Improving the thermal properties of fluoroelastomer (Viton GF-600S) using acidic surface modified carbon nanotube

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Abstract

Acid surface modified carbon nanotube (MCNT)-, Carbon nanotube (CNT)-filled fluoroelastomer (FE) and unfilled-FE were prepared (MCNT/FE, CNT/FE and FE). The compounds were subjected to thermogravimetric analysis (TGA) and heat air aging, and characterized by Energy Dispersive X-Ray (EDX). Results showed that MCNT improved the thermal properties of FE, resulting in a larger amount of FE and char remaining in the temperature range of 400-900 °C relative to unfilled FE and CNT/FE. The MCNT/FE TGA curve shifted towards higher temperatures compared to CNT/FE and FE. The same results also revealed that higher percentages of FE were undegraded or less degraded especially near MCNT in the temperature range of 400-540 °C. Energy Dispersive X-Ray (EDX) results indicated that the percentage of carbon and fluorine in the residue of TGA scans, up to 560 °C, of MCNT/FE were the same as CNT/FE, and were higher than FE. EDX results of TGA residue (run up to 900 °C) showed that most of the undegraded FE, which was not degraded at temperatures below 560 °C, was degraded from 560 °C to 900 °C in both MCNT/FE and CNT/FE, with the char in MCNT/FE being more than that in CNT/FE. EDX analysis of thermal aged specimens under air showed that, with increasing aging time, a greater percentage of C, O and F was lost from the surface of filler/FE and FE. The order of element loss after 24 hour aging time was: MCNT/FE > FE > CNT/FE.

Keywords: nanocomposites, fluoroelastomer, acidic surface modified, carbon nanotube, thermal properties, thermal aging.

1. Introduction

Viton is a synthetic rubber and fluoroelastomer (FE). Viton\textsuperscript{a} GF-600S is a terpolymer of hexafluoro-propylene, vinylidene fluoride and tetra-fluoroethylene with a cure site monomer. It is a peroxide cure, 70% fluorine FE\textsuperscript{1-2}.

Normally, in the formulation of FE, carbon black (CB) is used. Replacing CB with surface modified CNT (MCNT) is expected to improve the thermal properties of FE, which is very effective in making O-rings especially for the oil and gas industries\textsuperscript{3-5}. MCNT is also expected to improve thermo-oxidative degradation resistance of FE composites when subjected to thermal aging conditioning in air.

MCNT improves the properties of fluoropolymers, such as crystallinity, electrical response, mechanical properties, viscoelastic behavior, etc, and therefore their thermal stability. This finding has been reported by a number of researchers and examples of literatures and the reasons for the changes in these properties are mentioned bellow.

It has been reported\textsuperscript{6} that the use of multi-walled carbon nanotubes (MWCNTs)-embedded in fluorinated rubber leads to a rubber nanocomposite with cellulation structure. This structure possesses improved thermal resistance. Pham et al.\textsuperscript{7} observed that with increased MWCNT loading in FE/MWCNTs nanocomposites, there is a steady increase in the decomposition temperature. Interactions between CNT and fluoropolymers and the effect of surface area of CNT on the active interfacial area between them were investigated by a number of researchers, for examples: A strong interaction between fluoropolymers and CNT was reported by He et al.\textsuperscript{8}. The high aspect ratio of nanotubes also leads to a great increase in the active interfacial area between CNT and the fluoropolymer chain as reported by Chae et al.\textsuperscript{7}. An energetic relationship between the surface of CNT and fluoropolymer has been reported by Levi et al.\textsuperscript{9}.

CNT can change the mechanical and thermo-mechanical properties of polyvinylidene fluoride (PVDF)\textsuperscript{9-11} or the mechanical properties of polytetrafluoroethylene (PTFE)\textsuperscript{12,13}. CNT can also change the degree of crystallinity of fluoropolymers and, therefore, their properties\textsuperscript{14-18}.

Sementsov et al.\textsuperscript{11} reported that in nanocomposites of PTFE and MWCNT, the concentration and nature of oxygen containing MWCNT surface groups influences the strength parameters of the composite material. Wen et al.\textsuperscript{17} prepared (PVDF)/MWCNTs nanocomposites and studied the melt viscoelastic behavior of the composites. The results showed that PVDF/MWCNTs-g-OH (OH grafted) nanocomposites exhibit more significant solid-like behavior than PVDF/MWCNTs nanocomposites. Huang et al.\textsuperscript{16} showed that the interfacial interaction between the single walled carbon nanotubes (with hydroxyl groups) and PVDF has an effect on inducing crystallinity. The thermal stability of the PTFE composites is also enhanced by the presence of CNT as reported by Park et al.\textsuperscript{18}. The temperatures recorded at maximum decomposition rate were affected by the type of surface modification of MWCNT. The thermograms
for PTFE/MWCNT composites also shifted towards high temperatures.

