



EFFECTS OF JP-4 FUEL ON GRAPHITE/EPOXY COMPOSITES

B. L. White Structural Concepts Branch Structures and Dynamics Division

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FOREWORD

This report describes an in-house investigation conducted by the Structural Concepts Branch (FIBC), Structures and Dynamics Division, Flight Dynamics Laboratory, Air Force Wright Aeronautical Laboratories, Wright-Patterson Air Force Base, Ohio, under Project 2401, "Structural Mechanics", Task 240103, "Advanced Structures for Military Aerospace Vehicles", Work Unit 24010338, "Preliminary Design of Aircraft Structures." Mr. Billy L. White, AFWAL/FIBCA, served as Project Engineer and test fixture designer; Mr. Robert T. Achard, AFWAL/FIBCC, supervised the fabrication of the composite specimens; and Mr. Harold D. Stalnaker, AFWAL/FIBE, was the Test Engineer.



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SECTION I

INTRODUCTION

During the past decade, composite materials have developed into current state-of-the-art materials commanding an increasing percentage of the structural materials used in high performance aircraft. Aircraft are now being flown, such as the F-18 and AV8B, that contain composite materials in primary structure throughout the aircraft. These materials are subject to numerous environmental effects including prolonged contact with liquids such as jet fuel.

It is well known that the epoxies used as a matrix in fibrous composite structures absorb moisture. This moisture causes a reduction in certain structural properties of the matrix. Questions still arise as to the possible degradation of composite structural properties due to jet fuel penetration into the epoxy matrix. The resistance of the composites to JP-4 fuel depends largely on the chemical resistance of the resin, and on the presence of defects such as cracks, voids, resinrich regions and "dry" regions. Such defects can arise during the fabrication process or from subsequent service usage.

In the past, several investigations have been conducted where the composite materials were simply subjected to immersion into fuels to determine if any deterioration could be detected. In all cases except for a few test points, the data has shown little or no degradation in material properties due to contact with fuels (References 1 through 4).

SECTION II

OBJECTIVES

The primary objective of this program was to determine if structural properties of graphite-epoxy composite laminates could be degraded by subjecting the material to JP-4 fuel under pressure for an extended period of time. Also, an investigation was conducted to determine if fuel tank sump water would deteriorate the laminate material properties. A secondary aim was to observe whether the laminates exposed to pressurized fuel would absorb enough fuel for a measurable weight gain.

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SECTION III

SCOPE

Tests have been conducted previously in which composites were simply immersed in JP-4 for long periods of time. No degradation in material properties was found as a result of these tests. Unlike this approach, the specimens in this program had pressure applied to one face of the laminate, tending to force the fuel into the laminate. To subject the test specimens to pressurized fuel required the design, analysis, and fabrication of an innovative test fixture. This fixture allowed the composite panels to act as fuel tank walls without being over-stressed in bending when high fuel tank pressures were applied for long periods of time. (Figure 1)

The investigation consisted of three extended evaluations in which a total of 305 specimens were tested in various modes. During the initial phase, three laminate panels were fabricated using Hercules AS 3501-5A graphite/epoxy material system with a stacking sequence of $[0_2/90_2/+45]_c$. Two of the panels were installed in the fuel tank test fixture and the third one was used as a control specimen for baseline data. The fuel tank specimens were subjected to 20 psi pressure for six weeks. In the second test phase, four panels were fabricated from the same material systems as the first test but had 16 plies with orientation of $(\pm 45^{\circ})_{Ac}$. Two of the panels were placed in the test fixture for six months under a fuel pressure of 40 psi. One panel was placed next to the test fixture for ambient environment exposure and one panel was placed in an oven at 140°F as a control specimen. The third phase test was essentially the same as the second phase except a mixture of JP-4 fuel and 3% artificial fuel tank sump water was placed in the tast fixture to determine sump water effects on material properties.

Following the extended test period in each test phase, the laminate panels were removed from their respective test areas and were cut into 20 tensile test specimens of 1 inch width and 10 inches length. Additionally, three of the tensile test specimens from each panel were used for preparing specimens for flexural strength and short beam shear tests.



SECTION IV

TEST FIXTURE

To accomplish the objective of the program, an innovative fuel tank test fixture had to be designed, analyzed, and built so that large composite laminate panels could be subjected to pressurized JP-4 fuel. (Figure 2) The test fixture was designed with a center plate cut out to contain fuel. The two outside plates of the fixture were designed with cutouts permitting the composite panels to be exposed on the inside to pressurized fuel and on the outside to the atmosphere. These cutouts had posts every two inches to support a 23 x 10 inch test panel and to prevent excessive deflection while under pressure. (Figure 3)

An extensive analysis of the fuel tank design was made prior to fabrication to assure that the fuel tank fixture would n t fail due to internal pressure. The weakest area of the fixture was found to be the posts between the cutouts in the plates. The posts were analyzed for fixed end boundary conditions. AISI 4130, low carbon steel material with a tensile yield stress (F_{ty}) of 70,000 psi and an ultimate tensile stress (F_{tu}) of 90,000 psi was sed in the initial analysis of the last fixture (Reference 6). The system was analyzed for a test pressure of 100 psi. For this pressure, the stresses were found to be 10,816 and the MS was 5.47. Thus, the fuel tank fixture was designed to contain pressures much higher thar the maximum 40 psi used during the programs.

