpentane	573.70	2.5	2.9	3.4	5.1	6.6	8.4	1.06	5.80
4-Methyl-t-pentene-2	575.47	6.5	8.7	11.3	18.1	28.0	41.1	0.08	0.28
3-Methylpentane	585.52	2.6	2.9	3.2	4.0	5.6	7.5	0.60	2.50
2-Methylpentene-1	590.19	2.3	2.7	3.2	3.8	5.4	7.4	0.11	0.45
Hexene-1	591.06	3.1	4.2	5.5	5.9	8.2	10.9	0.06	0.26
n-Hexane	600.00	2.0	2.4	2.9	3.6	5.1	6.9	0.33	2.52
t-Hexene-3	602.83	1.9	3.2	4.9	7.4	11.8	17.7	0.08	0.35
t-Hexene-2	605.44	2.5	2.8	3.2	4.5	6.4	8.9	0.16	0.71
2-Methylpentene-2	607.86	2.4	3.0	3.8	6.1	13.1	24.1	0.22	0.97
3-Methyl-c-pentene-2	610.54	2.1	2.5	2.9	5.3	6.9	8.7	0.11	0.48
3-Methylcyclopentene	611.61	3.0	4.6	6.8	10.1	12.8	15.9	0.02	0.10
c-Hexene-2	614.67	2.6	3.3	4.1	4.8	5.4	6.1	0.09	0.40
3,3-Dimethylpentene-1	620.91	1.9	2.5	3.1	4.3	5.7	7.5	0.17	0.75
2,2-Dimethylpentane	624.17	3.3	4.3	5.5	4.4	8.2	13.7	0.01	0.09
Methylcyclopentane	625.86	2.2	2.6	3.1	4.5	6.4	8.7	0.37	2.35
2,4-Dimethylpentane	630.60	1.6	2.5	3.8	4.2	5.3	6.6	0.20	1.94
2,2,3-Trimethylbutane	634.86	3.2	7.1	13.1	12.5	20.4	31.1	0.02	0.08
3,4-Dimethylpentene-1	642.87	5.5	14.7	30.6	9.7	22.3	43.0	0.01	0.03
4,4-Dimethyl-c-pentene-2	646.65	3.8	5.6	8.0	6.1	11.0	17.9	0.01	0.11
2,4-Dimethylpentene-1	647.67	3.8	6.7	10.8	11.9	13.5	15.3	0.01	0.04
1-Methylcyclopentene	648.71	1.9	2.7	3.7	7.9	8.7	9.6	0.17	0.82
Benzene	649.92	2.6	3.6	4.8	5.5	9.0	13.7	0.17	1.58
5-Methylhexene-1	655.56	5.3	7.5	10.1	18.0	30.4	47.6	0.01	0.22
Cyclohexane	657.81	2.7	3.7	4.9	8.2	14.8	24.3	0.07	0.90
2-Methyl-t-hexene-3	661.03	3.2	6.3	11.0	16.8	23.4	31.6	0.03	0.14
2-Ethyl-3-Methylbutene-1	662.60	4.1	8.9	16.5	89.5	117.0	149.7	0.01	0.04
4-Methylhexene-1	663.81	2.5	4.5	7.2	11.7	14.7	18.3	0.02	0.09
4-Methyl-t/c-hexene-2	666.23	2.1	3.2	4.7	5.4	7.3	9.5	0.05	0.28
2-Methylhexane	667.61	1.6	2.2	2.9	5.1	6.1	7.2	0.39	1.09
2,3-Dimethylpentane	668.84	1.6	2.3	3.2	5.4	6.4	7.5	0.33	3.16
3-Methylhexane	675.89	1.9	2.8	4.0	4.7	5.6	6.7	0.37	1.08
3-4-Dimethyl-c-pentene-2	679.46	3.6	5.9	9.1	11.6	23.4	41.2	0.02	0.14
1c,3-dimethylcyclopentane	681.68	1.7	2.7	4.0	5.4	7.5	10.1	0.11	0.56
1t,3-Dimethylcyclopentane	684.37	1.7	2.8	4.4	5.4	7.7	10.5	0.08	0.48
3-Ethylpentane	685.98	3.1	3.7	4.3	5.6	8.3	11.9	0.08	0.26
2,2,4-Trimethylpentane	688.48	2.4	3.2	4.1	7.4	11.4	16.7	0.10	11.26

