Article

Photocatalytic Cleavage of β-O-4 Ether Bonds in Lignin over Ni/TiO₂

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Abstract: It is of great importance to explore the selective hydrogenolysis of β-O-4 linkages, which account for 45–60% of all linkages in native lignin, to produce valued-added chemicals and fuels from biomass employing UV light as catalyst. TiO₂ exhibited satisfactory catalytic performances in various photochemical reactions, due to its versatile advantages involving high catalytic activity, low cost and non-toxicity. In this work, 20 wt.% Ni/TiO₂ and oxidant PCC (Pyridinium chlorochromate) were employed to promote the cleavage of β-O-4 alcohol to obtain high value chemicals under UV irradiation at room temperature. The Ni/TiO₂ photocatalyst can be magnetically recovered and efficiently reused in the following four consecutive recycling tests in the cleavage of β-O-4 ether bond in lignin. Mechanism studies suggested that the oxidation of β-O-4 alcohol to β-O-4 ketone by oxidant PCC first occurred during the reaction, and was followed by the photocatalysis of the obtained β-O-4 ketone to corresponding acetophenone and phenol derivates. Furthermore, the system was tested on a variety of lignin model substrates containing β-O-4 linkage for the generation of fragmentation products in good to excellent results.

Keywords: lignin; β-O-4; photocatalyst; Ni/TiO₂; oxidant PCC

1. Introduction

While fossil fuels were widely regarded as the primary source for chemicals and energy, the fraction of chemicals and fuels obtained from renewable resources, such as biomass, can be expected to be good alternatives in the future [1–5]. Lignin contains complex natural aromatic subunits (sinapyl, coniferyl, coumaryl), and diverse types of linkages (β-O-4, α-O-4 and 4-O-5) [6,7]. Among these types of ether bonds, β-O-4 is the most abundant linkage in lignin, resulting in a variety of studies focused on the cleavage of the β-O-4 bond employing lignin dimeric model compounds (Figure 1a) [8,9]. In the past decades, tremendous efforts have been devoted into the degradation of lignin by Hartwig [10], Baker [11], Dyson [12], Ellman [13], Barta [14], etc.
was carried out in the practical and operationally simple two-step degradation of lignin from the same process \[39\]. When irradiated with ultraviolet light (wavelength \(< 400\)nm and energy band \(= 3.2\) eV), an electron in the valence band is excited and advances to the conduction band for the generation of a

In general, lignin depolymerization processes always involve the utilization of either transition-metal catalysts \[15\] or noble-metal catalysts \[16\]. However, those approaches still remain significant challenges that have never been dealt with. Particularly, a typical lignin degradation process requires highly demanding reaction conditions (elevated temperature, in the presence of H\(_2\), etc.) \[17,18\]. Therefore, it has become increasingly important to develop an efficient catalytic system to address these issues \[19\]. Recently, the photocatalytic degradation of lignin—which has various advantages such as milder reaction conditions, simple reaction process without further purification, filtration, or solvent changes \[20–22\]—has received increasing attention, as it is regarded as a potential alternative to traditional process. For example, Stephenson et al. explored an efficient and innovative two-stage lignin degradation method, in which, \[4-Acetamido-TEMPO\]BF\(_4\) mediated the benzylic oxidation in the first step and followed by a photoredox-catalyzed reductive C-O cleavage utilizing \([\text{Ir}(ppy)_2(dtbbpy)]PF_6\) as the photocatalyst \[23\]. Based on the previous study, a bimetal catalytic strategy was carried out in the practical and operationally simple two-step degradation of lignin from the same group involving Pd-catalyzed benzylic oxidation and photoredox-catalyzed \((\text{Ir}(ppy)_2(dtbbpy)]PF_6\) reductive fragmentation for the efficient cleavage of the \(\beta-O-4\) ketone obtained from the first step to generate lower-molecular-weight aromatic building blocks \[24\]. Besides \([\text{Ir}(ppy)_2(dtbbpy)]PF_6\), a variety of photocatalysts showed an excellent ability in cleaving C-O bonds in lignin, including \(\text{fac-Ir}(ppy)_3\) \[25\], \([\text{Ir}(ppy)_2(dtbbpy)]PF_6\) \[26\], \([\text{Ir}d^F(CF_3)_2ppy_2(dtbbpy)]PF_6\) \[27\]. However, these catalysts needed to be stored under anhydrous and anaerobic conditions, and could not be recovered for the next run. Semiconductor, such as TiO\(_2\), attracted the attention of many researchers due to its versatile advantages involving stable catalytic performance and low cost \[28–30\]. It has been widely reported that the introduction of transition metal \[31–33\] or noble metal nanoparticles \[34–36\] to TiO\(_2\) support is beneficial for electron transfer in the photochemical reaction process. For instance, Farnood et al. demonstrated a novel Bi and Pt co-modified TiO\(_2\) catalyst for the photo-oxidation of lignin under solar light for the generation of guaiacol, vanillin acid, vanillin and 4-phenyl-1-buten-4-ol \[37\]. Srisasivimon et al. employed lignin-based carbon to modify TiO\(_2\) for the generation of a composite photocatalyst (TiO\(_2\)/lignin), then supported Pt was prepared and exhibited excellent catalytic activity for the production of high value chemicals from lignin \[38\]. However, noble metal-based catalysts will inevitably add to the cost and always suffer from detaching issues; the development of non-noble metal based photocatalysts and the recovery of photocatalysts still meet huge challenges. Therefore, it is of great significance to develop a simple and efficient catalyst to address this issue.