The test fixture was designed to eliminate sealing problems to the extent possible. Recesses were cut into the end plates to seal the composite test panels and specimen retainers were designed to hold them securely against the tank walls. Buna-N rubber material was cut to shape and placed around the fuel tank plate boundaries to prevent leakage of the system.

A pressurization system was designed which would prevent any possibility of over-pressurization during the course of the test. The pressurization system incurporated a small cylinder with a piston that





was connected by a shaft to dead weights as shown in Figure 2. The cylinder filled with fuel when the fuel tank was filled. The piston compressed the fuel when the dead weight was applied, thus restricting the maximum system pressure. Over-pressurization could occur only by the application of more weight. A pressure gage was installed between the cylinder and the fuel tank for monitoring the system pressure.

The possibility of fuel leaking during the long test periods was taken into consideration when the test fixture was designed and the location of the facilities was selected. The fuel tank test fixture was designed to hold only 1.37 gallons of JP-4 fuel and the pressurization system held approximately 0.1 gallons of fuel. Consequently, if a leak had occurred, very little fuel would have been present for a possible fire. The conditions necessary for combustion were not permitted. The JP-4 fuel was enclosed in a sealed pressurized system, the test building was well ventilated, and no ignition sources were present. Also, the fuel tank test fixture was grounded to prevent the possibility of static electricity buildup.

SECTION V

TEST LAMINATE SIZING

To determine the most efficient size test fixture, several different size composite test panels were analyzed with 10 different laminate schedules of 8 to 16 plies each. Problem areas encountered during the design of the composite panels included: (1) larger panels, if unsupported would have excessive deflection, which could result in sealing problems, (2) small panels would have small deflections, but would not provide an adequate size or quantity of tensile test specimens; and (3) specimens from 7 panel that was very thick could not be failed in the test machine.

The analysis of several panels with different thicknesses and laminate orientations indicated that the 12 ply laminate panel with a lay-up of $[0_2/90_2/\pm 45]_s$ was the most suitable for the initial test. The results of the analysis indicated that a 20 psi test pressure would deflect the panel, supported by the test fixture cutout bars, .0037 inches at the center with a margin of safety of 8.39. This was an adequate MS to preclude possible failure due to the test pressure. The analyses of the graphite/epoxy composite panels were made utilizing two computer programs. The "Finite Element Plate Bending Analysis" program provided panel deflection data and resultant moments at the maximum deflection point (Reference 8). These resultant moments were input into the "Laminate Point Stress Analysis (SQ-5)" program which provided the margin of safety for each panel analyzed (Reference 9).

SECTION VI TEST PROGRAM

1. TEST NO. 1

Some consistent initial evaluation, a 12 ply composite panel with a stacking sequence of $[0_2/90_2/\pm 45]_s$ was selected to be tested at a fuel tank pressure of 20 psi for a six-week duration. Three laminate panels with dimensions of 23 x 10 inches were fabricated utilizing Hercules AS 3501-5A graphite/epoxy material. (Figure 4) The three laminates were cured in a single autoclave run and NDI was performed on each panel to determine if any voids or flaws were present. No significant defects were evident from the C-scan or X-ray inspections. The resin content was measured to be 30.5% by weight. Fiberglass (0⁰/90⁰ "S" glass) tabs were bonded to the laminate grip sections prior to insertion into the 140⁰ drying oven. The drying oven was used to assure a minimum moisture content at the start of the test period.

Two of the test panels (1A and 1B) were removed from the oven and placed in the fuel tank test fixture with one side subjected to JP-4 fuel at a 20 psi pressure and the other side to the ambient environment. The third panel (1C) was used as a control specimen exposed to the ambient environment. This panel was mounted on a metal plate with drying desiccant placed between the composite laminate and the plate. Consequently, the outside surface was subjected to the humid atmosphere and the inside was kept dry. Therefore, one side of the test panels were all exposed to the ambient environment. As a result, any reduction found in tensile strength could be attributed to the fuel exposure. The fuel tank test fixture and the control panel were placed in a remote building for the six week test period. The humidity and temperature of the area were recurded each afternoon. The temperature recorded during the test pericd ranged from $24^{\circ}F$ to $76^{\circ}F$, and the humidity ranged from 49% to 100%.

The three panels were removed from the drying oven and weighed prior to the start of the test period and weighed again after removal from the test fixture at the end of the six week test period.

÷μ







The before and after test weights were:

* · · ·	weight		
Specimen No.	Before	After	Change
14 (Suel Soaked)	777.4	777.5	0.1 (.01%)
IA (FUE? Soaked)	780.0	780.4	0.4 (.05%)
18 (ruer soared)	778.2	778.3	0.1 (.01%)
IC CARD LEAL WITT OF MILLION			

Further evaluation of moisture content was made since the panels were so large and the weight change so small that the accuracy of the available scales was questionable. The ends were cut off each panel and weighed. They were placed in a drying oven at 140° F for 30 days, removed and weighed. The results were:

	Weight	(Grams)	
Specimen NO.	Before	After	Change
14	10,7263	10.6995	0.0268 (0.25%)
19	5.6273	5.6155	0.0188 (0.21%)
. 10 	13.6639	13.6290	0.0349 (0.26%)

The results show that all three specimens had approximately 0.25% decrease in weight. It is evident from the small weight change that the resistance of the composite epoxies to JP-4 fuel is very good.

