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**Article**

**Catalytic Fast Pyrolysis of Kraft Lignin over Hierarchical HZSM-5 and Hβ Zeolites**

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**Abstract:** The hierarchical HZSM-5 and Hβ zeolites were prepared by alkaline post-treatment methods adopting Na$_2$CO$_3$, TMAOH/NaOH mixture, and NaOH as desilication sources, respectively. More mesopores are produced over two kinds of zeolites, while the micropores portion is well preserved. The mesopores formed in hierarchical Hβ zeolites were directly related to the basicity of the alkaline solution, indicating that Hβ zeolite is more sensitive to the alkaline post-treatment. The hierarchical HZSM-5 and Hβ zeolites are more active than the parent one for catalytic fast pyrolysis (CFP) of Kraft lignin. Hierarchical zeolites retained the function of acid catalysis, while additionally creating larger mesopores to ensure the entry of bulkier reactant molecules. The increase of the condensable volatiles yield can be attributed to the improvement of the mass transfer performance, which correlates well with the change of mesoporous surface area. In particular, the condensable volatiles yield for the optimized hierarchical Hβ reached approximately two times that of the parent Hβ zeolites. In contrast to the parent HZSM-5, the optimized hierarchical HZSM-5 zeolite significantly reduced the selectivity of oxygenates from 27.2% to 3.3%.

**Keywords:** catalytic fast pyrolysis; Kraft lignin; hierarchical zeolites; HZSM-5; Hβ

1. Introduction

Lignocellulose is the most abundantly available raw material on the earth for the production of biofuels. It is mainly composed of carbohydrate polymers (cellulose, hemicellulose), and a cross-linked aromatic polymer (lignin). Bio-oils derived from lignocellulose pyrolysis are characterized by high oxygen content, high acidity, and high viscosity, which significantly limit its utilization in the existing petrochemical infrastructure [1]. Removing oxygen from raw materials efficiently is usually a challenge for the catalytic fast pyrolysis (CFP) of lignocellulose to produce a high-energy density fuel with good combustion performance [2]. Generally, pyrolysis vapors can be upgraded by catalytic conversion over solid acid catalysts such as zeolites to generate products of lower oxygen content [3].

As the most recalcitrant component of lignocellulose, lignin has a high resistance to chemical attacks such as pyrolysis [4]. Monomeric phenolic compounds are the primary pyrolysis products of lignin and easily recombine to produce oligomeric compounds [5]. Lignin-derived phenols, which are strongly adsorbed on zeolite acidic sites, could act as a coke precursor and form a large number of cokes [6]. In CFP process, coke formation not only leads to rapid deactivation caused by blocking zeolite pores, but also declines the carbon conversion efficiency. Compared with the other two components, although lignin itself is composed of aromatic polymer structure, it has the lowest aromatics yield and highest coking under the same CFP conditions [7].
Microporous zeolites such as HZSM-5 are known to be effective deoxygenation catalysts, which can convert pyrolysis vapors of lignocellulose directly to olefins and aromatics [7–14]. However, due to size exclusion or pore blockage, HZSM-5 was invalid in the treatment of heavier feeds such as syringyl monolignols derived from CFP of lignin [13]. The diffusion and mass transfer resistance of molecules in and out of micropores often limit the space utilization in microporous zeolites; it is necessary to optimize the accessibility of zeolites to these components. Aluminosilicates possess a much larger pore size; however, CFP of lignin over this mesoporous material yields a similar aromatics and coke as HZSM-5 zeolite [15]. The relationship between the acidity and porosity of mesoporous aluminosilicates with product selectivity during CFP of lignin is hard to establish. The introduction of hierarchical zeolite, which has interconnected mesopores and micropores, is a promising approach for solving this problem [16–19]. Hierarchical zeolite retained the function of acid catalysis, while additionally creating larger mesopores to ensure the entry of bulkier reactant molecules. Puertolas had shown that the aromatics fraction for bio-oils conversion was correlated with the accessible Brønsted acid sites at the mesoporous surface for hierarchical zeolites [10]. Pinard had demonstrated that mesopores can be seen as “highways”, where big molecules can diffuse to more accessible Brønsted sites on micropores [11]. A hierarchical zeolite presents much higher selectivity to mono-aromatics and stability upon coke deposit than microporous zeolite.

