MICROCOPY RESOLUTION TEST CHART
NATIONAL BUREAU OF STANDARDS 1961-A
Antimisting Kerosene: Evaluation of Improved FM-9 Additive

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June 1986

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Optimization of FM-9 dissolution rate to attain both acceptable mist suppression and degradation properties within 15 to 20 minutes of in-line blending was carried out by Imperial Chemical Industries (ICI). This report discusses the evaluation of the FM-9 with improved dissolution rate and consequently better degradability characteristics for compatibility with engine filters and fuel control system. This additive was identified as a candidate additive to be used in a Control Impact Demonstration (CID) test. The major findings of the investigations are:

1) Quality of the FM-9 slurry meets the metering and dispersion requirements for single stage in-line blending.  
2) The dissolution rate at ambient fuel temperatures (15-20°C) is sufficient, and produces AMK fuel with adequate fire suppression in 30 minutes after blending.  
3) Freshly blended fuel can be degraded without difficulty with a specific power requirement of less than 30 kWsL⁻¹.  
4) Contamination with large amounts of bulk water leads to formation of gelled emulsion on the interface. The water vapor condensation on AMK surfaces produced a string-like second phase.  
5) Low temperature evaluation indicated no phase separation and gel formation problems.  
6) Significant loss in pumpability performance with FM-9 AMK was observed at both room and low temperature using the Cessna 441 boost pump. However, the deterioration of pumpability performance was not as significant with AMK as compared with Jet A using the DC-10 boost pump.

**Key Words**

Aircraft Fires, Aircraft Safety, Safety Fuels, Antimisting Fuel

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Acknowledgements

This work presents the results of one phase of research carried out at Jet Propulsion Laboratory, California Institute of Technology, Contract NAS7-918 Task Order RE152, Amendment 298, sponsored by Department of Transportation/Federal Aviation Administration Technical Center, Atlantic City Airport, New Jersey, under Agreement No. DTFA03-8000215. The authors extend their gratitude to Messrs Bruce Fenton, G. Klueg, and W.T. Westfield for many valuable technical suggestions throughout this program. We are also grateful to Messrs Wayne Bixler, Stan Kikkert, and R. Smither for their assistance in design, fabrication, assembly, and acquisition of the experimental data.
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Executive Summary

During the past few years, studies by the Federal Aviation Administration (FAA) and other government agencies have shown that the hazards from aircraft crash fires might be significantly decreased if an antimisting kerosene (AMK) fuel could be utilized. The addition of polymeric additive at low concentrations to jet fuels is known to suppress mist formation and ignition of the fuels under circumstances often encountered in survivable aircraft crash landings. An antimisting additive, FM-9TM has been developed by Imperial Chemical Industries (ICI) and is available under the trade name AVGARD. This material when dissolved in jet fuels imparts a strong time-dependent threshold type shear-thickening behavior. In case of fuel spillage from a ruptured fuel tank during an aircraft crash, the fuel misting is prevented. Simulated aircraft crash landing fuel spillage tests have indicated that fuel misting can be sufficiently suppressed, and the ignition and the subsequent fireball formation can be greatly reduced or eliminated.

Optimization of FM-9 dissolution rate to attain acceptable mist suppression and degradation properties within 15 to 20 minutes of inline blending was carried out by ICI. This report discusses the evaluation of FM-9 variant which has better dissolution rate and consequently better degradability and compatibility with engine filters and fuel control system and was identified as a candidate additive to be used in a Control Impact Demonstration (CID) test.
1.0 INTRODUCTION

During the past few years, studies by the Federal Aviation Administration (FAA) and other government agencies have shown that the hazards from aircraft crash fires might be significantly decreased if an antimisting kerosene (AMK) fuel could be utilized (Reference 1). The addition of polymeric additive at low concentrations to jet fuels is known to suppress mist formation and ignition of the fuels under circumstances often encountered in survivable aircraft crash landings. An antimisting additive, FM-9M has been developed by Imperial Chemical Industries (ICI) and is available under the trade name AVGARD. This material when dissolved in jet fuels imparts a strong, time-dependent, threshold type, shear-thickening behavior. In case of fuel spillage from a ruptured fuel tank during an aircraft crash, the fuel misting is prevented. Simulated aircraft crash wing fuel spillage tests and large scale ground-to-ground crash tests have indicated that fuel misting can be sufficiently suppressed, and the ignition and the subsequent fireball formation can be greatly reduced or eliminated.

Studies have indicated that the optimum method of making antimisting fuel is by a single-stage blending at the aircraft fueling point. Results reported in References 2, 3 and 4 have shown that the FM-9 additive, dispersed in a carrier fluid, could be blended into aviation kerosene to give a fuel which has adequate fire resistance 15 to 20 minutes after blending.

Work on optimization of FM-9 dissolution rate was carried out by ICI and led to the development of several FM-9 variants with improved dissolution characteristics. This report discusses the evaluation of FM-9 variant which has a better dissolution rate and, consequently, better degradability and compatibility with engine filters and the fuel control system. This additive is the latest development in the FM-9 variants formulations and was used in the Controlled Impact Demonstration (CID). The period of performance for the work reported herein was from August, 1983 thru November, 1984.

2.0 MATERIALS, EXPERIMENTAL PROCEDURES AND AMK CHARACTERIZATION TESTS

2.1 Materials

The antimisting additive FM-9 used in this program is a proprietary fuel additive developed by ICI. The FM-9 is a high molecular weight polymer with specifically designed properties for use with jet fuels. The additive is in the form of a free-flowing powder which is formulated with carrier fluids into a dispersion called AMK slurry. This slurry is available from ICI Americas, Inc.

In 1983, JPL evaluated approximately 40 batches of FM-9 variants, including four batches of additives which were used on the large-scale wing-spillage test facility at the FAA Technical Center, Atlantic City, N.J. The results of this evaluation were reported separately (Reference 9). This FM-9 variant additive was prepared and formulated by ICI as a 25-percent polymer solid loading in the carrier fluid. Only the results from the evaluation of this formulation are presented.

Lots JCK 14-247-1 and JCK 16-95-1 were extensively evaluated. These slurries were prepared by ICI Americas Inc. in Wilmington, Delaware.
The base fuel used in this program was Jet A aviation kerosene that came mainly from two sources: ICI, as lot RMH 30328 in 55-gallon drums; and Texaco/Martin Aviation Terminal, Burbank Airport, California. The fuel was stored as received in 55-gallon drums. Most of the tests were done using the Texaco Jet A and lot RMH 30328 was used as a control. The Texaco Jet A fuel water content stabilized in the 70 to 80 ppm range. Some of the properties of these base fuels (as received) relevant to AMK are presented in Table 1.

2.2 Experimental Procedure and AMK Characterization

2.2.1 AMK Blending Assembly and Procedure

The in-line blending setup which was used to produce AMK is presented in figure 1.