Immediately following the removal from the test fixture and after weighing, the test panels were cut into tensile specimens (Figure 5). Great care was taken to assure that the test specimens were cut in the 0° ply direction. A diamond cutoff wheel was used to cut each panel into 20 tensile test specimens one inch wide and 10 inches long (1 x 10). Three of the tensile test specimens(Numbers 1, 10, and 19) from each panel were saved for cutting into short beam shear and flexural strength test specimens (Figures 6 and 7).











Seventeen of the tensile test specimens from each of the three panels were tested to failure at ambient temperature in an Instron test machine. The specimens were loaded at a crosshead speed of 0.05 inches per minute. The results of the tensile tests were inconclusive as to whether or not the JP-4 fuel degraded the graphite-epoxy matrix since the laminate had a fiber dominated failure. The strength at the initial failure load was calculated using a nominal area for the fiber controlled laminate of .063 in² (12 ply x 0.00525 in./ply). The average failure strengths were 103,238; 98,502 and 97,621 (psi) for panels 1A, 1B, and 1C respectively (Figure A-1 and Table A-1).

Interlaminar shear stresses were determined by the short beam shear (SBS) test which is used primarily as a laminate quality check method. The short beam shear test method normally uses unidirectional laminates with the fibers running parallel to the length of the specimen. Although the test laminates were not specifically tailored to the SBS tests, it was decided in this program to use them for comparison purposes only to investigate effects of fuel exposure. However, the data generated is not recommended to be used as structural allowable values. Six SBS test specimens were cut from each of the three laminates. The specimen has the dimensions of 0.250 inch wide and 0.700 inch long. The specimen configuration is shown in Figure 6.

The short beam shear test fixture untilized a 3-point load assembly. The specimen supports were adjusted to a span-to-depth according to the relationship:

 $\frac{S}{T} = 4$

where S = span, inch

T = specimen thickness, inch

Therefore, with a specimen nominal thickness of .063 inch, a span of .25 inch was used between the specimen supports. The tests were conducted with the Instron test machine crosshead speed of 0.05 inch per

minute. The short beam shear strength at failure was calculated according to the following equation:

 $\tau = \frac{3P}{4A}$ where τ = Short beam shear strength, psi

P = Total load at failure, lbs

A = Cross sectional area, in^2

The average shear strength of the two fuel soaked laminates, 1A and 1B, were 11,286 and 10,971 psi respectively and the control (ambient environment) laminate, 1C, was 11,425 psi (Figure A-2 and Table A-2).

Six flexural test specimens were p[•] pared from each of the three panels IA, IB, and IC. Flexural testing was performed principally for checking the laminate material quality rather than for establishing basic mechanical properties. This is a convenient method for checking laminate quality since it simultaneously applies tension, compression, and horizontal shear. The test specimen was a straight sided, rectangular cross section beam 3.0 inches long and 0.5 inch in width. The specimen configuration is shown in Figure 7.

Flexural testing was conducted using a four-point loading method with the specimen support span of 2.0 inches. Each specimen was loaded to failure in an Instron test machine at a crosshead speed of 0.05 inch per minute. The ultimate flexural strength was calculated by the following equation:

$$\sigma = \frac{3PL}{4bt^2}$$

where:

 τ = stress in outer fiber at failure, psi

P = maximum load carried by the specimen, lbs

L = major span, inch

b = width of specimen, inch

t = thickness of specimen, inch

Comparison of the flexural strengths of the three panels shows very little difference in the values. The ambient environment exposed specimen, 1C, flexural strengths average value was 165,438 psi. The `wo fuel-soaked specimen strengths were 166,730 pci (1A) and 164,441 psi (1B) (Figure A-3 and Table A-3).

2. TEST NO. 2

1-

It was evident from the first test phase that the failure of the specimen was fiber controlled and the effects of JP-4 fuel on the matrix could not be determined with any reliability. It was decided that the next test should utilize a $\pm 45^{\circ}$ laminate schedule so the failure would be matrix rather than fiber controlled.

For the second test phase, the fuel tank pressure was increased to 40 psi and the test duration increased to six months in an attempt to force the fuel through the composite matrix. Analysis of $a \pm 45^{\circ}$ stacking sequence indicated that a 16 ply laminate would be required to prevent excessive deflection when exposed to the 40 psi test pressure. Also, four panel specimens were fabricated utilizing Hercules AS 3501-5A graphite-epoxy material rather than three used during the first test. The extra panel was used as a control panel and was placed in a 140°F drying oven for the duration of the test. The four panels were cured in a single autoclave run and NDI was performed on each panel to determine if any voids or flaws were present. No defects were evident from the C-scan and X-ray inspections. The resin content was measured to be 31% by weight. Fiberglass tabs were bonded to the grip sections prior to inserting the four panels in the 140°F drying oven. The drying oven was used to assure a minimum moisture content at the start of the test period.

Two of the test panels (2A and 2B) were removed from the oven and placed in the fuel tank test fixture with inside subjected to JP-4 fuel at a 40 psi pressure and the outside to the atmosphere. The third panel (2C) was used as a control specimen exposed to the ambient environment. This panel was mounted on a metal plate with drying desiccant placed

between the composite laminate and the plate. This resulted in the outside surface being subjected to the atmosphere and the inside surface staying dry. The fourth panel (2D) remained in the $140^{\circ}F$ drying oven for the six month test period. The outside air temperature at the test site ranged from $51^{\circ}F$ to $92^{\circ}F$ and the humidity ranged from 39% to 100% during the test period.