The most feasible preparation method for the hierarchical zeolite is a desilication of a microporous zeolite using an alkaline solution [16–19]. In this study, the influence of different alkaline desilication sources on the formation of intracrystalline mesopores through hydrolysis of the skeleton atom was investigated. The hierarchical HZSM-5 and \( \text{H}\beta \) zeolites with moderate acidity and pore structure were successfully prepared by an optimized alkaline treatment. The CFP of lignin over hierarchical HZSM-5 and \( \text{H}\beta \) zeolites was analyzed on-line by Py-GC/MS technique, by evaluating the impact of mass transfer performance on the yield of condensable volatiles and the deoxidation effect.

2. Results

In order to create more open mesopores in the microporous zeolites, four different treatment methods were investigated. The alkaline post-processing adopted \( \text{Na}_2\text{CO}_3 \) (Treatment A), TMAOH/NaOH mixture (Treatment B), and NaOH (Treatment C) as desilication sources respectively, reacting at 65 °C for 1.5 h. A sequential mild acid treatment with HCl was performed after desilication of zeolites with NaOH (Treatment D), in order to remove Lewis acid sites generated after mesoporosity introduction.

2.1. Nitrogen Physisorption Characterization

Table 1 summarizes the texture properties of HZSM-5 zeolites obtained by different post-treatment methods. The parent HZSM-5 is characterized by typical microporous material. The mesoporous surface area is only 18 \( \text{m}^2/\text{g} \), which is negligible compared to the microporous surface area of 395 \( \text{m}^2/\text{g} \). For HZSM-5 after different alkaline treatments, the mesoporous surface area all approximately increase to about 34 \( \text{m}^2/\text{g} \), while the corresponding microporous surface area decreases synchronously. A further mild HCl washing after desilication with NaOH makes little change to mesopores, but the microporous surface area was restored to 394 \( \text{m}^2/\text{g} \) for HZSM-5-D. The changes of mesoporous volume and microporous volume in each sample were in accordance with the corresponding trends of specific surface area. Figure 1 shows the pore size distribution of HZSM-5 zealites obtained by different post-treatment methods, which was calculated according to the adsorption branch of isotherms using the BJH method. It is obvious that the microporous size distribution of parent HZSM-5 is very concentrated. Alkaline treatment has no effect on the location of the distribution maximum for micropore, but the intensity decreases. For hierarchical HZSM-5, there are a small amount of mesopores formed in a pore width of 2–6 nm. Consistent with the specific surface area, the HZSM-5-D possess slightly more micropores than parent HZSM-5.
Table 1. Physicochemical properties of parent HZSM-5 and the hierarchical HZSM-5 obtained by different post-treatment methods.

<table>
<thead>
<tr>
<th>Sample</th>
<th>$S_{\text{Meso}}$ (m$^2$/g)</th>
<th>$S_{\text{Micro}}$ (m$^2$/g)</th>
<th>$V_{\text{Meso}}$ (cm$^3$/g)</th>
<th>$V_{\text{Micro}}$ (cm$^3$/g)</th>
<th>Rel. Crystal. (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HZSM-5</td>
<td>18</td>
<td>395</td>
<td>0.04</td>
<td>0.17</td>
<td>100</td>
</tr>
<tr>
<td>HZSM-5-A</td>
<td>34</td>
<td>363</td>
<td>0.07</td>
<td>0.15</td>
<td>85</td>
</tr>
<tr>
<td>HZSM-5-B</td>
<td>33</td>
<td>351</td>
<td>0.07</td>
<td>0.15</td>
<td>79</td>
</tr>
<tr>
<td>HZSM-5-C</td>
<td>35</td>
<td>369</td>
<td>0.08</td>
<td>0.16</td>
<td>91</td>
</tr>
<tr>
<td>HZSM-5-D</td>
<td>34</td>
<td>394</td>
<td>0.08</td>
<td>0.17</td>
<td>81</td>
</tr>
</tbody>
</table>

