Uranyl-Photocatalyzed Hydrolysis of Diaryl Ethers at Ambient Environment for the Directional Degradation of 4-O-5 Lignin

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ABSTRACT: Uranyl-photocatalyzed hydrolysis of diaryl ethers has been established to achieve two types of phenols at room temperature under normal pressure. The single electron transfer process was disclosed by a radical quenching experiment and Stern−Volmer analysis between diphenyl ether and uranyl cation catalyst, followed by oxygen atom transfer process between radical cation of diphenyl ether and uranyl peroxide species. The 18O-labeling experiment precisely demonstrates that the oxygen source is water. Further application in template substrates of 4-O-5 linkages from lignin and 30-fold efficiency of flow operation display the potential application for phenol recovery via an ecofriendly and low-energy consumption protocol.

KEYWORDS: lignin, phenol recovery, photocatalysis, oxygen atom transfer, flow operation

Lignin, one of the largest renewable resource of arenes in nature, is a kind of macromolecule polymerized from aromatic monomers through C−C bonds (e.g., β-5, β-β, β-1, and 5-5 linkages) and C−O bonds (e.g., α-O-4, β-O-4, and 4-O-5 linkages) (Scheme 1a).1 The recovery of aromatic chemicals from lignin would help solve the problems caused by accelerating consumption of fossil resources and corresponding environmental crisis, in which activation of C−O linkages is one of the pivot points.2 The 4-O-5 linkage is the most challenging part owing to its higher bond dissociation energy (BDE = 77.74 kcal/mol) in comparison to α-O-4 (BDE < 57.28 kcal/mol) and β-O-4 (BDE < 69.35 kcal/mol) (Scheme 1b).3,4

Typical studies mainly focused on hydrogenolysis and hydrolysis for C−O bond cleavage in diphenyl ether (DPE: template substrate for 4-O-5 linkage) (Scheme 1c). A series of elegant strategies by Hartwig, Grubbs, Mauriello, and others have been developed for selective hydrogenolysis with or without the assistance of transition-metal catalysts. 5−13 Hydrolysis of DPE was reported by Katritzky et al. 14 and Lercher and co-workers15−17 for cleaving the 4-O-5 linkage with great significance in lignin-first biorefinery,18 which is thermodynamically challenging with the reaction energy of −7.1 kcal/mol compared to −14.2 kcal/mol in hydrogenolysis.19 Recently, the degradation of DPE was realized by Li et al. through esterification under light with an acridinium photocatalyst with two molecules of phenols obtained via the following hydrolysis.20 Meanwhile, the cleavage of C−C21 and β-O-422,23 linkages of lignin were achieved with the development of photocatalysis.
Based on our previous study on uranyl species, the ligand-to-metal charge transfer (LMCT) mode and superior oxidative ability \([E_{\text{ox}} = 2.60 \, \text{V vs SCE}}\)\(^{29-32}\) supply great potential for DPE activation \([E_{\text{ox}} = 1.88 \, \text{V vs SCE}}\).\(^{33}\) Herein, the photocatalyzed hydrolysis of diaryl ethers was established to afford two kinds of phenols via uranyl catalysis with visible light stimulation at room temperature and normal pressure (Scheme 1d).

We commenced our study with DPE and uranyl nitrate hexahydrate as the photocatalyst stimulated with 460 nm light at room temperature, in which water was added as the oxygen source and diverse acids as coactivators (Table 1, entries 1–3).

