Hydrogen spillover assisted by oxygenate molecules over nonreducible oxides

Mingwu Tan¹, Yanling Yang¹, Ying Yang¹, Jiali Chen¹, Zhaoxia Zhang¹, Gang Fu¹, Jingdong Lin¹, Shaolong Wan¹, Shuai Wang¹, & Yong Wang²

Spontaneous migration of atomic hydrogen species from metal particles to the surface of their support, known as hydrogen spillover, has been claimed to play a major role in catalytic processes involving hydrogen. While this phenomenon is well established on reducible oxide supports, its realization on much more commonly used non-reducible oxides is still challenged. Here we present a general strategy to enable effective hydrogen spillover over nonreducible SiO₂ with aid of gaseous organic molecules containing a carbonyl group. By using hierarchically-porous-SiO₂-supported bimetallic Pt-Fe catalysts with Pt nanoparticles exclusively deposited into the micropores, we demonstrate that activated hydrogen species generated on the Pt sites within the micropores can be readily transported by these oxygenate molecules to Fe sites located in macropores, leading to significantly accelerated hydrodeoxygenation rates on the latter sites. This finding provides a molecule-assisted approach to the rational design and optimization of multifunctional heterogeneous catalysts, reminiscent of the role of molecular coenzymes in bio-catalysis.
Hydrogen spillover has been coined to denote an important effect involved in heterogeneous catalysis\(^1\)--\(^3\), in which atomic H species produced from H\(_2\) dissociative adsorption on a catalytic site spontaneously migrate to a different type of site that cannot readily dissociate H\(_2\) molecules under the same condition\(^4\)--\(^5\). This effect not only accounts for many phenomena observed in hydrogenation/hydrogenolysis processes\(^6\)--\(^7\), but also makes it possible to decouple the catalytic functions between H\(_2\) dissociation (to form active H atoms) and subsequent hydrogenation with these atomic H species, rendering the control of these catalytic sequences separately and precisely towards more efficient synthesis of chemicals\(^8\)--\(^14\).

Extensive studies have been devoted to investigating the mechanism of hydrogen spillover on oxide supports\(^15\)--\(^19\). On one hand, it is commonly accepted that atomic H species can migrate along surfaces of reducible oxide (e.g., TiO\(_2\) and WO\(_3\)) via coherent proton-electron movements\(^9\)--\(^16\), in which the H atom gives its electron to a reducible metal cation of the oxide support and diffuses in the form of proton. On the other hand, the possibility of hydrogen spillover on a nonreducible oxide (e.g., SiO\(_2\) and Al\(_2\)O\(_3\)) has been debated for a long time\(^17\)--\(^19\), and only until recently it was found by combined experimental and theoretical evidences that the mobility of H atoms on a nonreducible Al\(_2\)O\(_3\) support is orders of magnitude slower and restricted to a much shorter distance (less than sub-nanometer) than the case for a reducible TiO\(_2\) support\(^9\)--\(^15\). Although some studies suggest that the presence of sufficient acidic surface OH groups or introducing defect sites into the oxide surface may promote the hydrogen transfer process\(^10\)--\(^11\), a general and robust strategy is still highly desired for achieving an effective hydrogen spillover on nonreducible oxides with a low density of acid sites or defects, which are widely used as catalyst supports in industry because of their excellent thermochemical stability, low cost, and tunable acid-base property.

In our previous work\(^19\), Pt-Fe bimetallic catalysts (denoted as Pt@-Fe@SiO\(_2\)) with Pt and Fe nanoparticles encapsulated in the micropores and macropores of a hierarchically structured SiO\(_2\) support, respectively, showed activity for hydroydeoxygenation (HDO) of pyrolysis bio-oil vapor on Fe sites with H\(_2\) generated from steam reforming on Pt sites. However, questions such as how the H species are stabilized and how the H species “shuttle” or communicate between the spatially isolated Pt and Fe sites remained unanswered. Here, using HDO of lignin-derived guaiacol to produce value-added aromatic hydrocarbon products (e.g., benzene, toluene, and xylenes, denoted collectively as BTX) as a model reaction, we unveil that an efficient hydrogen spillover from the Pt sites to the Fe sites occurs over the SiO\(_2\) support with aid of gaseous oxygenate molecules containing a carbonyl functional group such as aldehydes, ketones, and esters (illustrated in Fig. 1), which leads to significantly accelerated HDO rates and is reminiscent of molecular coenzymes that assist in enzyme activity in a similar manner\(^20\).