The four panels were removed from the drying oven and weighed prior to start of the test period. They were once again weighed after the six month test period. The before and after test weights were:

		Weight	(Grams)		
_Spe	cimen No.	Before	After	Chang	e
2A	(Fuel Soaked)	901.1	902.9	+1.8	(.20%)
28	(Fuel Soaked)	895.6	897.0	+1.4	(.16%)
2C	(Ambient Environment)	901.9	902.0	+0.1	(.01%)
2D	(Oven Control)	898.3	897.6	-0.7	(08%)

The weight change data is questionable since it is improbable that the available scales could accurately measure the small change in the panel weights. It does indicate that the composite matrix never absorbed an exorbitant quantity of fuel.

The four test panels were each cut into 20 tensile test specimens of on inch width and ten inches in length. As in the first test, great care was taken to assure that the specimens were cut in the 0° direction. Three of the cut specimens from each panel (numbers 2, 10, and 18) were saved for use as flexural strength and short beam shear test specimens (Figures 6 and 7).

Sixty-eight tensile test specimens, 17 from each panel, were tested to failure at ambient temperature in an Instron test machine. Prior to testing, each specimen was measured for thickness and width in three places for determining specimen area. The failure stress of each

specimen was calculated using the average area and failure load. A mean stress, standard deviation, and coefficient of variation were calculated for the 17 specimens. The standard deviation for all data sets was less than one percent of the mean stress except the oven control panel, which was 1.17 percent. The mean failure strengths from the four panels shows a very small difference in value. The oven-dried control panel specimen strengths were slightly higher than the fuel-scaked and ambient-environment exposed panel specimens, which were essentially the same. The mean failure strength for the oven control panel specimens was 27,421 psi. The failure strength for the two fuel-exposed panel specimens and the exposed ambient environment panel specimens were 26,964 psi, 26,944 psi, and 26,982 psi respectively (Figure A-1 and Table A-4).

Interlaminar shear stresses were determined by the short beam shear (SBS) test as was described in Test No. 1. This test utilized specimen numbers 2, 10, and 13 of each panel for preparing SBS specimens (Figure 6). Three SBS coupons were cut from each specimen. The SBS test used a 3-point load assembly with supports adjusted to a span-to-depth ratio of four. A span of approximately 0.35 inch between the specimen supports was used for the 16 ply SBS specimen, which had a nominal thickness of 0.084 inch.

The results of the short beam shear tests, in which nine specimens from each panel were tested to failure, show that the ambient environment exposed panel (2C) had the greatest shear strength, whereas, the two fuel soaked panels (2A and 2B) had essentially the same shear strength as the oven control panel (2D). The mean shear strengths of the two fuel exposed panel specimens were 11,867 and 11,558 psi; the oven control panel had a strength of 11,868 psi and the ambient environment exposed panel shear strength was 12,415 psi (Figure A-2 and Table A-5).

The flexural test specimens were prepared from specimen numbers 2, 10, and 18 from each of the four panels (2A, 2B, 2C, and 2D) (Figure 7). The testing and flexural strength calculations were rerformed as in Test No. 1. The fuel-exposed specimens had average flexural strengths of

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45,161 ps1 and 44,834 ps1 and the ambient-environment-exposed and oven dried specimens average flexural strengths were 46,432 ps1 and 46,140 ps1 respectively (Figure 10 and Table 6).

3. TEST NO. 3

The third test phase consisted of the same type material, Hercules AS 3501-5A, and laminate schedule, $\pm 45^{\circ}$, as Test No. 2 with some variations in procedures. Since very little change was evident in Test No. 2 of the fuel-soaked and ambient-environment-exposed panels, it was decided to (1) determine the effects of sump water on the graphite-epoxy material; (2) instrument some of the tensile test specimens; (3) perform only tensile tests; and (4) fabricate two extra panels and test them prior to the start of the fuel soak tests.

For the third test, a 16 ply composite laminate with a stacking sequence of $(\pm 45^{\circ})_{45}$ was selected to be evaluated at a fuel tank pressure of 40 psi for a six month duration. Also, six panels were fabricated rather than the four used during the second test. The two extra panels were dried in an oven for four weeks at 140° F and tensile tested prior to the start of the fuel soak test to determine if the tensile properties had a significant change during the test period. The six panels were cured in a single autoclave run and NDI was performed on each panel to determine if any voids or flaws were present. No significant defects were evident from the C-scan and X-ray inspections. The resin content was measured to be 28.8% by weight. The panels were placed in a 140°F drying oven to assure a minimum moisture content at the start of the test period.

Two test panels (3A and 3B) were removed from the oven, weighed, and placed in a fuel tank test fixture. The third panel (3C) was used as a control specimen exposed to the ambient environment. The fourth panel (4D) remained in the drying oven for the six month test period. The fifth and sixth panels (4E and 4F) were cut into 20 tensile test specimens each. The tensile test specimens, numbers 2, 10, and 18 were instrumented on both sides with high elongation strain gages. Tensile tests were conducted on the 40 test specimens.

