Property and Structure Characterization of High Molecular Weight Polyacrylonitrile Polymers Initiated by 2,2'-Azobis (2-methyl propionamide) Dihydrochloride Using Aqueous Deposited Polymerization Technique

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Abstract. Powdery high molecular weight polyacrylonitrile (HMW-PAN) polymers were successfully synthesized by aqueous deposited polymerization (ADP) method with 2,2'-azobis (2-methyl propionamide) dihydrochloride (AIBA) as the initiator and itaconic acid (IA) as comonomer. The utilization of IA into the ADP system was conducive to relax the proceeding of thermo-oxidation process. Differential scanning calorimeter (DSC) curves of obtained PAN copolymers showed the doublet peaks character under air atmosphere, attributing to the exothermic cyclization and oxidative reactions during heat-treatment process. With the utilization of IA in the reaction system, crystallinity and grain sizes of the PAN polymers decreased. The stretching vibration of nitrile group (C≡N) at 2240cm⁻¹ was the strongest peak in both Fourier transform infrared spectra (FTIR). And the stretching vibration and bending of CH₂ were the stronger peaks in FTIR. Amido group (NH₂), carbonyl group (C=O) and C-O single bond existed in FTIR of PAN homopolymer and copolymers, indicating occurrence of C≡N hydrolysis during polymerization process except carboxyl group (COOH) introduced by IA.

1. Introduction

High molecular weight polyacrylonitrile (HMW-PAN) copolymer is one of the best precursors to prepare high quality carbon fibers [1-3]. During the preparation process of HMW-PAN copolymers, some vinyl comonomers containing flexible functional groups have been focused by many investigators, such as acrylic acid (AA), itaconic acid (IA), methacrylate (MA), vinyl acetate (VAc), and so on [4-10]. And their thermal properties and structural characters had a direct effect on PAN copolymer spinning dopes. Therefore, the introduction of a few comonomers greatly enhanced the internal mobility of PAN’s segments, depressed the onset of cyclized temperature during oxidation, and improved the final spinnability of PAN dopes. Compared with traditional homogeneous solution polymerization, HMW-PAN can be prepared by aqueous polymerization system, because of none chain transfer reactions and partially-solubility of acrylonitrile (AN) in water (H₂O) system. As one kind of heterogeneous solution polymerization, there have been some reports about aqueous deposited polymerization (ADP) of AN. But many complex initiating systems containing alkali metallic ions were usually employed to initiate their copolymerizations. This is not beneficial to manufacture high quality PAN-based precursors for carbon fibers [11-15]. So it is necessary to search an appropriate initiator for the production of HMW-PAN copolymers. By now, there have been some reports about the important water-soluble initiator ammonium persulphate with none alkali metallic ions [16-19]. But there have been few reports about the water-soluble azo initiator, which can also initiate the PAN’s copolymerization through the water-soluble radical in water phase. And HMW-PAN can be also obtained using the ADP technique. In this paper, as a single organic and water-soluble initiator without alkali metallic ions, 2,2'-azobis (2-methyl propionamide) dihydrochloride (AIBA) was used as the initiator, which is a relatively moderate initiator to control the polymerization process and thermal properties of PAN copolymers [20-22]. And a commonly used comonomer-itaconic acid (IA) was copolymerized with AN using the ADP technique. Furthermore, there are also few reports about detailed property and structure characterization of powdery HMW-PAN polymers directly obtained by this initiator. As a result, thermal properties of PAN polymers obtained were investigated by thermal analyzer. And their physical and chemical structures were discussed by wide-angle X-ray diffraction (WAXD) and Fourier transform infrared spectra (FTIR).

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2. Experimental Section

2.1 Original Materials
AN (Zhengzhou Paini Chemical Reagent Factory, China) and IA (Tianjin Guangfu Fine Chemical Research Institute, China) were analytically purity used as the reaction monomers without further treatment. AIBA (Shanghai Kaiyang Biological Technology Co., Ltd, China) with 98% purity was directly used. And deionized H₂O was adopted as the reaction medium during the polymerization process.

2.2 ADP of PAN
Powdery PAN homopolymer and PAN copolymer with different weight fractions of IA (0~10wt%) in the feed, i.e. Homo-PAN and various PAN(AN-co-IA)s, were synthesized under nitrogen (N₂) atmosphere using a three-necked flask with deionized H₂O at 60°C. The synthesized samples were described as Homo-PAN, PI-1, PI-2, PI-3, and PI-4, corresponding to 0wt%, 1wt%, 2wt%, 4wt%, 10wt% IA in the feed, respectively. While the concentration of AIBA was 1wt% based on the total monomer concentration at 22wt%. Good agitation was used to ensure that the reaction heat could be thrown off in the polymerization process. The reaction time lasted 2h. And then the precipitated products were washed successively with water and methanol, and dried to a constant weight under vacuum at 50°C.

