MECHANICAL PROPERTIES OF FIRE-RETARDANT GLASS FIBER-REINFORCED POLYMER MATERIALS WITH ALUMINUM TRI-HYDRATE FILLER

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ABSTRACT

Aluminum Tri-hydrate (ATH) can be effectively used to increase fire resistance of Fiber-Reinforced Polymer (FRP) materials. This paper studies the effect of ATH filler on mechanical properties of Glass FRP (GFRP) material, based on compression, tension, shear and flexural test results from three types of GFRP materials with the amount of 0% (control), 25%, and 50% ATH filler by weight of the resin. It was found that the control was the strongest for all tests except for flexure, which is 3% lower than the flexural strength of 25% ATH sample. The compressive strength dropped 19% and 25% for 25% and 50% ATH loadings, respectively, compared to the control. For shear and tensile strengths, the 25% ATH sample acted similarly to the control, but the 50% ATH sample had a significantly lower strength. For stiffness, changing the additive amount from 0% to 50% had only small changes for compression, tension, and flexure. It can be concluded that adding ATH generally decreases the strength and makes FRP more brittle. The performance of a 25% ATH loading is comparable to the control except compression, while a 50% ATH loading has a more significant effect on the mechanical properties of the GFRP. The data presented in this paper can be used to develop fire-resistant FRP systems.

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1. INTRODUCTION

Fiber-Reinforced Polymer (FRP) consists of high-strength fibers (glass, carbon, aramid, boron, etc.) embedded in a polymer resin (polyester, vinyl ester, resin, etc.). It is high strength (from the fibers), durable (from the resin), and lightweight. In the last two decades, FRP has gained acceptance and popularity in civil infrastructure applications, including FRP bridge decks, internal FRP reinforcement for concrete and prestressed concrete structures, strengthening of reinforced concrete, masonry, timber and metal with externally bonded FRP, etc. [7].

Numerous studies have also been performed to determine the behavior of FRP for building applications, mostly focused on Carbon FRP (CFRP) and Glass FRP (GFRP). Among others, Mukhopadhyaya et al. [19] studied whether FRP could be used in conjunction with concrete and concluded that it could perform well as a flexural reinforcement for concrete beams. Mirmiran et al. [18] studied the effects that wrapping hybrid concrete beam-columns with FRP would have on the strength and ductility of the beams. Davalos et al. [12] investigated the durability and shear strength by testing the bond strength between the FRP and the wood under both wet and dry conditions. Chowdury et al. [11] looked into the remaining structural strength in building systems after a fire event. Herwig and Motavalli [15] studied the axial behavior of reinforced concrete beams that had been externally strengthened with either lightweight concrete or with unbounded GFRP wrapping, i.e., there was no adhesive or other connecting mechanism between the GFRP wrapping and the concrete. Pantelides et al. [21] investigated the performance of concrete panels reinforced with synthetic fibers, mild steel, and GFRP when subjected to blast
loading. They also investigated the possibility of using a hybrid GFRP/steel concrete panel in order to develop composite action between a concrete-foam composite. The highlight of these efforts was the publications from the Committee of ACI 440, in particular, ACI 440.2R-08 [1], which provided a guide for the design and construction of externally bonded FRP systems for strengthening concrete structures.

However, unlike other civil infrastructure applications, buildings have stringent requirements on flame spread, smoke and toxicity, and fire ratings. Foster and Bisby [14] tested the bond strength between FRP and concrete and FRP to FRP via testing the ultimate tensile strength, elastic modulus, and failure strain at a range of temperatures. Their study showed that GFRP’s bond strength begins to decrease dramatically after 300°C likely due to the glass transition temperature of the polymer being surpassed. Petersen [22] conducted a detailed study on the requirements from International Building Code [16] for FRP to be used for buildings. Based on their findings, FRP with regular resins cannot meet the requirements of flame spread, smoke and toxicity, and fire ratings. These issues are generically addressed in ACI 440.2R-08 [1] as: “FRP-strengthened structures should comply with all applicable building and fire codes. Smoke and flame spread ratings should be determined in accordance with ASTM E 84. Coatings can be used to limit smoke and flame spread.” For fire, ACI 440.2R-08 [1] further states that “due to the low temperature resistance of most fiber-reinforced polymer materials, the strength of externally bonded FRP systems is assumed to be lost completely in a fire.”

