

Designation: D 5134 – 98 (Reapproved 2003)

An American National Standard

Standard Test Method for Detailed Analysis of Petroleum Naphthas through n-Nonane by Capillary Gas Chromatography¹

This standard is issued under the fixed designation D 5134; 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.

INTRODUCTION

Despite the many advances in capillary gas chromatography instrumentation and the remarkable resolution achievable, it has proven difficult to standardize a test method for the analysis of a mixture as complex as petroleum naphtha. Because of the proliferation of numerous, similar columns and the endless choices of phase thickness, column internal diameter, length, etc., as well as instrument operating parameters, many laboratories use similar *but not identical* methods for the capillary GC analysis of petroleum naphthas. Even minute differences in column polarity or column oven temperature, for example, can change resolution or elution order of components and make their identification an individual interpretive process rather than the desirable, objective application of standard retention data. To avoid this, stringent column specifications and temperature and flow conditions have been adopted in this test method to ensure consistent elution order and resolution and reproducible retention times. Strict adherence to the specified conditions is essential to the successful application of this test method.

1. Scope

- 1.1 This test method covers the determination of hydrocarbon components of petroleum naphthas as enumerated in Table 1. Components eluting after n-nonane (bp 150.8°C) are determined as a single group.
- 1.2 This test method is applicable to olefin-free (<2% olefins by liquid volume) liquid hydrocarbon mixtures including virgin naphthas, reformates, and alkylates. Olefin content can be determined by Test Method D 1319. The hydrocarbon mixture must have a 98 % point of 250°C or less as determined by Test Method D 3710.
- 1.3 Components that are present at the 0.05 mass % level or greater can be determined.
- 1.4 The values stated in SI units are to be regarded as the standard.
- 1.5 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. Specific warning statements are given in Section 7.

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2. Referenced Documents

- 2.1 ASTM Standards:
- D 1319 Test Method for Hydrocarbon Types in Liquid Petroleum Products by Fluorescent Indicator Adsorption² D 3700 Practice for Obtaining LPG Samples Using a Floating Piston Cylinder³
- D 3710 Test Method for Boiling Range Distribution of Gasoline and Gasoline Fractions by Gas Chromatography³ D 4057 Practice for Manual Sampling of Petroleum and Petroleum Products³

3. Summary of Test Method

3.1 A representative sample of the naphtha is introduced into a gas chromatograph equipped with a methyl silicone bonded phase fused silica capillary column. Helium carrier gas transports the vaporized sample through the column in which the components are separated. Components are sensed by a flame ionization detector as they elute from the column. The detector signal is processed by an electronic data acquisition system or integrating computer. Each eluting peak is identified by comparing its retention index to a table of retention indices and by visual matching with a standard chromatogram. The table of retention indices has been established by running

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 on Hydrocarbon Analysis.

² Annual Book of ASTM Standards, Vol 05.01.

³ Annual Book of ASTM Standards, Vol 05.02.

TABLE 1 Typical Retention Characteristics of Naphtha Components

Note—The abbreviations N and P refer to unidentified naphthenes and paraffins respectively.