Carabineiro et al.\textsuperscript{[19,20]} prepared CNT/PVDF composites using modified CNT samples. According to their research results, surface modifications of MWCNTs had an effect on the electrical response and the degree of crystallinity of the CNT/PVDF nanocomposites. Similarly, Xu and Yang\textsuperscript{[21]} prepared FE/MWCNT composite films using different surface modified CNTs. The better mechanical properties of CF4 plasma-modified CNT (FCNTs) composites over other studied CNTs is due to the better dispersion and enhanced chemical compatibility from introducing electron-rich fluorine atoms and also from hydrogen bonding.

The effect of thermal aging conditioning on FE in air and/or other environments were also verified by researchers. For example: the physical and chemical changes associated with degradation on the surface, near the surface and bulk of the FE seals in engine oil and additives were investigated by Smith et al.\textsuperscript{[22]}. Degradation of these materials was reported to be limited to the near surface region of the samples to a depth of less than approximately 50 μm. The process resulted in the release of either F\textsubscript{2} or HF.

Research on the verification of the effect of MCNT on the amount of residues in TGA runs of MCNT/FE nanocomposites is rather rare. In addition, comparison of the amount of residue in TGA runs of MCNT/FE, CNT/FE and FE detailing elemental composition is seldom published. Therefore, this investigation is unique in the sense that knowledge of the amount of residue and the composition of the elements in the residue gives an insight into the ability of MCNT to improve the thermal stability of CNT/FE and FE composites. Furthermore, the presence of elemental fluorine on the surface of MCNT/FE aged in air indicates whether MCNT can improve the resistance of the nanocomposites to thermo-oxidative degradation or not.

In the present work, MCNT was used as a filler for Viton® GF-600S with the aim of improving the thermal stability and thermal aging resistance of FE. The thermal properties of the composites MCNT/FE were assessed by Thermogravimetric Analysis (TGA) and compared vis-à-vis those of CNT filled and unfilled FE composites in our previous work\textsuperscript{[23]}. Energy Dispersive X-Ray (EDX) was also used to characterize the elemental composition of the degraded FE and char after TGA runs at temperatures below 900 °C. This was to ascertain whether the incorporation of MCNT could preserve more of the degraded FE and char compared to CNT/FE or unfilled FE. EDX was also employed to verify the amount of Zn in the residue and to confirm a possible reaction with MCNT. Thermal aging conditioning in air together with EDX was also conducted to investigate the elemental composition of the surfaces of MCNT/FE as a function of aging time.

2. Experimental

2.1 Materials and compounding procedure

The materials used were Viton GF-600S fluoroelastomer, FE; organic peroxide, Luperox 101 XL-45; acid surface modified carbon nanotube MCNT (TNMC8, -COOH Content:0.49wt%), carbon nanotube CNT (TNM8) both with an outside diameter > 50 nm, inside diameter:5-15nm, purity > 95%, and length 10-20 μm; zinc oxide; and triallylisocyanurate, TAIC supplied by ERIKS Sdn. Bhd. (Malaysia), Arkema Sdn. Bhd. (Malaysia), Chengdu Organic Chemicals Co. Ltd. (Chinese Academy of Sciences, China), Texchem Materials Sdn. Bhd. (Malaysia) and Liu Yang San Ji Chemical Trade Co. Ltd. (China) respectively. Three formulations were compounded, MCNT filled FE (MCNT/FE), CNT filled FE (CNT/FE) and unfilled FE (FE). The amounts of FE, organic peroxide, zinc oxide and TAIC in these compounds were 70.0, 2.1, 2.1 and 2.1 g respectively. For MCNT/FE and CNT/FE, 7.0 g filler was used.

Mixing FE with additives was done using a laboratory scale two roll mill with a roll temperature of 48 °C. FE in the above mentioned composition was supplied to the open roll. A uniform band was formed while three rolling cuts from each side of the mill were made, so that the polymer would be uniform and sufficiently warmed up. In the next stage, pre-blended ZnO and TAIC were added uniformly into the gum and three rolling cuts from each side of the mill were made. After setting the roll distance to 1.1 mm, MCNT was then fed in. The compound was then tightly milled ten times. The roll distance was then adjusted to 1.1 mm, and the peroxide was added and, after the final five to six rolled up end passes, the mixture was supplied to the open roll and sheeted. After 24 hours, re-milling was done with a roll temperature of 26 °C. A similar procedure was used for CNT/FE and FE.

