Study on the Isomerization Reaction of Turpentine Oil Catalyzed by the Acid-treated HangJin 2# Clay

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Abstract- In order to reduce costs and improve the activity of catalyst in turpentine oil isomerization reaction, two solid acid catalysts (b and c) were prepared by using sulfuric acid and hydrochloric acid treated HangJin 2# clay (a), respectively. On the basis of sulfuric acid-treated HangJin 2# clay (b) as carrier, SO4²-/HangJin 2# clay (d) and PO4³-/HangJin 2# clay (e) catalysts were prepared by dipping sulfuric acid and phosphoric acid in catalyst b, respectively. Five catalysts (a, b, c, d and e) were characterized by TPD, TG-DTA and BET. Moreover, the isomerization reaction of turpentine oil catalyzed by a, b, c, d, and e were also investigated. The results revealed that the catalytic activities as follow: b > c, e > d.

Keywords- HangJin 2# clay; solid acid; catalyst; turpentine oil; isomerisation.

I. INTRODUCTION

The isomerization reaction of turpentine oil is one of the most important chemical reactions in turpentine deep processing, which usually catalyzed by solid acid catalyst [1], molecular sieve [2], clay[3], etc.. Solid acid catalysts show highly active in turpentine oil isomerization reaction, but there still exist some problems such as complicated preparation process, high cost and short service life. Although molecular sieve show large specific surface area, simple preparation and recycle, problems remain in terms of low selectivity and easy carbon deposition. In recent years, Clay has received widespread attention owing to their low price, high activity, recycle and simple preparation [4].

In recent years, it has been found that HangJin 2# clay shows high catalytic activities and as potential applications in catalysis[5]. Jingfang Wei et al. found that sulfuric acid and hydrochloric acid treated HangJin 2# clay show larger BET and more stability with the conversion of the esterification reaction up to 99%[6]. Yuli Zhang and co-workers have reported a SO4²-/Fe-Al-activated clay catalyst with a yield of 45% after using in turpentine hydration reaction [7]. Limei Bai et al. prepared a SO4²-/SnO2-HangJin 2# clay catalyst with the reaction conversion rate up to 99%[8]. However, so far work on the preparation of HangJin 2# clay catalysts and its application in the isomerization reaction of turpentine oil have not been reported.
In this paper, we report the preparation, characterization and application of four new HangJin 2\textsuperscript{a} clay base catalysts.

II. EXPERIMENTAL

A. Preparation of Catalysts

Activated clay: A certain amount of HangJin 2\textsuperscript{a} clay was mixed with hydrochloric acid, sulfuric acid at solid-liquid ratio of 1:2.5 (g/mL) and kept stirring and reflowing for 4 hours at 90 °C. The filter cake was washed by distilled water, and dried under 110 °C for 3 hours and then triturated.

Prepared based on the impregnation method. The activated clay was mixed with sulfuric acid or phosphoric acid respectively at solid-liquid ratio of 1:5 (g/mL) and kept stirring and reflowing for 4 hours at room temperature. Filtration: filter cake in a drying box for drying, grinding and sieving. Then it was rubbed through a 100 mesh sieve, and calcinated for 3 hours in the muffle furnace at 350 °C.

B. Physical Measurements

Thermogravimetric analyses (TGA) were performed on an automatic simultaneous thermal analyzer (DTG-60, Shimadzu) under air at a heating rate of 20°C/min between ambient temperature and 800 °C.

Infrared spectra were recorded (4000-400 cm\textsuperscript{-1}) as KBr disks on a Shimadzu IR-8400 spectrometer.

NH\textsubscript{3}-TPD analysis was performed on multifunctional adsorption equipment, High purity nitrogen as the carrier gas, Ammonia is adsorbed gas, in 10°C/min heating rate from room temperature up to 600°C.