Table 2 summarizes the texture properties of Hβ zeolites obtained by different post-treatment methods. The specific surface area of parent Hβ is larger than HZSM-5, and contains a certain amount of mesopores itself (86 m$^2$/g). Different alkaline treatments process had varying impacts on the formed mesopores. There is only a slight increase in the mesoporous surface area of Hβ-A and Hβ-B. The mesoporous surface area of Hβ-C is twice as large as parent Hβ, while the microporous surface area decreased correspondingly. The mesoporous surface area of Hβ-D further increased, while the microporous surface area remained at 421 m$^2$/g. As can be seen from pore size distribution in Figure 2, the hierarchical Hβ still retains a large amount of micropores, while the newly-formed mesopores are not very concentrated. For Hβ-A and Hβ-B, the additional mesopores are mainly distributed in the range of 2–4 nm, while for Hβ-C and Hβ-D, the pore size distribution of the additional mesopores is shifted to larger pore diameters (3–6 nm and 4–9 nm, respectively).

Table 2. Physicochemical properties of parent Hβ and the hierarchical Hβ obtained by different post-treatment methods.

<table>
<thead>
<tr>
<th>Sample</th>
<th>$S_{\text{Meso}}$ (m$^2$/g)</th>
<th>$S_{\text{Micro}}$ (m$^2$/g)</th>
<th>$V_{\text{Meso}}$ (cm$^3$/g)</th>
<th>$V_{\text{Micro}}$ (cm$^3$/g)</th>
<th>Rel. Crystal. (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hβ</td>
<td>86</td>
<td>545</td>
<td>0.22</td>
<td>0.21</td>
<td>100</td>
</tr>
<tr>
<td>Hβ-A</td>
<td>116</td>
<td>535</td>
<td>0.28</td>
<td>0.21</td>
<td>70</td>
</tr>
<tr>
<td>Hβ-B</td>
<td>117</td>
<td>555</td>
<td>0.32</td>
<td>0.22</td>
<td>80</td>
</tr>
<tr>
<td>Hβ-C</td>
<td>171</td>
<td>450</td>
<td>0.36</td>
<td>0.18</td>
<td>67</td>
</tr>
<tr>
<td>Hβ-D</td>
<td>232</td>
<td>421</td>
<td>0.37</td>
<td>0.16</td>
<td>72</td>
</tr>
</tbody>
</table>
The hierarchical HZSM-5 zeolites all possess the typical diffraction peak original features of the parent one, indicating the crystal structure was basically maintained after alkaline treatments. At the same time, the intensity of the diffraction peak decreased to a certain extent, due to the decrease of the relative crystallinity for hierarchical zeolites. The relative crystallinity was obtained according to the intensity of the strongest diffraction peak, assuming that the relative crystallinity of the parent zeolites was 100%. For example, after desilication using NaOH solution, the relative crystallinity of Hβ-C was only 67% of the parent Hβ.

2.2. Structural XRD Characterization

Figure 2 shows the XRD patterns for parent HZSM-5 and the hierarchical HZSM-5 obtained by different post-treatment methods. Figure 3 shows the XRD patterns for parent Hβ and the hierarchical Hβ obtained by different post-treatment methods. The hierarchical zeolites all possess the typical diffraction peak original features of the parent one, indicating the crystal structure was basically maintained after alkaline treatments. At the same time, the intensity of the diffraction peak decreased to a certain extent, due to the decrease of the relative crystallinity for hierarchical zeolites. The relative crystallinity was obtained according to the intensity of the strongest diffraction peak, assuming that the relative crystallinity of the parent zeolites was 100%. For example, after desilication using NaOH solution, the relative crystallinity of Hβ-C was only 67% of the parent Hβ.
The high temperature peaks can be ascribed to strong Brønsted acid sites. It had been previously reported that Brønsted acid sites mainly exist over hierarchical zeolites formed in the partial desilication [18]. Na\(^+\) ions existed in the cationic sites of zeolite during alkaline treatment process, which could be removed by following ion exchanging with NH\(_4^+\); thus, the Brønsted acidity could be recovered.