The degradation of organics over TiO\(_2\)-supported catalysts is essentially a free radical reaction process \[39\]. When irradiated with ultraviolet light (wavelength \(< 400\)nm and energy band \(= 3.2\) eV), an electron in the valence band is excited and advances to the conduction band for the generation of a

**Figure 1.** (a). Representative structure of lignin and beta-O-4 linkage in lignin, (b). Schematic diagram of photoelectric transfer effect on TiO\(_2\).

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photo-generated electrons (e\textsuperscript{−}), leaving photo-generated holes (h\textsuperscript{+}) in the valence band (Figure 1b). The obtained photo-generated holes (h\textsuperscript{+}) have an excellent ability to oxidize organic chemicals attached to the surface of TiO\textsubscript{2} or oxidize OH\textsuperscript{−} to yield OH radical in the first place, and subsequently oxidize the organics for small molecule compounds [40–42]. In this work, TiO\textsubscript{2}-supported nickel catalyst was introduced into the selective oxidation of β-O-4 ketone model compounds, resulting in the generation of value-added aromatics. Afterwards, a two-step process was successfully achieved for the photocatalytic cleavage of β-O-4 alcohols. In the first step, β-O-4 alcohol was oxidized into β-O-4 ketone by PCC oxidant; then, the cleavage of the β-O-4 bond happened over Ni/TiO\textsubscript{2}. Through the screening of light sources, solvents and oxidations, herein we reported a super mild reaction condition (30W UV, iPrOH, and PCC) for the cleavage of the β-O-4 bond in lignin model compounds. Finally, the possible reaction mechanism was also proposed. The basic physicochemical properties were investigated by means of XRD, TEM, and XPS analyses.

2. Result and Discussion

2.1. Catalyst Characterization

XRD spectra of pure TiO\textsubscript{2} and 20 wt.% Ni/TiO\textsubscript{2} are presented in Figure 2i. The XRD pattern of pure TiO\textsubscript{2} was clearly shown in the pattern (a). The characteristic peaks of anatase and rutile crystals of TiO\textsubscript{2} could be obviously discovered, in which peaks of 2θ = 53.9° and 62.9° marked by Miller indices (210) and (002) belonged to the rutile crystal, and the other peaks of 2θ = 25.4°, 37.9° and 48.2° marked by Miller indices (101), (004) and (200) belonged to the anatase TiO\textsubscript{2} [43]. Moreover, the peak at 27.5° was too weak to be observed which belonged to rutile crystal. In the pattern of 20 wt.% Ni/TiO\textsubscript{2}, all the characteristic peaks of anatase and rutile crystals of TiO\textsubscript{2} could be clearly found, and three characteristic peaks at 2θ of 44.6°, 52.0° and 76.6°, marked by Miller indices (111), (200) and (220), could also be observed at relatively high angles, indicating the presence of metallic nickel. No NiO species were observed in the XRD pattern, which proved that metallic nickel and TiO\textsubscript{2} could be stably stored without oxidation in air before the reaction process [44]. The dispersion and average particle size of the photocatalyst Ni/TiO\textsubscript{2} were characterized by TEM, which was also presented in Figure 2 (Figure 2ii and iii). It could be clearly seen that small particles, representing clusters of metallic Ni, were homogenously dispersed on the surface of TiO\textsubscript{2} particles, and the average size of Ni particles was 13.45 nm or so.