The analysis of BET using American Mike company production of ASAP-2010, samples under vacuum pretreatment to 423K, nitrogen adsorption in 77.35K (Liquid nitrogen).

C. Catalytic Reactions

A mixture of turpentine oil (10 mL) and 3% of catalyst was stirred at 150 °C for 3 hours under air. After the reaction, the catalyst was separated by filtration. The filtrate was dried over Na\textsubscript{2}SO\textsubscript{4} and filtered. The products were analyzed by GC-MS analysis (Shimadzu GCMS-QP5050A equipped with a 0.25 mm × 30 m DB-WAX capillary column). The conversion and selectivity of the product (pinene) were calculated based on Area Normalization Method\textsuperscript{[9]}.

\[
\text{Conversion of pinene} = \frac{\text{Variation of pinene}}{\text{Content in raw material pinene}} \times 100\% \\
\text{Camphene selectivity} = \frac{\text{Camphene content}}{\text{The conversion rate of pinene}} \times 100\%
\]

III. RESULTS AND DISCUSSIONS

A. Turpentine Isomerization reaction catalysis.

The result of turpentine isomerization reaction catalyzed by three different catalysts (a, b and c) was shown in Table 1. From the table 1, compared to HangJin 2\textsuperscript{a} clay (a), the catalytic activities of acid treated HangJin 2\textsuperscript{a} clay (b and c) were obviously improved with the conversion of pinene up to 99.09% and 100%, respectively. Moreover, the BET of b and c were also significantly improved, up to 126.58 and 146.54 m\textsuperscript{2}/g, respectively. Compared to a, the principal product (camphene) catalyzed by b and c up to 35.75 and 32.41%, respectively. In summary, the acid treated HangJin 2\textsuperscript{a} clay (b and c) show larger BET and higher catalytic activity than a maybe due to the pore structure of b and c were opened after acid treatment, which causing the catalyst and the solution full contact. However, catalyst b shows higher activity than c with higher yield of principal product and lower yield of by-products.

| Sample | BET(m\textsuperscript{2}/g) | Conversion of pinene(%) | The selectivity of isomerization(%) |
|--------|------------------|------------------------|-----------------------------------|
| a      | 62.62            | 13.98                  | 55.60                             |
| b      | 126.58           | 99.09                  | 40.58                             |
| c      | 146.54           | 100.00                 | 36.22                             |

(a)HangJin 2\textsuperscript{a} clay (b)sulfuric acid treated HangJin 2\textsuperscript{a} clay (c)hydrochloric acid treated HangJin 2\textsuperscript{a} clay

The TPD spectra of catalysts (a, b, c) are shown in Fig. 1., the desorption peak of the weak acid sites move to the high temperature after treating the HangJin 2\textsuperscript{a} clay using acid. Compared to hydrochloric acid treated HangJin 2\textsuperscript{a} clay, which the stripping peak of weak/strong acid center move to 200°C /520°C, the sulfuric acid treated HangJin 2\textsuperscript{a} clay appeared in 125°C /510°C. The results reveal that the acidity of the weak acid will be improved after acid treatment of HangJin 2\textsuperscript{a} clay. Moreover, the acid content of the weak acid center of the catalyst help the turpentine isomerization reaction.

Figure 1. The TPD spectra of acid treated HangJin 2\textsuperscript{a} clay

(a)HangJin 2\textsuperscript{a} clay (b)sulfuric acid treated HangJin 2\textsuperscript{a} clay (c)hydrochloric acid treated HangJin 2\textsuperscript{a} clay

The TG curves of catalysts (a, b, c) are shown in Fig. 2. All of three catalysts show two weight loss steps. The first corresponding to the release of adsorbed water and crystal water is observed from 25-200°C. Then weight-loss step occurred between 400-800°C which corresponds to the release of the load acid. The weightlessness ratio of...
b larger than e, indicates that the acid load of b higher than c. It has been found that the acid load content play an important role in the number and acid strength of catalysts, which are in good agreement with the Table 1. However, catalyst a performs low catalytic activity is not agree with the high weight-loss between 400-800°C, which the weight-loss mainly from the surface of the acid load of catalyst. Thus, the acid strength of catalysts is one of the main reasons to affect the activity of catalyst. Moreover, the reaction activity of catalysts is not only related with the catalyst acidity, but also related to the distribution of acid sites. Meanwhile, catalyst activity strongly depends on the weak center, especially when the catalysts were used in the isomerization reaction of turpentine [10].

