Synthesis of Pyromellitic Dianhydride (PMDA) and Tetraoctyl Pyromellitate (TOPM) by Alkali Oxygen Oxidation of Anthraeite and Esterification of PMDA with Isooctyl Alcohol

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Abstract: Pyromellitic dianhydride (PMDA) was prepared by an alkali oxygen oxidation of anthracite and selectively decarboxylation of pyromellitic acid (PA). Tetraoctyl Pyromellitate (TOPM) was synthesized by the esterification of PMDA with isooctyl alcohol. The influence of temperature and time on the yield of PMDA was discussed. The effect of the amount of catalyst, weight ratio of PMDA and isooctyl alcohol on the percent conversion of TOPM was also investigated. The results indicate that worthy chemicals such as PMDA and TOPM can be prepared from coal.

1 INTRODUCTION

It is very important to develop effective methods for utilizing anthracite, which are most abundant fossil resources. Benzene polycarboxylic acid (BPCA) and its ramification are important materials of organic chemistry, and they are basic materials for the high value-added product [Song C,2002][Liu Fang-Jing, 2013][Wang Wenhua, 2013]. They usually produced from oxidation of aromatic hydrocarbon under the condition of high temperature, high pressure, etc. The drawback of the method is the high production cost, inactivation of catalyst and serious environment pollution, etc. To overcome the drawback, several methods have been proposed, and preparation of BPCA from coal is one of such methods [Calemma V,1994][Miurak, 2000][Miura, 2001].

Many aromatic and heteroatom containing compounds can be converted into monomers for high performance polymers, and many of these monomers can be obtained by oxidation of corresponding alkyl aromatics.

Polish Academy of Sciences Institute of Organic Chemistry studied decarboxylation dynamics using benzene-pentacarboxylic acid and benzene-hexacarboxylic acid as materials in the early 1970's. The reaction is under room temperature, and in the mixed medium of concentrated H$_2$SO$_4$ and KHSO$_4$. Benzene-pentacarboxylic acid and benzene-hexacarboxylic acid firstly dehydrated to form symmetrical dianhydride under 300 °C, and then converted into pyromellitic dianhydride through selectively decarboxylation when the temperature higher than 300 °C. In 1986, Salbut et al. investigated pyromellitic acid production from coal in the two-stage process of oxidation and decarboxylation. Chemical ways of aromatic polycarboxylic acids conversion to such valuable byproducts as phthalic, isophthalic or pyromellitic acids (PA) are presented [Salbust P D, 1986][Schlenk H, 1960]. In 1994, Dalian university of technology explored different coal type's Efficiency of controlled coal oxidation in an alkaline medium for the aim of preparing PA. The results indicate that 7~8% (daf) pyromellitic acid can be obtained from Yangquan anthracite.

The aim of this study was to through the two steps: alkali oxygen-oxidation and selectively decarboxylation for preparing PA. After that, used PA, which prepared from coal, and isooctyl alcohol to prepare tetraoctyl Pyromellitate (TOPM), a kind of plasticizer. Extend the downstream of coal and achieve the aim of coal's clean utilize. The structure of Pyromellitic dianhydride (PMDA) and Tetraoctyl pyromellitate (TOPM) were characterized by Fourier transform spectrometer (FT-IR),
1H nuclear magnetic resonance spectra (1H-NMR) and 13C nuclear magnetic resonance spectra (13C-NMR).

2 EXPERIMENTAL SECTION

2.1 Materials

Jingmei anthracite, which purchased from Yunnan was grounded to less than 0.35 mm particles then dried for 24 h before used. The detailed analysis of Jingmei anthracite is shown in Table 1 and Table 2, respectively.

| Table 1. Proximate analyses (wt %) of Jingmei anthracite |
|-----------------|------|------|------|------|
| Mad             | 2.98 | 19.88| 4.83 | 72.29|

| Table 2. Ultimate analyses (wt %) of Jingmei anthracite |
|-----------------|------|------|------|------|
| C               | 69.32| 2.94 | 0.92 | 26.82|
| H               |      |      |      |      |
| N               |      |      |      |      |
| O_{diff}        |      |      |      |      |
| S_{tot}         |      |      |      | 1.99 |

Concentrated H$_2$SO$_4$ (98%), KHSO$_4$, dilute hydrochloric acid, potassium hydroxide, isoctyl alcohol, tetrabutyl titanate, active carbon, activated clay, butanone and 2-pentanone are all purchased from Alpha Chemical Co., Ltd. Zhengzhou., and were used as received without further purification.

2.2 Alkali-oxygen oxidation of coal in aqueous solution of potassium hydroxide

A 50 g of Jingmei anthracite, 50 g of potassium hydroxide, and 160 g of distilled water were put into a 1.5L autoclave. The autoclave was heated to 280 °C and a pressure of 6.0 MPa O$_2$ (cold) was used. The reaction time was 1.0 h while the reactor was agitated at an average of 500 cycles/min to keep reactants well mixed. The reaction products were filtered with distilled water to remove insoluble residue (cinder + mineral) after the reaction was over.