![Figure 1. IN-LINE BLENDING APPARATUS](image)

The blending system consists of a slurry injection port, a pump, and the mixing element (static mixer). The entire system was made from off-the-shelf components with the exception of the fuel tanks. The injection port was part of the B-D Luer-Lock automatic syringe refill kit. The pump drive module was a high flow rate, explosion-proof unit, Model RP-F, manufactured by (FMI) Fluid Metering Inc., Oyster Bay, N.Y. The RP-F unit employs a 1/4-HP motor with model RP-F-2 pump head module. The pump head was made of 316 stainless steel with sintered carbon for cylinder liner material. The pump has a maximum flow rate of 16 gph and a maximum pressure rating of 100 psi. The pump has a simplified positive displacement mechanism based on a valveless pumping mode and was recommended for handling semi-solid fluids and heavy slurries. The main component of the blending system consists of a Static Mixer® manufactured by the Kenics Corp. The device is simply a straight 1/4-inch stainless steel tube, 9 inches long with a series of fixed, helical elements enclosed within the tubular housing. The helical design of the central element causes a transverse flow to arise normal to the pipe axis. As a consequence, fluid near the center of the pipe is rotated out toward the circular boundary, and vice versa. Radial mixing and multiple flow separation was achieved in this manner. The unit is an in-line mixer having no moving parts and no external power requirements; in addition, the unit is amenable to quick changes, has low cost of operation, and hardly requires any maintenance. The components of the in-line blending system were assembled using flexible PVC tubing which gives some see-through capabilities to the system.
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<td>392</td>
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<td>469</td>
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<td>95%</td>
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<td>492</td>
<td>483</td>
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<tr>
<td>E.P.</td>
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<td>518</td>
<td>512</td>
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<td>REC.%</td>
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<td>98.5</td>
<td>98.5</td>
<td>98.5</td>
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<td>98.5</td>
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<tr>
<td>RES.%</td>
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<td>1.5</td>
<td>1.5</td>
<td>1.5</td>
<td>1.5</td>
<td>1.5</td>
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<tr>
<td>LOSS.%</td>
<td>-0-</td>
<td>-0-</td>
<td>-0-</td>
<td>-0-</td>
<td>-0-</td>
<td>-0-</td>
</tr>
<tr>
<td>Freezing Point, °C, (D-2386)</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-43</td>
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<tr>
<td>Viscosity cSt @ -20°C (D-445)</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>5.92</td>
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<tr>
<td>Flash Point, °F, (D-56)</td>
<td>120</td>
<td>116</td>
<td>-</td>
<td>125</td>
<td>122</td>
<td>135</td>
</tr>
</tbody>
</table>

*Water Separation Index, Modified (ASTM D-2550)
In brief, the AMK blending operation consisted of placing the appropriate weight of slurry in a 50 ml B-D Plastipak™ Luer-Lok tip disposable syringe and then locking the syringe into the injection port. Care was taken that the slurry did not make contact with the fuel, since wetting of the slurry with jet fuel at this stage causes premature swelling of the slurry at the wetted surface, and result in the formation of transparent gel which makes the subsequent dispersion of the polymer particles very hard. With valves #1, #2, and #3 closed, half the required amount of jet fuel was placed in the base fuel tank and the other half was placed in the AMK tank. In a typical run 1.5 kg of Jet A was used in the base fuel tank, 36.0 gm of slurry was used in the syringe and 1.5 kg of Jet A was placed in the AMK tank (1 gallon polyethylene bottle). After the pump was turned on, valve #1 is opened. With the opening of the valve, the slurry from the syringe was carefully injected in the fuel line.

The AMK was collected in the tank and allowed to equilibrate for the desired amount of time. The AMK tank was gently stirred for 15 to 20 seconds at the start to allow mixing of the Jet A fuel. It should be noted that the end of the blending was always considered the start of the polymer equilibration process.

After each batch, the system was cleaned by circulating jet fuel through the system with valve #1 closed and valves #2 and #3 opened. In addition to the small-scale (one liter/minute) blender, some of the batches were tested for their dissolution properties using JPL's 5-10 gpm blender. This blender was similar to the one designed and built at JPL for blending large amounts of AMK for evaluation of the FM-9 variants at the FAA Technical Center, Atlantic City, N.J. A detail description of this blender can be found in reference 4.

The AMK blending was monitored by fire test, filter ratio, cup test, and by following the turbidity of the fuel with time. For the large scale runs the solid content of the fuel was also measured.

2.2.2 Screen Filter Ratio Test and Orifice Flow Cup Test (ICI Cup Test)

A filter ratio device (standardized by the United States/United Kingdom AMK Technical Committee) was utilized as the primary method of measuring viscosity properties. The details of this test are given in Appendix B and the description of the filter ratio device is given in Appendix C.

2.2.3 Flammability Comparison Test Apparatus (FCTA) and JPL's Mini Wing Shear Fire Test

The FCTA, shown schematically in figure 2, is described in detail in References 5 and 6. Air is released from a pressure vessel through a sonic orifice into a straight tube, where it atomizes a small jet of fuel. The spray issues through a conical diffuser into ambient air and is ignited by a propane torch. The fuel is delivered by a single stroke displacement pump, and issues through an upstream facing elbow with an inside diameter of 0.52 cm. The inside diameter of the straight mixing tube is 2.66 cm. The air mass flow is controlled by varying the air pressure and the fuel mass flow is controlled by a constant speed actuator that regulates the fuel pump. Once the air pressure and speed control are set by the operator, the operation of the apparatus is controlled by an automatic sequencing switch. Appendix E describes the JPL operating procedure for FCTA test.
The primary method to test the development of freshly blended AMK was done by the mini wing shear fire test. For the test, a measured amount (one gallon) of fuel is released from a two-inch (I.D.) pipe in front of a two-inch (I.D.) cylinder (flame holder) in an airstream produced by an open-jet wind tunnel.

![Diagram of the flammability comparison test apparatus](image)

Figure 2. SCHEMATIC DIAGRAM OF THE FLAMMABILITY COMPARISON TEST APPARATUS

An oxyacetylene torch is used as an ignition source located two inches downstream of the cylinder. The airstream velocities used for the fire test were 120 (61.7 m/s), 130 (66.8 m/s), and 140 (72.0 m/s) knots. This velocity was measured with a pitot tube located upstream of the nozzle exit plane. The flammability of the freshly blended fuel was compared to the flammability of ICI-prepared equilibrated AMK. It was assumed that the ICI prepared fuel will pass the FAA's large-scale wing spillage fire test. The length of the flame for the samples was visually observed to determine a rating of "pass," "fail," or "marginal." To follow the development of freshly blended AMK, one gallon samples of the fuel were tested for fire protection at various times after blending, and the time at which the fuel received a "pass" rating at 130 knots was also recorded. An additive batch, with an acceptable dissolution rate will get a "pass" fire test rating within 15 to 20 minutes after blending. It should be pointed out that this was one of the criteria for the evaluation of the antimisting additive dissolution rate.

2.2.4 Sample Degradation

The degradation of the samples was done in a blender with a 5-cup [1.25 liter] container. The sample size was always kept the same (300 ml) and samples were degraded for 30 seconds at 22°C at the highest speed (liquefy). The degraded samples were characterized by filter ratio tests and were always done within one minute after the sample was degraded. It is very important that the time after degradation at which the samples were characterized is kept always the same (one minute), especially for freshly blended samples where the additive in some cases is not fully equilibrated. In these cases, the undissolved polymer is not degraded during the 30-second degradation period and continues to dissolve.
If the FR test is done past the one-minute period, very high filter ratios can be obtained. The results of this test are presented as FRₜ where t is the time in minutes after blending, the degradation was performed. The equilibrated AMK fuel gives FRₜ values of 3-4 under these conditions. Based on this value, if FRₜ is less than 5, the AMK fuel is considered to have good degradability characteristics (and dissolution); if more than 10, it is poor; and between 5-10, is marginally degradable.

The degradability of equilibrated and freshly blended AMK fuel was evaluated also by degrading the samples using a continuous-flow single pass degrader which utilized a pressure drop across a needle valve. The schematic of the apparatus is included in Figure 3. The degrader operated at 4000 psi pressure drop and the degradability was evaluated in terms of the filterability of the degraded fuel. The filterability of the sample was monitored for at least 30 minutes and up to one hour. AMK fuel has an acceptable degradability if 20-30 minutes after blending no filter plugging is observed with 4000 psi pressure drop across the needle valve and a 1 gpm/in² volume flux through the filter (325 mesh (40 μm)) stainless steel screen) at 20°C inlet fuel temperature.

As in the blender degradation discussed above, one should be careful during characterization and interpretation of the FR data. It should be pointed out that in the degrader apparatus (Figure 3), as in the engine fuel system, the fuel was passed immediately through the filters. If one does the degradation and the filtration separately with some time in between, the partially equilibrated fuel may give rise to filtration difficulties.

For degradation of the equilibrated AMK fuel, JPL’s inline degrader-filtration apparatus (Figure 3) gave a FR values in the 1.2 to 1.08 range. Detailed description of the apparatus and the degradation procedure can be found in Reference 7.