Methane	Compound	Retention Time, min	Adjusted Retention Time, min	Kovats Retention Index @ 35°C	Linear Retention Index
Einane	Methane	3.57	0.00	100.0	
Propane					
2-Bulane	Propane	3.84	0.27	300.0	
2-Dimethylpropane	Isobutane	4.14	0.57	367.3	
Isopentane	<i>n</i> -Butane	4.39	0.82	400.0	
n-Pentane	2,2-Dimethylpropane	4.53	0.96	415.5	
2.2-Dimethylputane (Isopentane	5.33	1.76	475.0	
Cyclopentaine	<i>n</i> -Pentane	5.84	2.27	500.0	
2,3-Dimethylputane	2,2-Dimethylbutane				
### Aberty/pentane	• •				•••
Methylopentane					
### A					
2.2 Dimethylopentane	* 1				
Methycyclopentane					•••
2.4-Dimethylybuthane 11.68 8.11 630.3 Benzene 13.29 9.72 641.1 Serzene 13.29 9.72 641.1 Cyclobexane 14.19 10.62 658.3 Walthylhoxane 15.20 11.63 667.8 2.3-Dimethylopentane 15.05 11.73 669.1 1. F.Dimethylocylopentane 15.05 11.78 669.1 1. F.Dimethylocylopentane 16.81 12.81 070.2 1. F.Dimethylocylopentane 17.82 13.85 686.1 2. F. Trimethylocylopentane 17.44 13.87 686.1 2. F. Trimethylocylopentane 17.57 14.00 687.0 1. F. Trimethylocylopentane 17.57 14.00 687.0 1. J. Firmethylocylopentane 22.53 18.96 718.6 1. J. Firmethylocylopentane 22.55 18.95 718.6					•••
2.2.3 Timethylustane 12.09 8.52 635.4					•••
Benzene					
3.34 10.27 654.8					
Syciotesane					
2.4 Methylpeanane	7				
2,3-Dimethylycopentane	·				
1.1-Dimethykyclopentane					
Methylpkaxane 16.18 12.61 676.2					
cis-1,3-Dimethylocyclopentane 16.88 13.31 681.8					
Rans-1,3-Dimethyloyclopentane					
SEthylopentane					
trans-12-Dimethylcyclopentane 17.57 14.00 687.0 2.2.4-Trimethylppatane 19.43 15.86 700.0 Methylcyclopentane e					
17.80				687.0	
B-Heptane				688.7	
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3.3-Dimethylhexane	2,4-Dimethylhexane	25.47			
1.trans-2.cis-3-Trimethylocyclopentane					
1,12ms-2,cis-3-Immethylcyclopentane	All				•••
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cis-1,2-Ethylmethylcyclopentane + 2,3,5-Trimethylhexane 42.55 38.98 817.7 2,2-Dimethylheptane 43.20 39.63 822.0 cis-1,2-Dimethylcyclohexane 43.43 39.86 823.6 2,2,3-Trimethylhexane + 9N 43.76 40.19 825.8 2,4-Dimethylheptane 43.88 40.31 826.6 4,4-Dimethylheptane + 9N 44.09 40.52 828.0	· ·				
cis-1,2-Ethylmethylcyclopentane + 2,3,5-Trimethylhexane 42.55 38.98 817.7 2,2-Dimethylheptane 43.20 39.63 822.0 cis-1,2-Dimethylcyclohexane 43.43 39.86 823.6 2,2,3-Trimethylhexane + 9N 43.76 40.19 825.8 2,4-Dimethylheptane 43.88 40.31 826.6 4,4-Dimethylheptane + 9N 44.09 40.52 828.0	•				
cis-1,2-Dimethylcyclohexane 43.43 39.86 823.6 2,2,3-Trimethylhexane + 9N 43.76 40.19 825.8 2,4-Dimethylheptane 43.88 40.31 826.6 4,4-Dimethylheptane + 9N 44.09 40.52 828.0	•	42.55	38.98		
2,2,3-Trimethylhexane + 9N 43.76 40.19 825.8 2,4-Dimethylheptane 43.88 40.31 826.6 4,4-Dimethylheptane + 9N 44.09 40.52 828.0	2,2-Dimethylheptane	43.20	39.63		822.0
2,4-Dimethylheptane 43.88 40.31 826.6 4,4-Dimethylheptane + 9N 44.09 40.52 828.0	cis-1,2-Dimethylcyclohexane	43.43	39.86		823.6
4,4-Dimethylheptane + 9N 44.09 40.52 828.0					
Ethylcyclohexane + n-Propylcyclopentane 44.36 40.79 829.8					
	Ethylcyclohexane + n-Propylcyclopentane	44.36	40.79		829.8