Although providing coating as suggested in ACI 440.2R-08 [1] can improve the fire performance, it is costly. A more cost-effective way to address this problem is to add fire-retardant fillers to the resin, as discussed below. The resulting modified resins might even be cheaper than regular resins since some fillers are often less expensive than regular resins.
One of the additives investigated was rice hull ash (RHA). Due to the high silica content in the RHA, it was theorized that RHA could be a good fire retardant when added to a polymer system. Chand et al. [9] investigated the implications of using RHA as an additive to polyester composites. The RHA was added to the resin using volume fractions and after curing, tensile and impact strength was measured. Because RHA is weak in tension, it was found to decrease the tensile strength of RHA-loaded samples. Also it was found that RHA decreases the impact strength of the system using the IZOD impact test. Unfortunately, when RHA is generated at low temperatures it has a black color, which absorbs heat and therefore is not conducive to utilization in green environments. Chakraverty et al. [8] aimed at completing the following objectives through their research: (1) to determine the effects of various acid treatments on process of removing metallic impurities from the RHA, and (2) to determine the effects of acid treatments and different furnace temperatures on the time required to obtain completely white RHA. While it was found that the acid wash did not affect the silica structure of the RHA, it also did not affect the time for complete combustion of the RHA in order to produce a white silica product.

Another additive that was investigated was nanoclay (NC). Nazare et al. [20] investigated the use of NC in polyester resin to reduce smoke generation and improve the fire resistance of the system. They found that the addition of NC helps reduce the flammability because the dispersion of clay in the polymeric matrices produces a nanocomposite structure which allows for reduced flammability as well as improved mechanical properties. In addition, the NC forms a char layer, which helps insulate the FRP below. It was noticed that for higher clay concentrations (>5% clay) in combination with flame retardants, the crosslinking reaction, or the development of the nanocomposite structure in the nanoclays, was noticeably slower or halted completely. In addition to this problem, the resin also became more plastic and had an increased cure time when
the clay concentrations were high. From X-ray diffraction, it was found that little to no 
nanocomposite was formed. From the cone calorimetry test, it was found that the time to 
ignition was improved by 102 seconds. Therefore, nanoclay is a viable option for fire resistance 
in the GFRP.

Wu et al. [23] looked into using NC in conjunction with epoxy adhesives to provide fire 
resistant properties for the epoxy. This study investigated various flame spread and high-
temperature exposure tests. They found that NC can greatly improve the flame retardancy of 
epoxy systems at filling levels of 2%-3%. They also found that NC which was distributed into 
the epoxy using a mixer was better at improving fire properties than NC distributed by hand.

Intumescents were studied by Kandola et al. [17] to see how they changed the burning 
mechanism of composites with polyester resin. Intumescents are often found in paints and 
ceramics, and therefore may be incorporated into a building system. They found that 
intumescents significantly decreased the flaming behavior of the resins tested. However, the 
decrease was not as significant as expected. Also, they found that the intumescents did not help 
the structural integrity of the composite. Therefore, they suggested that intumescents be used as a 
protective coating on thick laminates.

Dholakiay [13] investigated the effects of the following four different additives: (1) 
hydroxyapatite or calcium phosphate, (2) zinc borate, (3) class C fly ash, and (4) antimony 
trioxide. They were tested for their fire resistance using LOI, TGA and IR spectroscopy. The 
materials are considered self-extinguishing if the LOI is greater than 26, and all of the 
composites tested had an LOI in the range of 25-26. It was also found that as the amount of 
fillers increases, so did the fire resistance of the composite. They concluded that while fly ash 
was a good additive, hydroxyapatite and zinc borate were better for increasing LOI. From the
TGA, it was found that composites containing filler have a better thermal stability. Mechanical properties were tested for the samples using Rockwell hardness test and flexural strength tests. From these tests, it was found that the mechanical properties increased as the filler content increased. The exception to this was for antimony trioxide and fly ash.

Liang et al. [27] studied mechanical properties and flame retardancy of polypropylene (PP) composites filled with Microencapsulated Red Phosphorus (MRP) and Magnesium Hydrate [Mg(OH)₂]/Aluminum Hydrate [Al(OH)₃]. They found that the increase of the weight fraction of MRP increases the index of limit oxygen, the rank of smoke density, the Young’s modulus and tensile elongation; but decreases the horizontal burning speed and tensile yield and fracture strengths.

Correia et al. [25] conducted an experimental program to study the behavior of GFRP profiles subjected to fire. Calcium silicate board, vermiculite/perlite based mortar and intumescent coating were used to provide fire protections, which was found to be effective in reducing the temperature throughout the profiles. The protected GFRP pultruded profiles achieved a fire resistance of 60-76 minutes, comparing to 38 minutes for unprotected ones. The addition of water cooling system could further increase the fire resistance to 120 minutes.

Chen et al. [29] developed a new glass fiber/bismaleimide (GF/BMI) composite that could improve the flame retardancy while achieving higher mechanical properties and lower dielectric loss. A new flame retardant resin (BDDP) was added to glass fibers. It was found that the interlaminar shear strength of the GF/BDDP composites were 1.4-1.9 times of those of GF/BMI composites and the flame retardancy was improved from UL 94 V-2 to V-0 grade.

Zhang et al. [28] investigated the mechanical properties and flame retardancy of wood-fiber/polypropylene (PP) composites. Cone colorimeter data (CONE) and LOI results showed that
Ammonium Polyphosphate (APP) and silica are effective flame retardant materials for wood-fiber/PP composites.