The surface element compositions of 20 wt.% Ni/TiO\textsubscript{2} photocatalyst were analyzed by XPS. The survey scan and XPS patterns of Ti2p and Ni2p were presented in Figure 3. The general spectra (Figure 3iii) exhibited the presence of respective metals. In Figure 3i, the binding energies at 852.5 eV and 869.0 eV might ascribe to Ni\textsuperscript{0}(2p\textsubscript{3/2}) and Ni\textsuperscript{0}(2p\textsubscript{1/2}), respectively, and the binding energies at 855.0 eV and 874.5 eV belonged to the main line of Ni\textsuperscript{2+}(2p\textsubscript{3/2}), and Ni\textsuperscript{2+}(2p\textsubscript{1/2}), which indicated the presence of both metallic Ni and NiO on the surface of TiO\textsubscript{2}. It could be seen from Figure 3ii, that the peaks at 458.5 and 464.2 eV were ascribed to Ti\textsuperscript{4+}(2p\textsubscript{3/2}) and Ti\textsuperscript{4+}(2p\textsubscript{1/2}) in TiO\textsubscript{2}, respectively [43].
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Figure 2. (i) XRD patterns of TiO₂ (a) and 20 wt.% Ni/TiO₂ (b), (ii) TEM image of 20 wt.% Ni/TiO₂, (iii) Particle size of Ni on the TiO₂ particles.

Figure 3. XPS spectra of the Ni 2p for 20 wt.% Ni/TiO₂ (i), and Ti 2p for the 20 wt.% Ni/TiO₂ (ii) and 20 wt.% Ni/TiO₂ (iii).

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2.2. Optimization of the Reaction Condition

Above all, the amount of Ni/TiO₂ could affect the efficiency of the cleavage of β-O-4 ether. 2-phenoxy-1-phenylethan-1-one (1a) was selected as a model compound, and the results were described in Table 1. As expected, the conversion increased with the increasing catalyst amount (from 0 to 30 wt.% Ni/TiO₂), and the changes of acetophenone and phenol yields indicated the same trend. Meanwhile, 20 wt.% Ni/TiO₂ could achieve a total conversion of 2-phenoxy-1-phenylethan-1-one in the photocatalytic system (Table 1, entry 1). Therefore, 20 wt.% Ni/TiO₂ was employed in the following experiments.

Table 1. Optimization of the amount of Ni/TiO₂ catalysts 1a.

| Entry | Catalysts | Solvent | T. (°C)/t. (h) | Con. (%) | Yield (%) |
|-------|-----------|---------|----------------|----------|-----------|
| 1     | TiO₂      | DMF     | r.t./12        | 56       | 40        |
| 2     | 10 wt% Ni/TiO₂ | DMF | r.t./12  | 76       | 66        |
| 3     | 20 wt% Ni/TiO₂ | DMF | 180/12 | 100      | 82        |
| 4     | 30 wt% Ni/TiO₂ | DMF | r.t./12 | 100      | 83        |

a Reaction conditions: 1a (100 mg), Catalyst (20 mg), DMF (2.5 mL); b T. = reaction temperature, t. = reaction time, Con. = conversion, r.t. = room temperature, the conversion and yields were determined by GC/MS with n-dodecane as the internal standard.

