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E-190

## NACA MR Hos. E5F20, E5I12

#### MATIONAL ADVISORY COMMITTEE FOR AEROFAUTICS

MEMORANDUM REPORT

for the

Air Technical Service Command, Army Air Forces

SYOKING CHARACTELISTICS OF VARIOUS FUELS AS DETERMINED

BY OPEN-CUP AND LABORATORY-BURNER SMOKE TESTS

By Earl R. Ebersole and Henry C. Barnett

#### ILTRODUCTION |

At the request of the Air Technical Service Command, Army Air Forces, the Cleveland laboratory of the MACA is conducting tests in an I-15 combustion chamber to obtain performance data on a variety of hydrocarbon fuels.

Inasmuch as the smoking tendency of jet-propulsion fuels is of interest, a laboratory program has been conducted to obtain data on the smoking behavior of fuels. The results of open-cub smoke tests of 25 hydrocarbon fuels and of two series of prepared blends are presented. In order to determine the effect of fuel-air ratio as well as hydrocarbon class on the smoking behavior of fuels, 21 hydrocarbons were tested in a laboratory burner at fuel-air ratios ranging from 0.062 to 0.119. The results of these controlled-burning smoke tests are also presented. Correlation with data from full-scale jetprovulsion tests will be necessary to determine the applicability of the results.

#### APPARATUS AND TES" PROCEDURE

<u>Open-cup tests</u>. - The apparatus used in the open-cup tests is shown schematically in figure 1. A 3-milliliter sample of the fuel to be tested was placed in the open cup and ignited with an open flame. Snoke passing up the chimney absorbed a portion of the light beam from the light source. The effect of this light absorption on the photoelectric cell was registered by the recording microammeter. Before each test, the recording microammeter was adjusted to zero, if necessary, by varying the resistance in the lamp circuit. A check run was made on each fuel. The readings in microamperes were converted to vercentage light absorbed by means of the calibration curve shown in figure 2. In the calibration of the recording instrument the light beam was interrupted with screens of varying mesh. The transmission through these screens was measured in a standard spectrophotometer using light with a wavelength of approximately 6000 A. The percentage light absorbed was then plotted against microamperb readings to obtain the calibration curve.

Each of 25 hydrocarbon fuels was burned in an open cup in order to determine snoking characteristics. In each test the burning rate of the fuel was determined by measuring the time required to burn a 3-milliliter sample.

The fuels tested included representative hydrocarbons of the paraffinic, cycloparaffinic, olefinic, and aromatic classes as well as mixed fuels. Kerosene and the same kerosene from which aromatics had been removed were included in the mixed fuels tested. Two series of binary blends consisting of toluene in hydrogenated triisobutylene and triisobutylene in hydrogenated triisobutylene were tested over the range of concentrations from 0 to 100 percent.

Laboratory-burner tests. - The burner and the auxiliary apparatus used in the laboratory-burner tests are shown schematically in figure 3. Primary air was passed through the preheater at a constant flow rate of  $4.1\pm0.1$  liters per minute. Secondary air was introduced at the base of the burner at a constant flow rate of  $10.4\pm0.2$  liters per minute. The fuel to be tested was placed in a graduated burette, which was modified to give low and constant flow rates. (See reference 1.) The fuel was then admitted to the primary air stream and was vaperized and mixed with the primary air in the preheater at a temperature of  $180^{\circ}$  to  $200^{\circ}$  F above the boiling point of the hydrocarbon. After the fuel was ignited through the small hole in the side of the chimney, the hole was scaled off by means of a slip ring.

The flow rate of the air was maintained at  $14.5 \pm 0.3$  liters per minute for all fuel-air ratios. The fuel-air ratio was varied by changing the flow rate of the fuel from the graduated burette. The rate of fuel flow was measured by determining to the nearest 0.01 minute the time required for a given volume (0.825 ml) to flow from the burette.

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Each of 21 fuels including representative hydrocarbons of paraffinic, cycloparaffinic, olefinic, aromatic, and dicyclic classes was tested for smoking tendency at various fuel-air ratios. Smoke particle size was not measured in these tests.

#### DISCUSSION OF RESULTS

<u>Open-cup tests</u>. - The results of the open-cup smoke tests are presented in table I. Smoke is reported as percentage light absorbed. Total smoke from a 3-milliliter sample was determined by measuring the area under the curve traced by the recording microammeter. Values of average smoke were computed by dividing the total smoke by the burning time. Values of peak smoke were obtained by visually averaging the deflections in the highest portion of the curve.

The data in table I show that, with the exception of the cycloparaffins, the fuels of a given hydrocarbon class gave readings within a range of 8 percent light absorbed and that the argumatics gave higher readings than the other classes. The greater smoking tordency of the argumatics can be further shown by the tests on kerosene and dearonatized kerosene. Kerosene gave a peak value of 21 percent light absorbed as compared with 5 percent for dearomatized kerosene.