2.3 Measurements
Conversions of the polymerizations were tested using weight method. Viscosity average molecular weights (Mv) of Homo-PAN and PAN(AN-co-IA)s were determined by Ubbelohde viscometer method at (30±0.2)°C using dimethylsulfoxide (DMSO) solvent. The intrinsic viscosity [η] equation was calculated by one-point method as follows:

\[ [\eta] = \frac{2(\eta_p - \eta_i)}{c} \]

where the needed PAN weight was about 0.04~0.05g in 50mL DMSO.
TA Instruments Co. SDT Q600 (USA) thermal analyzer was used to get simultaneous thermal analysis (STA) curves of different PAN polymers, including differential scanning calorimeter (DSC) and thermal gravity analysis (TGA) curves. Their STA curves with the temperature range from room temperature to 450°C were tested at a 10°C/min heating rate under air atmosphere. Also the DSC curves under N₂ atmosphere were measured using PerkinElmer DSC 8500 (USA) thermal analyzer with the aluminum crucible under aluminum cover, whose temperature range was from room temperature to 400°C. Wide-angle X-ray diffraction (WAXD) of Homo-PAN and PAN(AN-co-IA)s between 5~50° were measured at a 3.6°/min step rate under room temperature by PANalytical X'Pert Pro analyzer (Holland) All the WAXD measurements were well done under the same conditions. Crystallinity index (Xc) was calculated by the integral area method according the references [23-25]. And the crystalline planar spacing d and crystallite size (Lc) were determined by Scherrer equation and Bragg equation [25, 26].
FTIR of Homo-PAN and PAN(AN-co-IA)s were recorded on PerkinElmer Frontier (USA) spectrometer for 16 times. The transmission mode was used on finely pure powdery samples using direct compression method without KBr. The testing range was from 400cm⁻¹ to 4000cm⁻¹.

3. Results and discussions

3.1 Acquirement of HMW-PAN polymers
As detailed in Table 1, the Mv values measured by the method above are approximately above 16×10⁴. It is evidently investigated that the synthesized PAN polymers almost had the higher molecular weights than the results in the references [12,13,26]. The main reason is that H₂O was adopted as reaction medium in ADP process. Due to the presence of H₂O, chain transfer reactions were prevented, which could increase average molar weights of PAN polymers. While in the homogeneous solution polymerization process of AN and vinyl comonomers, chain transfer reactions to reaction solvents, such as DMSO, dimethylformamide (DMF) and so on, would lower molecular weights of the final PAN copolymers [12,13,26]. Furthermore, relative researches and monomer reactivity ratios of AN and IA have been reported by Zhao et al [16,17], which is deduced that the IA comonomer has the higher reactivity than AN monomer. So, the samples PI-1 and PI-2 with 1wt% and 2wt% IA in the feed had the higher Mv values than the Homo-PAN. When the IA amount reached 4wt%, the IA amount in the PAN main chains became much more enough to prevent the active chain growth. And the Mv value of sample PI-3 became lower. While the conversions of samples PI-2 and PI-3 had become higher than samples Homo-PAN and PI-1. However, only the 14.09% products were prepared with its Mv value at 16.18×10⁴ about sample PI-4 copolymer with 10wt% IA. Thus, more IA comonomer in the feed was not conducive to produce HMW-PAN copolymer with high conversion.

<table>
<thead>
<tr>
<th>Samples</th>
<th>AN/IA(w/w)</th>
<th>Conversion (%)</th>
<th>Mv (×10⁴)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Homo-PAN</td>
<td>100/0</td>
<td>76.39</td>
<td>40.60</td>
</tr>
<tr>
<td>PI-1</td>
<td>100/0</td>
<td>16.18</td>
<td>16.18</td>
</tr>
<tr>
<td>PI-2</td>
<td>98/2</td>
<td>66.10</td>
<td>55.94</td>
</tr>
<tr>
<td>PI-3</td>
<td>96/4</td>
<td>82.91</td>
<td>52.93</td>
</tr>
<tr>
<td>PI-4</td>
<td>90/10</td>
<td>83.12</td>
<td>41.74</td>
</tr>
</tbody>
</table>