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The fuel tank test fixture was filled with JP-4 field of 20104 2000 water. The artificial sump water was prepared by General Cynonics, Cert Worth Division, from the formula given in the Materials Laboratory Technical Report entitled. "Clastomers for Fuel Systems Containing Micro-Organisms-Centrolling Additives". (Reference 7 and Lable Add), the wrificial fuel tank sump water would not readily blend with the ode; fuel. The solution had to be mixed in a sonic vibrator. The 34 solution of sump water and JP-4 fuel was placed in the test fixture and tank was pressurized to 40 psi for the six month test period. The outside air temperature of the test site ranged from 46°F to 37°F and, the humidity ranged from 43% to 32% during the test period.

The four panels were removed from their test area after six coulds and weighed. The before and after test weights were:

		Weight	[Gr 305]		
Foorimon NO.		ilefore	After	Chi	e F Marina
28 (Fun) Soaked)		853.7	855.6	2.1	(0.253)
JA (ruel Souked)		850.4	852.4	`. 0	(0.747)
38 (File) 209kent	(nmont)	852.5	854.1	1.6	(0.194)
3C (Ambient Envir	Ormency	063 A	851.1	-0.3	(0.034)
30 (Oven Control)		036.4			

The weight change data is questionable since it is imprchable that the available scales could accurately measure the small change in the panel weights. It does indicate that the composite matrix never absorbed an exorbitant quantity of fuel.

A total of 120 tensile specimens were tested to failure in lest No. 3. Twenty specimens were prepared from each panel. Three test specimens from each panel were instrumented with strain gages and the data plotted in Figure A-4. A mean stress, standard deviation, and coefficient of variation were calculated for each set of data. Just standard deviation for all data sets was less than two precent of the mean stress with the oven control data having the largest coefficient of variation, which was 1.76. The results show that the fuel-surp water

exposed specimens mean strengths had a decrease in value when compared to the other laminates. The fuel-sump water soaked panels mean failure strengths were 25,176 psi and 25,308 psi. The ambient environment exposed panel and the oven control panel mean strengths were 26,129 psi and 27,568 psi respectively. The mean tensile strengths of the two new panels, which were tested prior to the six month test period, were 26,829 psi and 27,039 psi (Figure A-1 and Table A-7).

SECTION VII

CONCLUSIONS AND RECOMMENDATIONS

The objective of this program was to determine if graphite-epoxy composite material properties could be degraded by exposing the laminate to pressurized JP-4 fuel. The study dealt primarily with the tensile testing of fuel and ambient environment exposed laminates and oven dried laminates to determine relative strength values. Also, a test condition was set up to evaluate the effects of fuel tank sump water on the composite properties. The conclusions from this effort were:

1. Composite laminates will not absorb a significant amount of JP-4 fuel, after exposure to JP-4 fuel at a 40 psi pressure for six months.

2. The results of Test No. 1 show no degradation of the laminate strength due possibly to having fiber dominated failures rather than matrix controlled failures. The mean tensile strength of the fuel soaked laminates was actually greater than that of the ambient environment exposed laminates. The short beam shear and flexural strength average results were essentially the same for all laminates.

3a. The test results of Test No. 2 show no significant degradation in the strength of the test laminates. The tensile strength results show that the fuel exposed and ambient environment exposed laminates had the same average strength. The oven control panel had a slightly higher average failure strength than the other specimens although it was still within 2%.

b. The short beam shear test results show the ambient environment exposed laminate had a shear strength higher than the fuel exposed and oven dried laminates. Short beam shear strengths of the fuel exposed and oven dried laminates were essentially the same.

c. The flexural strength of the ambient environment exposed and oven control specimens was approximately 4% greater than the fuel soaked flexural strength values. No conclusions can be made from this test due to the small sample size and the large deviation in failure loads.

2. 2. 1.

4. Test No. 3 data shows that the fuel plus artificial sump water mixture reduces the strength of the composite materials. The fuel-sump water soaked tensile failure loads are consistenly lower than the other laminate failure loads, as shown in Figure 11. The mean failure load of the new material test specimens was 2176 lbs. The mean failure load of the fuel-sump water test specimens was 2060 lbs.

5. This study has shown that there is little or no degradation of composite material properties due to JP-4 fuel. The undamaged composite matrices used in this investigation were very resistant to JP-4 fuel. It is apparent from this study that further investigations are required to determine if fuel sump water has a significantly detrimental effect on composite material properties.

The following recommendations are made to further investigate the compatibility of composite fuel tanks with fuel and sump water:

1. Investigations should be conducted on matrix damaged composite fuel tanks since the resistance of the composites to JP-4 fuel depends largely on undamaged resins.

2. Additional studies should be made to determine if fuel tank sump water significantly reduces composite material properties beyond the normal reduction due to moisture absorption.