TABLE 1 Continued

Compound	Retention Time, min	Adjusted Retention Time, min	Kovats Retention Index @ 35°C	Linear Retention Index
2-Methyl-4-Ethylhexane	44.74	41.17		832.4
2,6-Dimethylheptane + 9N	44.95	41.38		833.8
1,1,3-Trimethylcyclohexane	45.21	41.64		835.5
Unidentified C9-Naphthene	45.56	41.99		837.8
2,5-Dimethylheptane + 9P	45.92	42.35		840.3
3,5-Dimethylheptane + 3,3-Dimethylheptane + N	46.09	42.52		841.4
Unidentified C9-Naphthene	46.31	42.74		842.9
Unidentified C9-Naphthene	46.55	42.98		844.5
Ethyl Benzene	47.15	43.58		848.5
Unidentified C9-Naphthene	47.37	43.80	•••	850.0
Unidentified Naphthene + 2,3,4-Trimethylhexane	47.53	43.96		851.0
Unidentified Naphthenes	47.78	44.21		852.7
Unidentified Naphthene + Paraffin	48.13	44.56		855.1
<i>m</i> -Xylene	48.49	44.92		857.5
p-Xylene	48.63	45.06		858.4
2,3-Dimethylheptane	48.93	45.36		860.4
3,4-Dimethylheptane C + N	49.10	45.53		861.6
3,4-Dimethylheptane ^C	49.29	45.72		862.8
Unidentified Naphthene	49.41	45.84		863.6
4-Ethylheptane + N	49.65	46.08		865.2
4-Methyloctane	50.10	46.53		868.3
2-Methyloctane	50.26	46.69		869.3
Unidentified Naphthene	50.41	46.84	•••	870.3
Unidentified Naphthene	50.73	47.16		872.5
3-Ethylheptane + N	50.96	47.39		874.0
3-Methyloctane	51.15	47.58		875.3
Unidentified Naphthene	51.35	47.78		876.6
o-Xylene + 1,1,2-Trimethylcyclohexane	51.54	47.97	•••	877.9
Unidentified Naphthene + 2,4,6-Trimethylheptane	51.74	48.17		879.2
Unidentified Naphthene	52.12	48.55		881.8
Unidentified Paraffin	52.24	48.67		882.6
Unidentified Naphthenes	52.56	48.99		884.7
Unidentified Naphthene	52.85	49.28		886.7
Unidentified Naphthene + Paraffin	53.06	49.49	L)	888.1
Unidentified Naphthene	53.26	49.69		889.4
Unidentified Naphthene	53.46	49.89	•••	890.8
Unidentified Naphthene	54.02	50.45	•••	894.5
Unidentified Naphthene	54.40	50.83	•••	897.1
n-Nonane	54.84	51.27		900.0
Unidentified Naphthene	54.98	51.41		900.9

^A Extrapolated from n- C_6 and n- C_7 . See A1.1.3.

reference compounds under identical conditions or by gas chromatographic—mass spectrometric (GC/MS) analysis of reference samples under the same conditions, or both.

3.2 The mass concentration of each component is determined by area normalization with response factors. Peaks eluting after n-nonane are summed and reported as C_{10+} .

4. Significance and Use

- 4.1 A knowledge of the hydrocarbon components comprising a petroleum naphtha, reformate, or alkylate is useful in valuation of crude oils, in alkylation and reforming process control, in product quality assessment, and for regulatory purposes. Detailed hydrocarbon composition is also used as input in the mathematical modeling of refinery processes.
- 4.2 Separation of naphtha components by the procedure described in this test method can result in some peaks that represent coeluting compounds. This test method cannot attribute relative concentrations to the coelutants. In the absence of supporting information, use of the results of this test method for purposes which require such attribution is not recommended.

5. Interferences

- 5.1 If present, olefinic hydrocarbons with boiling points less than 150°C will be separated and detected along with the saturates and aromatics. Some of the olefins will coelute with saturates or aromatics and give erroneously high concentrations for those components.
- 5.2 Alcohols, ethers, and other organic compounds of similar volatility can also interfere by coeluting with saturate or aromatic hydrocarbons thereby causing erroneously high values to be determined.