3.2 STA Analysis of PAN Polymers
Different DSC curves of powdery Homo-PAN and different PAN(AN-co-IA)s under air and N₂ atmosphere are listed in Figure 1(a) and Figure 1(b). The parameters obtained from the exotherms including the initiation temperature (Ti, the temperature when the exothermic reactions begin), the termination temperature, i.e., the
final temperature ($T_f$, the temperature when the exothermic reactions end), their difference ($\Delta T = T_f - T_i$), the exothermic peak temperature ($T_p$, the evident exothermic peak in DSC curves), the evolved heat ($\Delta H$) and the velocity of evolving heat ($\Delta H/\Delta T$) are described in Table 2.

DSC exotherms of different PAN polymers are due to the oxidative reactions and cyclization reactions under different atmospheres [1-3, 13]. The obtained exotherm parameters also reflect different reaction characters. The $T_i$ values for all the P(AN-co-IA) were much smaller than Homo-PAN, indicating that a greater release of the preoxidation process of PAN-based precursors, whose weight loss zone and slow weight loss zone, which were stages, named as micro-scale weight loss zone, intense exothermic reactions of all PAN polymers included three reaction stages, named as micro-scale weight loss zone, intense exothermic reactions and cyclization reactions during heat-treatment process. And the three stages were combinated into the preoxidation process of different PAN polymers. Furthermore, with the increase of IA amounts, much more intricate changes were found in DSC exotherms of different P(AN-co-IA), which was probably due to the superposition of the two peaks. The doublet peaks might be due to the exothermic cyclization and oxidative reactions during heat-treatment process. And the reactions might happen more or less simultaneously. However, initiation of oxidation reaction occurred before the onset of cyclization. The lower-temperature exothermic peak was attributed to the primary oxidation (including dehydrogenation) and cyclization reactions. And the higher-temperature exothermic peak was attributed to the secondary oxidation reactions leading to the polymer chain scission. The evolution of HCN, CO$_2$ and CO also happened in this region. There is no essential difference to the experimental results by the other many workers [13, 28-30].

While in the N$_2$ atmosphere, the different PAN polymers had higher $T_i$ values, lower $T_p$, $T_f$ and $\Delta T$ and $\Delta H$ values than those in air atmosphere. The main reason was due to strong oxidation reactions under air atmosphere of different PAN polymers. In addition, the Homo-PAN had the minimum $\Delta T$ and the maximum $\Delta H/\Delta T$. At the same time, the broader (higher $\Delta T$) and the smoother (lower $\Delta H/\Delta T$) exotherm were found in different PAN copolymer than in Homo-PAN. It is indicated that the evolving heat was concentrative and expeditious in Homo-PAN. The IA comonomer slowed the rate of exothermic reaction. It is obvious that the IA comonomer has role of relieving exothermic reactions of PAN polymers.

### Table 2. Parameters of DSC curves of Homo-PAN and different P(AN-co-IA)

<table>
<thead>
<tr>
<th>samples</th>
<th>atmosphere</th>
<th>$T_i$ (°C)</th>
<th>$T_f$ (°C)</th>
<th>$T_p$ (°C)</th>
<th>$\Delta T$ (°C)</th>
<th>$\Delta H$ (J·g$^{-1}$)</th>
<th>$\Delta H/\Delta T$ (J·g$^{-1}$·°C$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Homo-PAN</td>
<td>220</td>
<td>312</td>
<td>375</td>
<td>155</td>
<td>194</td>
<td>27.06</td>
<td></td>
</tr>
<tr>
<td>PI-1</td>
<td>Air</td>
<td>212</td>
<td>317</td>
<td>380</td>
<td>174</td>
<td>4968</td>
<td>28.55</td>
</tr>
<tr>
<td>PI-2</td>
<td>199</td>
<td>249</td>
<td>319</td>
<td>186</td>
<td>4920</td>
<td>26.45</td>
<td></td>
</tr>
<tr>
<td>PI-3</td>
<td>187</td>
<td>255</td>
<td>327</td>
<td>202</td>
<td>917</td>
<td>19.39</td>
<td></td>
</tr>
<tr>
<td>PI-4</td>
<td>267</td>
<td>271</td>
<td>275</td>
<td>8</td>
<td>445</td>
<td>55.63</td>
<td></td>
</tr>
</tbody>
</table>