2.3. NH\(_3\)-TPD Characterization

Figure 5 shows the NH\(_3\)-TPD patterns of parent H\(\beta\) and the hierarchical H\(\beta\) obtained by different post-treatment methods. Figure 6 shows the NH\(_3\)-TPD patterns of parent H\(\beta\) and the hierarchical H\(\beta\) obtained by different post-treatment methods. Two ammonia desorption peaks appear in all zeolites. The low temperature peak is derived from the weakly adsorbed ammonia molecules via hydrogen bonding, instead of the acid sites [19]. The high temperature peaks can be ascribed to strong Brønsted acid sites. It had been previously reported that Brønsted acid sites mainly exist over hierarchical zeolites formed in the partial desilication [18]. Na\(^+\) ions existed in the cationic sites of zeolite during alkaline treatment process, which could be removed by following ion exchanging with NH\(_4^+\); thus, the Brønsted acidity could be recovered.
Different alkaline treatments processes did not change the low temperature peak of the hierarchical HZSM-5. The high temperature peak remained essentially in the original position, indicating that alkaline treatment has tiny effect on the strength of the Brønsted acid sites. The peak area slightly decreased for HZSM-5-B and HZSM-5-C, indicating partial desilation reduces the number of Brønsted acid sites. The NH$_3$-TPD patterns of the hierarchical Hβ exhibit similar desorption peaks to the parent one, indicating the alkaline treatment of Hβ Zeolites may not significantly decrease the acidity. For the hierarchical zeolites, we also noted that the loss of relative XRD crystallinity did not correlate well with the amount of acid sites, especially for HZSM-5-C and Hβ-A. The characteristic tetrahedrally coordinated aluminosilicate framework of zeolites results in Brønsted acid sites. In the harsh treatment condition of strong alkaline environment, larger mesopores are created by a combination of amorphization and framework atom leaching, which both affect the amount of acid sites. The relationship between the amount of acid sites determined from NH$_3$-TPD and relative crystallinity is more complicated.
2.4. Catalytic Fast Pyrolysis of Kraft Lignin

Lignin is very chemically inert, due to its complex three-dimensional network structure formed by different non-phenolic phenylpropanoid units, which were linked with a variety of C–O and C–C bonds [2,4,5]. Pyrolysis can destroy these phenyl-propane units in the polymer lattice. As shown in Figure 7, the pyrograms of lignin over parent HZSM-5 and Hβ Zeolites were compared with those of non-catalytic fast pyrolysis. The yield of the condensable volatiles during Py-GC/MS experiments is semi-quantitatively evaluated by the sum of the absolute peak area of the total ion chromatogram peak (calibrated with the precise lignin sample weight), while the selectivity of a certain category of compound is represented by the percentage of peak area measured in total area. The light gases derived from the deoxygenation of oxygenates during CFP of lignin, mainly CO₂, CO, and other non-condensable gas, are not included in the sum of the absolute peak areas of condensable liquid products. As reported in previous studies by Bokhoven [15], the yield of light gases was about 5% when the lignin was pyrolyzed at 650 °C on different zeolites catalysts. Because the lignin has a high resistance to pyrolysis, the polymer structure can only be depolymerized into aromatic monomers at the pyrolysis temperature of 600 °C in this work; thus, the C–C bond will not be further broken to produce smaller molecules of light gases. It has also been reported previously that the yield of light gases produced by lignin pyrolysis is much lower than that of cellulose and hemicellulose [20]. Therefore, in the pyrolysis conditions of this work, pyrolysis products are mainly the condensable volatiles and coke correlating with each other. The products from lignin pyrolysis are very complex. It becomes very difficult to measure individual component. The retention time and the characteristic ion fragments of each compound in the total ion current chromatogram are shown in Table S1. Non-CFP of lignin mainly produces oxygen-containing species with very few aromatic hydrocarbons. During the process of lignin pyrolysis, Bokhoven verified that depolymerized lignin products undergo successive reactions to form phenol alkoxy ketones, phenol alkoxy, phenols, aromatic alkoxy, and, finally, aromatic hydrocarbon [14,15]. The presence of zeolites greatly changed the composition of pyrolysis products, leading to improved deoxygenation of the high oxygen-containing compounds into aromatics. The GC-detected condensable pyrolysis products were divided into 4 categories: oxygenates (predominantly phenols, guaiacols, and syringols), monocyclic aromatics (BTX), naphthalene, and polycyclic aromatic hydrocarbons (PAH). The retention time and the characteristic ion fragments of each compound in the total ion current chromatogram are shown in Table S2.