6. Apparatus

6.1 Instrumentation—A gas chromatograph capable of column oven temperature programming from 35°C to 200°C in 1°C/min increments is required. A heated flash vaporizing injector designed to provide a linear sample split injection (for example, 200:1) is also required for proper sample introduction. The associated carrier gas controls must be of adequate precision to provide reproducible column flows and split ratios in order to maintain analytical integrity. A hydrogen flame

^B Extrapolated from n- C_8 and n- C_9 . See A1.2.3. Indiands/sist/327ee2c8-e4fd-4a87-bec5-7494ad35505b/astm-d5134-982003

^C Stereoisomers.

ionization detector designed for optimum response with capillary columns (with the required gas controls and electronics) must meet or exceed the following specifications:

 $\begin{array}{lll} \mbox{Operating temperature} & 100^{\circ}\mbox{C to } 300^{\circ}\mbox{C} \\ \mbox{Sensitivity} & >0.015\mbox{ C/g} \\ \mbox{Minimum detectability} & 5\times 10^{-12}\mbox{ g carbon/second} \\ \mbox{Linearity} & >10^{7} \end{array}$

- 6.2 Sample Introduction System—Manual or automatic liquid syringe sample injection to the splitting injector may be employed. Devices capable of 0.2 μ L to 1.0 μ L injections are suitable. It should be noted that inadequate splitter design or poor injection technique, or both, can result in sample fractionation. Operating conditions which preclude fractionation should be determined in accordance with Section 11.
- 6.3 *Electronic Data Acquisition System*—Any data acquisition and integration device used for quantitation of these analyses must meet or exceed these minimum requirements:
 - 6.3.1 Capacity for at least 250 peaks/analysis.
- 6.3.2 Normalized area percent calculation with response factors.
- 6.3.3 Identification of individual components by retention time.
 - 6.3.4 Noise and spike rejection capability.
 - 6.3.5 Sampling rates for fast (<1 s) peaks.
 - 6.3.6 Positive and negative sloping baseline correction.
 - 6.3.7 Peak detection sensitivity for narrow and broad peaks.
 - 6.3.8 Perpendicular drop and tangent skimming as needed.
- 6.4 Capillary Column—This test method utilizes a 50-m (0.21-mm inside diameter) fused silica capillary column⁴ with bonded (cross-linked) methyl silicone phase and a film thickness (d_f) of 0.5 µm. Other columns with these nominal dimensions may be suitable. However, all columns must meet the criteria set out in Section 10 for efficiency, resolution, and polarity.

7. Reagents and Materials atalog/standards/sist/327ee2c8

- 7.1 Carrier Gas, helium, 99.99 % pure. (Warning—Compressed gas under high pressure.)
- 7.2 Fuel Gas, hydrogen, 99.9 % pure. (Warning—Extremely flammable gas under pressure.)
- 7.3 *Make-up Gas*, helium or nitrogen, 99.99 % pure. (**Warning**—Compressed gases under higher pressure.)
- 7.4 *n-Heptane*, 99+ mol %. (**Warning**—Flammable. Harmful if inhaled.)
 - 7.5 *Methane*—(Warning—Extremely flammable gas.)
- 7.6 2-Methylheptane, 99+ mol %. (Warning—Flammable. Harmful if inhaled.)
- 7.7 4-Methylheptane, 99+ mol %. (Warning—Flammable. Harmful if inhaled.)
- 7.8 2-Methylpentane, 99+ mol %. (Warning—Extremely flammable. Harmful if inhaled.)
- 7.9 *n-Octane*, 99+ mol %. (**Warning**—Flammable. Harmful if inhaled.)
- ⁴ The sole source of supply of the columns (designated HP-PONA) known to the committee at this time is Hewlett-Packard Company, Avondale, PA. If you are aware of alternative suppliers, please provide this information to ASTM International Headquarters. Your comments will receive careful consideration at a meeting of the responsible technical committee, ¹ which you may attend.