TABLE A1.3 *Continued*

Component	Average RI	r _{min}	r _{est}	r _{max}	R _{min}	R _{est}	R _{max}	C _{min}	C _{max}
3-Methyl-c-hexene-3	694.82	3.2	6.5	11.6	7.3	12.9	20.8	0.03	0.17
t-Heptene-3	698.39	2.6	3.2	3.8	5.0	8.1	12.3	0.11	0.67
n-Heptane	700.00	2.5	3.4	4.5	7.7	10.8	14.7	0.21	1.06
3-Methyl-c-hexene-2	702.30	1.5	2.5	4.0	4.6	7.2	10.6	0.13	0.75
3-Methyl-t-hexene-3	702.99	2.2	4.1	6.7	8.1	10.1	12.3	0.05	0.26
t-Heptene-2	704.58	2.6	4.2	6.4	5.9	7.1	8.4	0.06	0.34
3-Ethylpentene-2	705.96	4.7	8.3	13.4	14.5	17.9	21.8	0.03	0.16
c-Heptene-2	708.82	1.9	3.2	5.0	7.0	7.8	8.7	0.12	0.63
2,3-Dimethylpentene-2	712.07	2.4	3.7	5.5	7.8	9.1	10.5	0.06	0.57
O29	715.67	6.8	11.5	18.0	15.7	22.6	31.3	0.01	0.08
1c,2-Dimethylcyclopentane	717.13	2.4	3.7	5.3	9.5	11.6	13.9	0.05	0.20
Methylcyclohexane	717.89	2.8	3.4	4.0	4.1	5.9	8.2	0.11	1.20
2,2-Dimethylhexane	720.70	4.2	7.2	11.3	10.2	15.4	22.3	0.02	0.10
2,5-Dimethylhexane	730.05	2.5	3.0	3.5	4.9	6.2	7.7	0.16	1.12
2,4-Dimethylhexane	731.84	3.5	4.3	5.2	6.5	9.0	12.1	0.29	1.39
1c,2t,4-Trimethylcyclopentane	737.11	3.0	4.2	5.6	6.9	7.9	9.0	0.03	0.17
2,3,4-Trimethylpentane	746.83	2.3	3.8	6.0	5.8	7.8	10.3	0.08	4.26
I1	747.91	2.2	6.2	13.4	10.1	21.1	38.1	0.09	0.59
Toluene	751.77	1.9	2.7	3.8	10.8	13.5	16.5	1.99	10.34
2,3-Dimethylhexane	757.87	2.2	3.7	5.7	5.0	6.9	9.2	0.22	1.23
1,1,2-Trimethylcyclopentane	760.33	7.6	13.5	21.8	13.6	25.7	43.4	0.02	0.26
O44	761.73	4.2	9.4	17.7	12.7	20.5	30.8	0.03	0.24
2-Methylheptane	764.14	3.5	4.9	6.6	4.8	6.1	7.5	0.15	0.63
4-Methylheptane	765.62	4.1	6.2	9.1	7.1	9.1	11.5	0.05	0.29
3-Methyl-3-ethylpentane	766.62	5.4	7.1	9.3	11.0	15.8	21.8	0.05	0.10
3-Methylheptane	771.78	2.1	3.4	5.3	4.4	5.1	5.9	0.13	0.71
1c,2t,3-Trimethylcyclopentane	772.98	3.4	4.4	5.7	5.9	7.4	9.1	0.06	0.29
3-Ethylhexane	773.76	6.9	10.8	15.9	13.1	23.8	39.1	0.01	0.07
1t,4-Dimethylcyclohexane	774.89	9.8	18.0	29.8	22.2	49.5	93.1	0.01	0.11
2,2,5-Trimethylhexane	782.93	3.1	4.3	5.7	5.7	8.0	10.8	0.14	2.21
3c-Ethylmethylcyclopentane	784.35	5.7	12.1	22.1	9.7	22.2	42.5	0.05	0.21
3t-Ethylmethylcyclopentane	786.55	7.4	9.7	12.4	8.4	22.1	46.0	0.02	0.11
2t-Ethylmethylcyclopentane	787.86	8.9	11.4	14.2	12.6	26.7	48.7	0.02	0.12
1t,2-Dimethylcyclohexane	792.77	3.7	6.7	11.0	7.0	10.1	14.0	0.04	0.29
t-Octene-4	794.21	12.7	14.5	16.6	16.7	21.9	28.1	0.02	0.18