![Figure 7](image_url)

**Figure 7.** The pyrograms (Py-GC/MS) of catalytic fast pyrolysis of Kraft lignin over parent HZSM-5 and Hβ Zeolites. Experimental conditions: pyrolysis temperature of 500 °C, Kraft lignin 0.3 mg, catalyst-to-lignin ratio of 10:1.
The acid sites of zeolites can promote deoxygenation, decarboxylation, and decarbonylation, as well as cleavage, oligomerization, alkylation, isomerization, cyclization, and aromatization, to further convert the depolymerization products of lignin. Over the parent HZSM-5 and H\textbeta{} zeolites, there are significant differences in the selectivity of condensable volatiles. Using HZSM-5 catalyst, 27.2\% of oxygenates still exist in the condensable volatiles, mainly in trimethoxybenzene. On H\textbeta{} zeolite, oxygenates almost disappeared completely, only 1\%. The deoxygenation effect of H\textbeta{} is much higher than that of HZSM-5. Considering the difference in pore structure between the two zeolites, more lignin-derived oxygenates can diffuse into the larger pores of H\textbeta{}, even larger molecules trimethoxybenzene; thereby, most of oxygenates can be effectively converted. It has also been reported previously that HZSM-5 is invalid for the conversion of bulkier lignin monomers due to size exclusion or pore plugging, while H\textbeta{} is very effective for deoxygenation of lignin-derived oxygenates [13]. Zeolite strong acid sites are not the unique factor to achieve high conversion and selectivity to aromatics. The two most essential parameters are diffusion and the active site.

The yield of the condensable volatiles for the CFP over parent HZSM-5 zeolite is much higher than that of non-CFP of lignin. The reason is tentatively speculated that the microporous zeolites help to stabilize reactive intermediates derived during lignin pyrolysis, which otherwise readily recombine to form char. On the other hand, H\textbeta{} is not as effective as HZSM-5 in improving the yield of the condensable volatiles. The yields of condensable volatiles on parent H\textbeta{} zeolite are even lower than those of non-CFP. It is tentatively speculated that the reason for this is that less coke is present on the surface of HZSM-5, which is mainly derived from large size PAH molecule. It has been reported previously that the amount of coke derived from CFP of lignin is related to the pore size, and the larger pore size of the H\textbeta{} zeolites is favorable for the formation of coke occurring from polymerization reactions [13]. For HZSM-5 zeolite, the formation of large condensed molecules from side reactions was inhibited due to the smaller pore size. For H\textbeta{} zeolite, the concentration of PAH in the condensable volatiles is twice as much as HZSM-5. There is a considerable amount of 3 and 4 ring aromatics produced over H\textbeta{} zeolite, indicating that these large PAH are not hindered by the steric hindrance.