2.2 Curing and post curing

Curing of the FE compound was done in molds (15 cm × 15 cm × 1 mm and 18 cm × 18 cm × 2 mm) in a heated press, at 177 °C under a pressure of 10 MPa for 7 min. The post curing was done in an oven at 232 °C for two hours. The conditions for curing and post curing were recommended by the supplier.

2.3 Thermal gravimetric analysis (TGA)

25-35 mg of post-cured sample was subjected to TGA runs from room temperature to 900 °C. This was carried out on a Perkin-Elmer thermal analysis system, model Pyris Diamond TG/DTA, at a scan rate of 20 °C.min\textsuperscript{-1} and under nitrogen atmosphere of 20 ml.min\textsuperscript{-1}. In addition, TGA runs from room temperature to 560 °C were carried out under the same conditions and then cooled to room temperature at a scan rate of 100°C.min\textsuperscript{-1}. The same procedure was repeated with the temperature raised to 900 °C. The specimens were designated according to temperature, for example MCNT/FE-560 °C, where the temperature after the specimen designation is the highest temperature of the TGA run. The same procedure was applied for 3.5-5 mg of pure MCNT.

2.4 Thermal aging

Thermal aging was carried out according to ASTM D-573 for 24, 48 and 72 hours at 250 °C. The samples (5 mm × 5 mm × 2 mm) were named FE, FE-24, FE-48 and FE-72, where the numbers following FE indicate the aging time in hours. Similar types of abbreviation were used for the MCNT/FE and CNT/FE aged samples.
2.5 Energy Dispersive X-Ray (EDX) and Field Emission Scanning Electron Microscopy (FESEM) image analyses

EDX analysis was conducted using FESEM Philip XL-40 (UK) coupled with EDX. The residue obtained from the heating-cooling TGA scans was first applied on an aluminum stub using double sided copper tape (3M Company), then coated with gold for EDX elemental analysis to determine the elements of C, O, F, Si, Ca and Zn.

Elemental compositions of the surfaces of the aged and unaged samples were obtained using EDX. Un-aged and aged samples, 5 mm x 5 mm x 2 mm in dimension, from the molded post cured FE, CNT/FE and MCNT/FE rubber sheets were used for EDX analysis. The major elements characterized were C, O, F and Si. The surface images for some of the samples were obtained by FESEM. In FESEM imaging of MCNT/FE, the razor cut (cross-section) surface was used. Specimens for EDX and FESEM imaging analysis were applied on a copper tape and gold coated as mentioned above. However, for FESEM images of residues obtained from MCNT/FE-560 °C and MCNT/FE-900 °C, no gold coating was used.

3. Results and Discussion

3.1 TGA

Figure 1 shows the TGA thermograms of MCNT, FE, MCNT/FE and CNT/FE. The extracted data from these plots are presented in Table 1. In Table 1, $T_{\text{onset}}$, $T_{5\%}$, $T_{10\%}$ and $T_{50\%}$ are the onset temperature at which the compound started to lose weight (degrade), and temperatures at which the compound lost 5%, 10% and 50% weight respectively. $T_{\text{onset}}$, $T_{5\%}$, $T_{10\%}$ and $T_{50\%}$ increased for MCNT/FE compared to CNT/FE and FE. As can be seen from Figure 1 the MCNT/FE curve shifted towards higher temperatures and the thermal stability improved compared to CNT/FE and FE. As can be seen from Figure 1 the MCNT/FE curve shifted towards higher temperatures and the thermal stability increased compared to CNT/FE. Figure 1 shows the amount of this temperature (thermal stability) shift versus weight%. Figure 1 shows that for the weight% of 100 to 40% the amount of thermal stability shift is almost constant (≈10 °C). However, for weight% between ≈20 to 40% the thermal stability shift increases suddenly (≈20ºC). This increase in thermal stability shift indicates that in this weight% range compared to the 100 to 40 weight% range, FE in MCNT/FE showed more thermal stability improvement relative to FE in CNT/FE.