1c,2c,3-Trimethylcyclopentane	797.25	3.5	6.0	9.4	6.3	9.2	12.9	0.07	0.51
1t,3-Dimethylcyclohexane	798.80	6.8	9.3	12.4	11.9	16.3	21.7	0.02	0.10
n-Octane	800.00	2.2	3.6	5.5	6.5	15.7	30.9	0.14	0.75
Octene-2	804.40	4.9	8.2	12.7	11.1	20.1	32.9	0.03	0.23
I2	806.39	4.7	10.9	20.9	8.7	19.7	37.4	0.05	0.36
i-Propylcyclopentane	808.06	4.1	10.7	22.0	8.4	19.8	38.4	0.03	0.18
2,3,4-Trimethylhexane	818.10	2.5	5.0	8.9	5.1	7.2	9.8	0.05	0.37
N2	819.93	7.7	11.5	16.4	6.8	17.4	35.7	0.01	0.06
N3	822.29	8.0	13.3	20.7	11.3	23.2	41.5	0.01	0.09
2,3,5-Trimethylhexane	827.51	3.9	5.4	7.1	7.5	31.4	82.4	0.07	0.12
1,1,4-Trimethylcyclohexane	832.56	8.6	17.1	29.9	21.7	36.0	55.6	0.03	0.26
2,2,3-Trimethylhexane	834.96	8.4	13.1	19.2	14.8	20.3	27.0	0.05	0.09
2,5-Dimethylheptane	842.63	6.8	8.8	11.1	7.8	12.3	18.2	0.11	0.18
Ethylbenzene	854.65	2.2	3.2	4.4	7.2	10.6	14.9	0.62	2.62
1,3-Dimethylbenzene	864.22	2.6	3.3	4.2	9.7	12.5	15.7	1.55	6.66
1,4-Dimethylbenzene	865.20	3.6	4.2	5.0	10.4	14.1	18.5	0.62	2.97
I5	870.95	6.6	11.5	18.3	13.8	28.4	50.9	0.02	0.13
4-Methyloctane	873.81	4.4	7.6	12.0	5.9	11.2	18.9	0.05	0.20
2-Methyloctane	874.76	4.6	8.2	13.3	6.0	10.4	16.6	0.07	0.35
3-Ethylheptane	879.11	8.5	13.4	20.1	27.7	38.7	52.5	0.02	0.09
3-Methyloctane	880.24	5.1	8.5	13.0	8.7	15.5	24.9	0.07	0.29
1,2-Dimethylbenzene	883.47	2.1	3.2	4.7	8.8	11.5	14.8	0.83	3.85
I6	885.34	10.2	17.0	26.4	23.3	44.5	75.8	0.02	0.06
I7	886.38	6.5	9.0	12.0	7.8	21.1	44.7	0.05	0.32
N22	895.99	7.9	16.5	29.6	16.7	29.4	47.2	0.03	0.22
N23/c-nonene-2	897.24	5.7	15.9	34.2	26.1	48.7	81.6	0.02	0.15
I10	898.70	4.5	12.3	26.0	9.3	31.1	73.3	0.04	0.44
n-Nonane	900.20	3.9	6.4	9.8	8.6	10.3	12.2	0.06	0.34
i-Propylbenzene	912.28	3.2	5.0	7.3	8.3	15.1	24.9	0.04	0.33
N27	914.45	3.6	12.3	29.2	9.0	21.1	40.8	0.02	0.14
I12	921.30	4.4	11.0	22.2	8.7	21.2	42.2	0.03	0.34
2,4-Dimethyloctane	924.39	6.1	12.1	21.3	16.8	26.0	38.1	0.03	0.11
2,6-Dimethyloctane	930.83	7.0	14.6	26.4	15.4	27.7	45.3	0.02	0.10
2,5-Dimethyloctane	932.66	5.6	11.2	19.6	15.3	22.0	30.4	0.04	0.13
3,3-Dimethyloctane	942.30	4.3	10.4	20.6	7.5	17.5	34.0	0.03	0.11
n-Propylbenzene	946.33	2.8	5.0	8.1	7.6	11.9	17.7	0.21	0.77
3,6-Dimethyloctane	948.31	7.6	14.9	25.7	18.1	38.9	71.4	0.01	0.04
1,3-Methylethylbenzene	954.42	3.7	5.2	7.2	7.6	10.2	13.3	0.81	2.61
1,4-Methylethylbenzene	956.22	3.5	5.3	7.7	5.1	7.7	11.1	0.32	1.19