In addition, the influence of light source on the photocatalysis of β-O-4 ketone was conducted. To evaluate whether the light source played a significant role in the cleavage of C-O bond in the lignin, comparative tests were carried out in different light sources (darkness, sunlight, 30 W UV, and 30 W blue LED). As expected, the transformation of 1a failed in the darkness and no acetophenone and phenol were observed in GC/MS (Table 2, entry 1). Meanwhile, sunlight catalytic process achieved the same results and 100% recovery of 1a was observed (Table 1, entry 2). The above phenomena in Table 2 (entry 1 and entry 2) were reasonable, as previous studies reported that the lignin degradation reaction generally needed to be performed under elevated temperature, high pressure and even catalyzed by homogeneous and/or heterogeneous catalysts. However, the transformation of 1a at 180 °C under sunlight achieved a little improvement with a yield of 11% and 8% toward acetophenone and phenol, respectively (Table 2, entry 3). In comparison, it was amazing to find that the cleavage of β-O-4 C-O bond was remarkably expedited under UV irradiation, while the yield of acetophenone and phenol increased sharply to about 82% and 80%, respectively, indicating an excellent efficiency for the cleavage of β-O-4 ketone bond in lignin (Table 2, entry 4). Apart from darkness, sunlight and UV, blue LED was also selected as the light source in the 20 wt.% Ni/TiO₂ catalytic system. Similar to sunlight, blue LED also seemed to not be suitable for the photocatalyzed cleavage of C-O bond in lignin (Table 2, entry 5). Due to the relatively long wavelength of blue LED (>400 nm), the electron in the valence band could not be excited and subsequently advanced to the conduction band for the generation of a photo-generated electrons (e⁻) to leave a photo-generated holes (h⁺) [45], thus the reaction gave a very poor β-O-4 ketone conversion rate (Table 2, entry 5). The comparison results between different light sources suggested that the wavelength was a key influencing factor in the photolysis of β-O-4 model catalyzed by TiO₂ supported catalysts. Therefore, we found 20 wt.% Ni/TiO₂ performed the better catalytic activity for the cleavage of β-O-4 ketone under UV irradiation (30 W).
With the 20 wt.% Ni/TiO₂ photocatalytic system under 30 W UV irradiation in hand, the influence of solvents on the cleavage of β-O-4 models was explored subsequently. The results in Table 3 indicate that 20 wt.% Ni/TiO₂ exhibited an outstanding catalytic activity for the C-O bond cleavage, but the conversion and yields varied in different polar and apolar solvents. In the apolar solvent system, the photocatalytic process displayed a low conversion rate and low product yields (Table 3, entry 1 and 2), probably due to the lower solubility in solvent. Hydrogenation-donor solvents such as iPrOH and methanol were believed to favor the cleavage of β-O-4 models in lignin, while a conversion rate of nearly 100% was achieved in 1a, accompanied by remarkably high yields of acetophenone and phenol (Table 3, entry 3 and 4). When carried out in DMF, the reaction process also showed an excellent conversion rate (100%) and yields (82% of acetophenone and 80% of phenol). When the reaction was conducted in acetonitrile or acetone, improved conversion rate was observed, which were higher than 60% (Table 3, entry 6 and 7). However, the yields of acetophenone and phenol were quite low (all below 38%), indicating a poor selectivity. H₂O was expected to be a green and environmentally-friendly solvent in a variety of chemical reactions. However, relatively lower conversion rate (15%) and yields were achieved (6% toward acetophenone and 5% toward phenol) owing to poor solubility, limiting the occurrence of the photocatalytic reaction in water (Table 3, entry 8). In general, the catalytic performance of the solvents followed the below order: iPrOH > methanol > DMF > acetone > acetonitrile > n-hexane > cyclohexane > H₂O. All of this taken together, the results above reflected that iPrOH was the optimal solvents for the C-O bond cleavage over 20 wt.% Ni/TiO₂.

2.3. Scope of the Substrates

In order to continue the exploration of the 20 wt.% Ni/TiO₂ photocatalytic activity, a variety of substrates including β-O-4 linkage were chosen as model compounds under the optimal conditions. It could be clearly seen in Scheme 1 that, the main products in the above reactions were acetophenone and phenol derivates, suggesting the occurrence of the selective cleavage of the C-O bond in those β-O-4 model compounds. Substrates without any substituted groups on the benzene ring 1a could achieve a 100% conversion rate, to give 88% of acetophenone and 82% of phenol in iPrOH. Methoxy-substitutions of O-terminus aryl led to no apparent decrease in yields compared to 1a. It seemed that steric effect had no obvious influence in our photocatalytic system. Hence, sing-ortho-substituted ketone 1b and bis-ortho-substituted ketone 1c afforded ketone products 2b (78%), 2c (80%) and phenol products 3b (77%), 3c (78%), respectively, after 12 h. In addition, coumaryl-based substrates (1d–1f) were subsequently explored in the Ni/TiO₂-catalyzed photocatalytic system. It was gratifying to discover that methoxy-substituted ketones 1d–1f were tolerated with no apparent decrease in yields, affording phenol 3d, guaiacol 3e and syringol 3f in 80%, 77% and 72%, respectively. Moreover, different leaving

