Catalytic Hydrogenolysis of Substituted Diaryl Ethers by Using Ruthenium Nanoparticles on an Acidic Supported Ionic Liquid Phase (Ru@SILP-SO$_3$H)

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Abstract Catalytic hydrogenolysis of diaryl ethers is achieved by using ruthenium nanoparticles immobilized on an acidic supported ionic liquid phase (Ru@SILP-SO$_3$H) as a multifunctional catalyst. The catalyst components are assembled through a molecular approach ensuring synergistic action of the metal and acid functions. The resulting catalyst is highly active for the hydrogenolysis of various diaryl ethers. For symmetric substrates such as diphenyl ether, hydrogenolysis is followed by full hydrodeoxygenation producing the corresponding cycloalkanes as the main products. For unsymmetric substrates, the cleavage of the C–O bond is regioselective and occurs adjacent to the unsubstituted phenyl ring. As hydrogenation of benzene is faster than hydrodeoxygenation over the Ru@SILP-SO$_3$H catalyst, controlled mixtures of cyclohexane and substituted phenols are accessible with good selectivity. Application of Ru@SILP-SO$_3$H catalyst in continuous-flow hydrogenolysis of 2-methoxy-4-methylphenoxybenzene is demonstrated with use of commercial equipment.

Key words ruthenium nanoparticles, acidic SILP, multifunctional catalysis, diaryl ether cleavage, hydrogenolysis, deoxygenation, lignin

The catalytic reductive cleavage of C–O bonds in aryl ethers is an important transformation in synthetic organic chemistry, with possible applications including the synthesis of fine chemicals and the production of fuels from biomass. Breaking aromatic C–O bonds remains challenging, however, because of their high stability especially in diaryl ethers (bond dissociation energy = 82.5 kcal mol$^{-1}$). As a result, significant efforts have been dedicated to the development of homogeneous and heterogeneous catalysts for efficient hydrogenolysis of C–O linkages in diaryl ethers. Multifunctional catalytic systems are required to control the complex reaction network comprising the primary cleavage with parallel or secondary hydrogenation and deoxygenation processes (Scheme 1).

In the field of heterogeneous catalysis, third-row metal catalysts based on Ni,$^{5,6}$ Co,$^7$ and Fe$^8$ were proved to be active for the hydrogenolysis of aromatic C–O bonds, providing high selectivities towards the production of aromatics and phenols. However, these catalysts typically possess limited activities and stabilities under the reaction conditions required for the transformation.$^{6b,7}$ Ru-based catalysts are also known to be highly active for the cleavage of aromatic C–O bonds, typically associated with hydrogenation of the aromatic moieties.$^{5,9}$ Generally, the catalytically active materials for hydrogenolysis of diaryl ethers comprise a combination of metal and acid functionalities.$^{5g,9b,d,10}$ The bifunctional materials can lead to further hydrodeoxygenation of the primary aromatics produced to form...
completely saturated cycloalkanes.† To date, the influence of the substrate's substitution pattern on the product selectivity of the hydrogenolysis remains, however, fairly unexplored.¶

In the present work, ruthenium nanoparticles embedded in sulfonic acid-functionalized supported ionic liquid phases (Ru@SILP-SO$_3$H) are used as catalysts for the hydrogenolysis of substituted symmetric and unsymmetric diaryl ethers. A molecular approach to the preparation of bifunctional Ru@SILP-SO$_3$H catalysts was developed previously in our group and synergistic effects of the metal and acid components were demonstrated.\textsuperscript{12} We now report that the materials are highly active for the hydrogenolysis of aromatic C–O bonds in various substrates, allowing controlled tandem integration with hydrodeoxygenation or hydrogenation of the resulting aromatic moieties. The substrate scope includes substitution patterns as found in lignin feedstock, and controlled mixtures of cyclohexane and phenol derivatives can be produced under continuous-flow operation.

The synthesis of Ru@SILP-SO$_3$H was achieved following a procedure previously reported by our group.\textsuperscript{12a,b} In brief, the sulfonic acid-functionalized supported ionic liquid phase (SILP-SO$_3$H) was prepared through the condensation of the acidic ionic liquid [1-(4-sulfobutyl)-3-(3-triethoxysilyl-propyl)imidazolium][NTf$_2$] on dehydroxylated SiO$_2$ with a loading of 0.6 mmol$_\text{IL}$ g$^{-1}$SiO$_2$.\textsuperscript{11} The generation of ruthenium nanoparticles on the SILP-SO$_3$H phase involved the impregnation and in situ reduction of [Ru(2-methylallyl)$_2$(cod)] under an atmosphere of H$_2$ (100 bar) at 100 °C. A Ru loading of 0.27 wt% was determined by ICP, matching well the theoretical value of 0.32 wt%. TEM analysis of the resulting Ru@SILP-SO$_3$H material evidenced the formation of well-dispersed Ru NPs with a mean particle size of 1.7 ± 0.4 nm (Figure 1). Characterization details are summarized in Tables S1 and S2 in the Supporting Information.

