Chemical Characterization of Stabilized and Carbonized Polyacrylonitrile (PAN) Fibers Treated with Oleic Acid.

Shahrul Nizam Md Salleh1,a, Mohamad Zaki Abdullah2,b, Azmi Abdul Wahab3
1,2,3Mechanical Engineering Department,Universiti Teknologi PETRONAS
Bandar Seri Iskandar, 31750 Tronoh, Perak, Malaysia

Abstract. Polyacrylonitrile (PAN) fiber is the best precursor for carbon fibers due to high carbon content after heat treatment. After the polymer was spun into fibers, the fibers will undergo pretreatment process with chemical solution known as post spinning treatment. Post spinning will directly affect conversion of PAN fiber to carbon fiber. Oleic acid was used as post spinning treatment chemical solution to PAN fibers. The pretreated PAN fiber will be heated at 250°C and 800°C. The fibers were studied using Fourier Transform Infra-Red (FTIR), X-ray Photoelectron Spectroscopy (XPS) and DSC to study the chemical change during heat treatment. PAN fibers treated with oleic acid have reduced the cyclization energy and increase oxygen and carbon content leading to high performance carbon fibers.

1 Introduction.

Carbon fiber was used in high performance composite due to excellence weigh to strength ratio. Carbon fiber reinforced composite became more important in aerospace, civil engineering, military, and motorsports, along with other competition sports [1-2]. Carbon fibers can be derived from pitch based, polyacrylonitrile (PAN) polymer or rayon based chemicals. However most high performance carbon fiber was converted from polyacrylonitrile (PAN) based polymer due to high carbon content after heat treatment process [3]. Polyacrylonitrile (PAN) polymer was spun to form polyacrylonitrile (PAN) fiber and treated with solution known as post spinning treatment. Post spinning were divided into three categories, such as modification through coating, impregnation with chemicals (catalytic modification) and drawing/stretching with plasticizer [4,5,6]. Previous study shows fatty acid derivatives react as coating agent to reduce the entangling, fluffy, fusion and fiber to fiber adhesion of the PAN precursor fibers during thermal stabilization process [7]. In this paper an attempt to prove that fatty acid without any derivative can give similar effect to fatty acid derivatives coating type post spinning modification process. The post spinning modifications indirectly affect and ease the stabilization in several ways such as reducing the activation energy of cyclization, decreasing the stabilization exotherm, increasing the speed of cyclization reaction and also improving the orientation of molecular chains in the fibers [8]. Recently, the stabilization process is found to play an important role in converting PAN fiber to an infusible stable ladder polymer that converts C=N bonds to C≡N

a Corresponding author: shahruln@outlook.com
b Corresponding author: zaki_abdullah@petronas.com.my

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bonds and to develop crosslink between molecules of PAN which tend to operate at high temperatures, with minimum volatilization of carbonaceous material. The thermal stability of the stabilized fiber is attributed to the formation of the ladder structure due to cyclization of the nitrile groups in acrylic molecule and started from 180°C to 300°C. Carbonization was an aromatic growth and polymerization in which the fiber would undergo heating process at a high temperature up to 800-3000°C, typically to a 95% carbon content. Carbonization at 1000°C will produce carbon fiber in low modulus type and intermediate modulus or type II carbon fiber will produce at up to 1500°C.[8]

2 Experimental.

The PAN fiber was produced by Fortafil Fibre Inc. with 50000 (50k) s filament per tow was used in the study. The fibers have 17 micron average diameter.

2.1 Chemical treatment process.

10% oleic acid was dissolved in acetone with 10% weight by weight ratio. The PAN fiber was soaked into the solution for about 10 minutes and left to dry. The coated fibers was heated by batch in 250 ºC and 800 ºC.

2.2 Stabilization of fibers.

Stabilization is a heat treatment process starting from 200ºC to 300 ºC. In this study 250 ºC temperature was chosen for PAN fibers treated and untreated with oleic acid. Both samples were heat treated in tube furnace with 1 ºC /min heating rate.

2.3 Carbonization of fibers.

The stabilized PAN fibers will undergo a carbonization process started at 800 ºC with heating rate at 5 ºC /min in an inert atmosphere to produce carbon fiber. The treated and untreated samples with oleic acid were placed together in the tube furnace.

2.4 FTIR study.

The samples were characterized using attenuated total reflectance (ATR) FTIR model Nicolet iS5 with iD5 ATR attachment.

2.5 Differential scanning calorimetry(DSC) study.

The samples were characterized using DSC model 821 with 10 ºC /min heating rate from 30 ºC to 400 ºC. Nitrogen flow was set at 1 cm³ /m

2.6 X-ray Photoelectron Spectroscopy (XPS) study.

X-ray photoelectron spectra were recorded using a Omicron XPS spectrometer with a Mg Kα X-ray source(1254eV). Wide scan was perform on the sample before narrow scanning on carbon and oxygen element. The data collected from the instrument were analyzed by CasaXPS software.
3 Result and Discussion.