### Table 2. Optimization of light sources for the photocatalytic reaction of 1a a.

| Entry | Light Source | Solvent | T. (°C)/t. (h) b | Con. (%) b | Yield (%) b |
|-------|--------------|---------|-----------------|------------|-------------|
| 1     | darkness     | DMF     | r.t./12         | 0          | 0           |
| 2     | sunlight     | DMF     | r.t./12         | 0          | 0           |
| 3     | sunlight     | DMF     | 180/12          | 28         | 11          |
| 4     | 30W, UV      | DMF     | r.t./12         | 100        | 82          |
| 5     | 30W, blue LED| DMF     | r.t./12         | 0          | 0           |

a Reaction conditions: 1a (100 mg), 20 wt.% TiO₂ (20 mg), DMF (2.5 mL); b T. = reaction temperature, t. = reaction time, Con. = conversion, r. t. = room temperature, the conversion and yields were determined by GC/MS with n-dodecane as the internal standard.
groups such as α-acetoxy group instead of phenol compared with 1d exhibited a significant role in our photocatalyzed system, delivering trace amounts of 1-(4-methoxyphenyl)ethan-1-one. The same results were achieved when 1h was employed in the process and nearly 100% recovery of substrate 1h was observed in the GC/MS. Meanwhile, 1i was carried out in our reaction system and 32% yield of ketone derivative and 52% yield of phenol derivative were obtained. Finally, the Ni/TiO2-photocatalytic system was almost not responsible for the fragmentation of the corresponding β-O-4 alcohol (1j) under the optimal reaction condition and almost no target ketone and phenol products were observed. Therefore, this was again confirmed that previous reports could not achieve one-pot degradation of β-O-4 alcohol models, instead, the process always involved two steps: noble metal catalyzed oxidation process and photoredox-catalytic reductive fragmentation of the β-O-4 linkage. The dominant reason for this phenomenon was that the homolytic bond dissociation enthalpy of β-O-4 ether bond has been theoretically predicted to decrease from 62.2 to 55.9 kcal/mol after one oxidation process, leading to the fragmentation of β-O-4 ketone much easier than β-O-4 alcohol [46].

**Table 3.** Optimization of different solvents for the cleavage of 1a.

| Entry | Catalyst | Solvent | T. (°C)/t. (h) | Con. (%) | Yield (%) |
|-------|----------|---------|----------------|----------|-----------|
| 1     | Ni/TiO2  | n-hexane| r.t./12        | 26       | 15        |
| 2     | Ni/TiO2  | cyclohexane| r.t./12        | 20       | 12        |
| 3     | Ni/TiO2  | iPrOH   | r.t./12        | 100      | 88        |
| 4     | Ni/TiO2  | methanol | r.t./12        | 100      | 83        |
| 5     | Ni/TiO2  | DMF     | r.t./12        | 100      | 82        |
| 6     | Ni/TiO2  | acetone | r.t./12        | 60       | 32        |
| 7     | Ni/TiO2  | acetone | r.t./12        | 69       | 38        |
| 8     | Ni/TiO2  | H2O     | r.t./12        | 15       | 6         |

*Reaction conditions: 1a (100 mg), 20 wt.% Ni/TiO2 (20 mg), Solvent (2.5 mL); b T. = reaction temperature, t. = reaction time, Con. = conversion, r.t. = room temperature, the conversion and yields were determined by GC/MS with n-dodecane as the internal standard.