The catalytic performance of Ru@SILP-SO$_3$H for the hydrogenolysis of diaryl ethers was assessed starting with diphenyl ether (1) as model substrate. According to the reaction network shown in Scheme 1, the catalytic transformation of 1 under hydrogen pressure can lead to the formation of several products, such as benzene (1a), phenol (1b), cyclohexane (1c), cyclohexanol (1d), and uncleaved mono- and disaturated ethers (1e and 1f) (Table 1). Gratifyingly, the material proved very active leading to full conversion at a substrate/Ru loading of 500:1 under a standard set of reaction conditions [170 °C, H$_2$ (100 bar), 2 h; Table 1, entry 1]. Cyclohexane (1c) was obtained with a selectivity (S) of 62% indicating a significant contribution of hydrodeoxygenation from cyclohexanol (1d, S = 28%). The only observable side products were the saturated ethers 1e + 1f (S = 10%). Hydrogenolysis of dicyclohexyl ether (1f) under similar reaction conditions gave only 37% conversion, confirming that the main pathway for ether cleavage originates from the aromatic level (Table S3). Extending the reaction time for the conversion of 1 to 16 hours led to almost complete transformation into the fully saturated cyclic product 1c (S = 94%; Table 1, entry 2).

Several control experiments were carried out in order to validate the synergistic action of the metal-acid functionalities of the supported material (Table 1, entry 3–7). The results show that the Ru-free acidic support SILP-SO$_3$H is not active for the cleavage of 1 under the reaction conditions considered, as expected (Table 1, entry 3). While the acid-free Ru@SILP is active for the hydrogenolysis of 1, large amounts of saturated ethers (1e + 1f) (S = 17%) are produced and cyclohexanol (1d) (S = 46%) is not efficiently deoxygenated to cyclohexane (1c) (S = 37%) (Table 1, entry 4). Employing a physical mixture of the two individual materials Ru@SILP and SILP-SO$_3$H resulted in only marginal improvement of the cleavage and deoxygenation activities (Table 1, entry 5). Use of a dispersion of Ru NPs in the nonsupported IL-SO$_3$H resulted in formation of 1c as the major product, but the reaction reached only 34% conversion accompanied by agglomeration of the nanoparticles, in agreement with previous reports\textsuperscript{12b} (Table 1, entry 6 and Figures S1, S2). The catalytic activity and selectivity of Ru@SiO$_2$ under these conditions were very low (Table 1, entry 7). Even after 16 hours the yield of the cleavage products could not
be further increased (Table S4). In addition, TEM analysis of Ru@SiO₂, Ru@SILP and Ru@SILP-SO₃H before and after catalysis (16 hours reaction time) evidenced severe aggregation in the case of Ru@SiO₂, whereas Ru@SILP and Ru@SILP-SO₃H were stable (Figure S3–S6). This demonstrates the importance of the ionic liquid layer in the SILP to stabilize the Ru NPs under these reaction conditions. Taken together, these results clearly demonstrate the synergistic bifunctional catalysis of metal and acid sites and the stabilizing effect of the IL moiety on the silica support.