Pham et al.[4], used a SEM micrograph of the ash residue collected after 90% degradation of the compound and showed that less or partial degradation of the polymer chains attached to the nanotubes occurred whereas extensive degradation occurred in the bulk. Therefore, the higher thermal stability shift for weight% between ≈20 to 40% represents the part of FE which is near or attached to MCNT. All of the above mentioned results show that FE in MCNT/FE is more thermally stable than FE in FE under N$_2$. Furthermore in MCNT/FE, FE near MCNT is considerably more thermally stable than FE in the bulk. FE near MCNT in MCNT/FE to a larger extent is more thermally stable than both FE near CNT and also FE in the bulk in CNT/FE.

It can be concluded that a highly filled FE with MCNT will have a high percentage of FE near or attached to MCNT which can have thermal stabilities up to 540 °C. Therefore

<table>
<thead>
<tr>
<th>Specimen</th>
<th>$T_{\text{onset}}$ (°C)</th>
<th>$T_{5%}$ (°C)</th>
<th>$T_{10%}$ (°C)</th>
<th>$T_{50%}$ (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FE</td>
<td>423.4</td>
<td>442.6</td>
<td>470.7</td>
<td>497.8</td>
</tr>
<tr>
<td>CNT/FE</td>
<td>421.1</td>
<td>446.2</td>
<td>475.4</td>
<td>505.1</td>
</tr>
<tr>
<td>MCNT/FE</td>
<td>433.1</td>
<td>457.7</td>
<td>485.4</td>
<td>513.7</td>
</tr>
<tr>
<td>MCNT</td>
<td>627.7</td>
<td>693.6</td>
<td>748.4</td>
<td>-</td>
</tr>
<tr>
<td>CNT</td>
<td>594.5</td>
<td>640.2</td>
<td>692.9</td>
<td>-</td>
</tr>
</tbody>
</table>

Figure 1. Heat and cool TGA curves of pure MCNT, FE and filler/FE: (a) zoomed and (b) unzoomed.

Figure 2. Thermal stability shift curve calculated from TGA curves of FE and filler/FE.
it is predicted that a highly filled FE with MCNT can stay up to 540 °C.

Figure 2 also shows the amount of this thermal stability shift versus weight% for MCNT/FE relative to FE. Again the MCNT/FE curve shifted towards higher temperatures and the thermal stability improved compared to FE. Figure 2 shows that, for the weight% of 100 to 50%, the amount of thermal stability shift is almost constant (≈13-15 °C). However for weight% between ≈20 to 50% the thermal stability shift increases gradually (≈15-33 °C). This increase in thermal stability shift shows that in this weight% range compared to the 100 to 50 weight% range, FE in MCNT/FE has more thermal stability improvement relative to FE in unfilled FE.

Once more, based on Pham et al.\textsuperscript{\[4\]} work and the above results it can be concluded that greater thermal stability shift for weight% between ≈20 to 50% will be for the part of FE which is near or attached to MCNT. All of the above mentioned results show that FE in MCNT/FE is more thermally stable than FE in unfilled FE under N\textsubscript{2}. Furthermore in MCNT/FE, FE near MCNT is considerably more thermally stable than FE in the bulk. FE near MCNT in MCNT/FE to a large extent is more thermally stable than FE in unfilled FE.

By comparing the thermal stability improvement of FE in MCNT/FE and CNT/FE, both relative to FE in unfilled FE, it can be seen that for all weight% (100 to ≈20%) the thermal stability improvement of FE in MCNT/FE is considerably more than that of CNT/FE.

3.1.1 TGA: weight of residual

Figure 1 also shows that with the incorporation of MCNT, a greater amount of the FE remained and produced char in the temperature range of 400-540 °C compared to CNT/FE and FE. This amount was calculated as follows.

Table 2. The percentage of filler remaining in pure filler and filler/FE at different temperatures.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>MCNT</th>
<th>MCNT/FE</th>
<th>CNT</th>
<th>CNT/FE</th>
</tr>
</thead>
<tbody>
<tr>
<td>520</td>
<td>100.00</td>
<td>8.40</td>
<td>100.00</td>
<td>8.4</td>
</tr>
<tr>
<td>530</td>
<td>99.94</td>
<td>8.39</td>
<td>100.00</td>
<td>8.4</td>
</tr>
<tr>
<td>540</td>
<td>99.87</td>
<td>8.39</td>
<td>99.70</td>
<td>8.4</td>
</tr>
<tr>
<td>560</td>
<td>99.67</td>
<td>8.37</td>
<td>99.30</td>
<td>8.3</td>
</tr>
<tr>
<td>600</td>
<td>99.04</td>
<td>8.32</td>
<td>97.90</td>
<td>8.2</td>
</tr>
<tr>
<td>650</td>
<td>97.21</td>
<td>8.17</td>
<td>94.50</td>
<td>7.3</td>
</tr>
<tr>
<td>700</td>
<td>94.39</td>
<td>7.93</td>
<td>89.30</td>
<td>7.5</td>
</tr>
<tr>
<td>800</td>
<td>82.52</td>
<td>6.93</td>
<td>66.60</td>
<td>5.6</td>
</tr>
<tr>
<td>875</td>
<td>74.15</td>
<td>6.23</td>
<td>54.80</td>
<td>4.6</td>
</tr>
</tbody>
</table>