Figure 8 shows the yield of the condensable volatiles for lignin CFP over HZSM-5 zeolites obtained by different post-treatment methods. Compared with parent HZSM-5, the yield of the condensable volatiles on hierarchical HZSM-5 zeolites increased slightly, which is basically in accordance with the increase of mesoporous surface area. As shown in Figure 9, the alkaline desilication treatment improved considerably the selectivity in aromatics hydrocarbon over hierarchical HZSM-5 zeolites. Compared with pure HZSM-5, the selectivity in naphthalene and PAH for hierarchical HZSM-5 zeolite is increased without affecting the yields in BTX. The most significant distinction for hierarchical HZSM-5 zeolite is the sharp reduction of oxygenates in the condensable volatiles, especially for HZSM-5-D, only 3.3\%. Oxygenates are significantly reduced, while the amount of aromatic hydrocarbons increased. The formation of mesopores after desilication enhances a faster diffusion of reactants into the active surface and of products out of the zeolite particles. The introduction of mesopores in hierarchical HZSM-5 zeolite makes the larger oxygenates molecules diffuse into the pores and become effectively deoxygenated under the catalysis functions of Brønsted acid sites. As the main active sites for biomass pyrolysis vapors conversion, the remained Bronsted acid sites mainly presented in the micropores are more accessible in the hierarchical HZSM-5 zeolite. The deoxidation of oxygenates derived from lignin pyrolysis was effectively improved, which could be tentatively ascribed to the suppression of mass transfer resistance over hierarchical HZSM-5 zeolite. We speculated that the obtained deoxidation product can quickly diffuse out of the pore, avoiding the formation of unnecessary coke due to molecules condensation.
Figure 8. The total yield of the condensable volatiles from catalytic fast pyrolysis of Kraft lignin over parent HZSM-5 and the hierarchical HZSM-5 obtained by different post-treatment methods. Experimental conditions: pyrolysis temperature of 500 °C, Kraft lignin 0.3 mg, catalyst-to-lignin ratio of 10:1.

Figure 9. The selectivity of various volatile products from catalytic fast pyrolysis of Kraft lignin over parent HZSM-5 and the hierarchical HZSM-5 obtained by different post-treatment methods. Experimental conditions: pyrolysis temperature of 500 °C, Kraft lignin 0.3 mg, catalyst-to-lignin ratio of 10:1.

Figure 10 shows the yield of the condensable volatiles for lignin catalytic fast pyrolysis over Hβ zeolites obtained by different post-treatment methods. The hierarchical Hβ zeolite is much more active than the parent one. The yield of the condensable volatiles gradually increases from Hβ-A to Hβ-D, which correlates well with the change of mesoporous surface area. In particular, the condensable product yield for Hβ-D is 2 times that of the parent Hβ. The increase of the yield can also be attributed to the improvement of the mass transfer performance, because the alkaline desilication treatments are more effective for the formation of mesopores on hierarchical Hβ zeolite. Similar to the case of
hierarchical zeolite HZSM-5, the introduction of mesopores mainly promotes the diffusion of molecules in and out of the active sites during the pyrolysis process. The yield of condensable volatiles over parent Hβ is lower than that of parent HZSM-5. Considering that the parent HZSM-5 contains hardly any mesopore, the mesopores present in the parent Hβ itself had tiny effect on inhibiting carbon deposition. It is speculated that the additional mesopores in hierarchical Hβ zeolite may be interconnected with micropores. The hierarchical pore structures can improve the utilization of micropores, due to the promoted mass transfer performance and the more accessible surface Brønsted acid sites. As shown in Figure 11, the selectivity in oxygenates for hierarchical Hβ zeolites is slightly increased without affecting the yields in aromatic hydrocarbons. The improved activity of the hierarchical zeolite is ascribed to the synergistic effect between the presence of Brønsted sites and accessible mesopores.

Figure 10. The total yield of the condensable volatiles from catalytic fast pyrolysis of Kraft lignin over parent Hβ and the hierarchical Hβ obtained by different post-treatment methods. Experimental conditions: pyrolysis temperature of 500 °C, Kraft lignin 0.3 mg, catalyst-to-lignin ratio of 10:1.