To gain further insight into the influence of the substitution pattern at aromatic moieties for the hydrogenolysis with Ru@SILP-SO₃H, reactions of symmetric (Table 2) and unsymmetric (Table 3) diaryl ethers were performed under slightly adjusted reaction conditions (200 °C, 100 bar H₂, 3 h). Under these conditions, diphenyl ether (1) reacted nearly quantitatively to cyclohexane (1c) (S = 97%) through the hydrogenolysis/hydrodeoxygenation sequence, with only minor amounts of cyclohexanol (1d) (S = 1%) and benzene (1a) (S = 2%) as the side products (Table 2, entry 1). The presence of methyl groups in para position of the phenyl groups (4,4'-dimethyldiphenyl ether, 2) did not affect the activity and selectivity of the catalyst, producing methylocyclohexane (2c) as sole product (S = 99%) (Table 2, entry 2). The introduction of methoxyl groups in ortho position (2,2'-dimethoxydiphenyl ether, 3) also led to nearly full conversion of the substrate (96%). However, a mixture of products including aromatics [anisol (3a, S = 3%), guaiacol (3b, S = 12%)] and saturated compounds (3c + 3d + 1c + 1d) was formed (Table 2, entry 3). These results indicate that the cleavage of the diaryl ether group was efficiently achieved, but the resulting aromatic monomers 3a and 3b were only partially hydrogenated to their saturated analogues 3c and 3d. This indicates the hindrance of the hydrogenation of the aromatic products by the methoxy substituents. In line with previous findings, the production of 1c as the main product occurred through the hydrogenolysis of the methoxy group in 3c and 3d to form the corresponding alcohols and methane, followed by hydrodeoxygenation of the hydroxyl group.¹²b

On the basis of this observation, we further investigated the influence of the substitution pattern focusing on unsymmetric diaryl substrates possessing a methoxy group in ortho position. Several unsymmetric diaryl ethers were synthesized (see experimental procedures for details¹³) and subjected to hydrogenolysis by using Ru@SILP-SO₃H as catalyst under the previously established reaction conditions (Table 3). In all cases, cyclohexane was obtained as the major component together with phenols and their secondary products. The hydrogenolysis of the monosubstituted 2-methoxyphenoxycyclohexane (4) led to significant hydrodeoxygenation of the phenolic compounds, leading to overstoichiometric amounts of cyclohexane and only minor amounts of the aromatic products (Table 3, entry 1). The amount of aromatic products increased significantly to reach synthetically useful values for substrates 5 and 6, which possess an additional methyl group in meta and para position, respectively (Table 3, entries 2 and 3). High conversions were achieved and the selectivities for cresol 5a (37% selectivity, 41% isolated yield) and of iso-cresol 6a (S = 42%) were nearly equivalent to those of cyclohexane. This trend continued for the sterically hindered substrates 7 and 8, for which the substituted phenols 7a and 8a were obtained in high selectivities in an approximate 1:1 ratio with cyclohexane (Table 3, entries 4 and 5). Only small amounts of other possible side products were formed with these substrates.

### Table 1 Hydrogenolysis of Diphenylether (1) with Use of Ru@SILP-SO₃H and Control Experiments

<table>
<thead>
<tr>
<th>Entry</th>
<th>Catalyst</th>
<th>X (%)</th>
<th>Selectivity S (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>1a</td>
<td>1b</td>
</tr>
<tr>
<td>1</td>
<td>Ru@SILP-SO₃H</td>
<td>&gt;99</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>Ru@SILP-SO₃H</td>
<td>&gt;99</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>SILP-SO₃H</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>Ru@SILP</td>
<td>&gt;99</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>Ru@SILP + SILP-SO₃H</td>
<td>&gt;99</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
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<td>34</td>
<td>11</td>
</tr>
<tr>
<td>7</td>
<td>Ru@SiO₂</td>
<td>&gt;99</td>
<td>0</td>
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</table>

*Ru@SILP-SO₃H (37.5 mg, 0.0012 mmol Ru), 2 h, 170 °C, H₂ (100 bar), substrate (0.6 mmol, 500 equiv), decalin (0.5 mL).

b 16 h.

c Scale-up by factor 4. Conversion and selectivity determined by GC-FID by using hexadecane as an internal standard.
**Table 2** Hydrogenolysis of Symmetric Diaryl Ethers with Ru@SILP-SO$_3$H$^a$

<table>
<thead>
<tr>
<th>Entry</th>
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<th>Selectivity S (%)</th>
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<tr>
<td>3</td>
<td><img src="image3.png" alt="image" /></td>
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<td>3</td>
</tr>
</tbody>
</table>

$^a$ Ru@SILP-SO$_3$H (37.5 mg, 0.0012 mmol Ru), 3 h, 200 °C, H$_2$ (100 bar at rt), substrate (0.6 mmol, 500 equiv), decalin (0.5 mL). Conversion and selectivity determined by GC-FID with use of hexadecane as the internal standard.