The results in Table 3 support the finding that, in MCNT/FE, a greater amount of FE remained and produced char preserve, particularly in the temperature range of 520-540 °C, compared to CNT/FE and FE. Therefore, the presence of MCNT enhances the thermal stability of FE, resulting in a higher percentage of the nanocomposite remaining undegraded at higher temperatures compared to CNT in FE or unfilled FE. Besides, in MCNT/FE and CNT/FE, most of the char was stable up to 900 °C.

3.1.2 EDX analysis: residual elemental

The composition of the residue from the heating-cooling TGA run cycle, as mentioned in the experimental section, was characterized by EDX. The TGA curves of the samples are shown in Figure 3. In the heating-cooling run, the cooling was done with the highest possible cooling rate in order to prevent further degradation and to keep the same composition of the remaining materials at 560 °C and at 900 °C upon cooling. Table 4 shows the elemental compositions of the residues in heating-cooling TGA runs for the compounds under study which were extracted from the EDX results (Figures 4 and 5). Figures 4 and 5 also show FESEM images of MCNT/FE-560 °C and MCNT/FE-900 °C residues respectively.

From Figure 3 it can be seen that, at low temperatures, after a cooling scan of the MCNT/FE-560 °C sample, 19.18% of the nanocomposite remained. If this amount

The percentage of “MCNT remaining” within the nanocomposite can be calculated by multiplying the percentage of MCNT obtained from the TGA data of pure MCNT at that temperature by 8.4 (the initial percentage of MCNT in FE). These amounts are shown in Table 2. From Figure 1 and Table 2, it can be seen that pure MCNT lost less weight at temperatures up to 900 °C than pure CNT. Again, in the TGA of the nanocomposite, at each temperature, the percentage of “undegraded FE + char + mineral additives” can be calculated by subtracting the percentage of “MCNT remaining” in the nanocomposite (Table 2) from the percentage of nanocomposite remaining in the TGA curve extrapolated in Figure 1 at that temperature. The results of these calculations are shown in Table 3. This amount, minus the initial mineral filler (2.5%), gives the percentage of “undegraded FE + char” as recorded in the same table. The same calculations were done for CNT/FE. However, in Table 3 the possible reactions of fillers at high temperatures are not considered.

Table 3. The percentage of “undegraded FE + char + mineral additives” and “undegraded FE + Char” remained in the compounds at different temperatures.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>“undegraded FE + char + mineral additives”</th>
<th>“undegraded FE + Char”</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FE</td>
<td>MCNT/FE</td>
</tr>
<tr>
<td>520</td>
<td>-</td>
<td>30.58</td>
</tr>
<tr>
<td>530</td>
<td>-</td>
<td>17.63</td>
</tr>
<tr>
<td>540</td>
<td>-</td>
<td>12.19</td>
</tr>
<tr>
<td>560</td>
<td>-</td>
<td>10.54</td>
</tr>
<tr>
<td>600</td>
<td>7.90</td>
<td>9.65</td>
</tr>
<tr>
<td>650</td>
<td>7.00</td>
<td>8.84</td>
</tr>
<tr>
<td>700</td>
<td>6.10</td>
<td>8.24</td>
</tr>
<tr>
<td>800</td>
<td>4.90</td>
<td>7.65</td>
</tr>
<tr>
<td>875</td>
<td>4.30</td>
<td>7.30</td>
</tr>
</tbody>
</table>
is multiplied by weight% of each element in Table 4, the weight% of that element based on the initial nanocomposite can be obtained. The same procedure was carried out for all specimens. The results of these calculations are presented in Table 5.