Another phenomenon deduced from DSC curves was the doublet peaks character in different P(AN-co-IA) under air atmosphere. While in case of Homo-PAN, the doublet character was not very clear. It could be relevant to the different reaction mechanism during heat-treatment process of different PAN polymers. However, the increase of IA amounts, much more intricate changes in DSC exotherms of different P(AN-co-IA), which was probably due to the superposition of the two peaks. The doublet peaks might be due to the exothermic cyclization and oxidative reactions during heat-treatment process. And the reactions might happen more or less simultaneously. However, initiation of oxidation reaction occurred before the onset of cyclization. The lower-temperature exothermic peak was attributed to the primary oxidation (including dehydrogenation) and cyclization reactions. And the higher-temperature exothermic peak was attributed to the secondary oxidation reactions leading to the polymer chain scission. The evolution of HCN, CO$_2$ and CO also happened in this region. There is no essential difference to the experimental results by the other many workers [13, 28-30].

### 3.3 TGA Analysis of PAN Polymers

TGA curves of powdery Homo-PAN and different P(AN-co-IA) are shown in Figure 2. Obviously, weight loss reactions of all PAN polymers included three reaction stages, named as micro-scale weight loss zone, intense weight loss zone and slow weight loss zone, which were related to small molecules loss and intense exothermic reactions. And the three stages were combinated into the preoxidation process of PAN-based precursors, whose

![Figure 1](image-url)  
**Figure 1.** DSC curves of Homo-PAN and different P(AN-co-IA)s under air atmosphere from STA results and DSC curves of Homo-PAN and different P(AN-co-IA)s under N$_2$ atmosphere.
difference reflected the different degree and rate of the preoxidation reaction. According to the TGA curves, the initiation decomposition temperature ($T_{id}$) and different weight loss values obtained from the TGA curves including $w_1$(%), $w_2$(%) and $w_3$(%) corresponding to the three stages were listed in Table 3.

**Table 3. Parameters for TGA curves of Homo-PAN and different P(AN-co-IA)s**

<table>
<thead>
<tr>
<th>Samples</th>
<th>$T_{id}$ (°C)</th>
<th>$w_1$ (%)</th>
<th>$w_2$ (%)</th>
<th>$w_3$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Homo-PAN</td>
<td>303</td>
<td>0.5</td>
<td>12.9</td>
<td>22.3</td>
</tr>
<tr>
<td>PI-1</td>
<td>282</td>
<td>0.7</td>
<td>11.4</td>
<td>23.3</td>
</tr>
<tr>
<td>PI-2</td>
<td>277</td>
<td>1.1</td>
<td>11.5</td>
<td>23.0</td>
</tr>
<tr>
<td>PI-3</td>
<td>275</td>
<td>1.2</td>
<td>13.7</td>
<td>25.1</td>
</tr>
<tr>
<td>PI-4</td>
<td>132</td>
<td>1.7</td>
<td>10.3</td>
<td>34.5</td>
</tr>
</tbody>
</table>

It is investigated that the first stage occurred at the higher temperature to the Homo-PAN with none IA, with its largest weight loss slope seen from Figure 2. All P(AN-co-IA)s had ease exothermic reactions under air atmosphere, with its lower $T_{id}$ values. When the IA amount was less than 4wt%, the $w_1$ and $w_3$ weight loss values had low rise trend. Due to the introduction of IA comonomer in the feed, the increase of $w_1$, $w_2$ and $w_3$ were attributed to higher IA content in the PAN main chains, which might promote the pyrolysis reaction. At the same time, when the IA amount reached 10wt%, the $w_1$ and $w_3$ reached the maximum values. That is to say, sample PI-4 had much more weight loss at the first and third stage. However, the sample PI-3 had the highest $w_2$ value and the sample PI-4 with the lowest $w_2$ value. Higher IA content in sample PI-3 was benefit to the exothermic reaction, which made the $w_2$ value become higher. While the $T_{id}$ value of sample PI-4 was so lower that it made the exothermic reaction occurred much more early. And the sample PI-4 also had the lowest $M_v$ value. Therefore, the $w_2$ value became a little smaller. It was probably attributed to interactive affect of higher IA contents in the final P(AN-co-IA) polymers and their $M_v$ values [1-3, 13, 25-30].

**3.4 WAXD Analysis of PAN Polymers**

WAXD curves of Homo-PAN and different P(AN-co-IA)s measured under room temperature are depicted in Figure 3. It is shown that all the PAN polymers had the strongest diffraction peaks appearing at 2θ around 17°. They were corresponded to (100) crystalline plane of the pseudohexagonal cell or (200) reflections of the orthorhombic structure with their $d$ values equaling about 0.53nm [31-33]. It is noticed that the centers of the strongest diffraction peaks hardly changed with the increase of required IA amounts. Therefore, $d$ values of different PAN polymers basically remained unchanged. But the changes of peak intensity and peak area caused changes of crystallinity in different PAN polymers.