Figure 11. The selectivity of various volatile products from catalytic fast pyrolysis of Kraft lignin over parent Hβ and the hierarchical Hβ obtained by different post-treatment methods. Experimental conditions: pyrolysis temperature of 500 °C, Kraft lignin 0.3 mg, catalyst-to-lignin ratio of 10:1.
3. Materials and Methods

3.1. Materials and Chemicals

Commercial Kraft lignin (product number 471003, Sigma-Aldrich, Saint Louis, MO, USA), NaOH (≥96% AR assay, Aladdin, Shanghai, China), HCl (36–38% AR assay), Hβ zeolite (Si:Al = 20, Nankai University Catalyst Co. Ltd., Tianjin, China), HZSM-5 zeolite (Si:Al = 25, Nankai University Catalyst Co. Ltd., Tianjin, China) NH₄NO₃ (≥96% AR assay, Aladdin, Shanghai, China), Na₂CO₃ (99% AR assay, Aladdin, Shanghai, China), TMAOH (25% of water AR assay, Aladdin, Shanghai, China).

3.2. Catalyst Preparation

The hierarchical zeolites were prepared by alkaline post-processing method using Na₂CO₃, TMAOH/NaOH mixture, and NaOH as desilication sources, respectively. Typically, 4.0 g HZSM-5 or Hβ powders were added into following aqueous solutions.

Treatment A: 133 mL 0.1 M Na₂CO₃ alkaline aqueous solution;

Treatment B: 133 mL mixed alkaline solution (C₅MAOH/CHOH⁻ = 0.6), which contained TMAOH (5.8190 g, 25 wt %) and NaOH (0.4380 g);

Treatment C: 133 mL 0.1 M NaOH alkaline solution.

Then, the suspension was heated to 65 °C and stirred for 1.5 h. After being cooled down to room temperature, the solid was recovered by centrifugation and washed thoroughly with deionized water.

Treatment D: Selected NaOH-treated samples were treated in aqueous HCl solution (0.1 M, 100 cm³/g zeolite), and then the above stirring, cooling, centrifugation, and washing procedure were repeated.

The recovered solid sample was further exchanged three times with an aqueous solution of NH₄NO₃ (1.0 M, 100 mL), followed by drying at 80 °C for 12 h and calcination in air to obtain the H-format 550 °C for 4 h. The hierarchical HZSM-5 was denoted as HZSM-5-X, in which X was the corresponding alkaline post-processing method (treatment A or B or C or D) and so on.

3.3. Catalyst Characterization

N₂ adsorption-desorption isotherms were measured with Automatic physical adsorption instrument at 77 K (TriStar 3020, Micromeritics, Norcross, GA, USA). Prior to analysis, each sample was degassed under dynamic vacuum (VacPrep 061) at 300 °C for 4 h. Pore size distribution was determined from the adsorption branch of isotherm, using the Barrett-Joyner-Halenda (BJH) Analysis. The Brunauer-Emmett-Teller (BET) method was used to estimate the total surface area (0.01 < p/p₀ < 0.08). The t-plot method was adopted to distinguish mesopores from micropores.

X-ray powder diffraction (XRD) patterns of the prepared samples were measured on a Philips CM-1 (Cu Kα, λ = 0.1543 nm) powder X-ray diffractometer (Panalytical, Almelo, The Netherlands). Typically, the data was collected from 5° to 70° (2θ) for conventional wide-angle XRD patterns. The software X’Pert Highscore (Version: 1.0e, Panalytical, Almelo, The Netherlands, 2003) was used to perform microstructure analysis.

Temperature-programmed desorption of NH₃ (NH₃-TPD) was carried out in an automatic chemisorb analyzer (ChemiSorb 2720, Micromeritics, Norcross, GA, USA). One hundred milligrams of each sample was introduced into a U-shaped tubular quartz reactor. Each sample was first treated at 500 °C for 1 h and then cooled down to room temperature in He. NH₃ was absorbed at 120 °C for 30 min (5 mL/min) of 10% NH₃/He followed by a He purge. TPD experiments were performed under a flow of He (25 mL/min) from 120 to 700 °C at a constant heating rate (10 °C/min). The desorbed NH₃ was monitored with a thermal conductivity detector (TCD).