These results show that the percentage of carbon and fluorine in the residue of the TGA scan of MCNT/FE-560 °C are the same as CNT/FE-560 °C. This proves that MCNT is able to preserve the same amount of FE from degradation compared to the samples containing CNT at this temperature. Similarly, based on the EDX results and considering the work of Pham et al.\cite{4} as mentioned earlier, it can be seen that FE attached to or near MCNT at the higher temperature range exhibits a lower degradation tendency. Considering this and the fact that more carbon and fluorine remained in this nanocomposite residue after the TGA run, it can also be concluded that a greater amount of the FE remained undegraded or less degraded especially near MCNT. The amount of remaining fluorine represents the undegraded FE.

These results also show that the percentage of carbon and fluorine in the residue of the TGA scan of MCNT/FE-560 °C is higher compared to FE-560 °C. This proves that MCNT is able to preserve more FE from degradation compared to the samples containing unfilled FE.

Figure 3. Heat and cool TGA curves of FE and filler/FE up to 560 °C.

Figure 4. EDX spectra (a) and FESEM image (b) of MCNT/FE-560 °C.

Figure 5. EDX spectra (a) and FESEM image (b) of MCNT/FE-900 °C.
### Table 4. The weight% (Wt%) and Atomic% (At%) composition of the remaining composites in TGA runs.

<table>
<thead>
<tr>
<th></th>
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<tbody>
<tr>
<td>Wt %</td>
<td>27.16</td>
<td>50.48</td>
<td>69.78</td>
<td>84.46</td>
<td>68.98</td>
</tr>
<tr>
<td>At %</td>
<td>50.48</td>
<td>69.78</td>
<td>84.46</td>
<td>80.89</td>
<td>83.87</td>
</tr>
</tbody>
</table>

### Table 5. Amount of element remained based on initial weight of compound.

<table>
<thead>
<tr>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>3.42</td>
<td>13.38</td>
<td>13.13</td>
<td>10.69</td>
<td>6.26</td>
</tr>
<tr>
<td>F</td>
<td>0.85</td>
<td>1.75</td>
<td>1.76</td>
<td>2.00</td>
<td>0.20</td>
</tr>
<tr>
<td>Zn</td>
<td>2.08</td>
<td>2.75</td>
<td>1.76</td>
<td>0.92</td>
<td>2.00</td>
</tr>
<tr>
<td>Ca</td>
<td>0.27</td>
<td>0.28</td>
<td>0.16</td>
<td>0.15</td>
<td>0.20</td>
</tr>
<tr>
<td>Si</td>
<td>-</td>
<td>-</td>
<td>0.09</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

#### 3.1.3 EDX analysis: undegraded FE and char

At 560 °C, the “undegraded FE + char” for MCNT/FE-560 °C can also be calculated by subtracting the percentage of “MCNT remaining” in the nanocomposite (Table 2) from the percentage of “C + F” in the EDX analysis at 560 °C. Thus, the percentage of “undegraded FE + char” for MCNT/FE-560 °C is 6.76 and for CNT/FE-560 °C is 6.98%, again showing the same percentage of “undegraded FE + char” as in the case of using MCNT or CNT as filler for FE.

Table 2 also shows that when MCNT/FE-900 °C and CNT/FE-900 °C were run up to 900 °C, much less fluorine and carbon remained compared to MCNT/FE-560 °C and CNT/FE-560 °C respectively. These results indicate that most of the undegraded FE at temperatures below 560 °C was degraded from 560 °C up to 900 °C in both MCNT/FE and CNT/FE.

At 900 °C, the “char” for MCNT/FE-900 °C can also be calculated by subtracting the percentage of “MCNT remaining” in the nanocomposite (Table 2) from the percentage of “C + F” in EDX analysis at 900 °C. Thus, the percentage of “char” for MCNT/FE-900 °C is 3.29% and for CNT/FE-900 °C is 6.76%, which is 6.98%, again showing the same percentage of “undegraded FE + char” in the case of using MCNT or CNT as filler for FE.

#### 3.2 Verifying thermal stability in FE composites

The increased decomposition temperatures caused by using MCNT in FE are due to several reasons, some of which are the same as those mentioned for CNT in FE\[^{19}\]. Firstly, the presence of MCNT in FE makes the active centers of the FE main chains inactive, preventing degradation, therefore saving FE that is nearer to the MCNT surface. Secondly, the interactions between MCNT and FE result in increased physical and chemical cross-linking points, which prevent the degradation of the polymer chains. Besides, considering that the degradation of FE is a radical chain reaction, it is therefore susceptible to inhibition by reagents capable of trapping such radicals. Furthermore, the antioxidant nature of MCNT, attributed to its high electron affinity (≈ 2.65 eV), enables it to act as a radical scavenger. Consequently, MCNT helps to trap radicals and inhibits the degradation of FE nearer to it. Moreover, Endo et al.\[^{13}\] reasoned that the thermal stability of FE near MCNT is due to the presence of bound rubber to MCNT and concluded that this structure could prevent the decomposition of rubber at high temperatures by resisting molecular mobility.