Jeencham et al. [26] studied the efficiency of the following flame retardant fillers: APP, Mg(OH)₂, zinc borate (Zb), and a combination of APP, Mg(OH)₂ and Zb in sisal fiber/PP composite. It was found that the addition of APP or a combination of APP and Zb could improve the flame retardancy of sisal fiber/PP composites.

Rowen et al. [24] conducted Cone Calorimeter and ASTM E 84 flame and smoke tests to compare performances among GFRPs with standard resin, which acted as a reference, and resins with three different additives. Sample 1 contained pultruded brominated resin panel with polyester veil, sample 2 contained pultruded panel with alumina trihydrate (ATH) and intumescent veil, and sample 3 consisted of pultruded panel with ATH and expandable graphite intumescent veil. It was concluded that the third sample achieved the best performance based on cone tests, where Peak Heat Release Rate (PHRR) and Average Heat Release Rate (AHRR) were at 47 kW/m² and 21 kW/m², which were significantly lower than the values of 185 kW/m² and 85 kW/m² for sample 1. Improved performance was also observed based on ASTM E 84 test results, where sample 3 achieved Flame Spread Index (FSI) of 20 and Smoke-development Index (SDI) of 125 compared with FSI of 25 and SDI of 985 for sample 1.

Similar conclusions were drawn by Petersen [22]. They compared performances of a wide range of fillers, which included: ATH, a mixture of boric acid and rice hull ash (BA/RHA), coarse graded gypsum (CG), coarse graded limestone (CLS), a mixture comprised of 60% BA, 20% RHA, 10% limestone, and 10% ATH (Conc), fine graded gypsum (FG), fine graded limestone (FLS), nanoclays (NC), and rice hull ash (RHA). In addition, two controls were created with no additives to the resin, one with 1% MEKP hardener and the other with 5%
MEKP hardener. They concluded that by adding ATH to the resin in different amounts, the heat resistive properties of the FRP could be increased. The time to ignition for the 50% ATH was 6 seconds, which was a 4 second improvement from the control samples. Additionally, it was found that with the additional of ATH, the char layer was generated, the quantity of the smoke produced was low, and flame spread could be significantly decreased during fire tests.

As shown above, extensive studies has been conducted on the fire performance of FRP with fillers and shown that the fire resistance of the FRP can be increased by adding various additives to the resin. However, there is only limited study on the effect of the fillers on mechanical behavior, including stiffness and strength, of the FRP system. Other than the preliminary study from Dholakiay [13] as described above, Aziz et al. [6] investigated how modifying polyester resins with additives could change the mechanical properties, glass transition temperature, fracture surface, and bond performance of the systems. They found that adding kenaf, a plant fiber, could improve the mechanical behavior of the composites. However, to the authors’ knowledge, no study is available on the effect of ATH filler on the mechanical properties of the GFRP, which is the objective of this study. To achieve this, comprehensive tests were conducted, as described next.

2. EXPERIMENTAL PROGRAM

2.1 Testing Plan

The experimental program consisted of compression, tension, shear and flexural tests, conducted on FRP coupon samples made of resins with three different amount of ATH, namely 0% (control), 25%, and 50% ATH by weight, with five samples for each test.
2.2 Materials

404 isophthalic resin, A202 ATH, and E-glass Chopped Strand Mat (CSM) were used to manufacture the samples. The material properties are shown in Table 1.

2.3 Workability Test

In past research [13], it was found that as the concentration of additives increased in the resin, so did the fire resistance of the GFRP until the workability was compromised. Therefore, the workability limits need to be found for ATH in order to determine the optimal loading amount. A workability test was performed and the results are shown in Table 2, where samples mixed with ATH were rated on a scale of one to ten with ten being very runny, and one being completely unworkable. It was determined that four was the ideal workability for 51 g/m² (1.5 oz/yd²) of chopped strand mat given that at this workability the resin was still able to completely penetrate the fibers. These tests were conducted in parts per hundred (PPH). For example, 100 PPH means that the resin is 50% resin and 50% ATH. From Table 2, it can be seen that at 25% ATH, or 50 PPH, the resin had a workability of 8 and at 50% ATH, or 100 PPH, the resin had a workability of 4. Therefore, these two were selected for further testing along with a control which contained 0% ATH. Further information on the workability tests can be found from Petersen [22].

2.4 Sample Fabrication

Samples were cast using vacuum bagging methods. First, a layer of visqueen was placed down, followed by a breather cloth and peel ply. Next, one layer of 102 g/m² (3 oz/yd²) chopped strand fiber was laid down, and the appropriate amount of resin, by weight, was added. After the
resin had been applied, the system was pressed down to eliminate air from the system. The process with the resin and the fiber was repeated until the appropriate number of layers had been reached. After the FRP was cast, peel ply was placed on the top, and then another layer of breather cloth. Finally, the visqueen was folded on the top, and the bag was sealed. Before the vacuum bag was opened and the FRP was removed, the FRP was allowed to cure for a minimum of twenty-four hours. Examples of this process can be seen in Figures 1A and 1B.