Partial degradation of AMK fuel and subsequent characterization by FR test was used also to evaluate the unintentional degradability of the fuel. The degradability (as measured by FR) of ICI prepared equilibrated AMK fuel was used as a baseline control. The partial degradation to simulate unintentional degradation was done by pumping the fuel in one or more passes through the Kenics (K) Mixer using the in-line blending apparatus. Equilibrated FM-9 AMK fuel after one pass-through the static mixer gives an FR of 13 to 15. Such measurements on non-equilibrated fuel were not performed during this study.

2.2.5 Turbidity

The measurements of turbidity were done with model DRT-100 Turbidimeter manufactured by H.F. Instruments. The DRT-100 Turbidimeter is a continuous reading nephelometer which measures reflected light from scattered particles in suspension and direct light passing through a liquid. The resulting ratioed optical signal is stabilized and amplified to energize a meter. The instrument provides a linear readout of turbidity in nephelometric turbidity units (NTU).

2.2.6 Water Reaction Test

Visual observation of the interaction of water with AMK fuel showed strings which forms when water vapor is condensed on a cold fuel surface. This was done in a one liter "Pyrex," heavy wall, filtering flask. AMK fuel (400 cc)
was placed in the stoppered flask and the head space evacuated to about 3 inches Hg and sealed. The flask was then immersed halfway in CO₂/acetone bath at -30°C. After the temperature of the fuel reached -20°C, the flask was taken out from the bath and ambient air was allowed to enter the flask until ambient pressure was reached. This process took approximately 20 seconds. The fuel was gently swirled and then allowed to rest. Visual observations were then made of string formation due to polymer/water reaction; their relative amounts and lengths were noted. At these conditions, equilibrated (ICI) FM-9 AMK fuel will form small amounts of strings, and its behavior at these conditions was used as a control.

2.2.7 Low Temperature Gel Formation and Pumpability Test

The low temperature gel formation test was done in the apparatus described for the water reaction test. The AMK fuel was placed in the flask, the head space was inerted with dry nitrogen gas, closed, and then placed in CO₂/acetone bath at -30°C. After the fuel temperature reached -25°C, it was stirred using a magnetic stirrer. Stirring and cooling of the fuel continued for 10 minutes. The flask was then opened and fuel poured as fast as possible through a four-mesh stainless steel screen. The presence of transparent gel on the top of the screen, the relative amount of the gel and its behavior with time (warming) were visually noted. The test is a "pass" or "fail" depending on the collection of gel on top of the screen. The ICI equilibrated FM-9 AMK fuel under these conditions does not give any gel.

The impact of gel formation after exposure of the fuel to subzero temperatures was characterized by flammability (fire test). This was done as described in Section 2.2.3 using one gallon of fuel which has been cooled down to -25°C.

The low temperature pumpability performance of the AMK fuel was evaluated and compared with Jet A performance mainly by determining the pumping efficiency. This was done in the JPL low temperature facility shown in Figure 4 which consists of an Airborne IC12-17 (Cessna 441) centrifugal fuel boost pump mounted at the bottom of a jacketed 10-gallon fuel tank, equipped with hand stirrer, inlet and outlet for nitrogen gas, and a thermometer. The schematic of the apparatus is shown in Figure 5. The fuel was placed in the tank and the air above the fuel replaced with nitrogen gas in order to prevent moisture condensation at low temperatures. Using an acetone/dry ice mixture in the jacket and slow stirring, the fuel temperature was lowered to the desired temperature (-25°C to -30°C). The efficiency of the boost pump was determined by measurements of the flow rate, pressure rise (ΔP), and the input electric power to the pump. The pumping efficiency is defined as

\[ \eta = \frac{Q \Delta P}{VI} \times \text{Conversion Factor} \]

where

- \( Q \) = Volume flow rate;
- \( V \) = Input voltage
- \( I \) = Input current
- \( \Delta P \) = Pressure differential

The Airborne IC12-17 pump specifications for Jet A are: 15 psi at 4.4 GPM and 19 psi minimum at 2.4 GPM. The actual measurement gave: 15 psi at 5.2 GPM and 19 psi at 2.4 GPM. Figure 6 presents the pumpability criterion employed to evaluate the performance of AMK fuels. The following figures of merit were used:
Figure 4. LOW TEMPERATURE PUMPING FACILITY
Figure 5. SCHEMATIC OF AXI AL PUMPABILITY TEST APPARATUS
A. Maximum flow rate in GPM delivered by the pump at 10 psi.

B. Decrease in delivery pressure in psi associated with an incremental increase in flow rate (see the slope in Figure 6).

In addition to the above test the low temperature behavior of AMK was characterized using JPL's aircraft wing tank environmental simulator (Figure 7). The test tank size is 50 gallons and represents a cross-section of an aircraft outer wing tank. The tank is equipped with heat exchangers on the top and bottom wall and can operate at temperatures from -55 to +40°C. The procedure for conducting the test involved loading the tank with fuel and then by controlling the temperature on the upper and lower walls; to lower the fuel bulk temperature to the desired level. The fuel was held at the temperature for a predetermined time and then gravity discharged from the tank to determine the fraction of holdup or frozen, unpumpable fuel. The percent holdup at various temperatures for AMK was compared with the data obtained for Jet A fuel (used to blend AMK) under the same condition. Detailed description of the apparatus and the testing procedures can be found in Reference 8. Because of the complexity, the test was performed only for two batches of slurry.

Figure 6. PUMPABILITY CRITERION FOR AMK FUEL
2.2.8 Slurry Preparation and Characterization

The slurry batches were prepared by ICI as a 25 percent by weight FM-9 polymer powder in a carrier fluid. The size of the shipments varied from 5 to 40 lb and were received in 5-gallon plastic containers. The slurries were used at least one week after arrival at JPL. The slurries were first homogenized by using a stirrer with an overhead motor and then passed through a "Tyler" equivalent 14-mesh sieve to remove any large agglomerates or foreign matter. The large batches of slurry were then divided and stored in one-gallon plastic containers. Prior to blending, the slurry was again homogenized by tumbling the containers for several hours on a rotary tumbler.

During the course of this program, a simple test procedure to determine the presence of large polymer particle agglomerates was developed. Description of this procedure is presented in Appendix F. Presence of such agglomerates in the slurry results in blending problems such as polymer settling, formation of gel, and plugging of the fuel lines.

3.0 EXPERIMENTAL RESULTS AND DISCUSSIONS

The objective of this investigation was to characterize the improved FM-9 AMK additive using ICI prepared equilibrated AMK (see Appendix A) as the baseline control. Preliminary development batches of this additive were previously characterized at JPL. Five lots of this additive were received as powders and one in a slurry form. The results of their evaluation can be found in Reference 9.

The order in which a slurry batch was evaluated was as follows:

A. Slurry properties
B. Blending (single stage)
C. Fire suppression capabilities
D. Degradability (combustion and filtrability)
E. Unintentional degradation
F. Water reaction and low temperature behavior
G. Pumpability

Batches which failed one of the evaluation steps were not further evaluated.

3.1 Slurry Properties

Twelve (12) FM-9 slurries were evaluated in the order of which they were received at JPL.

All the slurries from the various batches, after homogenizing, produced thick, but free flowing, homogeneous liquids. Random slurries were tested for presence of large particles using the procedure in Appendix F. All slurries were practically free of particles larger than 100μ and one had less than 0.05 percent by weight of particles above 75μ. In several blending runs no pumping or blending problems were encountered. The only problem area observed (for all of the slurry lots) was the tendency to settle with time in storage. This resulted in the formation of a clear liquid phase on top and relatively hard layer of
solids at the bottom of the storage containers. This slurry settling rate was not determined but it was noticed that it began almost immediately on storage and a thin film of clear liquid appeared within 48 hours. The liquid on top, in the case of one slurry was decanted and measured. A 40 lb quantity of slurry produced 6 lb of clear liquid on top in 6 months of storage.