The FT-IR spectra of catalysts (a, b, c) are shown in Fig. 3. In the IR spectrum, the bands at 3600 cm⁻¹ may be due to the ν(-OH) stretching vibrations, the bands at 3420 cm⁻¹ may be due to the ν(H₂O) stretching vibrations. After the acid treated HangJin 2# clay, the bands of the ν(Fe₂O₃) stretching vibrations at 1460 cm⁻¹ was disappeared. The characteristic peaks between 1020 cm⁻¹ and 1150 cm⁻¹ are associated with the solid super acid. After the acid treated agent, the strength of characteristic peaks were increased. The characteristic peaks of b catalyst were stronger than a and e catalyst, which indicated that the b catalays performs superior activity.

B. The effects of Acid Dipping Activated Clay of Turpentine Isomerization reaction.

The result of turpentine isomerization reaction catalyzed by two different catalysts (d and e) was shown in Table 2.

From the Table 2, compared to catalyst b, the catalytic activities of the acid dipping SO₄²⁻/activated clay (d) and acid dipping PO₄³⁻/activated clay (e) show low conversion of pinene (17.00% and 28.53%) and low BET (78.87 and 9.80 m²/g), which maybe due to the pore structure were loaded by acid and reduce the BET of d and e. The results reveal that the BET play an important role in catalytic activities.

| Sample | BET(m²/g) | Conversion of pinene(%) | The selectivity of isomerization(%) |
|--------|-----------|-------------------------|-----------------------------------|
| b      | 126.58    | 99.09                   | 40.58 59.42                       |
| d      | 78.87     | 17.00                   | 57.85 42.15                       |
| e      | 9.80      | 28.53                   | 34.76 65.24                       |

(b) sulfuric acid treated HangJin 2# clay; (d) SO₄²⁻/activated clay; (e) PO₄³⁻/activated clay

The TG curves of catalysts (b, d, e) are shown in Fig. 4. All of three catalysts show two weight loss steps. The first corresponding to the release of adsorbed water and crystal water is observed from 25-200°C. Then weight-loss step occurred between 400-800°C which corresponds to the release of the load acid (obsd. 2.5%, 12.7% and 0.71%). Too much acid were loaded in catalyst d, result in the acid coated in the surface of catalysts and reduce the catalytic reactivity after high-temperature roasting[11]. The low weight-loss of catalyst e relate to the phosphor acid concentration and volume. It has been found that small amount of phosphoric acid or excessive amount of phosphoric acid were used will make the BET and catalytic activity decrease. On the basis of above mention, the amount of phosphoric acid plays a crucial role in catalytic activity.
The FT-IR spectra of catalysts (b, d, e) are shown in Fig. 5. In the IR spectrum, the bands at 1400 cm⁻¹ may be due to the \( \nu(S=O) \) stretching vibrations. The characteristic peaks between 1050 cm⁻¹ and 1150 cm⁻¹ were reduced. The characteristic peaks of d catalyst were stronger than e catalyst, which indicated that the d catalyst performs superior activity.

**IV. CONCLUSIONS**

In summary, we have successfully prepared four different catalysts based on HangJin 2# clay and were used in the isomerization reaction of turpentine, with the conversion of pinene up to 99%. The results reveal that the acid play an important role in catalytic activity. Moreover, the acidity of the weak acid sites was increased will improved the yield of the camphene main product.

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