In order to make the FRP samples, three 61 x 61 cm² (2 x 2 ft²) samples were cast with quantities of 0%, 25%, and 50% ATH by weight of the total amount of resin, i.e., if 50% was used, 50% of the system was resin and the other 50% was ATH. Using a volumetric fraction of the fiber (V_f) of 0.1877 following a previous study [10], for each 102 g/m² (3 oz/yd²) layer of fiberglass, the required weight of the resin was determined to be 894 g/m² (26.37 oz/yd²) per layer. Since the desired thickness of each panel was 6.4 mm (0.25 inches), the required number of plies was calculated to be four.

The resin additive mixture was created by first weighing out the amount of ATH to be used. Next, resin was measured and the premeasured ATH was added. The ATH was then carefully mixed into the resin in order to avoid creating air bubbles. Next, 5% MEKP hardener was added by weight of the total amount of resin. This was then also carefully mixed, and the resin and additive was poured onto the FRP and smoothed evenly over the entire surface. After the resin had been spread over the FRP, a roller was used to eliminate air pockets between layers of fiberglass. This process was repeated for each layer of FRP. The goal was to achieve a 6.4 mm (0.25 inches) thickness. However, the 50% ATH system was thicker, as will be shown next, because the resin was too thick to properly soak the fibers.
2.5 Sample details

After the composite had cured for a minimum of twenty-four hours, it was removed from the vacuum bag and placed under a fume hood until it was cut into the testing samples. From each 61 x 61 cm² (2 x 2 ft²) panel, five samples for each test were cut according to their ASTM standard measurements. Compression samples were cut to be 12.7 x 81.3 mm² (0.5 x 3.2 in²), as can be seen in Figure 2. Tension samples were 25.4 x 254 mm² (1 x 10 in²) tabs with 76.2 mm (3 inches) tabs at each end. After the 76.2 mm (3 inches) tabs, the samples were narrowed down to 12.7 mm (0.5 inches) wide, as shown in Figure 3. The samples for shear were 19.1 x 76.2 mm² (0.75 x 3.0 in²) tabs, with 45° notches cut from the middle of the span, as shown in Figure 4. Flexural samples were cut to be 50.8 x 381 mm² (2 x 15 in²), as shown in Figure 5.

2.6 Test Setup

All tests were conducted on an MTS 318.10 machine with a Flex Test SE control unit. Five samples were tested for each type of the test. Loading rates were specified by respective ASTM standards, as shown in Table 3. Loads and displacements were recorded by the MTS machine and a separate National Instruments Data Acquisition System (DAS) was used to record the strain.

2.7 Test Methods

2.7.1 Compression (ASTM D 695) [3]

For compression test, the samples were placed in the fixture as shown in Figure 6. The bolts were then hand tightened to the point that there was a limited amount of resistance. This was done because if the bolts were loose then the samples could buckle, but if they were too tight
then the fixture and the sample could form a steel-GFRP composite. The buckling case would provide data resulting in strength that was less than the actual strength of the composite, while the composite case would create a strengthened system, and therefore the values for the strength would be greater than the actual strength of the FRP.

### 2.7.2 Tension (ASTM D 638) [5]

For tension test, Syntech grips were used. The grips were able to clamp samples that were up to 6.4 mm (0.25 inches) thick. Therefore, the samples were filed to a thickness of 5.8 mm (0.23 inches) in order to allow the samples to fit into the grips and prevent failures from occurring at the grips. For testing, the grips were hand tightened around the samples until a force of 222 N (50 lbs) was applied, and then the test was run. The system for the tension testing can be seen in Figure 7.

### 2.7.3 Shear (ASTM C 1292) [4]

Each shear sample was inserted into the fixture as can be seen in Figure 8. Care was taken to align the sides of V-notches at the appropriate location on the shear fixture, so that the start of the bottom V-notch would line up with where the fixture started to slope. In addition, the front of the samples was flush with the front of the fixture.

### 2.7.4 Flexure (ASTM D 790) [2]

For flexure, a three point bending test was performed using a custom-made fixture with a span of 279 mm (11 inches), as shown in Figure 9. Therefore, the sample extended past the fixture 51 mm (2 inches) on each end. The load was applied at the center of the sample.
2.8 Strain Gage Placement

Strain gages were placed on the compression, tension and flexural samples. No strain gages were used for shear samples. However, accuracy was checked by comparing the results to previous studies. For each test, strain gages were attached to four of the five samples for each additive quantities (0%, 25%, and 50%). For compression, one strain gage was placed at the center of the sample, as shown in Figure 6. For tension, a strain gage was placed at the middle of each sample, as shown in Figure 10. For flexure, two strain gages were attached to the top and bottom at the center of each sample, as can be seen in Figure 11.