TABLE A1.3 *Continued*

Component	Average RI	r _{min}	r _{est}	r _{max}	R _{min}	R _{est}	R _{max}	C _{min}	C _{max}
1,3,5-Trimethylbenzene	961.92	3.7	5.5	7.7	5.4	8.3	12.1	0.39	1.21
2-Methylnonane	971.77	6.5	10.6	16.2	17.5	25.9	36.6	0.03	0.19
3-Methylnonane	977.26	7.5	13.5	22.1	23.5	41.0	65.5	0.03	0.16
I18	980.12	7.3	15.7	28.8	14.9	30.0	52.9	0.04	0.82
1,2,4-Trimethylbenzene	983.40	4.2	5.7	7.5	7.8	10.6	13.9	1.19	4.32
I21	987.40	6.9	18.6	39.3	16.9	49.6	109.2	0.03	0.27
I24	994.53	6.9	12.2	19.6	18.9	31.1	47.7	0.04	0.36
n-Decane	1000.20	7.5	9.2	11.1	12.1	17.9	25.3	0.03	0.25
N38	1003.39	9.7	19.8	35.2	25.8	47.6	79.3	0.01	0.12
1,2,3-Trimethylbenzene	1006.88	3.8	5.8	8.5	7.2	8.5	10.0	0.28	0.96
2,3-dihydroindene	1019.44	5.2	7.8	11.3	8.8	10.8	13.0	0.18	0.37
I30	1024.82	7.5	13.9	23.3	20.4	61.8	138.4	0.01	0.15
1,2-Methyl-i-propylbenzene	1027.73	17.3	19.5	22.0	66.4	99.4	141.9	0.02	0.08
?	1038.53	6.8	16.0	31.1	16.8	37.4	70.2	0.02	0.16
1,3-Diethylbenzene	1039.97	5.0	7.3	10.1	9.9	13.7	18.5	0.08	0.22
1,3-Methyl-n-propylbenzene	1042.60	4.0	6.4	9.5	6.7	13.5	23.6	0.30	0.68
1,4-Methyl-n-propylbenzene	1046.40	4.6	8.7	14.7	7.8	12.7	19.4	0.16	0.32
1,3-Dimethyl-5-ethylbenzene	1049.78	3.4	6.0	9.8	7.6	10.6	14.3	0.22	0.51
1,2-Methyl-n-propylbenzene	1057.87	6.3	9.3	13.2	20.9	30.3	42.1	0.11	0.20
		4.5	11.4	23.2	14.4	36.1	73.1	0.06	0.56
I38	1065.53	7.9	16.6	30.0	28.4	37.4	48.2	0.02	0.09
1,4-Dimethyl-2-ethylbenzene	1068.05	4.1	6.7	10.3	15.3	27.8	45.7	0.15	0.38
1,2-Dimethyl-4-ethylbenzene	1075.25	5.1	7.0	9.3	13.8	18.7	24.6	0.29	0.71
Undecene-1	1090.45	8.2	17.8	32.9	20.5	35.4	56.1	0.01	0.09
1,2-Dimethyl-3-ethylbenzene	1093.12	6.8	10.3	14.8	15.2	18.4	22.1	0.09	0.19
1,2-Ethyl-i-propylbenzene	1097.22	14.3	31.3	58.2	51.7	74.6	103.4	0.02	0.06
n-Undecane	1100.00	8.6	13.9	21.0	24.4	40.0	61.2	0.03	0.18
1,2,4,5-Tetramethylbenzene	1104.83	6.1	8.3	11.1	12.5	16.0	20.1	0.15	0.36
1,2,3,5-Tetramethylbenzene	1108.79	6.4	7.8	9.3	10.2	13.9	18.3	0.21	0.51
5-Methylindan	1127.35	6.6	9.2	12.4	7.5	9.3	11.4	0.06	0.34
I43	1131.42	9.1	15.0	23.0	18.8	30.8	47.1	0.02	0.35
4-Methylindan	1133.70	8.0	13.8	21.9	15.3	20.9	27.6	0.02	0.10
1,2-Ethyl-n-propylbenzene	1136.52	7.4	12.0	18.3	25.3	39.6	58.4	0.02	0.11
2-Methylindan	1138.11	5.6	7.7	10.2	6.4	8.9	12.0	0.08	0.34
1,3-Di-i-propylbenzene	1142.70	7.9	9.5	11.2	16.2	17.9	19.7	0.06	0.18
1,2-Di-i-propylbenzene	1153.16	13.9	22.4	34.0	21.3	36.0	56.2	0.01	0.06
?	1157.64	12.8	20.7	31.4	24.5	44.1	71.9	0.02	0.06
1,4-Di-i-propylbenzene	1159.52	11.1	18.6	28.8	18.6	35.6	60.8	0.01	0.09
Napthalene	1168.01	6.1	8.5	11.3	12.9	16.9	21.5	0.13	0.40
1,4-Ethyl-t-butylbenzene	1173.72	9.8	12.0	14.5	18.0	28.1	41.3	0.04	0.30
I48	1187.14	9.1	14.3	21.1	16.8	30.7	50.8	0.01	0.09
1,3-Di-n-propylbenzene	1188.64	9.9	14.4	19.9	18.9	24.2	30.4	0.02	0.08
A5	1190.24	11.5	19.2	29.8	26.7	30.3	34.2	0.02	0.06
A6	1198.52	13.7	23.3	36.7	28.1	39.5	53.5	0.01	0.05
n-Dodecane	1200.00	12.2	16.7	22.1	20.2	32.9	50.0	0.01	0.11
1,4-Methyl-n-pentylbenzene	1241.71	8.2	14.1	22.3	16.5	31.2	52.7	0.01	0.14
1,2,3,4,5-Pentamethylbenzene	1274.04	11.4	13.9	16.7	23.1	29.7	37.5	0.01	0.11
2-Methylnaphthalene	1282.57	7.6	11.1	15.4	17.5	22.3	28.0	0.05	0.50
1-Methylnaphthalene	1297.72	7.3	11.0	15.8	14.0	21.0	30.1	0.02	0.22