Table 1. Condition Optimization\(^{\text{a}}\)

| entry | H\(_2\)O (equiv) | acid (equiv) | solvent | 2a (%) |
|-------|-----------------|-------------|---------|--------|
| 1     | 20              | CF\(_3\)COOH | MeCN    | 26     |
| 2     | 20              | CH\(_3\)COOH | MeCN    | 33     |
| 3     | 20              | CCl\(_3\)COOH| MeCN    | 55     |
| 4     | 20              | CCl\(_3\)COOH| DCM     | NR     |
| 5     | 20              | CCl\(_3\)COOH| THF     | NR     |
| 6     | 20              | CCl\(_3\)COOH| acetonite | 79  |
| 7     | 40              | CCl\(_3\)COOH| acetonite | 68  |
| 8     | 10              | CCl\(_3\)COOH| acetonite | 84  |
| 9     |                 | CCl\(_3\)COOH| acetonite | 96(80) |
| 10\(\text{a} \) |                 | CCl\(_3\)COOH| acetonite | 83   |
| 11\(\text{d} \) |                 | CCl\(_3\)COOH| acetonite | 76   |
| 12    |                 | CCl\(_3\)COOH| acetonite | 48   |
| 13\(\text{f} \) |                 | CCl\(_3\)COOH| acetonite | NR   |
| 14\(\text{f} \) |                 | CCl\(_3\)COOH| acetonite | NR   |
| 15\(\text{g} \) |                 | CCl\(_3\)COOH| acetonite | NR   |

\(\text{a} \)General conditions: DPE (0.4 mmol), UO\(_2\)(NO\(_3\))\(_2\), 6H\(_2\)O (4 mol %), acid (2 equiv), solvent (2 mL), N\(_2\), RT, 48 h, blue light (460 nm), \(^1\)H NMR yields with CH\(_3\)Br\(_2\) as the internal standard. \(^2\)Isolated yields.

Encouragingly, the desired product phenol was afforded in 55% yield with trichloroacetic acid. Considering the solvent effect, acetone turned out to be the optimal one, while tetrahydrofuran and dichloromethane were not due to the tendency of free radical generation (Table 1, entries 4–6).\(^{34,35}\) Increasing the amount of water disturbed the transformation, while decreasing the water helped the reaction (Table 1, entries 7 and 8). It is worth noting that the transformation was greatly promoted even without additional water being supplied (Table 1, entry 9), in which water was supplied by the dimerization and dehydration of acetone. Two equivalents of trichloroacetic acid afforded the optimal results (Table 1, entries 10–12). Control experiments further demonstrated that UO\(_2\)(NO\(_3\))\(_2\), 6H\(_2\)O and blue light were essential (Table 1, entries 13 and 14). Compared to the uranyl cation, Ir(ppy)\(_2\)(dtbbpy)BF\(_4\) \([E_{1/2} = +0.66 \, \text{V vs SCE}}\)\(^{26}\) Ru(bpy)\(_3\)Cl\(_2\), 6H\(_2\)O \([E_{1/2} = +0.77 \, \text{V vs SCE}}\)\(^{26}\) or Eosin Y \([E_{1/2} = +0.78 \, \text{V vs SCE}}\)\(^{36}\) was inefficient for the transformation (Table 1, entry 15).

With the optimized conditions in hand, symmetrical diaryl ethers were first investigated as shown in Scheme 2. A broad range of diaryl ethers with electron-neutral, -deficient, and -rich groups at the para-position were well tolerated in this reaction, delivering the desired products with good to excellent yields. The transformation went smoothly for phenol 2b, though it was latent to be oxidized at the benzylic position. C–O bonds were successfully disconnected to obtain the corresponding phenols 2d and 2e with halogen substituents, which were never found in previous reports due to the C–X bond dissociation tendency.\(^{5,11}\) Remarkably, easily hydrolyzed carboxylic ester (2g) and cyano (2h) were preserved during the transformation. Diaryl ethers with substitutions at the meta-position afforded the corresponding phenols 2i–2l as well.