The pH of the filtrate was acidized by Concentrated H$_2$SO$_4$ (98%) to 3.8 and the humic acid was filtrated and throw away. The pH of the filtrate was acidized by Concentrated H$_2$SO$_4$ (98%) once again to 1.7, and the produced humic acid was also filtrated and throw away again. After that, the filtrate was extracted by butanone time and again, and the butanone was recovered by a vacuum rotary drier. The remaining concentration was dried to get BPCA.

2.3 Selectively decarboxylation of BPCA and the refining of PMDA

50 g BPCA, 150 g concentrated H$_2$SO$_4$, 350 g KHSO$_4$ (the weight ratio of BPCA, concentrated H$_2$SO$_4$ and KHSO$_4$ were 1:3:7) and 400 ml distilled water were charged into a 1000 ml three-necked flask while adding a little zeolite to the flask. The three-necked flask was equipped with an air set pipe and was put in a sand bath furnace, and heated the sand bath furnace to 300 °C and maintained for 2 h.

After that, cooling the reaction products slightly and filtered them, and then the filtrate was extracted by butanone for five times. The extraction was concentrated under vacuum rotary evaporation. The butanone was recovered, and the extraction was dried in a vacuum drying oven to get crude PA.

Adding 2-pentanone to crude pyromellitic acid under constant stirring, heated the solution to the boiling point of 2-pentanone. Then adding activated charcoal and the hot solution was filtration to remove insoluble impurities; the filtrate was cooled, and the pyromellitic acid crystal was precipitated. Using 2-pentanone washed the crystals several times and the crystals were heated in vacuum chamber at 260 °C for 3 h to get pure PMDA.
Fixing the reaction time at 2 h and performed the selectively decarboxylation of BPCA at 250 °C, 300 °C, 350 °C and 400 °C, respectively to study the influence of the reaction time to the yield of PA.

Fixing the reaction temperature at 350 °C and performed the selectively decarboxylation of BPCA at 15min, 30min, 45min and 60 min, respectively to study the influence of the reaction temperature to the yield of PA.

2.4 Synthesis of TOPM

1 mole pure PMDA, which prepared from coal, 5 mole isoctyl alcohol and 5 g tetrabutyl titanate were put into a 1000 ml three-necked flask equipped with thermometer, water knockout drum and reflux condensing tube. The mixture was heated from atmospheric temperature to 220 °C and controlled the reaction temperature to 220 ±10 °C. The generated water was periodically measured from the water knockout drum, and stopped the reaction when the acid number of the product under 0.2 mgKOH/g.

The three-necked flask was linking-up to a vacuum pump to pull off the remaining isoctyl alcohol until there was no isoctyl alcohol can be seen in the reflux condensing tube. And then, 1% active carbon of the products was added to decolorize. Afterwards, the products which through decolorize were put into a sand core funnel to carry out vacuum suction filter, and 10 cm activated clay was spread out in the sand core funnel as filter aid.

Fixing the reaction temperature at 220 °C, changed the weight ratio of PMDA and isoctyl alcohol at 1:1, 1:3, 1:5 and 1:7, respectively to investigate the effect of weight ratio of PMDA and isoctyl alcohol on the percent conversion of PMDA.

Fixing the reaction temperature at 220 °C and the weight ratio of PMDA and isoctyl alcohol at 1:5, changed the amount of tetrabutyl titanate catalyst at 1 %, 3 %, 5 % and 8 % (based on the total weight of PMDA and isoctyl alcohol) to investigate the effect of weight ratio of catalyst on the percent conversion of PMDA.

3 RESULTS AND DISCUSSION

3.1 Influence of temperature on the yield of PA

![Figure 1. Effect of temperature on the PA yield.](image)

Figure 1 gives the influence of temperature on the crude PA yield. When the time of the selectively decarboxylation of BPCA keep at 2 h, the yield of PA continue to show an upward tendency directly with temperature while the time of the selectively decarboxylation of BPCA keep at 2 h. The yield of PA reached its highest value: 43.98%. The yield of PA decreased when the temperature continue to rise.
The results illustrate that temperature has great influence on the yield of PA. The reaction speed is very slow and it needs a long time to selectively decarboxylate, and the yield of PA is quite low. Obviously, it doesn’t really amount to much. But then, too high temperature will make PA further decomposed, and it also lead to the yield of PA decrease. The most suitable temperature for selectively decarboxylation is 2 h.

3.2 Influence of reaction time on the yield of PA

Figure 2 shows the effect of reaction time on the PA yield. The selectively decarboxylation reaction of BPCA is insufficient when the reaction time is relatively short, and the yield of PA is rather low. It presents the tendency of increase with the extension of the reaction time. The yield of PA begins to decrease after it reaches its peak value.

It is generally supposed that the process of coal oxidation is: Coal → humic acid → coal acids → CO₂ + H₂O. The first stage engages in the surface of coal particles, and its speed is comparatively rapid. The second stage engages in the liquid phase, and its speed is slower than the first stage. Too long reaction time makes the speed of the third stage accelerate, then the PA is be further oxidized. The result is that the ultimate yield of PA decreases, and the results illustrate that 45 minutes is the most favourable time.