3. TEST RESULTS

3.1 Compression (ASTM D 695) [3]

3.1.1 Stiffness and Strength

For compression test, force-displacement curves as shown in Figure 13 were obtained from MTS machine, and stress-strain curves were developed from both the DAS and force-displacement curves from the MTS machine, as shown in Figure 12, where good correlations can be observed between the two systems. However, the strain gages stopped working before the samples failed. Therefore, the data from the MTS machine were used to calculate the strain to develop complete stress-strain curves.

It is noted that the force-displacement curves in Figure 13 show that the strength of 50% ATH sample is higher. This is because the 50% ATH samples were 2.4 mm (0.094 inches) and 2.6 mm (0.103 inches) thicker than the control and 25% ATH samples, respectively, as shown in Table 4. Therefore, the stress-strain curves from the MTS machine shown in Figure 14, which
take into account the section area differences, are better representative of the actual strength of the composites. Also, from this graph, the modulus of elasticity for each additive amount was calculated using a linear trend line. In order to add the trend line, first the initial section due to equipment settlement was removed, and then a best fit line was created, as shown in Figure 15 for a representative sample. The equation for the best fit line can be seen on the graph, where the slope is the modulus of elasticity. The modulus of elasticity for each group is reported in Table 4, and a comparison between the control, 25%, and 50% ATH including their error bars is shown in Figure 16, which indicates that the compressive modulus of elasticity is fairly similar from the control to the 50% ATH, with a maximum of 5% difference.

Table 4 summarizes the values for the maximum forces, stresses, and displacements for all the three additive amounts as well as those from previous research [10]. The maximum stress is higher than those from previous tests [10] since the resin and fiber are different, but they are still within a similar range. Similar to Figure 13, the area has not been taken account in the forces, and therefore indicates that 50% ATH has the highest ultimate load. But when adjusted for the differences in the cross section areas of the samples, it can be seen from Table 4 and Figure 17 that the composite with the highest maximum stress is the control with an ultimate stress of 178 MPa (25.8 ksi). The next strongest is the 25% ATH which has an ultimate stress of 145 MPa (21.0 ksi), followed by the 50% ATH which has an ultimate stress of 138 MPa (20.0 ksi). Therefore, there is a 19% reduction in ultimate stress from the control to the 25% ATH, and a 23% reduction from the control to the 50% ATH.

Although the maximum deflections are reported in Table 4, they cannot be directly compared and used to calculate the maximum strain since they include the initial settlement. However, because the load-displacement curves are linear excluding the initial part due to the equipment
settlement, i.e., the materials are elastic in compression, the maximum strain can be calculated as the maximum stress divided by the modulus of elasticity, with values shown in Table 4. It can be seem from Figure 18 that, comparing to control samples, the maximum strain drops 14% and 20% for the 25% ATH and 50% ATH samples, respectively, indicating that adding ATH generally makes the samples more brittle in compression.

3.1.2 Failure Mode

There were three crack patterns that occurred in all the three ATH amounts, as can be seen in Figure 19, in which the cracks have been highlighted with a pink dye for visibility. The crack patterns did not seem to have significant impact on the maximum stress or displacement of their respective composite groups.

3.2 Tension (ASTM D 638) [5]

3.2.1 Stiffness and Strength

Similar to compression test, force-displacement curves were created from the data obtained from the MTS machine, and stress-strain curves were developed from both the MTS machine and the DAS. As shown in Figure 20, good correlations can be observed between the two systems, but the strain gages stopped working before the samples failed. Therefore, the displacement data from the MTS machine were used to calculate strain in the following analysis.

Unlike the results from the compression test, there was no thickness change between the control, the 25% ATH, and 50% ATH samples since they were all cut to have a constant thickness in order to fit in the tension grips, as shown in Table 5. Therefore, the force-displacement curves, as shown in Figure 21, and the stress-strain curves, as shown in Figure 22,
are similar, both correctly representing that the 50% ATH sample is significantly weaker than the other two composites. Also, from Figure 21 and Figure 22, it can be seen that the control and the 25% ATH sample act in similar fashions, with similar slopes and the maximum loads and stresses.

Using the same methods as described for compression tests, modulus of elasticity can be calculated, as shown in Table 5 and Figure 23. It can be seen that all samples are fairly similar, with the 25% ATH sample 6% higher and the 50% ATH sample 5% lower than the control samples.

Table 5 compares the differences in strength among all three laminates and those from previous tests. Figure 24 presents the difference in maximum stress with error bars for the various additive amounts. It can be seen that the control is able to hold the most load and stress. The maximum stresses of the 25% and 50% ATH samples are 9% and 47% lower than the control samples, respectively.