Subsequently, the cleavage of C–O bond with unsymmetrical diaryl ethers was tested, starting with diaryl ethers possessing unilateral groups. Phenol derivatives bearing electron-withdrawing substitutions (2f–2h and 2m) at the para-position along with phenol were readily obtained with good efficiency as expected. Phenol 2n with the methoxy group substituted at the meta-position was also compatible. Remarkably, substrates with ortho-position substituted groups could be degraded to corresponding products (2o–2r). Furthermore, diaryl ethers bearing different substitutions varied on both sides of the aromatics were successfully transformed. Good yield was obtained when both parts of the diaryl ether possessed electron-donating groups (Scheme 2, 22). The substituted diaryl ethers with electron-deficient and -rich groups on both sides could be converted to corresponding phenols in middle to good yields (Scheme 2, 23–27). Good to excellent yields could be obtained when both parts of the diaryl ether possessed electron-withdrawing groups (Scheme 2, 28, 29). It is noteworthy that nitroen (herbicide) and triclosan (antibiotic) with multiple substituents were transformed smoothly (Scheme 2, 30, 31), showing the potential for the degradation of pesticide waste. For our lignin degradation target, models of 4-O-5 linkage in lignin were evaluated. For trisubstituted substrates, 2z1 and 2zn were obtained in 48% and 54% yield, respectively. The cleavage of the C–O bond of the pentasubstituted one, whose presence was confirmed in lignin by 2D NMR spectroscopy,\(^{37,38}\) was achieved at room temperature and normal pressure for the first time. 2z1, widely used for food additives and spices, was obtained in 14% yield. And 2z2, the midbody for Tinib anticancer drugs, was obtained in 22% yield.

To further demonstrate the application potential of the strategy, a flow device was designed for the transformation of lignin templates (Scheme 3). Compared to tube operation, the desired products were obtained in up to 30 times efficiency with a flow device, in which the reaction mixture was pumped into light permeable PTFE tube wrapping around a high reflective aluminum bar, surrounded by 60 tandem LED lamps (430 nm), despite the residue volume of the flow pipeline being about 4.7 mL.

Mechanistic studies were carried out to understand the process. First, radical quenching experiments with 2,2,6,6-tetramethyl-1-piperidinedinyloxy (TEMPO) and butylated hydroxytoluene (BHT) suggested the radical property of the system (Scheme 4a). The uranyl cation was approved to serve as the photosensor via UV–vis absorption at 424 nm (Scheme 4b), and it was also confirmed to interact with DPE directly via Stern–Volmer analysis (Scheme 4c). Based on the above results, this transformation was initiated with excited uranyl cations through a single electron transfer (SET) process between DPE under blue light irradiation.
$^{18}$O labeling experiments unambiguously demonstrated the oxygen source from water (Scheme 4d). According to the previous study, uranyl peroxide complexes were obtained from uranyl photolysis of water, which is responsible for the oxygen atom transfer (OAT) process. Then, the bias of C–O bond breaking was investigated (Scheme 4e). Whether diaryl ethers possessed a tert-butyl or cyano group at the para-position with the other side unsubstituted or a tert-butyl group on one side and a cyano group on the other, both phenols were labeled with $^{18}$O, indicating that the cleavage of the C–O bond took place on both bond 1 and 2 as shown in Scheme 4e. Moreover, the ratio of the different phenols depicted the cleavage trend on the bias of the low electron cloud density side. A Hammett plot was carried out to illustrate the relationship between the reaction rate and substituent effect (Scheme 4f). The $\rho$-value of +0.5096 for reactions showed that the electron-withdrawing groups promoted the transformation and the decisive step was the process of negative charge accumulation.

Therefore, a possible reaction pathway is depicted in Scheme 5. The uranyl photoredox catalysis was stimulated by blue light through the LMCT process, generating $^*$UO$_2$$^{2+}$. The SET process between $^*$UO$_2$$^{2+}$ and diaryl ether produced UO$_2$$^+$ along with radical cation A, which was attacked by the uranyl peroxide dimer afforded via water-splitting, yielding the radical cation of phenol B and phenyl oxygen anion C, accompanied by cleavage of the original C–O bond and formation of the new one. The protonation of C, together with the SET process between B and UO$_2$$^+$ [$E^{\circ}(B^*/B) = +1.56$ V].