APPENDIX TEST DATA

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1. UN
RESULTS-TEST
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TINGTE
TABLE A-1

L 1C NVIRONMENT)	STRESS	(FSI)	88571	80952	101429	98413	100397	100317	100476	100159	103333	109206	109683	95238	87302	92095	93016	98413	100476	97612	7454	7.63	
FANE (AMBIENT E	FAILURE LOAD	(TB)	5580	5100	6390	6200	6325	6320	6330	6310	6510	6880	6910	6000	5500	5802	5860	6200	6330	6149	469	7.63	
L 1B SOAKED)	STRESS	(ISI)	93333	101587	92222	113810	111746	111905	86984	95238	95286	95079	92857	98730	91905	106508	106391	92063	88905	98502	8535	8.66	
PANEL S	FAILURE LOAD	(TB)	5880	VU79	5810	7170	7040	7050	5480	6000	6003	2990	5850	6220	5790	6710	6702	5800	5601	6205	537	<u></u> х.00	
. 1A OAKED)	STRESS	(PSI)	99683	112778	106380	109556	108000	93650	105556	105873	100952	96873	111143	105556	106032	91746	105556	98413	97619	103238	6085	5.89	cness = 0.063 in
PANEL FUEL S	FAILURE LOAD	(TB)	6280	7105	6702	6902	6804	2900	6650	6670	6360	6103	7002	6650	6680	5780	6650	6200	6150	6504	viation 383	iration 5.89	inal 12 ply Thick
TENSILE SPECIMEN	.04		2	ŵ	4	5	Q	7	8	6	11	12	13	14	15	16	17	18	20	Mean:	Standard De	Coef. of Vé	Note: Nomi

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NO.	
RESULTS-TEST	
TEST	
SHEAR	
BEAM	
SHORT	
TABLE A-2	

PANEL 1C (AMBIENT ENVIRONMEN	EA FAILURE LOAD 1 ²) (LB)	172 244	1174 289	1173 275	177 294	1177 263	1173 229	1174 266	002 25.5	
	I ARE (IN	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
ED)	SHEAR STRENGTH (PSI)	11466	11063	9738	12119	10684	10757	10971	802.5	7.31
PANEL 1B (FUEL SOAK	FAILURE LOAD (LB)	266	267	235	286	255	251	260	17.3	·
	AREA (IN ²)	0.0174	0.0181	0.0181	0.0177	0.0179	0.0175	0.0177	0.0003	
	SHEAR STRENGTH (PSI)	10838	12659	11337	11571	10614	10695	11286	770.7	6.83
PANEL 1A (FUEL SOAKED)	FAILURE LOAD (LB)	250	292	260	270	242	241	259	19.5	
IEAR :IMEN IO.	AREA (IN ²)	0.0173	0.0173	0.0172	0.0175	0.0171	0.0169	. 0.0172	<mark>d.</mark> : 0.0002	. of
SH PEC		-	7	ę	4	ŝ	9	lean	tan ev.	oef ar.

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PANEL 1C MBIENT ENVIRONMENT)	HICK- WIDTH LOAD FLEXURAL NESS STRENGTH (IN) (IN) (LB) (PSI)	.069 0.499 261 164791 .069 0.499 279 176156 .069 0.499 279 165391 .069 0.501 269 165824 .069 0.505 269 153186 .069 0.504 238 153186 .069 0.504 238 153186 .069 0.501 266 167277 .069 0.501 262 167277 .069 0.501 266 167277 .069 0.501 266 7334 .069 0.511 266 167277 .069 0.511 266 167277 .069 0.511 266 167237 .069 0.511 266 167237 .060 13.6 7334
PANEL 18 (FUEL SOAKED)	THICK- WIDTH LOAD FLEXURAL NESS STRENGTH (IN) (IN) (LB) (PSI)	0.070 0.500 266 162857 0.070 0.504 272 165209 0.069 0.501 264 166020 0.069 0.503 264 165359 0.070 0.503 285 173449 0.079 0.503 285 173449 0.079 0.503 285 173449 0.069 0.500 244 153749 0.069 0.500 244 153749 13.3 266 164441 13.3 6355 3.86
FLEXURAL PANEL 1A SPECIMEN (FUEL SOAKED) NO.	THICK- WIDTH LOAD FLEXURAL NESS STRENGTH (IN) (LB) (PSI)	1 0.067 0.501 236 157404 2 0.067 0.504 258 172077 3 0.067 0.504 259 171716 4 0.069 0.501 254 159731 5 0.068 0.501 265 171586 6 0.069 0.503 268 167365 Mean: 257 166730 Stand. Dev. : 11.3 6546 Coef. of Var. : 3.92 3.92 3.92 3.92