The viscosity of these slurries ranged from 6000 to 15168 centipoise.

3.2 Blending and Fire Suppression Capabilities

AMK blended with these slurries and the subsequent testing of fire suppression characteristics was done as described in Section 2.2.1 and 2.2.3 of this report. The blending and the fire test were done at ambient temperature which varied from 10°C to 27°C depending upon the time of the year.

Besides the fire test, the blending properties of the slurries were evaluated by the filter ratio, the cup test and by following the turbidity of the AMK fuel with time. For the large scale runs the gum content (percentage solids) of the blended fuel was also determined.

The evaluation of the FM-9 slurries are presented in the table 2 and 3. The numbers in parentheses are the air velocity, (in knots) at which the fire tests were run.

**TABLE 2. EVALUATION OF FM-9 SLURRY**

<table>
<thead>
<tr>
<th>SLURRY LOT</th>
<th>FIRE TEST (FT) AT 120 KNOTS</th>
<th>FILTER RATIO (FR)</th>
<th>CUP TEST (CT) IN CC</th>
</tr>
</thead>
<tbody>
<tr>
<td>JCK 14-125</td>
<td>FT^60 = MAR-FAIL</td>
<td></td>
<td>CT^60 = 2.2</td>
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<tr>
<td>JCK 14-125</td>
<td>FT^40 = FAIL</td>
<td>FR^40 = 44.8</td>
<td>CT^20 = 2.9</td>
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<td>FT^15 = FAIL</td>
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<td>CT^30 = 2.7</td>
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<tr>
<td>RMH 1-242</td>
<td>FT = PASS MAR (130)</td>
<td>FR = 33</td>
<td>CT = 3.3</td>
</tr>
<tr>
<td>JCK 14-125</td>
<td>FT^20 = FAIL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JCK 14-125</td>
<td>FT^15 = FAIL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JCK 14-125</td>
<td>FT^17 = FAIL</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note 1: superscripts indicate time after blending in minutes.

Note 2: absence of superscript indicates that the data is for equilibrated AMK.

Note 3: Base Fuel: RMH 30328
<table>
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<tr>
<th>SLURRY LOT</th>
<th>FIRE TEST (FT) AT 120 KNOTS</th>
<th>FILTER RATIO (FR)</th>
<th>CUP TEST (CT) IN CC</th>
</tr>
</thead>
<tbody>
<tr>
<td>JCK 14-163-3</td>
<td>FT$^{20}$ = FAIL</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>FT$^{15}$ = FAIL</td>
<td></td>
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</tr>
<tr>
<td>RMH 1-242</td>
<td>FT = PASS</td>
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<td>CT = 3.5, 3.5</td>
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<tr>
<td></td>
<td>FT = MAR (130)</td>
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<tr>
<td>JCK 14-163-3</td>
<td>FT$^{30}$ = MAR</td>
<td>FR$^{50}$ = 49.0</td>
<td>CT$^{10}$ = 3.2</td>
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<td>CT$^{15}$ = 2.7, CT$^{40}$ = 2.4</td>
</tr>
<tr>
<td>JCK 14-163-3</td>
<td>FT$^{65}$ = PASS (120 at 10°C)</td>
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</tr>
<tr>
<td></td>
<td>FT$^{30}$ = PASS (120 at 10°C)</td>
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<tr>
<td>JCK 14-163-3</td>
<td>FT$^{20}$ HRS = MAR</td>
<td></td>
<td>CT$^{70}$ = 2.2</td>
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<td></td>
<td></td>
<td></td>
<td>CT$^{24}$ HRS = 2.0</td>
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<tr>
<td>JCK 14-163-3</td>
<td>FT$^{20}$ = FAIL</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>FT$^{30}$ = MAR</td>
<td></td>
<td></td>
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<td>FT$^{20}$ HRS = MAR</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>FT$^{44}$ = FAIL (130)</td>
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<tr>
<td></td>
<td>FT$^{60}$ = MAR-FAIL (130)</td>
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Note: Superscript indicates time after blending in minutes
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<tr>
<th>AIR VELOCITY m/sec</th>
<th>SETTING FUEL FLOW RATE</th>
<th>RPMH 1-241</th>
<th>RPMH 1-242</th>
<th>JCK 14-125</th>
<th>JCK 14-162-3</th>
<th>JCK 13-195</th>
<th>JCK 13-196</th>
<th>JCK 13-197</th>
<th>TIME AFTER BLENDING</th>
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</thead>
<tbody>
<tr>
<td>70</td>
<td>650 (10 ML/SEC)</td>
<td>F/F</td>
<td>F/F</td>
<td>F/F</td>
<td>F/F</td>
<td>F/F</td>
<td>F/F</td>
<td>F/F</td>
<td>15</td>
</tr>
<tr>
<td>70</td>
<td>450 (14 ML/SEC)</td>
<td>N/M</td>
<td>N/M</td>
<td>F/F</td>
<td>F/F</td>
<td>MP/F</td>
<td>M/M</td>
<td>M/M</td>
<td></td>
</tr>
<tr>
<td>70</td>
<td>650 (18 ML/SEC)</td>
<td>-</td>
<td>-</td>
<td>M/M</td>
<td>M/M</td>
<td>M/F</td>
<td>M/M</td>
<td>MP/MP</td>
<td>60</td>
</tr>
<tr>
<td>70</td>
<td>450 (18 ML/SEC)</td>
<td>-</td>
<td>-</td>
<td>M/M</td>
<td>M/M</td>
<td>M/M</td>
<td>M/M</td>
<td>M/M</td>
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</tr>
<tr>
<td>70</td>
<td>650 (18 ML/SEC)</td>
<td>-</td>
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<td>M/M</td>
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<td>MF/MP</td>
<td>M/M</td>
<td>M/M</td>
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<tr>
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<td>450 (14 ML/SEC)</td>
<td>-</td>
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<tr>
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<td>60</td>
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<tr>
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<td>450</td>
<td>-</td>
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<td>-</td>
<td>-</td>
<td>60</td>
</tr>
</tbody>
</table>

M - Marginal; F - Fail; MP - Marginal Pass
Note: Repeat results shown.
Figure 9. MONITORING OF JCK 14-247-2 SLURRY BLENDING WITH TURBIDIMETER
TABLE 5. EVALUATION OF FM-9 SLURRY

<table>
<thead>
<tr>
<th>SLURRY LOT AND BASE FUEL</th>
<th>FIRE TEST (FT) AT (KNOTS)</th>
<th>FILTER RATIO TEST (FR)</th>
<th>CUP TEST (CT) IN C.C.</th>
</tr>
</thead>
<tbody>
<tr>
<td>JCK 14-247-2 RMH 30328</td>
<td>FT(^{20}) = PASS (120)</td>
<td>FR(^{24}) HRS = 52</td>
<td>CT(^{30}) = 2.0</td>
</tr>
<tr>
<td>((\triangle)-in Fig. 9)</td>
<td>FT(^{20}) = PASS (130)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>FT(^{20}) = MARG (140)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JCK 14-247-2 RMH 30328</td>
<td>FT(^{50}) = PASS (140)</td>
<td></td>
<td>CT(^{24}) = 2.05</td>
</tr>
<tr>
<td>((\triangle)-in Fig. 9)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>JCK 14-247-2 TEX/BUR JET-A</td>
<td>FT(^{20}) = MAR (130)</td>
<td>FT(^{6}) HRS = 60</td>
<td>CT(^{25}) = 2.6</td>
</tr>
<tr>
<td>(January) ((\triangle)-in Fig. 9)</td>
<td>FT(^{28}) = PASS (130)</td>
<td></td>
<td>CT(^{40}) = 2.5</td>
</tr>
<tr>
<td>56 PM Blender</td>
<td></td>
<td></td>
<td>CT(^{5}) HRS = 2.3</td>
</tr>
<tr>
<td>REPEAT AS ABOVE</td>
<td>FT(^{30}) = PASS (130)</td>
<td></td>
<td>CT(^{28}) = 2.4</td>
</tr>
<tr>
<td>((\triangle)-in Fig. 9)</td>
<td></td>
<td></td>
<td>CT(^{45}) = 2.2</td>
</tr>
<tr>
<td>1 LPG Blender</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1-23-84 RMH 1-246 CONTROL AMK Equilibrated</td>
<td>FT = PASS (120)</td>
<td></td>
<td>CT = 3.8</td>
</tr>
<tr>
<td>2-9-84 RMH 1-246 CONTROL AMK Equilibrated</td>
<td>FT = PASS (130)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Additional fire tests not presented in Table 3 indicated that even when equilibrated, the rating of this material at 120 knots was marginal at the most.