3.3 Influence of weight ratio of PMDA and isoocetyl on the percent conversion of PMDA

Figure 3 shows the effect of weight ratio of PMDA and isoocetyl on the percent conversion of PMDA by looking at Figure 3. The higher of the weight ratio of PMDA and isoocetyl leads to the higher of
the percent conversion of PMDA. That is to say, higher weight ratio of PMDA and isooctyl is beneficiafor the percent conversion of PMDA. But too high weight ratio of PMDA and isooctyl has rendered the pressure of the reaction rise, and it involves the difficulty of isooctyl reflux. It turns the percent conversion of PMDA decrease.

3.4 Influence of weight ratio of catalyst on the percent conversion of PMDA

We can recognize from Figure 4 that the more quantity of catalyst adding involve higher percent conversion of PMDA. But the percent conversion of PMDA has not obviously increased when the adding of catalyst exceeded 5%. Besides, more catalyst adding increases the difficulty of production post-treating.

3.5 Analysis of PMDA and TOPM

The structure of PMDA was confirmed by $^1$H-NMR and $^{13}$C-NMR, and the structure of TOPM was confirmed by $^1$H-NMR and FT-IR.

FT-IR spectral data were obtained on a Digilab FTS-20E system by using KBr disc techniques. The mixture of 0.8 to 1.2 mg (+ 0.01 mg) of sample and 95-105 mg (+0.01 mg) of KBr was finely ground for at least 15 minutes to assure the optimum measurement and then pressed under vacuum with a load of 8000 kg.

The $^1$H-NMR spectra were recorded in deuteriochloroform solutions in the usual way. As for $^{13}$C-NMR, the sample were recorded in (CD$_3$)$_2$SO solutions, and a relaxation agent, chromium trisacetylacetonate (0.1 M) was added in an inverse gated decoupling system to get quantitative $^{13}$C-NMR spectroscopy.

As for the $^1$H-NMR of PMDA, there is only one kind of proton: 8.0 (Ar-H) appears at 8.73 ppm. The sharp peak at 2.5 ppm could be assigned to (CD$_3$)$_2$SO solvent peak and the small impurity peak appears at 8.27 ppm.

For the $^{13}$C-NMR of PMDA, the carbonyl carbon atom exhibits at 161.9 ppm; the tertiary carbon atom shows at 137.6 ppm and the quaternary carbon atom appears at 124.8 ppm. The heights of the spectrum are broadly in line with 2 : 2 : 1, which reflected the type and number of carbon atoms in PMDA. Some petty impurity peaks display at 121.7, 133.1, 139.4, 161.3 and 166.8 ppm.

As for the FT-IR of TMOP, The -CH$_3$- and -CH$_2$- absorption show intense peak at 2800 to 3000 cm$^{-1}$, and the characteristic bands of 1730 cm$^{-1}$ is the (-C=O) stretching vibration. The vibration of asymmetric bending is shown at 1459 cm$^{-1}$ and the strong sharp peak at 1104 cm$^{-1}$ is attributed to (-OCH$_2$CH$_2$-) stretching vibration. The (-C=C-) stretching is shown a sharp intense peak at 1643 cm$^{-1}$. The C=C stretching is shown as a sharp intense peak at 1622 cm$^{-1}$, 1023 cm$^{-1}$ and 952 cm$^{-1}$ for C-H plane deformation vibration of =CH$_2$.This 1602 and 1502 cm$^{-1}$ absorptions are at-
tributed to the existence of benzene nucleus. The FTIR proves that the synthesized monomer has the anticipated structure.

For the $^1$H-NMR of TOPM, the signal at $\delta = 8.6$ ppm assigned to the (Ar-2H). The chemical shifts at $\delta = 4.28$ ppm and $\delta = 4.21$ ppm resulting from the ether (CH$_2$-O-CH$_2$-, 8H) groups. $\delta = 1.51$, 1.34, 1.00, 0.92, 0.88 ppm were the high coupling peak at the structure of -CH-CH$_2$-CH$_2$-CH$_2$-CH$_3$.

From FT-IR measurements and $^1$H-NMR analysis, it can confirm that synthetic TOPM have the expected structure.

4 CONCLUSION

1. The authors have prepared pure PMDA and TOPM from anthracite, and survived better reaction conditions for the two compounds. Studied multifarious factors influenced on the yield of the two compounds.

2. Under optimal conditions, the yield of PMDA is 14.7% based on the weight of BPCA and 11.2% based on the weight of daf. coal. The yields of TOPM is 11.21% based on the weight of BPCA and 6.57% based on the weight of daf. coal.

3. The optimal conditions for PMDA production: 300 °C, 45 min and the PMDA purity can be further improved by recycling the mother liquid of 2-pentanone recrystallization. The optimal conditions for TOPM production: 220 °C, 45 min, and the weight ratio of PMDA and isooctyl alcohol is 1:5.

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