Wang et al.\[^{25}\] reported that in PVDF/SiO2- graphene oxide (GO) composites there is a strong interaction between many of the functional groups in the GO and PVDF chains. In particular, the C-F bond in PVDF and carboxyl or hydroxyl groups of graphene oxides (GO) in the PVDF/SiO2-GO composites form hydrogen bonds, and this supports the enhancement of interfacial interaction. The same hydrogen bonding can be formed between functional groups in MCNT and C-F of FE. Ma and Larsen\[^{26}\] also reported that nitric acid treated single wall nanotubes (SWNT) compared to untreated MWNT have better physical surface affinities with PVDF. The same compatibility can be mentioned here between MCNT and FE compared to CNT and FE. Therefore, there is a strong interaction, which is that the hydrogen bonding and compatibility described above, between MCNT and FE, and these interactions are not present between CNT and FE. More interactions between MCNT and FE compared to CNT and FE result in increased physical and chemical cross-linking points, which prevent more degradation of the polymer chains in the case of MCNT/FE. Furthermore, these interactions will increase the amount of bounded rubber to MCNT in MCNT/FE compared to CNT in CNT/FE and,
as explained above, more bounded rubber will increase the thermal stability in the case of MCNT/FE.

3.3 Effect of thermal aging on residual elemental composition

Figure 6 illustrates representative figures of EDX obtained for aged and un-aged samples. The elemental weight percent extracted from these figures is presented in Table 6. This table shows the surface composition of the composites before and after aging. The major elements in the composites are C, O, F and Si. The initial loss in element is defined as the percentage lost due to aging for different periods of time. The percentage loss can be calculated as follows: For 100 g of un-aged FE (Table 6), there were 34.26, 4.99, 59.82 and 0.92 g of C, O, F and Si respectively. After 72 hours of aging, if \( C_{\text{rem}}, O_{\text{rem}}, \) and \( F_{\text{rem}} \) g of C, O and F respectively remained and \( C_{\text{los}}, O_{\text{los}}, \) and \( F_{\text{los}} \) g of C, O and F respectively were lost. Then;

\[
C_{\text{rem}} + C_{\text{los}} = 34.26 \text{ g}; \\
O_{\text{rem}} + O_{\text{los}} = 4.99 \text{ g and} \\
F_{\text{rem}} + F_{\text{los}} = 59.82 \text{ g}
\] (1)

Since Si is not lost during aging, the amount of Si after 72 hours of aging is 0.92 g. If T is the total weight of the remaining elements, then; \( T = C_{\text{rem}} + O_{\text{rem}} + F_{\text{rem}} + 0.92 \text{ g} \)

After 72 hours of aging, the weight percentage of Si is 2.32% (Table 6), therefore; \( \frac{0.92}{T} = 0.0232 \), and thus \( T = 39.66 \text{ g} \). Using T and the percentage of C (Table 6), after 72 hours of aging (30.89%), \( C_{\text{rem}} \) can be calculated as; \( 0.3089 = \frac{C_{\text{rem}}}{T} \). Thus, \( C_{\text{rem}} = 0.3089 \times 39.66 = 12.25 \text{ g} \). Therefore, the percentage of \( C_{\text{los}} \) is,

\[
C_{\text{los}}(\%) = \frac{W_{\text{los}} \cdot C_{\text{los}}}{W_{\text{los}} \cdot \text{initial} C} \times 100(\%) = \frac{22.01}{34.26} \times 100(\%) = 64.24\% \] (2)

This means 64.24% of the initial C was removed from the surface after 72 hours of aging at 250 °C. The same calculations were done for F and O in all other aged composites. The results of these calculations are presented in Table 7. These results show that for each composite, with increasing aging time, the percentage loss of C, O and F increased. This is because, as the aging time increased, more degradation including dehydrofluorination, defluorination and carbon oxygen reactions occurred especially at the surface[23]. The results also show that at each aging time, for CNT/FE the loss of elements was lower than for MCNT/FE. Therefore, CNT can preserve a larger portion of the elastomer surface from degradation compared to MCNT.