s,

TABLE A-3 FLEXURAL STRENGTH TEST RESULTS-TEST NO. 1

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20.00

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(1c	STRESS (PSI)	27431 27553 27553 27553 27553 27569 27699 27693 27693 27693 27693 27615 27615 27615 27615 27615	27421 322.1 1.17
PANEL 2D TEN CONTRO	FAILURE I OAD (LB)	2381 2381 2417 2417 2420 2435 2435 2435 2435 2435 2435 2436 2436 2436 2436 2436 2436 2436 2436	2412 22.76
NO)	AREA (1N ²)	.0363 .0877 .0879 .0878 .0878 .0893 .0882 .0882 .0882 .0882 .0882 .0882 .0882 .0882 .0882 .0882	.0000
)NMENT)	STRESS (PSI)	27085 26623 26993 27100 27100 27121 27313 27095 27095 27095 27121 26970 26970 26937 26733 26736 26733 26736 26953	26982 256.2 0.949
PANEL 2C NT ENVIRC	FAILURE LOAD (LB)	2267 23267 23370 23370 23369 23369 23369 23357 23369 23357 23357 23357 23357 23357 23357	2357 28.58
(AMBIE	AREA (IN ²)	.0337 .0874 .0878 .0876 .0871 .0871 .0871 .0875 .0886 .0886 .0875	.035
(G	STRESS (PSI)	27350 27125 27017 26904 26896 26880 26880 26880 26880 26880 26880 26880 26880 26830 26652 27000 27371	26944 212.3 0.788
PANEL 2B UEL SOAKE	FAILURE LOAD (LB)	2334 2334 23354 23354 23354 23354 2334 233	2353
(F	AREA (IN ²)	.0868 .0873 .0875 .0871 .0877 .0877 .0870 .0870 .0870 .0870 .0870 .0870	.0873
(9	STRESS (PSI)	27503 27378 26923 26923 26954 26667 26667 26667 26667 26667 26653 26710 26633 26710 26633 26796 26735 26796 26735 27123 27123	26964 259.9 0.964
PANEL 2A UEL SOAKF	FAILURE LOAD (LB)	2390 2325 2325 2337 2335 2337 2335 2337 2335 2337 2335 2337 2335 2337 2337	2350 23.09
LE IEN (FI	AREA (1N ²)	.0%69 .0%62 .0%73 .0%62 .0%873 .0%82 .0%873 .0%873 .0%873 .0%73 .0%68 .0%68 .0%68 .0%68 .0%68 .0%68	.0872 .00089 of Var.
TENSIL SPECIM NO.		2616655321937000000	Mean Stand. Dev. Coef.

TABLE A-4 TENSILE LEST ABSULIS-TEST NO. 2

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STELLAR STELLAR RU.	r	PANEL 2A	(ED)	(۲	PAKEL 28 UEL SCARE	H	(ANBIE	PANEL 20 NT ENVIRC	(INGAN)	COV.	PANEL 201 EN CONTRO	C
	ABEA (13 ²)	FAILURE LOAD (LB)	E SHEAR STRENGTH (PSI)	AREA (15 ²)	FAILURE LUAD (LB)	SHEAR STRENGTH (F31)	AREA (13 ²)	FAILURE Load (LB)	SHEAR STRENCTH (PSI)	AREA (1 ^{N²)}	FAILURE LOND (LB)	SHE UR STRENGTH (PST)
1	.0222	352	11867	.0232	353	11418	.0223	126	12473	.0229	<u> 1</u> 94	12930
2	.0234	352	11274	.0227	36.8	12171	.0225	361	12053	1520-	35.0	10337
•	.0226	377	12514	.0227	351	11576	.0218	329	11325	.0233	349	11223
.4	.0236	358	11398	.0246	366	11159	.0218	186	13090	.0236	367	11679
Ś	.0225	364	12137	.0233	366	11798	-0219	361	12360	1520.	332	11422
9	.0226	360	11964	4620.	351	11267	.0217	334	11532	. 0234	367	11775
٢	.0225	376	12391	.0222	543	11576	.0216	382	13256	.0217	367	12671
•0	C223	353	11872	.0238	352	11105	.0213	390	13696	.0225	373	64421
6	-02.30	349	11383	.0228	363	11953	.0216	344	11950	.0226	355	11775
Mean	.0228	360	11867	.0232	357		.0218		12415	1620.	367	11868
Stand Dev.	·000	3.6	148.2	.000	60 60 60	363.7	,000 .	21.8	800.4	0100.	14.7	683.7
Coef. o	f Var.		3.74			3.14			6.44			5.76

TADLE A-3 SHORT BEAM SHEAR TEST RESULTS-TEST NO.

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FLEXURAL SPECIMEN NO.		PANEL (FUEL S	2A Oaked)			PANE (Fuel	L 2B Soaked)	•
	THICK- NESS (IN)	(II)	LOAD (LB)	FLEXURAL STRENGTH (PSI)	THICK- NESS (IN)	(NI)	LOAD (LB)	FLEXURAL STRENGTH (PSI)
1 2 7	.0870 .0864	.5017 .5023	115 113	45426 45204	.0863 .0863	.5004 .5006	601 111	43871 44658 42033
3	.0863	.5029	112	44800	.0802	anne.	114	7/60%
Mean	.0866	.5023	113	45161	.0863	.5005	111	44834
Stand Dev.	.0004	•0006	1.5	288	1000.	1000.	1.5	613
Coef. of Var.				.637				2.36
FLEXURAL SPECIMEN		PANEI (AMBIENT EN	, 2C WIRONMENT)			PANE (oven c	IL 2D Control)	
	THICK-	HLCIM	QVOI	FL EXURAL STRENGTH	THICK- NESS	HIDIM	LOAD	FLEXURAL
	(NI)	(NI)	(TB)	(IS4)	(NI)	(NI)	(81)	(ISI)
7	.0869	.5006	120	47615	.0873	.5011	115	45169
ΝŴ	.0881 .0872	.5007	120 115	45363 45363	.087 8 .0871	.5006	123 115	47810 45440
Mean	.0874	. 5005	118	46432	.0874	.5007	117	46140
Stand Dev.	.0006	£000°	2.9	1130	.0004	.0004	4.6	1452.8
Coef. of Var.				2.43				3.14