**Scheme 1.** Cleavage of various β-O-4 ketones over 20 wt% Ni/TiO2-photocatalytic system. *a Reaction conditions: 1a-1j (100 mg), 20 wt.% Ni/TiO2 Catalyst (20 mg), iPrOH (2.5 mL). The yields were determined by GC/MS with n-dodecane as the internal standard.
In the past few years, a two-step method for the fragmentation of β-O-4 alcohol to valued-added aromatics had been widely explored. Stephenson and his team firstly developed a room-temperature lignin degradation strategy with a chemoselective benzylic oxidation [4-Acetamido-TEMPO]BF₄, followed by the reductive C-O bond cleavage over Ir-based catalyst [23]. In 2017, Stephenson et al. proposed an electrocatalytic oxidation method, coupled with a photocatalytic catalyst (Ir catalyst) catalyzed cleavage for β-O-4 bond in lignin [47]. Subsequently, Stephenson et al. developed a novel and operationally simple two-step lignin degradation method, involving Pd-catalyzed aerobic oxidation and visible-light photoredox-catalyzed reduction for the efficient cleavage of β-O-4 alcohol in 2019 [24]. Taking all of these into consideration, we assumed whether it could be achieved in one-pot (Scheme 2).

According to the special ability of Ni/TiO₂ for the transformation of β-O-4 ketone to corresponding acetophenone and phenol, we attempted the mechanical mixing of oxidant and Ni/TiO₂ for the C-O bond cleavage, and found that it was hardly efficient for the cleavage of β-O-4 alcohols. Herein, three oxidants ([4-Acetamido-TEMPO]BF₄, H₂O₂, PCC) were employed as oxidants in the reaction under optimal conditions. Unfortunately, it seemed that it was impossible to achieve the cleavage of C-O bond in β-O-4 alcohols by adding the oxidants (herein [4-Acetamido-TEMPO]BF₄, H₂O₂ and PCC) and photocatalyst (herein 20 wt.% Ni/TiO₂) in a one-pot process (Table 4, entry 1–3). This was probably due to the interference with the Ni/TiO₂ photocatalyst. As is well-known, PCC is a perfect oxidant to transfer primary and secondary alcohols to aldehydes and ketones in DCM, and we question whether we could take PCC, 1k and photocatalyst Ni/TiO₂ in DCM in one-pot. Disappointingly, one case using PCC as an oxidant and DCM as a solvent gave a perfect conversion of 1k (100%), but failed to yield acetophenone and phenol (Table 4, entry 4). When the experiment was carried out in two steps, β-O-4 alcohol was firstly oxidized by PCC, and the obtained β-O-4 ketone was transferred to yield acetophenone (66%) and phenol (61%) (Table 4, entry 5). We also suspect that solvents played a crucial role in this reaction. Hence, a mixture solvent (iPrOH: DCM=1:1) was tried under the optimal reaction condition and the same results were achieved—no acetophenone and phenol were detected. We suspected that the most likely reason was that the oxidant and reductant are mixed together, and that they could react with each other with no useful outcome. (Table 4, entry 6). Taken together, we continued to explore a two-step approach to achieve the hydrogenolysis of the β-O-4 alcohols of lignin-derived compounds (PCC oxidation followed by Ni/TiO₂ photocatalysis).

![Scheme 2. Two-step method for the fragmentation of β-O-4 alcohol to valued-added aromatics.](image-url)
Inspired by the excellent performance of two-step strategy, that is PCC oxidation in the first step followed by 20 wt.% Ni/TiO₂ photocatalysis under the optimal reaction condition for the efficient cleavage of the β-O-4 bond. Then, the hydrogenolysis of C-O bond was further studied in more detail by varying the substituent groups on the benzene ring (Scheme 3). Owing to the successful two-step strategy for the transformation of β-O-4 alcohol to corresponding aromatics. However, the yields of aromatics exhibited only a slight decrease in Scheme 3, due to the loss of substrates in the two-step process. When no substituted groups on the benzene ring, 1k could reach a high conversion to yield 66% of acetophenone and 61% of phenol. Methoxy substitutions of O-terminus aryl resulted in no apparent decrease in yields compared to 1k, which was similar to the above study of β-O-4 ketones. Therefore, sing-ortho-substituted alcohol 11 and bis-ortho-substituted alcohol 1m delivered ketone products 2l (62%), 2m (64%) and phenol products 3l (65%), 3m (58%), respectively, after 12 h at room temperature. Moreover, coumaryl-based system (1n–1p) was soon carried out in the two-step Ni/TiO₂-mediated photocatalytic strategy. It was satisfactory to find that methoxy-substituted alcohols 1n–1p were tolerated with no apparent decrease in yields, delivering phenol 3n, guaiacol 3o and syringol 3p in a yield of 66%, 58% and 49%, respectively. Based on the previous study of β-O-4 model compound (1j) in Scheme 2, we attempted the two-step strategy for the transformation of 1j. Unfortunately, oxidation PPC showed an excellent activity to turn α-hydroxyl group and γ-hydroxyl group to corresponding ketone and aldehyde, resulting in the failed generation of 2q. Above all, it could be verified that the two-step strategy was highly efficient for the transfer hydrogenolytic cleavage of the β-O-4 bond in a variety of lignin-derived model compounds.