The maximum strain can be calculated in the same way as compression test, with values shown in Table 5. It can be seen from Figure 25 that, comparing to control samples, the maximum strain drops 14% and 44% for the 25% ATH and 50% ATH samples, respectively, indicating that adding ATH also makes the samples more brittle in tension.

3.2.2 Failure Mode

For tension, the fracture type was consistent throughout all the resin types. A crack began in the center of the sample, and then propagated across the 12.7 mm (0.5 inches) mid-section until pull-apart occurred, as shown in Figure 26.
3.3 Shear (ASTM C 1292) [4]

3.3.1 Stiffness and Strength

Similar to compression tests, force-displacement curves were obtained from the data from the MTS machine for the shear tests, as shown in Figure 27. Due to time constraints and back order of the shear strain gages, strain gages were not used for shear samples; therefore no stress-strain curve was developed. However, in order to verify the data, the maximum stresses were calculated and compared to previous research as can be seen in Table 6, which also reports the maximum forces and displacements recorded during the test. Figure 28 shows the difference in the maximum stress among different additive amounts. As can be seen from Table 6 and Figure 28, the strengths of the control and 25% ATH samples were about the same, but there was a reduction in strength of 27% from the control to 50% ATH samples.

3.3.2 Failure Mode

The three different loading amounts all had slightly different failure types as can be seen in Figure 29. The control failed with no sign of delamination, while both 25% and 50% ATH samples had shear failures and also slight delamination failures. In addition, the 50% ATH sample had one sample that had a catastrophic failure. The failure started at the bottom of the samples at the V-notch, and then propagated up to the top of the sample.

3.4 Flexure (ASTM D 790) [2]

3.4.1 Stiffness and Strength

Similar to compression test, load-displacement curves were obtained from the data from the MTS machine. However, due to the nature of the flexural test, stress-strain curves were only
created from the strain data from the DAS combined with the stress data from the MTS and no stress-strain curve was developed solely from the MTS data. The stress-strain curves from the DAS can be seen in Figure 31, and the force-displacement curve from the MTS can be seen in Figure 30. Unlike elastic behavior observed from previous tests, it is noted that the 50% ATH samples assume a nonlinear behavior. This is probably because the samples are thicker, as shown in Table 7, and much stiffer. Any small amount of slippage at the support will affect the displacement.

Similar to the compression test, the strain gages stopped working before failure occurred. Therefore, the force-displacement curves were used to calculate the stiffness. The mid-span deflection of the sample under 3-point loading can be calculated as:

\[
\Delta = \frac{PL^3}{48EI}
\]  

(1)

where \( \Delta \) is the displacement from the MTS machine, \( P \) is the force from the MTS machine, \( L \) is the distance between supports on the sample, \( E \) is the modulus of elasticity, and \( I \) is the moment of inertia as:

\[
I = \frac{bh^3}{12}
\]  

(2)

where \( b \) is the width and \( h \) is the depth of the sample, as shown in Figure 5.

From the initial linear region in each load-displacement curve, slope of the curve, \( P/\Delta \) can be obtained. From Eq. (1), the flexural modulus of elasticity can be calculated using:

\[
E = \frac{P}{\Delta} \times \frac{L^3}{48I}
\]  

(3)
The results from these calculations are listed Table 7. A comparison of modulus of elasticity from the control to the other additive amounts can be seen in Figure 32, which shows that the control and 25% ATH samples are similar, but the modulus of 50% ATH is 11% lower than the control sample.

Table 7 summarizes the maximum forces, stresses, deflections and the modulus of elasticity. The maximum stresses with their error are shown in Figure 33. It can be seen that strength of the control and 25% ATH are about the same, with 3% difference. Comparing to control samples, there is a decrease of 21% for 50% ATH samples.

### 3.4.2 Failure Modes

The fracture types for all three quantities of ATH were fairly similar. The crack first formed at the middle of the span, and then propagated both upwards and across the sample, as shown in Figure 34 and Figure 35, where Figure 34 shows the bottom of the samples and Figure 35 shows the side of the samples. The failure initiated from the bottom of the sample and then spread up to the top. The fractures are dyed pink for better visibility.

### 4. DISCUSSION

As shown above, adding ATM generally decreases the strength of GFRP. Although the strength is decreased, it is still high enough (about 1/3 of the strength of the steel) to be used for structural applications. The difference is that more materials are required to take the same load. Additionally, the strength reported in this paper was based on the specific fiber volume fraction, $V_f$, adopted in this study. Higher strength can always be achieved with higher $V_f$. 
5. CONCLUSIONS

Three layups containing varying amounts of ATH ranging from 0%, 25% to 50% by weight were tested in order to evaluate the effects of ATH on the mechanical properties of FRP. The tests, including compression, tension, shear and flexure, were performed according to ASTM standards. The following conclusions can be drawn from this study:

1) Strengthwise, the control was the strongest for all of the tests except for flexure, which is 3% lower than the flexural strength of 25% ATH sample. For compression, there is a 19% and 25% reduction from the control to 25% and 50% ATH samples, respectively. For tension, shear, and flexure, the 25% ATH sample acted similarly to the control. However, the 50% ATH sample had a significantly lower maximum stress.