TABLE A-6 FLEXURAL STRENGTH TEST RESULTS-TEST NO. 2

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ENT)	STRESS	26452 26200 25783	26098 26717	26103 26445 26445	26412 25543 25966 26370	26190 25908 26156 26576	25903 25568 26188 26273	26129 324 1.24
PANEL 3C ENT ENVIRONM	FAILURE LOAD	2140 2117 2140	2140 2140	2140 2130 2150	2115 2115 2150 2165	2145 2140 2150 2150	2150 2117 2120 2115	2138 14.5
(AMB)	AREA	.0809 .0808 .0830	.0820 .0801 .0832	.0813 0813 0813	.0814 .0828 .0828	.0819 .0826 .0822 .0809	.0830 .0828 .0312 .0805	.03177 .0009
ER)	STRESS (PSI)	25743 25362 25275	25000 24940 24237	25593 25561 25636	25287 25277 25371	25401 25494 25091 25746	25181 25571 24970 25616	25308 349 1.38
PANEL 3B EL/SUMP WAT	FAILURE LOAD (LB)	2080 2067 2070	2080 2065 2065	2050 2050 2040	2028 2050 2050	2060 2065 2070	2090 2079 2070 2080	2064 15.3
(FU	AREA (IN ²)	.0808 .0815 .0819	.0832 .0828 .0852	.0801 .0802 .0802	.0802 .0811 .0808	.0810 .0810 .0823 .0804	.0830 .0813 .0829 .0812	.0816
TER.)	STRESS (PSI)	25304 24563 25092	24939 24940 25398	25776 25214 25340	25266 25339 25213 25213	25398 25398 25184	24939 25191 24908 25123	25176 248 0.986
PANEL 3A JEL/SUMP WAY	FAILURE LOAD (LB)	2080 2024 2050	2070 2070 2075	2075 2060 2050	2044 2055 2070 2060	2070 2075 2055	2043 2040 2040	2057 14.9
(F	AREA (IN ²)	.0822 .0824 .0817	.0830 .0817	.0805 .0817 .0809	.0809 .0811 .0821	.0818 .0817 .0816	.0811 .0819 .0812	.0817 .0006 ar.
TENSILE SPECIMEN NO.		- 10 6 4 *		6 8 7	10 * 11 13	112	18 * 20	Mean Stand Dev. Coef. of V

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TABLE A-7 TENSILE TEST RESULTS-TEST NO. 3

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* - Strain Gaged Specimens

ENSILE PECIMEN NO.	_	PANEL 3D (OVEN CONTROI	.		PANEL 3E (NEW)			PANEL 3F (NEW)	
	AREA	FAILURE LOAD	STRESS	AREA	FATLURE LOAD	STRESS	AREA	FAILURE LOAD	STRESS
	(IN ²)	(TB)	(PSI)	(1N ²)	(TB)	(ISI)	(IN ²)	(TB)	(ISI)
	.0825	2220	26909	.0810	2125	26235	.0808	2165	26795
*	.0814	2217	27232	.0812	2135	26293	.0809	2112	26106
.	0180.	2200	27160	.0811	2200	27127	.0808	2160	26733
u to	.0819	2180	26618	.0817	2205	26989	.0814	2190	26904
n v	1280.	2200	26797	.0807	2200	27261	.0800	2160	27000
0 1	6080.	2190	27071	.0810	2195	27099	.0812	2180	26847
~ 0	4T80.	2150	26781	.0807	2220	27509	.0802	2200	27431
0 0	.0806	2250	27915	.0812	2175	26786	.0811	2195	27065
ן אר י	.0804	2240	27861	.0805	2175	27019	.0310	2195	27099
*	.0802	2217	27643	.0818	2090	25550	.0800	2120	26500
11	.0800	2230	27875	.0812	2150	26479	.0806	2185	27109
7 7	.0800	2230	27875	.0800	2165	27163	.0806	2200	27295
×	- 0806	2250	27916	.0813	2190	26937	.0802	2200	27431
4	.0804	2250	27985	.0816	2200	26961	.0801	2205	27528
	.0802	2260	28180	.0816	2195	26900	.0816	2200	26961
	.1813	2260	27798	.0816	2130	26715	.0808	2205	27290
+ 0+	1180.	2250	27743	.0814	2200	27027	.0792	2190	27652
0 1	0080.	2248	28100	.0808	2149	26597	.0797	2119	26587
100	.0803	2250	28020	.0807	2190	27139	.080.	2190	27273
7.0	CT00.	0622	27676	.0808	2165	26795	.0802	2180	27182
an	.0809	2228	27568	.0811	2172	26829	.0805	7716	92039
and Dev.	.000	27.0	484	7000	181	127	9000	- F C	
ef. of V			76 6		1.07		0000.	1.10	3/8
	• •		0/ · T		·	1.63			1.40
- Strain	Gaged Sp	ecimens							

TABLE A-8 ARTIFICIAL FUEL TANK SUMP WATER (Reference 7)

COMPOSITION	Z BY WEIGHT	METAL	CHLORIDE
		(Had)	(Mdd)
cac1 ₂	0.005	18	32
cdc1_2	0.100	067	310
MgC12	0.005	9	18
NaCl	0.010	20	30
ZnC12	0.001	4.7	5.2
CrC1 ₃ .6H ₂ 0	0.001	0.2	0.3
cuC12.2H20	1000.0	0.4	0.4
rec1 ₃	0.0005	1.7	3.3
MnC12.4H20	0.0005	1.4	1.8
NICL2.6H20	0.0001	0.2	0.3
PbC12	0.0001	0.7	0.3
		543.3	401.6

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Adjust to PH = 4.5 Using HNO₃

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