The fire test results for JCK 13-195, 196 and 197 of Table 4 slurries were similar to JCK 14-163-3. All three slurries got a "fail" rating at 120 knots, 15 to 20 minutes after blending and all got "pass" ratings at the same test conditions after equilibrating for 24 hours.

Because of the poor performance of these additive slurries the fire protection characteristics were double checked by FCTA test. The data presented in Table 4 confirmed the marginal fire protection properties of these batches. With the exception of some turbidity measurements presented in Figure 8 further evaluation of these batches was discontinued.

The evaluation of the FM-9 additive continued with lot JCK 14-247. It was marked lot 1 and 2, since it came in two pails (40 lb quantity each). It was considered the "final slurry lot" and was received in larger than the usual quantity to allow for large scale testing, e.g., blending, degradation and low temperature characterization.

Part of the data for this slurry is presented in Table 5 and Figure 9. It was found that filter ratio, cup and turbidity test values were about the same as the previously characterized slurries but the fire protection capability was significantly improved. These results were confirmed by data from larger blending runs (5-10 GPM blender) which also indicated good fire protection characteristics. Based on these results this AMK batch was further characterized for its degradability, pumpability and low temperature properties. These results are presented in Sections 3.3 through 3.6 of this report.

The JCK 14-247 slurry properties were evaluated again to compare them with newer batches of slurry additive which were received at JPL in July and later in November 1984.

The results presented in Table 6 and Figure 10 indicated partial deterioration of the flammability protection in comparison to the result obtained about 6 months earlier. The same slurry batch was tested again in November and the fire test results were the same as in January 1984. The ambient fuel temperatures in January and November were found to be about 10 to 15°C lower than those in July and August. These results demonstrate the influence fuel temperature had on AMK flammability characteristics. Fuels with "marginal" properties may pass the fire test at fuel temperatures of 10 to 15°C and get a marginal to fail rating at 22 to 25°C. Slurries which were rated as "good" when blended yield AMK which would get a "pass" rating in the entire range from 10 to about 30°C. At about 30°C fuel temperature even for these slurries, the fire test results become marginal and above 33°C they all fail the fire test. The data on two such slurries is given at the end of Table 5 as lot JCK 16-88-3 and JCK 16-95-1.

3.3 Degradability

The intentional degradability (restoration of base fuel properties) was assessed as described in Section 2.2.4. The results from the Hamilton blender degradation test and the subsequent characterization of the samples by the filter ratio test are presented in Table 7. Most of the work in this area was done
### TABLE 6. EVALUATION OF FM-9 SLURRIES

<table>
<thead>
<tr>
<th>SLURRY LOT AND BASE FUEL</th>
<th>FIRE TEST (FT) AT (KNOTS)</th>
<th>FILTER RATIO (FR) TEST</th>
<th>CUP TEST (CT) IN C.C.</th>
</tr>
</thead>
<tbody>
<tr>
<td>JCK 14-247-1 IN TEX/BURB JULY JET A 5 GPM BLENDER 0.28% SOLIDS</td>
<td>FT\textsuperscript{25} = FAIL (130) AT 26°C FT\textsuperscript{24} HRS = PASS (130) AT 21°C</td>
<td>FR\textsuperscript{40} = 49 FT\textsuperscript{9} DAYS = 53</td>
<td>CT\textsuperscript{18} = 3, CT\textsuperscript{30} = 2.3 CT\textsuperscript{22} = 2.5 CT\textsuperscript{9} DAYS = 2.2</td>
</tr>
<tr>
<td>JCK 16-88-3 IN TEX/BURB AUGUST JET A</td>
<td>FT\textsuperscript{25} = PASS-MAR (130) AT 28°C</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AS ABOVE IN JULY JET A</td>
<td>FT\textsuperscript{30} = PASS (130) AT 22°C</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JCK 14-247-1 IN TEX/BURB AUGUST JET A 5 GPM BLENDER 0.31% SOLIDS ( - in Fig. 10)</td>
<td>FT\textsuperscript{20} = MAR-FAIL (130) AT 22°C FT\textsuperscript{6} HRS = PASS (15°C) (130)</td>
<td>FR\textsuperscript{60} = 56</td>
<td>CT\textsuperscript{60} = 1.95</td>
</tr>
<tr>
<td>JCK 16-88-3 IN TEX/BURB AUGUST JET A 5 GPM BLENDER 0.31% SOLIDS</td>
<td>FT\textsuperscript{15} = FAIL (130) FT\textsuperscript{60} = PASS (130) ALL AT 24°C FT\textsuperscript{100} = PASS (130) AT 25°C FT\textsuperscript{80} = MARG (130) AT 31°C FT\textsuperscript{110} = FAIL (130) AT 33°C FT\textsuperscript{8} DAYS = PASS (130) AT 25.5°C</td>
<td>FR\textsuperscript{4} HRS = 80.5</td>
<td>CT\textsuperscript{40} = 2.1 CT\textsuperscript{24} HRS = 1.9</td>
</tr>
<tr>
<td>MIAMI SLURRY MOJAVE BLEND GE BLEND 0.31% SOLIDS</td>
<td>FT\textsuperscript{48} HRS = PASS (130)</td>
<td>FR\textsuperscript{34} HRS = 63</td>
<td>CT\textsuperscript{34} HRS = 2.0</td>
</tr>
</tbody>
</table>
### TABLE 6. EVALUATION OF FM-9 SLURRIES (CONT'D.)

<table>
<thead>
<tr>
<th>SLURRY LOT AND BASE FUEL</th>
<th>FIRE TEST (FT) AT (KNOTS)</th>
<th>FILTER RATIO (FR) TEST</th>
<th>CUP TEST (CT) IN C.C.</th>
</tr>
</thead>
<tbody>
<tr>
<td>JCK 16-95-2 TEX/BURB (AUGUST) JET A</td>
<td>FT&lt;sup&gt;30&lt;/sup&gt; = PASS (130) (15-25°C) FT = PASS (130) (15-31°C)</td>
<td>FR&lt;sup&gt;60&lt;/sup&gt; = 85±3</td>
<td>CT = 1.8 ± 0.1</td>
</tr>
<tr>
<td>TEX/BUR (SEPT) JET A 1 LPM AND 5 GPM BLENDERS</td>
<td>FT = PASS (160)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JCK 16-98-1 IN TEX/BUR(AUGUST) JET A</td>
<td>FT&lt;sup&gt;30&lt;/sup&gt; = MAR-FAIL (130) (20-25°C)</td>
<td>FR&lt;sup&gt;60&lt;/sup&gt; = 40±3</td>
<td>CT&lt;sup&gt;50&lt;/sup&gt; = 2.6 ± 0.2</td>
</tr>
<tr>
<td>TEX/BUR SEPT JET A 1 LPM RUNS</td>
<td>FT = MAR-PASS (130) (20-25°C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JCK 14-247-2 IN TEX/BUR AUGUST JET A</td>
<td>FT = PASS (130) 15°C FT = PASS (130) 24°C</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>REPEAT ABOVE WITH CID JET A</td>
<td>FT = PASS (130) 15°C FT = PASS (130) 25°C</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CID COMPOSITE SLURRY IN CID JET A, 1 LPM BLENDER</td>
<td>FT&lt;sup&gt;30&lt;/sup&gt; = PASS (130) 18°C FT&lt;sup&gt;60&lt;/sup&gt; = PASS (160) 18°C FT&lt;sup&gt;3&lt;/sup&gt; DAYS = (160) 18°C</td>
<td>FR&lt;sup&gt;60&lt;/sup&gt; = 55</td>
<td>CT&lt;sup&gt;45&lt;/sup&gt; = 2.0</td>
</tr>
<tr>
<td>AS ABOVE 5 GPM IN TEX/BURB (NOV.) JET A 0.296% SOLIDS</td>
<td>FT&lt;sup&gt;30&lt;/sup&gt; = PASS (130) 20°C</td>
<td>FR&lt;sup&gt;50&lt;/sup&gt; = 52 FR&lt;sup&gt;120&lt;/sup&gt; = 63 FR&lt;sup&gt;24&lt;/sup&gt; HOURS = 75.4</td>
<td>CT&lt;sup&gt;120&lt;/sup&gt; = 1.9 CT&lt;sup&gt;24 HR&lt;/sup&gt; = 1.9</td>
</tr>
</tbody>
</table>