2) The moduli of elasticity were compared for all three additive loadings, and it was found that changing the additive amount of ATH has insignificant effect on stiffness.

3) Adding ATH generally decreases the strength and makes the FRP more brittle.

4) Using 25% ATH by weight in conjunction with isophthalic does not have a significant effect of the stiffness and strength comparing to control samples except compression, while 50% ATH has more significant effect.

The data presented in this paper can be used to develop fire-resistant FRP systems. It should be pointed out the findings are based on the ATH and resin used in the study. Further study is recommended for other types of fire-retardant fillers. Other properties, such as impact properties, are also important for FRP material, which deserves further study.
6. ACKNOWLEDGEMENTS

We gratefully acknowledge the US Department of Energy for financial support. We appreciate the help from Dean Larry Stauffer (College of Engineering) and Chair Richard Nielsen (Department of Civil Engineering) from University of Idaho. We thank Dr. Lloyd Smith (Washing State University), Dr. Pizhong Qiao (Washington State University) and Dr. Edwin Odom (University of Idaho) for their testing fixtures and advice. We also thank Tony Parris, Nicolas Pena and Dallas Roberts for helping with the test.

7. REFERENCES

[1] ACI 440.2R-08. Guide for the Design and Construction of Externally Bonded FRP Systems for Strengthening Concrete Structures, American Concrete Institute, Farmington Hills, MI.


Table 1: Material Properties of the Resin and ATH

<table>
<thead>
<tr>
<th>Type</th>
<th>Tensile Strength (MPa)</th>
<th>Tensile Modulus (GPa)</th>
<th>Compressive Strength (MPa)</th>
<th>Density</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resin 404 Isophthalic Resin</td>
<td>50</td>
<td>37</td>
<td>88</td>
<td>1100 kg/m³</td>
</tr>
<tr>
<td>ATH A 202</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>561 kg/m³</td>
</tr>
<tr>
<td>E-Glass Fiber Chopped Strand Mat</td>
<td>2400</td>
<td>73</td>
<td>--</td>
<td>2600 kg/m³</td>
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</table>

Table 2: Workability Results

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<thead>
<tr>
<th>PPH</th>
<th>Workability</th>
<th>Comments</th>
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<tr>
<td>40</td>
<td>8</td>
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<tr>
<td>50</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>60</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>70</td>
<td>6</td>
<td></td>
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<tr>
<td>80</td>
<td>5</td>
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<tr>
<td>90</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>100</td>
<td>4</td>
<td>honey like</td>
</tr>
<tr>
<td>110</td>
<td>3</td>
<td>Un-pourable, still spreadable</td>
</tr>
<tr>
<td>120</td>
<td>3</td>
<td>very un-pourable, still semi-spreadable</td>
</tr>
<tr>
<td>130</td>
<td>2</td>
<td>waxy texture</td>
</tr>
</tbody>
</table>

Table 3: Loading Rates

<table>
<thead>
<tr>
<th>ASTM</th>
<th>Test</th>
<th>Loading Rate</th>
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<tbody>
<tr>
<td>Modified D 695</td>
<td>Compression</td>
<td>1.3 mm/min</td>
</tr>
<tr>
<td>D790</td>
<td>Flexure</td>
<td>1.3 mm/min</td>
</tr>
<tr>
<td>D638</td>
<td>Tension</td>
<td>5.1 mm/min</td>
</tr>
<tr>
<td>C1292</td>
<td>Shear</td>
<td>8.4 mm/min</td>
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Table 4: Maximum Force, Stress and Displacement for Compression

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Previous Research (Chen and Davalos [10])</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average:</td>
<td>STDEV:</td>
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<tr>
<td>Thickness (mm)</td>
<td>7.9</td>
<td>0.3</td>
</tr>
<tr>
<td>Max Load (kN)</td>
<td>16</td>
<td>1</td>
</tr>
<tr>
<td>Max Stress (MPa)</td>
<td>178</td>
<td>18</td>
</tr>
<tr>
<td>Max Δ (mm)</td>
<td>2.41</td>
<td>0.26</td>
</tr>
<tr>
<td>Stiffness E (MPa)</td>
<td>6758</td>
<td>535</td>
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<tr>
<td>Max Strain (mm/mm)</td>
<td>0.026</td>
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</tr>
<tr>
<td>25% ATH</td>
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<td></td>
</tr>
<tr>
<td>Thickness (mm)</td>
<td>7.6</td>
<td>0.3</td>
</tr>
<tr>
<td>Max Load (kN)</td>
<td>14</td>
<td>2</td>
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<tr>
<td>Max Stress (MPa)</td>
<td>145</td>
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<tr>
<td>Max Δ (mm)</td>
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<td>Stiffness E (MPa)</td>
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<td>Max Strain (mm/mm)</td>
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Table 5: Average Maximum Force, Stress and Displacement for Tension