**Scheme 2. Hydrolysis of Diaryl Ethers to Phenols**

| Diaryl Ethers | Acetone (2.0 mL), N$_2$, RT, blue light | Isolated Yields |
|--------------|----------------------------------------|----------------|
| 1            | 2a 80%                                 | 2f 72%         |
| 2            | 2b 64%                                 | 2g 69%         |
| 3            | 2c 69%                                 | 2h 59%         |
| 4            | 2d 49%                                 | 2i 62%         |
| 5            | 2a 70%                                 | 2j 62%         |
| 6            | 2g 61%                                 | 2k 52%         |
| 7            | 2f 60%                                 | 2l 61%         |
| 8            | 2d 60%                                 | 2m 65%         |
| 9            | 2n 69%                                 | 2o 65%         |
| 10           | 2p 59%                                 | 2q 61%         |
| 11           | 2a 70%                                 | 2r 72%         |
| 12           | 2g 61%                                 | 2s 52%         |
| 13           | 2f 60%                                 | 2t 61%         |
| 14           | 2d 60%                                 | 2u 65%         |
| 15           | 2g 61%                                 | 2v 61%         |
| 16           | 2a 70%                                 | 2w 62%         |
| 17           | 2g 61%                                 | 2x 52%         |
| 18           | 2f 60%                                 | 2y 61%         |
| 19           | 2d 60%                                 | 2z 74%         |
| 20           | 2a 70%                                 | 2a 62%         |

$^{18}$O labeling experiments unambiguously demonstrated the oxygen source from water (Scheme 4d). According to the previous study, uranyl peroxide complexes were obtained from uranyl photolysis of water, which is responsible for the oxygen atom transfer (OAT) process. Then, the bias of C–O bond breaking was investigated (Scheme 4e). Whether diaryl ethers possessed a tert-butyl or cyano group at the para-position with the other side unsubstituted or a tert-butyl group on one side and a cyano group on the other, both phenols were labeled with $^{18}$O, indicating that the cleavage of the C–O bond took place on both bond 1 and 2 as shown in Scheme 4e. Moreover, the ratio of the different phenols depicted the cleavage trend on the bias of the low electron cloud density side. A Hammett plot was carried out to illustrate the relationship between the reaction rate and substituent effect (Scheme 4f). The $\rho$-value of +0.5096 for reactions showed that the electron-withdrawing groups promoted the transformation and the decisive step was the process of negative charge accumulation.

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Scheme 3. Flow Reaction

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Reactor (R1) \[ R_1 \rightarrow 2 \]

Rate in flow (mmol/h) 0.170 0.150 0.080 0.070 0.050 0.040
Rate in tube (mmol/h) 0.013 0.005 0.003 0.003 0.003 0.003
Efficiency promotion (times) 12 35 30 27 22 17 13
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The desired product 2 was afforded and the catalyst was regenerated. In summary, C–O bond activation of diaryl ethers was achieved at room temperature and

vs SCE > E°(UO₂⁺/UO₂²⁺) = + 0.32 V vs SCE affords desired product 2 and regenerated the catalyst. In summary, C–O bond activation of diaryl ethers affording phenols was achieved at room temperature and

(vs SCE > E°(UO₂⁺/UO₂²⁺) = + 0.32 V vs SCE)
Scheme 5. Proposed Mechanism

normal pressure under blue light stimulation. Both symmetrical and unsymmetrical substrates were achieved with high compatibility. The C$_2$=O bond was activated via the SET process between the diaryl ethers and excited uranyl cation. The dimerization of uranyl transferred the oxygen atom from the water to generate easily oxidized phenols with the LMCT process of uranyl excitation. The synergistic mechanism of SET and OAT contributes a straightforward pathway for mild photocatalytic transformation. Further application of the 4-O-5 linkage model with a flow setup indicates the potential for upgrading the degradation of lignin. Further studies on uranyl catalyzed reactions are ongoing in our laboratory.

**ASSOCIATED CONTENT**

1. Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/10.1021/jacsau.1c00168.

Experimental procedures; NMR spectra and analytical data for all new compounds (PDF)

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**Notes**

The authors declare no competing financial interest.

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