In order to ascertain the effect of burning rate on the smoking tendency, average smoke (percentage light absorbed) was plotted against burning rate (grams per min) in figure 4. Both test run and check run for each fuel are plotted. The five-carbon ring cycloparaffins are not shown in this figure because data were obtained for only two commounds in this series. It can be seen in table I that considerable difference in smoking tendency was found between the five-carbon ring and the six-carbon ring cycloparaffins. The fuels tested (fig. 4) show that the type of hydrocarbon exerts a much greater influence on the smoking tendency than does the burning rate of the fuel.

Figure 5 shows the variation of the burning rate of the fuels with boiling point. As in figure 4 both test and check runs are plotted. For the fuels tested, the burning rate of hydrocarbons within a given class decreased with increasing boiling point. It is of interest that two fuels of different classes with different boiling points will burn at the same rate; for example, from the curves of figure 5, a paraffinic hydrocarbon with a boiling point of 176° F will burn at approximately the same rate as an aromatic hydrocarbon with a boiling point of  $305^{\circ}$  F.

In order to show the relation of smoking tendency and composition for fucls having equal boiling points, tests were made on blends of triisobutylene in hydrogenated triisobutylene. A linear relation was obtained. (See fig. 6.) Figure 7 shows the results of tests of two fuels (toluene and hydrogenated triisobutylene) having different boiling points.

Laboratory-burner tests. - The results of the laboratoryburner smoke tests are presented in table II together with the boiling point end the stoichiometric fuel-air ratio of each fuel tested. At the maximum fuel-air ratios that would support combustion in the burner no smoke was obtained for parafrinic, cycloparaffinic, or olefinic hydrocarbons with the exception of triisobutylene, which gave datectable smoke at a fuel-air ratio of 0.102.

Aromatic and dicyclic hydrocarbons began smoking at approximately their stoichiometric fuel-air ratios. Figures 8 and 9 show the variation of smoke (percentage light absorbed) with fuel-air ratio for aromatic and dicyclic hydrocarbons, respectively. Korosene is also included in figure 9. Within experimental error the smoking tendency appears to be a linear function of the fuel-air ratio for aromatic and dicyclic hydrocarbons. Comparison of figures 5 and 9 indicates that the smoking tendency of neuhthalenic hydrocarbons is greater than that of aromatic hydrocarbons.

## SUMMARY OF RESULTS

The data obtained from an investigation of the smoking characteristics of 25 hydrocarbons indicate that for uncortrolled burning in an open cup:

1. The smoking tendency of a hydrocarbon fuel is more dependent upon the type of hydrocarbon than upon its boiling point or burning rate.

2. The burning rate of hydrocarbons within a given class tended to decrease with increasing boiling point.

3. The smoking tendency of a commercial kerosene was about four times that of the same kerosene from which aromatics had been removed. Results obtained from controlled-burning smoke tests indicate that:

1. The smoking tendency of hydrocarbon fuels is dependent upon both the class of hydrocarbon and the fuel-air ratio.

2. Within experimental error the smoking tendency of aromatic and dicyclic hydrocarbons is a linear function of the fuel-air ratio.

The following table summarizes the data obtained:

Class	Boiling point (°F)	Hydrogen- carbon retio	Foak smoke (porcont light absorbed)	Burning rate (grams/min)
Paraffins	122-230	0.189-0.196	10-15	0.69-1.07
Six-carbon ring cyclopareffins	175-2.9	.1.7	8-13	.6285
Five-carbon ring cycloparaffirs	122-151	.167	2331	.97-1.13
Olofins	101-172	.167	40-47	.97-1.33
Arom: tics	230-323	.095111	6 <b>8</b> -75	.75-1.36

OPEN-CUP SMOLT TESTS

#### LABORATORY-BURGER SMOKE CESTS

Hydrocarbon class	Bciling- point range (°F)	Stoichiometric fuel-air ratio	Fuel-air-ratio range	Smoke (norcent light absorbed)
Paraffins	175-350	0.035-0.066	0.064-0.114	0
Cyclohexanes	178-270	.067	.073115	0
Olefins	133-350	.067	.062109	0-3
Aromatics	175-465	.070075	.063108	0-47
Dicyclics	365-605	.069077	.069119	3-80

Aircraft Engine Research Laboratory,

National Advisory Committue for Aeronautics,

Cleveland, Ohio, June 20, 1045; Sept. 12, 1945.

## REFERENCE

 Zentner, E. Thomas: Delivory of Liquids at Low and Constant Ratos. Ind. and Eng. Chem. (Anal. ed.), vol. 16, no. 7, July 21, 1944 pp. 471-472.