**Table 4. Parameters for WAXD curves of Homo-PAN and different P(AN-co-IA)s**

<table>
<thead>
<tr>
<th>Samples</th>
<th>2θ (°)</th>
<th>$d$ (nm)</th>
<th>FWHM (°)</th>
<th>$L_c$ (nm)</th>
<th>$X_c$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Homo-PAN</td>
<td>16.92</td>
<td>0.524</td>
<td>1.298</td>
<td>6.125</td>
<td>53.1</td>
</tr>
<tr>
<td>PI-1</td>
<td>17.10</td>
<td>0.519</td>
<td>1.272</td>
<td>6.251</td>
<td>48.9</td>
</tr>
<tr>
<td>PI-2</td>
<td>16.98</td>
<td>0.522</td>
<td>1.270</td>
<td>6.260</td>
<td>48.0</td>
</tr>
<tr>
<td>PI-3</td>
<td>16.95</td>
<td>0.523</td>
<td>1.273</td>
<td>6.245</td>
<td>47.5</td>
</tr>
<tr>
<td>PI-4</td>
<td>16.98</td>
<td>0.522</td>
<td>1.409</td>
<td>5.642</td>
<td>43.3</td>
</tr>
</tbody>
</table>

**3.5 FTIR Analysis of PAN Polymers**

FTIR of powdery Homo-PAN and P(AN-co-IA)s are described in Figure 4. It is obvious that the stretching vibration (ν) of nitrile group (C≡N) at 2240 cm⁻¹ was the
strongest peak in FTIR spectra. And the 2240 cm\(^{-1}\) band remained unchanged in the Homo-PAN and P(AN-co-IA)s. It is indicated that the AN units appeared the status of uninterrupted long sequences [34]. A shoulder-like absorption peak around 2190 cm\(^{-1}\) near the C≡N group might be due to the vC≡NH formed during the ADP process [35]. Besides, the stretching vibration around 2940cm\(^{-1}\) and bending vibration (δ) around 1450cm\(^{-1}\) of methylene group (CH\(_2\)) were the stronger peak in the two spectra. During ADP of AN and IA, C≡N would be changed into amide group (CONH\(_2\)) or carboxyl group (COOH) owing to hydrolysis under water bath heating [28,36]. Thus, a double broad absorption peak at 3628cm\(^{-1}\) and 3546cm\(^{-1}\) belonging to stretching vibration of NH\(_2\) and a weak absorption peak at 1621cm\(^{-1}\) belonging to bending vibration of NH\(_2\) could be exhibited in FTIR of Homo-PAN and P(AN-co-IA)s. Simultaneously, a weak and broad band at 3248cm\(^{-1}\) might belong to double-frequency band of δNH\(_2\) (DFB-δNH\(_2\)) at 1624cm\(^{-1}\).

4. Conclusion

Powdery HMW-PAN polymers were synthesized by ADP technique using AIBA as initiator. According to investigations of thermal properties and structure characters using TA, WAXD and FTIR, some conclusions can be obtained as follows:

The different PAN polymers synthesized had the higher molecular weights and higher conversions with the required IA amounts at 1wt%, 2wt% and 4wt%.

The use of IA into the polymerization system was available for relaxing the thermooxidation process of different PAN copolymers. Simultaneously, DSC curves of the PAN copolymers under air atmosphere revealed the doublet peaks phenomenon, which was not evident in Homo-PAN and in N\(_2\) atmosphere. And TGA curves of the PAN polymers were divided into three stages. More IA used in the ADP system made the P(AN-co-IA)s have lower initiation decomposition temperature and larger loss weight value.

(3) IA in the feed caused destructive effect to the crystal structure of PAN polymers, which made the crystalinity and grain sizes of the PAN polymers decrease. The nitrile group (C≡N) stretching vibration at 2240cm\(^{-1}\) and stretching vibration and bending of CH\(_2\) were very strong in both FTIR. In Homo-PAN, a weak absorption at 1730cm\(^{-1}\) was attributed to the C=O group stretching vibration because of hydrolysis of C≡N. However, it became stronger in different P(AN-co-IA)s with the increase of required IA dosages in the feed.

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References