Note: Solids were not determined for each blend unless stated.
using the JCK 14-247-2 slurry. With the exception of JCK 14-125 which showed a marginal degradability, the rest of the samples had good degradability.

The AMK degradability was also evaluated using a continuous flow single pass degrader which utilized pressure drop across the needle valve to degrade the fuel. The quality of the degraded fuel was evaluated in terms of its filterability. The description of the degrader, all the details of the test procedure and the test results are the subject of a separate report (Reference 7) and are summarized below.

The following samples were evaluated:

- JCK 14-247-2 AMK, freshly blended (20 minutes) or equilibrated.
- AMK blended in Miami (Florida); Convair-880 wing tank test sample from Mojave Airport; equilibrated
- AMK blended at Mojave Airport, Lot #7-5-84 tests equilibrated for 35 hours
- JCK 16-98-1 AMK
- JCK 16-95-2 AMK
- JCK 16-88-3 AMK equilibrated for 48 hours
- ICI Equilibrated AMK, Lot #RMH 1-241
- JCK 14-247-2 AMK, equilibrated for 6 hours and degraded at -22°C

With the exception of the last sample, the AMK fuel samples were successfully degraded without any filter plugging problems. The pressure drop across the filter was approximately 10 to 15 percent higher than Jet A fuel with degrader power requirement of 27.6 kw-s/liter (2.335 HP/GPM).

The filter ratio of the degraded samples were in the 1.1 to 1.3 range when measured 1 to 2 minutes after degradation. For equilibrated and then degraded AMK the filter ratio remained in that range. Filter ratio slowly increased with time to 10, 15, or in some cases even higher for freshly blended and degraded AMK samples.

The sample of AMK fuel which was degraded at -22°C gradually plugged the filter and the rate of plugging was 3.3 psi/minute. After modification of the degrader and with the introduction of a bypass loop the degrader performance markedly improved, enabling filtration of even low-temperature, freshly-blended and then degraded AMK fuel. (For further details of these findings, see Reference 7).

3.4 Unintentional Degradation

As previously indicated the unintentional degradation of the AMK fuel was evaluated by exposing the various samples to the same mechanical shear (one pass through the 1 liter per minute (LPM) Kenics blender) and then comparing the degree of partial degradation by filter ratio or fire test. The resistance to degradation of ICI blended equilibrated FM-9 AMK under these conditions has been the baseline control when evaluating the unintentional degradability of the various AMK batches. Past work done by FAA, JPL and Douglas Aircraft Company indicated that for FM-9 AMK fuel, an acceptable level of unintentional degradation in the wing tank as measured by filter ratio should be in the 15 to 20 range. One pass through the 1 LPM blender produced partially degraded fuel with filter ratio in that range. Due to limitation of funds and time
| SLURRY LOT                                      | FILTER RATIO (FR), CUP TEST (CT) OR NTU BEFORE DEGRADATION | DEGRADED FILTER RATIO FR\textdagger| |
|-----------------------------------------------|---------------------------------------------------|----------------------------------| |
| JCK 14-125 in RMH 30328 Jet A                 | FR$^{20}$ = 44.8, NTU$^{40}$ = 16 CT$^{30}$ = 2.7 | FR$\textdagger_{d}^{35}$ = 7.75 | |
| JCK 14-247-2 in RMH 30328 Jet A               | FR$^{24}$ HRS = 52, NTU$^{24}$ HRS = 5.8 CT$^{24}$ HRS = 2.0 | FR$\textdagger_{d}^{24}$ HRS = 3.27 | |
| JCK 14-247-2 5 GPM Blender in Tex/Burb. January Jet A | FR$^{6}$ HRS = 60, NTU$^{5}$ HRS = 6.2 CT$^{5}$ HRS = 2.3 | FR$\textdagger_{d}^{5}$ HRS = 3.4 | |
| JCK 14-247-2 in Tex/Burb. January Jet A       | FR$^{45}$ = 50.1, NTU$^{45}$ = 8.6 CT$^{45}$ = 2.2 | FR$\textdagger_{d}^{60}$ = 4.5 | |
| Control, RMH 1-246                            | CT = 3.3                                           | FR$\textdagger_{d}$ = 2.9       | |
| JCK 14-88-3 in Tex/Burb. July Jet A           | FR = 80.5, NTU = 3.5 CT = 2.0                      | FR$\textdagger_{d}$ = 4.1       | |
| JCK 14-247-1, 0.28% in Tex/Burb. July Jet A   | FR = 53, NTU = 4 CT = 2.2                          | FR$\textdagger_{d}$ = 4.27      | |
| 5 GPM Blend                                   |                                                    |                                  | |
| JCK 14-247-1, 0.31% as above                  | FR$^{60}$ = 56, NTU$^{60}$ = 5.0 CT$^{60}$ = 1.95 | FR$\textdagger_{d}^{60}$ = 4.4 | |
| Miami Slurry GE Blend in Mojave                | FR$^{34}$ HRS = 63 CT$^{34}$ HRS = 2.0             | FR$\textdagger_{d}^{34}$ HRS = 1.08 (5-10 hrs later) | |
|                                              |                                                   | FR$\textdagger_{d}^{34}$ HRS = 1.16 (4 hrs later) | |
| JCK 16-88-3, Tex/Burb. (July) Jet A, equilibrated | FR = 80.5, CT = 2.0, NTU = 3.5                      | FR$\textdagger_{d}$ = 4.1       | |
| JCK 14-247, Tex/Burb. (July) Jet A, 0.28% solids | FR = 53, CT = 2.2, NTU = 3.45                      | FR$\textdagger_{d}$ = 4.27      | |
| JCK 14-247 0.31% in Tex/Burb. (Aug.) Jet A    | FR$^{60}$ = 56, NTU$^{60}$ = 5.0 CT$^{60}$ = 1.95 | FR$\textdagger_{d}^{60}$ = 4.4 | |
constraints, solid contents of each blend was not determined.

In the case of the improved FM-9 AMK only two batches were tested for unintentional degradability and at the time of the experiment the ICI blended FM-9 AMK fuel was not available for comparison purposes. After one pass through the Kenics blender the fuel has higher filter ratios than FM-9 but failed the fire test, indicating excessive degradation. The data for the two batches is presented below in Table 8 and the JCK 14-247-1 data is not representative since the fuel is marginal even before degradation.