<table>
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<th>Previous Research</th>
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</thead>
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<tr>
<td></td>
<td>Average:</td>
<td>STDEV:</td>
</tr>
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<tr>
<td>Max Load (N)</td>
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<td>Max Stress (MPa)</td>
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<td>Max Δ (mm)</td>
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<tr>
<td>Stiffness E (MPa)</td>
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<tr>
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</tr>
<tr>
<td>25% ATH</td>
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<td></td>
</tr>
<tr>
<td>Thickness (mm)</td>
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<td>0.5</td>
</tr>
<tr>
<td>Max Load (N)</td>
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<td>Max Stress (MPa)</td>
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<td>5</td>
</tr>
<tr>
<td>Max Δ (mm)</td>
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<td>0.5</td>
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<tr>
<td>Stiffness E (MPa)</td>
<td>6330</td>
<td>815</td>
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<tr>
<td>Max Strain (mm/mm)</td>
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Table 6: Average Maximum Force, Stress and Displacement for Shear

<table>
<thead>
<tr>
<th>Thickness (mm)</th>
<th>Max Load (N)</th>
<th>Max Stress (MPa)</th>
<th>Max Δ (mm)</th>
<th>25% ATH</th>
<th>50% ATH</th>
</tr>
</thead>
<tbody>
<tr>
<td>7.5</td>
<td>7037</td>
<td>83</td>
<td>2.0</td>
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<td>7.8</td>
<td>7741</td>
<td>112</td>
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Table 7: Average Maximum Force, Stress and Displacement for Flexure

<table>
<thead>
<tr>
<th>Thickness (mm)</th>
<th>Max Load (N)</th>
<th>Max Stress (MPa)</th>
<th>Max Δ (mm)</th>
<th>Stiffness E (MPa)</th>
<th>25% ATH</th>
<th>50% ATH</th>
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<tbody>
<tr>
<td>7.3</td>
<td>871</td>
<td>132</td>
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<td>7741</td>
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<table>
<thead>
<tr>
<th>Thickness (mm)</th>
<th>Max Load (N)</th>
<th>Max Stress (MPa)</th>
<th>Max Δ (mm)</th>
<th>Stiffness E (MPa)</th>
<th>50% ATH</th>
</tr>
</thead>
<tbody>
<tr>
<td>10.0</td>
<td>1319</td>
<td>104</td>
<td>32.0</td>
<td>6877</td>
<td>50% ATH</td>
</tr>
</tbody>
</table>

Figure 1: Vacuum Bagging Process

(A) FRP sealing bag  (B) After sealing bag

Figure 2: Compression Dimensions

Figure 3: Tension Dimensions

Figure 4: Shear Dimensions
Figure 5: Flexure Dimensions

Figure 6: Compression Testing Fixture

Figure 7: Tension Testing Fixture and Failed Tension Sample
Support

Figure 8: Shear Testing Fixture and Failed Sample

Figure 9: Flexural Testing Fixture

Figure 10: Placement of Tension Strain Gage
Figure 11: Placement of Flexure Strain Gage

![Strain Gages]

Figure 12: MTS vs. DAS for Compression with 50% ATH

Figure 13: Force-Displacement Curves for Compression
Figure 14: Stress-Strain Curves for Compression

Figure 15: Stiffness Calculation
Figure 16: Comparison for Compressive Modulus of Elasticity

Figure 17: Max Compressive Stress with Error Bounds
Figure 18: Comparison of Maximum Strain for Compression

Figure 19: Failure Mode
Figure 20: MTS vs. DAS for Tension with 50% ATH

Figure 21: Force-Displacement Curves for Tension
Figure 22: Stress-Strain Curves for Tension

Figure 23: Comparison of Tensile Modulus of Elasticity
Figure 24: Max Tensile Stress with Error Bounds

Figure 25: Comparison of Maximum Strain for Tension

Figure 26: Failure Mode
Figure 27: Force-Displacement Curves for Shear

Figure 28: Max Shear Stress with Error Bounds
Figure 29: Failure Mode for (A) Control, (B) 25% ATH, (C) 50% ATH

Figure 30: Force-Displacement Curves for Flexure
Figure 31: Control Stress-Strain Curve from DAS

Figure 32: Comparison of Flexural Modulus of Elasticity
Figure 33: Comparison of Maximum Flexural Stress

Figure 34: Failure Mode, Bottom View of (A) Control, (B) 25% ATH, (C) 50% ATH

Figure 35: Failure Mode, Side View of (A) Control, (B) 25% ATH, (C) 50% ATH