- 7.10 *Toluene*, 99+ mol %. (**Warning**—Flammable. Vapor harmful.)
- 7.11 2,3,3-Trimethylpentane, 99+ mol %. (Warning—Extremely flammable. Harmful if inhaled.)
- 7.12 Column Evaluation Mixture, a qualitative synthetic mixture of pure liquid hydrocarbons with the following approximate composition: 0.5 % toluene, 1 % *n*-heptane, 1 % 2,3,3-trimethylpentane, 1 % 2-methylheptane, 1 % 4-methylheptane, 1 % *n*-octane in 2-methylpentane solvent.
- 7.13 *Reference Alkylate*,⁵ actual refinery alkylation product used to prepare Fig. 1. (**Warning**—Extremely flammable. Harmful if inhaled.)
- 7.14 *Reference Naphtha*,⁵ actual refinery stream used to prepare Fig. 2. (**Warning**—Extremely flammable. Harmful if inhaled.)
- 7.15 Reference Reformate,⁵ actual refinery reformer product used to prepare Fig. 3. (**Warning**—Extremely flammable. Harmful if inhaled.)

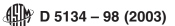
8. Sampling

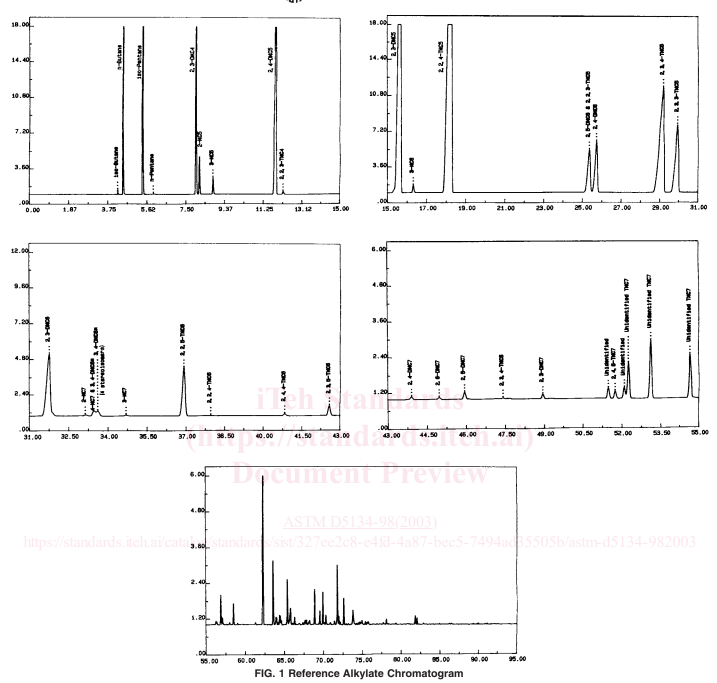
- 8.1 Hydrocarbon liquids (including naphthas) with Reid vapor pressures of 110 kPa (16 psi) or less may be sampled either into a floating piston cylinder or into an open container.
- 8.1.1 Cylinder Sampling—Refer to Test Method D 3700 for instructions on transferring a representative sample of a hydrocarbon 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 container immediately after drawing sample.
- 8.2 Preserve the sample by cooling to approximately 4°C and by maintaining that temperature until immediately prior to analysis.
- 8.3 Transfer an aliquot of the cooled sample into a precooled septum vial, then seal appropriately. Obtain the test specimen for analysis directly from the sealed septum vial, for either manual or automatic syringe injection.

9. Preparation of Apparatus

- 9.1 Install and condition column as per manufacturer's or supplier's instructions. After conditioning, attach column outlet to flame ionization detector inlet and check for leaks throughout the system. If leaks are found, tighten or replace fittings before proceeding.
- 9.2 Calibrate the gas chromatograph column oven temperature sensors using an independent, electronic temperature measuring device such as a thermocouple or platinum resistance temperature detector.
- 9.2.1 Place the independent temperature measuring probe in the oven in the region occupied by the column. Do not allow sensor to touch the walls of the oven.