**Table 3** Hydrogenolysis of Unsymmetric Diaryl Ethers with Use of Ru@SILP-SO$_3$H$^a$

<table>
<thead>
<tr>
<th>Entry</th>
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<th>Selectivity S (%)</th>
</tr>
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<tr>
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<td>3</td>
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<tr>
<td>2</td>
<td><img src="image5.png" alt="image" /></td>
<td>93</td>
<td>2</td>
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</table>

$^a$ Ru@SILP-SO$_3$H (37.5 mg, 0.0012 mmol Ru), 3 h, 200 °C, H$_2$ (100 bar at rt), substrate (0.6 mmol, 500 equiv), decalin (0.5 mL). Conversion and selectivity determined by GC-FID with use of hexadecane as the internal standard.
Table 3 (continued)

<table>
<thead>
<tr>
<th>Entry</th>
<th>Substrate</th>
<th>X (%)</th>
<th>Selectivity S (%)</th>
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<td><img src="image" alt="Structure 8" /></td>
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<td><img src="image" alt="Structure 6a" /> 8b</td>
</tr>
<tr>
<td></td>
<td></td>
<td>85</td>
<td>6</td>
</tr>
</tbody>
</table>

\(^a\) Ru@SILP-SO$_3$H (37.5 mg, 0.0012 mmol Ru), 3 h, 200 °C, H$_2$ (100 bar at rt), substrate (0.6 mmol, 500 equiv), decalin (0.5 mL).

\(^b\) Substrate (0.3 mmol, 250 equiv). Conversion and selectivity determined by GC-FID by using hexadecane as an internal standard.

\(^c\) Isolated yield.

The selectivities shown in Table 3 provide valuable information on the reaction network of the sequential catalytic transformations. The initial hydrogenolysis of unsymmetric diaryl ethers can proceed in principle through two different routes, which are illustrated in Scheme 2. The production of cyclohexane and substituted phenols in comparable quantities strongly indicates that the ether cleavage occurs preferentially adjacent to the nonsubstituted arene ring with high regioselectivity (Scheme 2, Route 2).

After cleavage of the diaryl ethers, the Ru@SILP-SO$_3$H catalyst hydrogenates quickly the benzene produced to form cyclohexane. The substituted phenols are hydrogenated significantly slower, leaving the substituted phenols as the second main products. The conservation of the phenols' substituent pattern in the aromatic products is consistent with previous literature reports, which suggest that the de-functionalization of phenols requires, first, their hydrogenation to the corresponding substituted cyclohexanols.\(^{12b}\)

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**Scheme 2**  Reaction pathway for the hydrogenolysis of unsymmetric diarylethers with use of Ru@SILP-SO$_3$H: example of 2-methoxy-4-methylphenoxybenzene
Once the hydrogenation occurs, subsequent hydrodeoxy-
genation reduces the oxygen content until, finally, the fully
saturated cyclic products are obtained very slowly.

The hydrogenolysis of 2-methoxy-4-methylphenox-
ybenzene (5) by using the Ru@SILP-SO₃H catalyst was inves-
tigated under continuous-flow conditions in order to evalu-
ate the product distribution as a function of time. In order
to demonstrate the compatibility of the procedure with
commercial equipment, the experiments were carried out
in an H-Cube Pro™ equipped with a Phoenix Flow Reactor™
oven for high temperature experiments. After optimi-
ization of the reaction conditions towards high yields of cre-
osol (5a), (Supporting Information, Tables S5–S7 for de-
tails), the performance of Ru@SILP-SO₃H was tested in a
six hours run (Scheme 3). A temperature of 170 °C, a pres-
sure of 40 bar H₂ (90 NmL min⁻¹), and a substrate flow of
0.3 mL min⁻¹ were chosen corresponding to a residence
time of τ = 12 s (see Supporting Information for further in-
formation). After an induction period (1–2 hours), the reac-
tion reached a maximum conversion of 96% with good se-
lectivity of creosol (5a) (34%) and alkanes (1c + 2c) (43%),
similar to what was obtained under batch conditions
(Table 3, entry 2). In addition, small amounts of hydroge-
nated ethers (5c) (S = 11%), alcoholic intermediates (1d + 2d
+ 5b) (S = 8%) and benzene (1a) (S = 2%) were detected.