3.1 FTIR Result.

Figure 1 show FTIR spectral comparison between PAN fiber before and after heated at 250°C. Before heated, PAN fiber show prominent peaks at 2924 and 2242cm⁻¹ due to stretching vibration of the methylene (CH₂) and the nitrile groups (CN) in acrylonitrile structure. In figure 1, 2242 cm⁻¹ of cyanide (-C≡N) which is main functional group of PAN fibers gradually decrease in concentration when heated to 250°C. In figure 1, it is clearly shown the existing of a new peak in 1580 cm⁻¹ region after heat treated to 250°C. 1580 cm⁻¹ was referred to double bonding of carbon (C=C) exist in cyclic form. The double bond only exists when heat treated at 250°C. This suggests that intermolecular or intramolecular cross-linking might have taken place forming the cyclisation reaction. For 1731 cm⁻¹, indicate the existent of carbonyl group in PAN fibers and was gradually decreased with increasing temperature. The carbonyl group was found in PAN fibers due to commoner exist in the polymer which is a common for PAN fibers manufacturing process in order to increase the plasticity effect to the fibers. Compound Methyl acrylate was often used as commoner in PAN polymer to produce the fibers. PAN fibers treated with oleic acid and heated at 250°C show interesting FTIR spectra pattern. Most of the peaks have significantly high intensity compared to untreated PAN fibers. At 2924 cm⁻¹, sharp and high intensity peak referred to methylene (CH2) was recorded in the spectra. This may due to methylene (CH2) coming from oleic acid attach to stabilized PAN fibers surface. Sharp and high intensity peak at 1709cm⁻¹ show existent of carbonyl (C=O) derived from carboxylic acid which is a functional group of oleic acid. PAN fibers treated with oleic acid and heated at 250°C also show additional peaks at 724 and 942 cm⁻¹ and was referred to rocking CH₂ and NO formation. Figure 2 show FTIR spectral comparison between PAN fiber before and after heated at 800°C. The FTIR spectra between PAN fibers coated and uncoated with oleic acid show almost similar pattern. Main
peaks at 2885 cm\(^{-1}\) due to CH from aldehyde of ketone group. 2116 cm\(^{-1}\) was referred to N=C-C group. However PAN fiber treated with oleic acid and heated at 800\(^\circ\)C has additional peak at 1570 cm\(^{-1}\) which was referred to N-H bending in amide group.

### 3.2 Differential scanning calorimetry (DSC)

![Figure 3. DSC of PAN fiber before and after treated with oleic acid.](image)

As shown in figure 3, the range of the exotherm for PAN fibers was from 217.7\(^\circ\)C to 309.7\(^\circ\)C a spread of 92\(^\circ\)C whereas for PAN Fibers treated with oleic acid range was from 175.5\(^\circ\)C to 372\(^\circ\)C, a spread of 196.5\(^\circ\)C. Pan fiber treated with oleic acid has lower exothermic peak at 283\(^\circ\)C as oppose to PAN fiber with exothermic peak at 295\(^\circ\)C. However, for Pan fiber treated with oleic acid, a shoulder appear in the graph, suggesting there are two exothermic peak that may be came from oleic acid treatment. The broadening of the exothermic peak is probably contributed to the oleic acid that initiates cyclization at a lower temperature through an ionic mechanism.

### 3.3 X-ray Photoelectron Spectroscopy (XPS)

The data received from Omicron XPS was analyzed by CasaXPS software. All binding energy are measured in electron volt (eV) and were referred to National Institute of Standards and Technology (NIST) X-ray Photoelectron Spectroscopy Database.

![Figure 4. XPS spectra of PAN fiber.](image)

Figure 4 show XPS spectrum of C1s for PAN fiber. After deconvulation, the spectra revealed 3 peaks at 281.20eV, 284.07 eV and 287.32 eV. The three peaks can be assigned to C-H, C-C and carbon attached cyanide carbon (CN). It shows the monomer of PAN polymer as shown in figure 5.
### Figure 5. Polyacrylonitrile polymer.