A1.10 Tables A1.4-A1.9 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 in-

cluded only spark ignition fuels and different results may be obtained with pure blending components.

TABLE A1.4 Benzene

Sample	Benzene (wt %)	
	D 5580	D 6730
2	1.52	1.58
6	1.05	1.12
8	1.10	1.15
10	1.13	1.19
13	0.14	0.17
14	0.62	0.69
Average	0.93	0.98

**D 6730 – 01****TABLE A1.5 Toluene**

Sample	Toluene (wt %)	
	D 5580	D 6730
2	4.3	4.5
6	2.1	2.0
8	10.1	10.3
10	5.0	5.2
13	3.3	3.3
14	4.4	4.7
Average	4.9	5.0

TABLE A1.6 Total Aromatics

Sample	Total Aromatics (wt %)		
	D 5580	PIONA ^A	D 6730
2	30.3	28.2	30.2
6	18.9	18.7	18.3
8	49.1	49.0	47.6
10	23.9	24.5	23.1
13	19.7	19.8	19.3
14	23.8	24.6	24.2
Average	27.6	27.5	27.1

^A Multidimensional PIONA**TABLE A1.7 Total Olefins**

Sample	Total Olefins (wt %)	
	PIONA ^A	D 6730
2	7.1	4.5
6	9.8	8.7
8	6.6	6.1
10	15.1	12.9
13	11.1	10.6
14	24.6	19.5
Average	12.4	10.9

^A Multidimensional PIONA**TABLE A1.8 Total Oxygenates**

Sample	Total Oxygenates (wt %)	
	PIONA ^A	D 6730
2 ^B	15.3	15.1
6 ^B	7.0	7.8
8 ^B	4.2	4.3
10 ^C	>8	10.5
13 ^B	20.5	20.2
14 ^B	2.8	2.9
Average	N/A	10.1

^A Multidimensional PIONA^B Major oxygenate = MTBE^C Major oxygenate = Ethanol

TABLE A1.9 Total Paraffins and Total Naphthenes

Sample	Total Paraffins (wt %)		Total Naphthenes (wt %)	
	PIONA ^A	D 6730	PIONA ^A	D 6730
8	35.6	38.0	2.2	2.7
10	41.1	45.5	5.6	6.5
13	42.6	46.0	1.3	2.1
14	34.1	41.3	5.9	9.3
Average	38.4	42.7	3.8	5.2

^A Multidimensional PIONA

APPENDIX

(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 methods.

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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.”

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