It was of great importance to understand the pathway for the photocatalytic degradation of β-O-4 ether bond in lignin. Generally, the hydroxyl substituted on the α-C was firstly turned to ketone through the PCC oxidation process (1j), and subsequently, the generated β-O-4 ketone underwent a C-O bond cleavage to deliver the desired keto and phenol fragmentation products (2–4) after hydrogen atom abstraction (5) and protonation (6). This was in accordance with Enright’s study, which put forward the oxidation and photochemical reduction of a lignin model substrate from benzyl alcohol to guaiacol and 4-methoxyacetophenone via a benzylic ketone intermediate [21]. In addition, the reusability of Ni/TiO₂, based on the recycling tests under the optimal reaction conditions, was later investigated by employing 1k conversion as a model reaction. The photocatalyst Ni/TiO₂ could maintain a good catalytic activity after four successive runs and almost no change in the yields of corresponding aromatics was observed. Apart from this, 20 wt.% Ni/TiO₂ could be recovered magnetically in the end (Figure 4), which could facilitate the recycling process.

Table 4. Optimization of the oxidants for the cleavage of β-O-4 alcohols in one-pot a.

| Entry | Oxidant | Solvent | T. (°C)/t. (h) b | Con. (%) b | Yield (%) b |
|-------|---------|---------|-----------------|-----------|------------|
| 1     | [4-Acetamido-TEMPO]BF₄ | iPrOH | r.t./12 | 12 | 0 | 0 | 0 |
| 2     | H₂O₂    | iPrOH  | r.t./12 | 0  | 0 | 0 | 0 |
| 3     | PCC     | iPrOH  | r.t./12 | 100 | 66 | 61 |
| 4     | PCC     | iPrOH  | r.t./12 | 10 | 0 | 0 | 0 |
| 5 c   | PCC     | iPrOH  | r.t./12 | 66 | 61 |
| 6 d   | PCC     | iPrOH:DCM=1:1 | r.t./12 | 0 | 0 | 0 | 0 |

a Reaction conditions: 1k (100 mg), 20 wt.% TiO₂ (20 mg), Oxidant PCC (200 mg), iPrOH (2.5 mL); b T. = reaction temperature, t. = reaction time, Con. = conversion, r.t. = room temperature, the conversion and yields were determined by GC/MS with n-dodecane as the internal standard. c Two-step strategy (oxidation of 1k firstly and followed by Ni-TiO₂ photocatalysis). d 1k was used as substrate and iPrOH:DCM=1:1.
3. Experimental

Mechanical mixing of oxidant and photocatalyst cannot achieve an ideal effect. Therefore, it is important to seek an efficient catalyst that can both oxidize alcohol and break the C-O bond.