### TABLE 8. UNINTENTIONAL DEGRADABILITY OF FM-9

<table>
<thead>
<tr>
<th>SLURRY LOT</th>
<th>FILTER RATIO (FR), CUP TEST (CT), FIRE TEST (FT) BEFORE DEGRADATION</th>
<th>CUP TEST, FILTER RATIO, AFTER KENICS DEGRADATION</th>
<th>FIRE TEST</th>
</tr>
</thead>
<tbody>
<tr>
<td>JCK 14-247-1, 0.28% in Tex/Burb., (JULY) Jet A, 5 GPM Blend</td>
<td>FR = 53 FT = Marginal CT = 2.2 FT = Mar/pass (23°C) FT = Fail (27°C)</td>
<td>One Pass FR = 27.1 CT = 3.1</td>
<td>Fail (27°C)</td>
</tr>
<tr>
<td>JCK 16-88-3 in Tex/Burb., (July) Jet A, 6 GPM Blend</td>
<td>FT = pass (130) FR = 80.5 CT = 1.9 FR = 32 CT = 2.7</td>
<td>One Pass</td>
<td>Mar-Fail (25°C)</td>
</tr>
<tr>
<td>JCK 16-95-2 in Tex/Burb. in (Sept.), Jet A</td>
<td>FT = Pass (130) FR = 85 CT = 1.8</td>
<td>One Pass</td>
<td>FT = Mar-Fail (130) FT = Fail (140)</td>
</tr>
</tbody>
</table>

In view of these results and the change in the equilibrated filter ratio range from 45 ±5 for FM-9 to 85 ±5 for the improved FM-9, the 15 to 20 filter ratio range for acceptable unintentional level of degradation cannot be valid. The unintentional degradation in the aircraft fuel system depends on the degradability of AMK fuel and needs to be determined for this FM-9 variant. The only data which gave some information in this area was the characterization of AMK fuels from the Convair 880 wing tank. This fuel was blended at the FAA Technical Center, Atlantic City Airport, New Jersey, and flown to Mojave Airport in California with a stopover in Cincinnati, Ohio. The data for this fuel is as follows:

<table>
<thead>
<tr>
<th>ATLANTIC CITY*</th>
<th>CINCINNATI SAMPLE</th>
<th>MOJAVE AIRPORT SAMPLE</th>
</tr>
</thead>
<tbody>
<tr>
<td>FR = 46.3</td>
<td>FR = 33.1</td>
<td>FR = 36.6</td>
</tr>
<tr>
<td>CT = 2.4</td>
<td>CT = 2.75</td>
<td>CT = 2.8</td>
</tr>
<tr>
<td>NTU = 17.2</td>
<td>NTU = 7.2</td>
<td>NTU = 7.0</td>
</tr>
<tr>
<td>FT = Pass (130)</td>
<td>FT = Mar-Pass (130)</td>
<td></td>
</tr>
</tbody>
</table>

*FAA tests 3 days after blending
3.5 Water Reaction and Low Temperature Behavior

The investigation of AMK fuel with water vapor at the fuel/water vapor interface was done following the procedure in Section 2.2.6. The tests were performed using AMK fuel prepared from JCK 14-247-2 slurry. No difference in the behavior of this AMK fuel as compared to FM-9 fuel were observed. The same batch of AMK fuel was used for evaluation of the low temperature behavior. The test procedure can be found in Section 2.2.7. No presence of gel on top of the 4-mesh stainless steel screen was detected after one gallon of AMK fuel at -25°C was poured through the filter. In a similar experiment using the same batch of AMK, the content (50 gallons) of the low temperature wing simulator at -40°C was passed through the same size screen and again no gel on top of the screen was observed.

The same fuel when tested for flammability at -350°C fuel temperature retained its fire protecting capability at 130 knots air velocity.

As previously indicated (Section 2.2.7) the low temperature behavior of FM-9 AMK fuel was extensively evaluated using JPL’S aircraft wing tank environmental simulator (Figure 7). The low temperature characteristics (percent holdup) of JCK 14-247-2 AMK fuel under these conditions was compared to the characteristics of Jet A and ICI blended equilibrated AMK. For further details see Reference 8. In summary, these results indicated that under these test conditions the differences between the holdup behavior of FM-9 equilibrated AMK fuel, Jet A and JCK 14-247-2 AMK fuels are insignificant.

3.6 Pumpability of FM-9 AMK Fuel

The pumpability of the improved FM-9 AMK fuel was evaluated following the test procedure and the pumpability criteria described in Section 2.2.7. These tests were performed in the Cessna Boost Pump Rig. The AMK fuel used in these tests were prepared using JCK 14-247-1 slurry and was evaluated at two temperatures (22°C ±1°C and -25°C ±1°C). The data is presented in Figure 11 for JCK 14-247-1 AMK and Figures 12 and 13 for ICI equilibrated AMK and for Jet A. Because of the significant pumpability loss observed with this AMK fuel the tests were repeated using AMK made from slurry lot JCK 16-95. The results in Figure 14 indicate even higher pumpability loss when compared to JCK 14-247 AMK.

Finally, the pumpability performance of the improved FM-9 AMK fuel was evaluated using a full-scale DC-10 boost pump. Due to the complexity and the large amount of fuel required, this test was done on only one slurry batch (CID composite). This particular batch was selected because it "passed" all other tests, e.g., blending, fire test, degradability, etc.

The DC-10 boost pump was located inside JPL’s wing tank environmental simulator (Figure 7). The pumpability test results are presented in Figure 15. The results indicated that the pumpability performance deterioration when switching from Jet A to AMK fuel was not as marked for the DC-10 boost pump as for the Cessna boost pump. This may be attributed to the differences in the pump design and size.
FIGURE 11. PUMPABILITY OF IMPROVED FM-9 AMK FUEL RELATIVE TO JET A AND AT COLD TEMPERATURES
FIGURE 12. PUMPABILITY OF FM-9 AMK FUEL AT ROOM TEMPERATURE
RELATIVE TO JET A USING CESSNA PUMP
Figure 13. Pumpability of FM-9 AMK fuel at low temperature relative to Jet A at cold temperatures using Cessna pump.
FIGURE 14. PUMPABILITY OF IMPROVED FM-9 AMK FUEL ALL TEMPERATURES USING CESSNA PUMP
FIGURE 15. PUMPABILITY OF IMPROVED FM-9 AMK FUEL ALL TEMPERATURES USING DC-10 PUMP
CONCLUSIONS

The principal conclusions of this report are as follows:

1. Laboratory evaluation of the improved FM-9 slurry indicated that the quality of the slurry meets the metering and dispersion requirements for single state, in-line blending of AMK fuel.

2. The dissolution rate of the improved FM-9 additive in Jet A at ambient fuel temperatures (15-20° C) is sufficient, and produces AMK fuel with adequate fire suppression characteristics within 30 minutes after blending.

3. Evaluation of the intentional degradability of the improved FM-9 using degrader/filterability apparatus indicated that freshly blended fuel can be degraded without difficulty and with a specific power requirement of less than 30 kWsL⁻¹.

4. The interaction of the improved FM-9 AMK fuel with bulk or vapor condensed water is similar to the reaction of FM-9 AMK with water. Contamination with large amounts of bulk water leads to formation of gelled emulsion on the interface. The water vapor condensation of AMK surfaces produced a string-like second phase that redissolves when pumped or heated.

5. The low temperature evaluation tests indicated that the fuel holdup behavior of the improved FM-9 AMK is similar to that of Jet A and FM-9 AMK. The evaluation also indicated that there were no phase separation and gel formation problems at low temperature.