For CNT/FE, up to 24 hours of aging, the initial percentage loss of each element slightly increased. Nevertheless, as the aging time increased, even a decrease in percentage loss was observed. This means that the slight degradation happened mainly in the first 24 hours and did not increase after that. For MCNT/FE, up to 24 hours of aging, the same trend as with CNT/FE was observed. However, as the aging time increased, there was a drastic increase in percentage loss. FE underwent thermal oxidation and dehydrofluorination during hot air aging[27]. Due to the dehydrofluorination reaction, HF was produced[27] and the carboxylic acidic groups on the surface of MCNT reacted with HF. Wang et al.[28] reported a similar reaction between dispersed graphene oxide (GO) and hydrofluoric acid (HF) during the synthesis of fluorinated graphene sheets (FGS). The authors mentioned that fluorine was grafted onto the basal plane of graphene. Furthermore, the oxygen-containing groups in GO played a major role in FGS formation, and the fluorination degree could be easily controlled by varying the reaction temperature, times and amounts of HF. They also stated that fluorination...
occurred at the original sp³ C sites in GO, that is, the C sites connected with the oxygen-containing groups. In the case of MCNT/FE, this reaction caused increased degradation particularly at aging times higher than 24 hours, leading to more loss of F and C from the surface compared to CNT/FE. Comparing the results of FE and MCNT/FE at aging times longer than 24 hours, it can also be seen that for FE without MCNT, the loss of elements was lower than for FE with MCNT. This could be due to the same reasons as those given above in the comparison of MCNT/FE and CNT/FE. The reasons adduced for the saving of a higher percentage of undegraded FE and char in the TGA runs are also valid for the thermal aging of MCNT/FE in the first 24 hours. Besides, the antioxidant nature of MCNT enables it to act as a radical scavenger especially for the oxide radicals produced due to the presence of air. This also helps to trap radicals, thereby inhibiting the degradation of nearby MCNT, and results in an increased percentage of undegraded FE.

A considerable depletion of F and C on the surface and to a depth of around 10–15 μm of the cross-section of the CB/FE seals aged in oil containing amine based dispersant at 150 °C was also reported by Smith et al. They also reported the catalytic effect of amine on the degradation of FE. The authors reported that polytetrafluoroethylene (PTFE) also began to soften and release F at around 200–220 °C. This may explain the apparent susceptibility to heating in air shown by elastomers with a high TFE content. A drastic decrease in C and F due to thermal aging in air at 200 °C from the surface of CB/FE (up to a depth of 5 nm) was also reported by Wang et al.

Figure 7 shows the FESEM image of MCNT/FE razor cut surfaces. The very good distribution and dispersion of MCNT in FE can be seen in this figure.

3.4 Other works on FE and fillers/FE

In our before study, we verified the crystallinity of some of the above mentioned FE and fillers/FE, by Dynamic mechanical analysis (DMA), Differential Scanning Calorimetry (DSC) and X-ray diffraction (XRD). Verifying thermal properties of fluoroelastomer using carbon nanotubes in presence of air and under nitrogen flow is the other study.
4. Conclusions

TGA shows that the thermal stability of FE in a N$_2$ atmosphere increased with the presence of MCNT relative to when either CNT was used or while FE was unfilled. A higher percentage of this composite in the form of undegraded FE or char remained within the temperature range of 400-900°C compared to CNT/FE or FE. The MCNT/FE TGA curve shifted towards higher temperatures compared to CNT/FE or FE. The same results also revealed that a higher percentage of FE was undegraded or less degraded especially near MCNT in the temperature range of 400-540 °C.

EDX analysis shows that the percentage of carbon and fluorine in the residue of TGA runs up to 560 °C of MCNT/FE was the same as CNT/FE and higher than that of FE. EDX of the residue of TGA runs up to 900 °C shows that most of the undegraded FE which was not degraded at temperatures up to 560 °C was degraded from 560° up to 900 °C in both MCNT/FE and CNT/FE. However, the char in MCNT/FE was more than that in CNT/FE at 900 °C.

The EDX results of aged specimens under air indicate that the percentage of C, F and O lost on the surface of filler/FE and FE increased with increasing aging time. Again, at 24 hour aging time, MCNT preserved a higher percentage of these elements compared to CNT. The order of percentage loss in elements after 24 hour aging time was MCNT/FE > FE > CNT/FE.

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6. References


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