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## TABLE I. - RELATIVE SNOKING CHARACTERISTICS OF

VARIOUS FUELS BURNED IN OPEN CUP

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Class of hydro- carbon	Compound	Peak smoke observed (percent light absorbed)	Average smoke (percent light absorbed)	Total smoke (area in sq in.)	Rate of burning (grams/ min)	Boiling point ( <sup>O</sup> F)	Hydrogen- carbon ratio
Paraffins	2,2-Dimethyl-	11	10	1.4	1.052	122	0.196
	2,3-Dimethyl-	11	11	2.0	.941	137	.196
	2,2,3-Tri-	15	10	2.2	.805	178	.192
	2,3-Dimethyl-	15	11 7	2.4	.772	193	.192
	pentane 2,2,3-Tri- methylpentane	10 13 11	6 10 8	1.4 2.5 2.0	.723 .679 .688	230	.189
Cyclo-	Cyclopentane	26	20	3.6	1.125	122	0.167
pererring	Nethylcyclo-	31 31	26 24	5.4	.965	161	.167
	Cyclohexane	8	8	1.8	.825	178	.167
	Methyl- cyclohexane	11 13	8 10	2.0	.755	212	.167
	Ethyl- cyclohexane	11 13	9 8	2.8 2.3	.617 .517	269	.167
Olefins	Trimethyl-	42	37 37	5.8	1.325	101	0.167
	2,3-Dimethyl-	44	37	6.4	1.223	133	.167
	2,3-Dimethyl-	40	33	6.9	1.026	164	.167
	Triptene	42 46 47	38 38 38	8.2 8.2	.968 1.063 1.063	172	.167
Cyclo- olefins	Methyl- cyclohexene	50 47	41 37	8.0 8.1	1.323 1.142	216	0.143
Aromatics	Toluene	71	50 49	10.7	1.363	230	0.095
	Xylene	68	52	18.3	.844	279 <b>-28</b> 8	.104
	1somers <u>n</u> -Propyl-	75 70	57 52	19.1	.893	320	.111
	benzene m-Ethyl-	70 70	54 51	19.7	.816 .763	323	.111
	toluene	72	55	22.7	.745		
Mixed	Virgin	21			0.698	115-298	
TAATS	Hot-acid	13			.652	174-257	
	Triiso-	35			******	340-350	
	Hydrogenated triiso-	16				335-350	
	butylene <sup>#</sup> Kerosene Kerosene (dearo- matized)	21 5			 NAT 1	302-486	ISORY

<sup>a</sup>Listed as mixed fuels because their purity is questionable.

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Class of hydrocarbon	Compound	Boiling point (°F)	Stoichi- ometric fuel-air ratio	Fuel-air- ratio range	Hydrogen- carbon ratio	Smoke (percent light absorbed)
Paraffins	Triptane	178	0.065	0.067-0.103	0.191	0
	Diisopropyl	137	.065	.064071	.195	0
	Neohexane	122	<b>- 065</b>	.070085	.195	0
	2,2,3-Trimethyl- pentane	230	•066	.074083	.188	0
	Hydrogenated triisobut <b>y</b> lene	335-350	•066	.083114	.181	0
Craloberanes	Croloberane	179	067	073- 093	167	
UJCI OMERANICO	Ethylcyclo- hexane	269	.067	.073115	.167	õ
Olefins	2,3-Dimethyl-	133	.067	.080096	.167	o
	2,3,3-Trimethyl-	172	.067	.062090	.167	0
	Triisobutylene	340-350	.067	.092109	.167	0-3
Aromatics	Benzene	176	.075	.063104	.083	0-37
	<b>Bthylbenzene</b>	277	.073	.076100	.104	3-34
	Isopropylbenzene	306	.072	.063100	.111	0-32
	1,2,4-Trimethyl-	337	.072	.076108	••••	3-47
	Triisopropyl- benzene	465	.070	.073107	.133	5-44
Dicyclics	1-Methyl- naphthalene	473	.077	.073114	.076	3-80
	Dimethyl- naphthalene	511	.075	.078102	.083	11-59
	Monoamyl- naphthalene	535 <b>-6</b> 08	.073	.069097	.101	9-50
	Tetralin	405	.073	.076108	.101	7-61
	Decalin	365-882	.069	.084119	.150	3-18
Mixed fuels	Kerosene	302-486		.060108		0-5

## TABLE I. - SMOKING CHARACTERISTICS OF HYDROCARBON FUELS

DETERMINED FROM CONTROLLED-BURNING TESTS

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Figure 2. - Calibration curve for photocell.

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Figure 3. - Schematic diagram of burner and auxiliary apparatus.



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Figure 8. - Variation of smoke with fuel-air ratio for aromatic hydrocarbons.

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Figure 9. - Variation of smoke with fuel-air ratio for dicyclic hydrocarbons and kerosene.

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