3.1. Materials

TiO$_2$ were purchased from Aladdin Industrial Inc. Shanghai, China and used without further treatment. N,N-dimethylformamide was obtained from Alfa Aesar Reagent Co., Ltd. (Shanghai, China). Ni(NO$_3$)$_2$·6H$_2$O was provided from Aladdin Industrial Inc. Shanghai, China. β-O-4 ketones (1a–1f) were synthesized according to the previous literature [48]. In a typical process of 2-phenoxy-1-phenylethan-1-one (1k) formation through a reductive method, in which NaBH$_4$ was employed.
(1a), a 500 mL round bottom flask equipped with a reflux condenser and a dropping funnel was charged with phenol (5.2 g, 55 mmol) and potassium carbonate (10.4 g, 76 mmol) in acetone (250 mL) and stirred at room temperature. To this solution, 2-bromoacetophenone (10.0 g, 50 mmol) in acetone (50 mL) was added dropwise over 10 min at room temperature. The resulting suspension was stirred at reflux for 5 h. After the reaction, the suspension was filtered and concentrated in vacuo. The crude product was purified by recrystallization from petroleum ether to obtain 2-phenoxy-1-phenylethan-1-one as a white solid (10.5 g, 49 mmol) in a 98% yield. Moreover, β-O-4 ketones (1k–1p) was synthesized through a reductive method, in which NaBH₄ was employed as the reductant. In addition, substrates 1g–1j was obtained from leyan.com.cn.

3.2. General Procedure for Ni/TiO₂ Catalyzed β-O-4 Model Compounds

In a typical catalytic reaction, 100 mg of 2-phenoxy-1-phenylethan-1-one or other derivates, 20 mg of Ni/TiO₂ catalyst, and 2.5 mL N,N-dimethylformamide were placed into a glass tube (10 mL). The photocatalytic reaction was performed under 30 W ultraviolet light (main wavelength at around 395 nm) in air at room temperature. At the end of the reaction, the mixture solution was filtered to collect the catalyst and the filtrate was analyzed by the Gas Chromatograph/Mass Spectrometer (GC/MS, Agilent 7890, Shanghai, China) using n-dodecane as an internal standard. The collected catalyst was washed with water and ethanol for three times and dried at 105 °C for the next run under the optimal reaction condition. The conversion and product yields in the liquid phase were calculated according to the following formula, respectively:

\[
\text{Conversion} = \frac{\text{mole of reacted substrate}}{\text{total mole of substrate feed}} \times 100\% \quad (1)
\]

\[
\text{Yield of acetophenone} = \frac{\text{mole of acetophenone}}{\text{total mole of substrate feed}} \times 100\% \quad (2)
\]

\[
\text{Yield of phenol} = \frac{\text{mole of phenol}}{\text{total mole of substrate feed}} \times 100\% \quad (3)
\]

3.3. Catalyst Preparation

Ni/TiO₂ was prepared by the traditional impregnation method. In a typical process of 20 wt.% Ni/TiO₂, Ni(NO₃)₂·6H₂O (0.2 g) and TiO₂ (2.0 g) were added in deionized water and stirred for 24 h. Then, the suspension was dried for 12 h at 105 °C in the oven. Subsequently, obtained grey solid was calcined at 500 °C for 2 h in the muffle furnace, and then reduced in a tube furnace under hydrogen atmosphere at 500 °C for 2 h.

3.4. Catalyst Characterization

Powder X-ray diffraction (XRD) was conducted on a Bruker D8 Advance X-ray powder diffractometer (Shanghai, China). Transmission electron microscopy (TEM) images were collected using a TEM Tecnai G2 20 (Thermo-VG Scientific, Shanghai, China). The X-Ray photoelectron spectroscopy (XPS) was examined on an ESCALAB-250 (Thermo-VG Scientific, Shanghai, China) spectrometer with Al Kα (1486.6 eV) irradiation source.

4. Conclusions

In summary, we have shown a mild and efficient two-step strategy that proceeded through the oxidation of β-O-4 alcohol to ketone first, then 20 wt.% Ni/TiO₂-photocatalysis for the transformation of β-O-4 ketone to obtain value-added chemicals. The Ni/TiO₂ photocatalyst not only had an excellent ability to catalyze the transfer hydrogenolytic cleavage of β-O-4 ether bonds in diverse substrates under UV irradiation (30 W), but could also be easily recovered magnetically from the reaction process for the next four recycling tests. Our photocatalytic system was also suitable for the cleavage of various β-O-4
alcohols in lignin, and the basic physicochemical characterization illustrated that the high activity of photocatalyst originated from the metallic Ni on the surface of TiO$_2$. It is worth noting that even though the two-step strategy for the cleavage of β-O-4 bond in lignin performed well, it still remains a great challenge to realize the hydrogenolysis in one-pot. This work may inspire more researches on the depolymerization of lignin using photocatalysts in one-pot.

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