6. Significant loss in pumpability performance with the improved FM-9 AMK was observed at both room and low temperature using the Cessna 441 boost pump. However, the deterioration of pumpability performance was not as significant with AMK as compared with Jet A using the DC-10 boost pump.
REFERENCES


APPENDIX A

BATCH-BLENDED AMK RECEIVED BY JPL
AMK-FM-9-0.30 PERCENT SOLIDS
<table>
<thead>
<tr>
<th>LOT NUMBER</th>
<th>RMH 1-160</th>
<th>RMH 1-172</th>
<th>RMH 1-177</th>
<th>RMH 1-195</th>
<th>RMH 1-205</th>
<th>RMH 1-231</th>
<th>RMH 1-232</th>
<th>RMH 1-233</th>
</tr>
</thead>
<tbody>
<tr>
<td>DATE SHIPPED</td>
<td>7/9/81</td>
<td>8/21/82</td>
<td>10/14/84</td>
<td>11/18/84</td>
<td>12/7/81</td>
<td>3/23/82</td>
<td>5/12/82</td>
<td>6/3/82</td>
</tr>
<tr>
<td>AMOUNT, LBS.</td>
<td>1980</td>
<td>990</td>
<td>660</td>
<td>330</td>
<td>330</td>
<td>330</td>
<td>330</td>
<td>2640</td>
</tr>
<tr>
<td>% SOLIDS</td>
<td>0.30</td>
<td>0.297</td>
<td>0.310</td>
<td>0.290</td>
<td>0.303</td>
<td>0.297</td>
<td>0.297</td>
<td>0.290</td>
</tr>
<tr>
<td>FLOW CUP ml/30 s</td>
<td>2.57</td>
<td>2.60</td>
<td>2.70</td>
<td>2.40</td>
<td>2.30</td>
<td>2.40</td>
<td>2.8</td>
<td>2.57</td>
</tr>
<tr>
<td>CLARITY</td>
<td>Clear</td>
<td>Clear</td>
<td>Clear</td>
<td>Clear</td>
<td>Clear</td>
<td>Clear</td>
<td>Clear</td>
<td>Clear</td>
</tr>
<tr>
<td>VISCOSITY @ 25°C</td>
<td>2.75</td>
<td>2.73</td>
<td>3.12</td>
<td>2.80</td>
<td>2.90</td>
<td>2.99</td>
<td>2.86</td>
<td>2.87</td>
</tr>
<tr>
<td>FILTER RATIO</td>
<td>N.A.</td>
<td>N.A.</td>
<td>59.2</td>
<td>44.0</td>
<td>48.2</td>
<td>67.0</td>
<td>51.0</td>
<td>38.5</td>
</tr>
<tr>
<td>LOT NUMBER</td>
<td>RMH 1-237</td>
<td>RMH 1-242</td>
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APPENDIX B

OPERATING PROCEDURE FOR FILTER RATIO TEST

Fuel temperatures for Jet A and AMK are 20 °C.

Apparatus: Filtration ratio apparatus as shown in Appendix C.

Type of filter used: 16-18 micron twilled Dutch weave stainless steel 165 x 1400 mesh cloth, warp diameter 0.07 mm and weft diameter 0.04 mm, pre-cut into discs of 44.5 mm diameter. The material is obtained from Tetco, Inc., 525 Monterey Pass Road, Monterey Park, CA 91754.

1. Make sure filter apparatus has been rinsed clean with Jet A and then drained. Residual AMK can influence the filter time of the next sample.

2. Place an unused filter on lower filter plate, positioning it in the center so that it overlaps the edge of the orifice.

3. Both ‘O’ rings should be properly seated. Align upper and lower filter plates the same way each time; attach lower to upper and apply screws, tightening them to the same tolerance each time.

4. Insert a rubber stopper in bottom orifice, choosing a size which does not contact the filter. Hold stopper steady until removal. Excess motion may induce gelation in the filter.

5. Tilt apparatus to diagonal and pour the reference Jet A slowly down side of tube.

6. Once tube is about 3/4 filled, return it to vertical, add fuel till it overflows into gallery.

7. Remove rubber stopper. Record time between timing reference points.

8. When apparatus has drained, replace stopper, tilt apparatus to diagonal and pour sample AMK slowly (90 seconds) down side of tube, not letting it hit bottom directly.


10. Wait 60 seconds (fuel relaxation time) before removing stopper. Remove it slowly and gently with a turning motion to avoid causing suction.

11. Record time between timing reference points.

12. Dismantle lower filter plate and discard used filter. Rinse and drain apparatus.
APPENDIX C

DESCRIPTION OF FILTER SCREEN DEVICE

1st TIMING REFERENCE POINT

2nd TIMING REFERENCE POINT

GLASS METAL JOINT

CLAMP

SOLDER FILLET

VITON-O-RING

O-RING GROOVE 2mm DEEP

1.0 mm SQUARE

37 mm DIAM. FILTER

80-86°
APPENDIX D

OPERATING PROCEDURE FOR ICI ORIFICE FLOW CUP TEST

CLEANING PROCEDURE:
1. Place cup in Jet A. Fill cup about halfway with Jet A.
2. Sonicate for 30 seconds in Jet A fuel; power rating at 7.
3. Blow until dry with 25 psi nitrogen (1/4" hose). It is important that the area around the orifice hole both inside and out is completely dry and void of any particles.

OPERATING PROCEDURE:
1. Suspend cup inside ring on ring stand; allow enough room below cup to permit introduction of graduated cylinder (preferably 10 cc).
2. Place finger over the hole, tilt cup slightly to one side. Pour in fuel sample allowing fuel to run down the sides of the cup rather than hitting the bottom directly.
3. Let fuel overflow into gallery.
4. Once cup is full, allow 30 seconds before releasing finger (fuel relaxation time).
5. Release finger at 30 second mark, recovering fuel in beaker beneath hole. Let the cup drain for another 30 seconds.
6. Again at the 30 second mark, simultaneously slide graduated cylinder in place of beaker, collect for another 30 seconds then remove graduated cylinder and replace beaker. Record the amount of fluid collected in cylinder to the nearest 0.10 milliters (cc).
7. Discard collected material and repeat cleaning procedure.
APPENDIX E

OPERATING PROCEDURE FOR FCTA TEST

A special run procedure, described below, was devised for the FCTA to obtain rapid relative flammability measurement for quality control tests only. This procedure yields a single-point flammability temperature measurement and is not intended to replace standard FCTA procedure. It was incorporated because of the need to carry out testing on a routine basis.

1. The speed control dial which controls the fuel injection rate is set and recorded. Control dial settings range from 90-100 corresponding to low to high flow rates.

2. The air accumulator tank pressure which determines the air flow rate is allowed to climb to 6.5 atm (95 lb in\(^{-2}\)). This reading is taken at the highest pressure reached during the run and occurs just as the air begins to flow through the nozzle.

3. Temperature measurements are made with a 0.76 mm diameter lead, chromel-alumel thermocouple. The probe is placed level with and 25 cm downstream of the exit flange tip. Thermocouple readings are made with a strip chart recorder set so that a 1 mm deflection (the minimum resolvable) corresponds to a 240 temperature change.

4. A series of runs is performed until these tests yield results consistent within the measuring precision of \(\pm 12^\circ\)C.
APPENDIX F

JPL PROCEDURE FOR AMK SLURRY PARTICLES SIZE EVALUATION

1. Place 100 grams of well homogenized slurry in a 2000 ml graduated "Griffin" beaker equipped with magnetic bar and a stirrer.

2. Slowly, with gentle stirring, dilute the slurry sample with 1500 ml of tap water. Continue stirring until the liquid is homogeneous and has the consistency of milk.

3. Pour the contents of the beaker through a 200 mesh (Tyler equipment) sieve and wash any material which remains on top of the sieve first with water and then with methyl alcohol. Place the sieve in drying oven at 50°C and dry to constant weight.

4. Collect the powder and record its weight. A slurry with less than 0.05% (w/w) of particles above 75 micron size and practically free of particles above 100 micron size is of acceptable quality.
## Appendix G
### Distribution List

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<td>Dr. Frank A. Albini</td>
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<td>A. Allcock</td>
<td>Department of Industry Abell House, Room 643</td>
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<tr>
<td>Dr. S. J. Armour</td>
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<td>Robert Armstrong</td>
<td>NASA ARC M.S. 254-1</td>
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<td>M. C. Ingham</td>
<td>Chevron Research Company P.O. Box 1627</td>
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<td>M. C. Ingham</td>
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<td>Bartlesville, OK 74004</td>
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<td>Dr. D. E. Boswell</td>
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<td>William A. Callahan</td>
<td>ARCO Chemicals Company 1500 Market Street Philadelphia, PA 19101</td>
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<tr>
<td>Paul Campbell</td>
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<td>Michael Cass</td>
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Glenn Austin Halfway House, 1685  
REPUBLIC OF SOUTH AFRICA

Stephen L. Imbrogno  
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