The conversion stayed within 86–96% for the period be-
tween two and six hours time on stream. ICP characteriza-
tion of the catalyst after six hours on stream did not show
any leaching of the Ru or the ionic liquid, and changes in the
catalyst’s textural properties were not detected by BET sur-
face area analysis. Characterization of the catalyst before
and after six hours on stream by using DRIFT-IR did not
show any significant change in the regions corresponding
to the characteristic absorption bands of the IL moiety
(2900–3200 cm⁻¹: C–H stretches of the imidazolium ring
and N-alkyl chain; 1450–1570 cm⁻¹: symmetric ring
stretches of the imidazolium moieties; Supporting Informa-
tion, Figures S7–9). In combination with the absence of
leaching, this suggests that the ionic liquid layer is stable
under the reaction conditions considered. In addition, the
size and dispersion of the Ru NPs did not change during the
reaction, as evidenced by TEM analysis (see Supporting In-
formation, Tables S1, S2, and Figure S10 for characterization
details). The Ru@SILP-SO₃H catalyst thus appears to be

![Scheme 3](image-url)
chemically stably under continuous flow conditions. Nevertheless, a small decrease in conversion and alkane yield after four hours suggests a loss of hydrogenolysis and hydrogenation activity with time. On the basis of previous findings, the loss of catalytic activity observed may be attributed to the accumulation of water on the catalyst during the reaction.\textsuperscript{12b,14}

The hydrogenolysis of diaryl ethers was accomplished by using a bifunctional catalytic system consisting of ruthenium nanoparticles embedded on an acidic supported ionic liquid phase (Ru@SILP-SO\(_3\)H). The molecular design of the catalyst allowed for a close proximity between the metal and acid sites, which was shown to be a requirement for high catalytic activity. Various diaryl ethers with different substitution patterns were used as the substrates to study the catalytic activity and selectivity of Ru@SILP-SO\(_3\)H. While diphenyl ether and 4,4′-dimethyldiphenyl ether were readily converted to the corresponding cycloalkanes in high yields, the presence of methoxy groups as substituents hindered the hydrodeoxygenation of the cleaved moieties. In case of unsymmetrical diaryl ethers, the cleavage of the C–O bond occurred adjacent to the unsubstituted phenyl ring regioselectively. This allows to achieve controlled formation of mixtures of methoxy-substituted phenols and cyclohexanes starting from substrates with substitution patterns as found in lignin-type feedstocks. Application of the Ru@SILP-SO\(_3\)H catalyst under continuous-flow conditions was demonstrated for the hydrogenolysis of 2-methoxy-4-methylphenoxybenzene. Production of a mixture of cyclohexane and cresol as the main products was achieved over six hours on stream with conversions and yields similar to those observed under batch conditions. These results further substantiate the potential of molecular approaches for the design and the synthesis of multifunctional catalytic systems based on metal nanoparticles immobilized on functionalized supported ionic liquid phases (M@SILP-func) to control complex reaction networks involving consecutive and parallel hydrogenation and hydrogenolysis steps.

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Supporting Information

Supporting information for this article is available online at https://doi.org/10.1055/s-0037-1611678.

References and Notes

\textsuperscript{(6)} (a) Von Stein, I.; Weigand, T.; Hermannsdorfer, J.; Kretschmer, W. P.; Motz, G.; Kempe, R. ChemCatChem 2013, 5, 11759.
\textsuperscript{(7)} (a) Worster, D. F.; Cartwright, R.; Rinaldi, R. ChemSusChem 2016, 9, 4567.


(13) Experimental Procedures

Safety warning: High-pressure experiments with compressed H2 must be carried out only with appropriate equipment and under rigorous safety precautions.

General: If not otherwise stated, the syntheses of the ionic liquids (ILs), the supported ionic liquid phases (SILPs), and the nanoparticles immobilized on SILPs (Ru@SILP and Ru@SILP-SO3H) were carried out under an inert atmosphere by using standard Schlenk techniques or inside a glovebox. After synthesis, ILs, SILPs, Ru@SILP, and Ru@SILP-SO3H were stored under an inert atmosphere. If not otherwise stated solvents were used after distillation without any further purification. For reactions under an inert atmosphere, solvents were additionally dried with molecular sieves (4 Å) and degassed by flushing solvents with argon. For catalysis decalin (from ACROS, 99% anhydrous) and iodobenzene (456 mg, 2.40 mmol, 10.0 mol%) were charged into an autoclave and used without purification. For catalysis decalin (from ACROS, 99% anhydrous) and iodobenzene (456 mg, 2.40 mmol, 10.0 mol%) were charged into an autoclave and used without purification.