![Polyacrylonitrile polymer molecule](image)

### After PAN fiber was heated at 250°C

<table>
<thead>
<tr>
<th>Name</th>
<th>Pos.</th>
<th>FWHM</th>
<th>Area</th>
<th>%Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>C-C/C-H</td>
<td>281.36</td>
<td>2.157</td>
<td>6249.37</td>
<td>27.72</td>
</tr>
<tr>
<td>ester carbon</td>
<td>286.00</td>
<td>4.457</td>
<td>4167.65</td>
<td>18.42</td>
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<tr>
<td>graphitic carbon</td>
<td>284.07</td>
<td>1.823</td>
<td>12169.55</td>
<td>53.86</td>
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</table>

### Carbon (PAN Fiber Treated with Oleic Acid and Heated at 250°C)

<table>
<thead>
<tr>
<th>Name</th>
<th>Pos.</th>
<th>FWHM</th>
<th>Area</th>
<th>%Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>graphitic carbon</td>
<td>284.96</td>
<td>1.560</td>
<td>4289.08</td>
<td>63.00</td>
</tr>
<tr>
<td>COO ester carbon</td>
<td>288.94</td>
<td>2.167</td>
<td>532.68</td>
<td>7.80</td>
</tr>
<tr>
<td>C-C or graphitic carbon</td>
<td>285.96</td>
<td>2.156</td>
<td>1989.33</td>
<td>29.20</td>
</tr>
</tbody>
</table>

### Figure 6. XPS spectrum of PAN fiber heated at 250°C(A) and oleic acid treatment (B)

After PAN fiber was heated at 250°C, C1s XPS spectrum show 3 peaks centered at 281.36 eV, 286.00 eV and 284.07 eV. All peaks are referred to C-H, ester carbon (COO-R) and C-C or graphitic carbon. The carbon content (C-C), of PAN fiber heated at 250°C is almost similar to PAN fiber which was calculated at 53.86% and 55.13% respectively. However ester type carbon was found due to cyclization of PAN polymer in the fiber. In figure 6(B), graphitic carbon (C-C) has slightly increased to 63%. This show oleic acid does help in cyclization process. Ester carbon content also increases compare to PAN fiber without oleic acid treatment. 288.89 eV peak show carbon from carboxylic acid, a functional group of oleic acid.

### After heated at 250 °C, the fiber was heated at 800 °C

<table>
<thead>
<tr>
<th>Name</th>
<th>Pos.</th>
<th>FWHM</th>
<th>Area</th>
<th>%Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>graphitic carbon</td>
<td>284.51</td>
<td>1.606</td>
<td>12188.73</td>
<td>53.63</td>
</tr>
<tr>
<td>NC=O</td>
<td>288.45</td>
<td>1.576</td>
<td>1089.56</td>
<td>4.78</td>
</tr>
<tr>
<td>ester C</td>
<td>283.08</td>
<td>2.232</td>
<td>3760.23</td>
<td>16.56</td>
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<tr>
<td>C-H</td>
<td>285.52</td>
<td>2.713</td>
<td>5690.97</td>
<td>25.02</td>
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</tbody>
</table>

### Carbon (PAN Treated with Oleic Acid and Heated at 800°C)

<table>
<thead>
<tr>
<th>Name</th>
<th>Pos.</th>
<th>FWHM</th>
<th>Area</th>
<th>%Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>graphitic carbon</td>
<td>283.98</td>
<td>2.770</td>
<td>27204.56</td>
<td>94.31</td>
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<tr>
<td>amide</td>
<td>287.99</td>
<td>2.392</td>
<td>1645.72</td>
<td>5.69</td>
</tr>
</tbody>
</table>

### Figure 8. XPS spectra of PAN fiber heated at 800°C (C) and oleic acid treatment (D)

After heated at 250 °C, the fiber was heated at 800 °C in inert environment. The C1s xps spectrum show 4 peaks after deconvolution. The peak represent graphitic carbon, C-C (284.51 eV), amide carbon, NC=O (288.45eV), ester carbon (285.52 eV) and C-H carbon. Carbonization started at 800°C to 1500°C to form low to intermediate modulus carbon fibers. Figure 8(C) show the existence of ester carbon that indicate carbonization process not fully occur. Graphitic carbon content shows no significant increase around 53.63%. However amide type carbon starts to exist on the surface of the fiber due to cyclization process. Figure 8(D) shows PAN fiber treated with oleic acid have higher carbon content at 94.31%. This indicate oleic acid have important role in carbon fiber conversion process.
4 Summary

PAN fiber coated with 10% w/w oleic acid play important role in lowering cyclization energy of PAN fiber to form carbon fibers. The coated PAN fiber could be fully preoxidized at lower temperature and reduce thermal stabilization temperature as indicate by the DSC. FTIR result show insertion of carboxylic acid to the polyacrylonitrile polymer has altered the structure, thus promoting early stabilization process that lead to thermal stability of the fibers before carbonization stage. This will assist aromatic growth of carbon structure with higher carbon content at carbonization stage. Higher carbon content of carbon fibers have higher mechanical properties as reported by previous studies.[9,10]

References