⁵ These qualitative reference samples are available from Supelco, Inc., Bellefonte, PA.

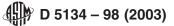


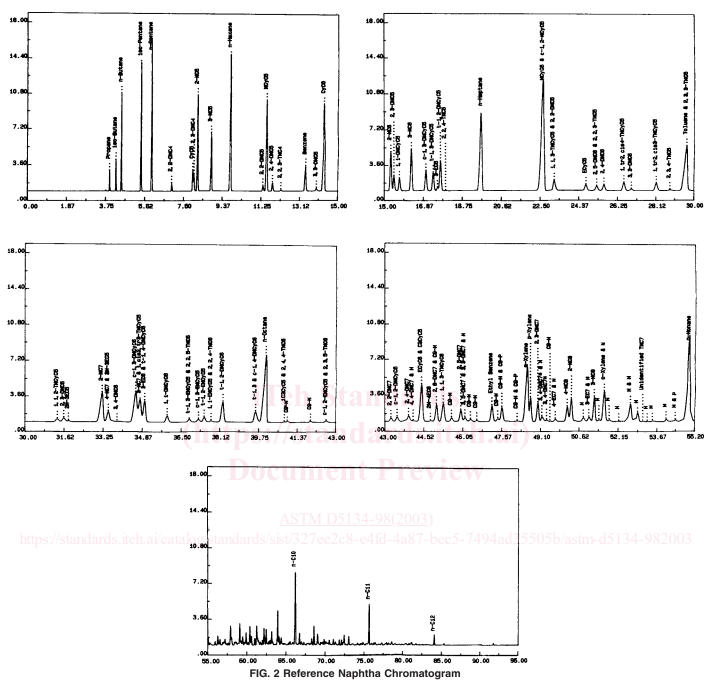


- 9.2.2 Set the oven temperature to 35°C and allow oven to equilibrate for at least 15 min, then observe the temperature reading.
- 9.2.3 If the reading of the independent temperature sensor is more than 0.5°C different from 35°C, follow manufacturer's instructions to adjust calibration of GC oven temperature.

Note 1—Differences of as little as 1° C can change the resolution of two closely eluting peaks (of dissimilar hydrocarbon types) enough to affect integration and quantitation while 2 to 3° C may cause those same peaks to be unresolved or even reverse their elution order.

- 9.3 Adjust the operating conditions of the gas chromatograph to conform to the list in Table 2. Turn on the detector, ignite flame, and allow the system to equilibrate.
- 9.4 Set carrier gas flow rate such that the retention time of toluene at 35°C is 29.6 \pm 0.2 min.
- 9.4.1 As a matter of practicality, it may be easier to first set an *approximately* correct flow rate, using methane gas injections. To do this, adjust the carrier gas flow (or column head pressure) until the retention time of methane on the 50-m column is 3.6 min.





9.4.2 Make final adjustments to flow rate so that toluene is retained for the specified 29.6 \pm 0.2 min. As this specification is critical to achieving reproducibility of retention times among different laboratories, care must be taken that the toluene does not overload the column and cause skewed peaks with resultant shifts in peak apex position. Injection of a 1 % toluene solution should preclude this possibility.

10. Column Evaluation

10.1 In order to establish that a column will perform the required separation, certain specifications must be met with respect to efficiency, resolution, and polarity. Determine the following data for new columns. Check older columns on a

periodic basis to ensure that column deterioration has not occurred. A column which does not meet these specifications is unsuitable for use.

10.2 Set oven temperature parameters for isothermal operation. Under isothermal conditions at 35°C, inject ~25 μL of methane and record the retention time. Also at 35°C, analyze the column evaluation mixture described in 7.12. Record the retention times and the peak widths at half height of each of the components.

10.2.1 Calculate efficiency of the column using Eq 1. The number of theoretical plates (n) must be greater than 225 000.

$$n = 5.545 \left(t_{\rm R} / W_{\rm h} \right)^2 \tag{1}$$