Synthesis of catalysts: The Ru@SILP, Ru@SILP-SO3H, and Ru@IL-SO3H were synthesized as previously reported and Ru@SO3H was synthesized accordingly.12a

Synthesis of substrates: The synthesis of diaryl ethers was carried out as previously reported.12a A round-bottom Schlenk flask (250 mL) was equipped with a magnetic stir bar, copper iodide (456 mg, 2.40 mmol, 10.0 mol%), picolinic acid (590 mg, 4.80 mmol, 20.0 equiv), and potassium phosphate (10.2 g, 48.0 mmol, 2.00 equiv). A second Schlenk flask (100 mL) was charged with the aryl iodide (24.0 mmol, 1 equiv), the phenol (28.8 mmol, 1.2 equiv), and anhydrous DMSO (50 mL). The solution of flask 2 was transferred into flask 1 under flowing Argon and the reaction mixture was stirred for 20 h at 100 °C, subsequently. After cooling down, the reaction mixture was diluted with a 1:1 mixture of a saturated aqueous solution of NaH2PO4 (200 mL) and H2O (200 mL). After addition with CH3Cl2 (3 × 200 mL) the combined organic phases were washed with a 5% aqueous solution of KOH (300 mL) and brine (300 mL). After drying the mixture with Na2SO4 the solvent was removed and the crude reaction mixture was preadsorbed on silica gel. The purification of the crude mixture was achieved with use of silica gel. Remaining iodine compounds caused the deactivation of the catalyst in subsequent catalytic reactions and therefore had to be separated carefully. This afforded in some cases purification by two subsequent chromatography columns. Full characterization of all substrates can be found in the Supporting Information.

2.6-Dimethoxy-4-methylphenoxymethane: Prepared according to the general procedure by using iodobenzene (50.0 g, 24.0 mmol) and 2.6-dimethoxy-4-methylphenol (4.8 g, 28.8 mmol). The crude product was purified by flash column chromatography (elucent: ethyl acetate/n-pentane 1:8). 2.6-Dimethoxy-4-methylphenoxymethane was obtained as a slightly off-white solid (4.46 g, 18.3 mmol) in 76% yield.12c H NMR (500 MHz, CDCl3): δ = 7.29–7.26 (m, 2 H), 7.02–6.98 (m, 1 H), 6.93–6.90 (m, 2 H), 6.51 (s, 2 H), 3.75 (s, 6 H), 2.42 (s, 3 H) ppm.13c NMR (125.7 MHz, CDCl3): δ = 158.77 (C), 153.23 (C), 135.65 (C), 129.84 (C), 129.39 (CH), 121.52 (CH), 114.87 (CH), 106.28 (CH), 56.33 (CH3), 22.24 (CH3) ppm. HRMS (EI): m/z 244.11.

Batch catalysis: In a typical experiment, Ru@SILP (37.5 mg, 0.0012 mmol Ru), substrate (0.6 mmol, 500 equiv), and decalin (0.5 mL) were combined in a glass insert and placed in a high-pressure autoclave. After purging the autoclave with H2, the reaction mixture was stirred at 170 °C in an aluminium heating cone under 120 bar H2 (pressurized at 100 bar H2 at rt). After the reaction, the autoclave was cooled in an ice bath, carefully vented, and the reaction mixture filtered before GC analysis with use of hexadecane as an internal standard. In some cases a gap in mass balance was observed because of the loss of cyclohexane in the headspace of the autoclave (scale-up decreased this error).

Isolated yield (5a): The catalysis was scaled up by a factor of 5. After reaction the decalin phase was isolated and the catalyst washed with decalin (1 × 5 mL). The combined organic phases were extracted with aqueous KOH (3 × 10 mL, 0.3 M). The KOH phase was washed with n-pentane (3 × 10 mL) and neutralized with aqueous HCl, subsequently. After neutralization, the aqueous phase was extracted with CH3Cl2 (4 × 20 mL). After drying the CH3Cl2 phase with MgSO4 and removal of the solvent a yellow-brownish oil (186 mg, 1.35 mmol) was obtained in 45% yield (See Supporting Information, Figure S11 for characterization).

Continuous flow catalysis: A 70 mm CatCart® (V packed react = 0.5126 mL) was filled with Ru@SILP-SO3H (439 mg, 0.0144 mmol Ru) and installed into the H-Cube ProTM. Prior to catalysis the catalyst was flushed with decalin (1 mL min–1 for 30 min). Then, the reactor was pressurized to 40 bar H2 (90 NmL min–1) and heated to 170 °C. When reaching stable reaction conditions, the substrate solution (0.025 M in decalin) was introduced into the system (0.3 mL min–1). The reaction was allowed to equilibrate for 30 min before the first samples were collected. The reaction mixture was analysed by GC analysis with use of hexadecane as an internal standard.