3.4. Py-GC/MS Test

Pyrolysis-gas chromatography-mass spectrometry (Py-GC/MS) study was performed at 500 °C using a micro-furnace system (PY-3030D, Frontier Laboratories, Fukushima, Japan) installed with a gas
chromatograph (GC) (7890A, Agilent Technologies, Santa Clara, CA, USA). A mass spectrometer (MS) (5975C, Agilent Technologies, Santa Clara, CA, USA) was used to analyze the products. The injector temperature was kept at 300 °C with a 1:100 split ratio. Analysis of condensable volatiles for lignin catalytic fast pyrolysis was performed using UA-5 Ultra ALLOY® Capillary Column (30 m x 0.25 mm and 0.250 µm film thickness, with a stationary phase consisting of 5% diphenyldimethyl polysiloxane) with Helium as carrier gas of 1.0 mL/min flow rate. The GC oven temperature program began with 40 °C (holding 3 min) to 280 °C with the heating rate of 5 °C /min. The final temperature was held for 4 min. The temperature of the GC/MS interface was held at 300 °C, and the mass spectrometer was operated in EI mode at 70 eV. The mass range from m/z 29 to 350 was scanned at 4.37 scans/s. Pyrolysis products were identified according to the NIST MS library and the literature data of bio-oils. In each experiment, about 0.3 mg Kraft lignin (and 3 mg HZSM-5 or Hβ zeolites for catalytic pyrolysis) were placed in a sample cup, and for each sample, experiments were conducted at least three times to confirm the reproducibility of the reported procedures.

4. Conclusions

Lignin is the most recalcitrant component of lignocellulose, which has a high resistance to pyrolysis. A large number of cokes derived from CFP of lignin usually lead to rapid deactivation of zeolites, which restricted the practical application of lignocellulose pyrolysis due to the loss of carbon conversion efficiency. The diffusion and mass transfer resistance of heavier feeds often limit the space utilization in microporous zeolites. The hierarchical HZSM-5 and Hβ zeolites, which have interconnected mesopores and micropores, were prepared successfully by alkaline desilication methods to optimize the accessibility to these big molecules. We noticed that there was an approximate mesoporous surface area for the hierarchical HZSM-5 zeolites under the treatment condition of different alkaline desilication sources. However, the surface area of mesopores formed in the hierarchical Hβ zeolites was directly related to the basicity of alkaline solution. The hierarchical zeolites all maintained the original crystallographic structure of the parent one. The CFP of lignin over the hierarchical HZSM-5 and Hβ zeolites was analyzed on-line by Py-GC/MS technique. The deoxidation of oxygenates derived from lignin pyrolysis was effectively improved without the formation of extra coke, which could be ascribed to the suppression of mass transfer resistance over hierarchical HZSM-5 zeolite. The yield of the condensable volatiles over hierarchical Hβ zeolite gradually increases, which correlates well with the change of mesoporous surface area. The improved activity of the hierarchical zeolites is ascribed to the synergistic effect between the presence of Brønsted sites and accessible mesopores.

Supplementary Materials: The following are available online at http://www.mdpi.com/2073-4344/8/2/82/s1, Table S1: The main products of non-catalytic fast pyrolysis of Kraft lignin determined by Py-GC/MS at 500 °C; Table S2: The main products of catalytic fast pyrolysis of Kraft lignin over HZSM-5 zeolite determined by Py-GC/MS at 500 °C.

Author Contributions: H.C. and J.H. conceived and designed the experiments; X.L. and G.X. performed the experiments; X.L. and Y.B. analyzed the data; Y.B. and H.C. wrote the paper.

Conflicts of Interest: The authors declare no conflict of interest.

References