### Results

**Synthesis and characterization of Pt@-Fe@SiO\(_2\) catalysts.** As reported in our previous study\(^18\), Pt@-Fe@SiO\(_2\) catalysts were synthesized via a dual template method with a nonionic polyglycol ether surfactant (Tergitol 15-S-5) and a cellulose extraction thimble used as the templates to generate micropores and macropores in SiO\(_2\), respectively (described in Supplementary Fig. 1 of the supporting information). The content of Pt in these catalysts varied within 0.5--2.0 wt%, while the content of Fe was kept at 10 wt%. These catalysts are denoted as xPt@-10Fe@SiO\(_2\) (x = 0.5--2.0) henceforth. For comparison, 1Pt@SiO\(_2\) with 1.0 wt% Pt encapsulated within the micropores of SiO\(_2\) and 10Fe@SiO\(_2\) with 10 wt% Fe mainly deposited within the macropores of SiO\(_2\) were also synthesized using this dual template method.

The micropores of the synthesized SiO\(_2\) support possess a size distribution mainly within 1.0--2.0 nm as determined by N\(_2\) physisorption isotherms (Supplementary Fig. 2a) and small angle X-ray powder diffraction (XRD) spectra (Supplementary Fig. 3a), and show a well-ordered wormhole-like feature according to transmission electron microscopy (TEM) images (Supplementary Fig. 4). Mercury intrusion porosimetry measurement further confirms the existence of macropores with a diameter ranging from 100 to 160 nm for the SiO\(_2\) support (Supplementary Fig. 2b). This hierarchically porous structure is well maintained after depositing the Pt and Fe nanoparticles onto the SiO\(_2\) support, consistent with insignificant changes of the N\(_2\) physisorption isotherms and average pore sizes (Supplementary Fig. 5).

XRD patterns of these Pt@-Fe@SiO\(_2\) samples show the diffraction peaks belonging to cubic Fe crystalline phases (e.g., \(2\theta = 44.6^\circ\)), whereas characteristic diffraction peaks associated with the Pt nanoparticles are undetectable reflective of their high dispersion (Supplementary Fig. 6). TEM images confirm uniformly dispersed Pt nanoparticles of around 1.6 ± 0.3 nm for the Pt@-Fe@SiO\(_2\) catalysts, irrespective of the presence of Fe species (Fig. 2a and b) and the Pt loading amount within 0.5--2.0 wt% (Supplementary Fig. 7), which reflects the confinement effect on the growth of these Pt nanoparticles within micropores. In contrast, the Fe nanoparticles deposited in the macropores show much larger sizes (e.g., 6.5 ± 0.7 nm for 1Pt@-10Fe@SiO\(_2\); 7.2 ± 0.7 nm for 10Fe@SiO\(_2\), Fig. 2a, c). H\(_2\)-temperature-programmed reduction (H\(_2\)-TPR) profiles of the oxide precursors for 10Fe@SiO\(_2\) and 1Pt@-10Fe@SiO\(_2\) show nearly identical temperatures (c.a. 441°C) for the reduction of Fe\(_2\)O\(_3\) (Fig. 2d), consistent with the spatial isolation between Pt and Fe nanoparticles in these Pt@-Fe@SiO\(_2\) catalysts. Otherwise, the promoted reduction of Fe\(_2\)O\(_3\) species is expected if they are in contact with Pt nanoparticles as found for the Pt-Fe/SiO\(_2\) catalyst (1 wt% Pt, 10 wt% Fe) synthesized via a conventional co-impregnation method, which exhibits the H\(_2\)-TPR peak at an apparently lower temperature (419°C, Fig. 2d). Moreover, these close reduction temperatures for the Fe\(_2\)O\(_3\) species also confirm that hydrogen spillover is inhibited on the SiO\(_2\) support used in this study.

### Rates and selectivity of guaiacol hydroydeoxygenation on Pt@-Fe@SiO\(_2\) Catalysts.

Catalytic performances of 1Pt@SiO\(_2\), 10Fe@SiO\(_2\), and 1Pt@-10Fe@SiO\(_2\) were compared in guaiacol HDO (450°C, 0.5 kPa guaiacol, 50 kPa H\(_2\), 4.0 mL s\(^{-1}\) g\(_{\text{cat}}\)\(^{-1}\) of space velocity), which has been widely applied as a model reaction for studying the upgrading of lignin-derived phenolics to aromatic hydrocarbons\(^21\)--\(^27\). As illustrated in Fig. 3, the C-O bond cleavage in guaiacol on a metal catalyst can lead to sequential formations of catechol, phenol, and benzene. Concomitant with these conversions, the methyl species generated in the catechol formation can recombine with phenol to yield anisole and cresol, which are grouped with phenol as the partially deoxygenated products (denoted here as PAC), while the corresponding methylated derivatives of benzene mainly include toluene and xylene and are grouped with benzene since they are all fully deoxygenated products (i.e., BTX). Alternatively, guaiacol, catechol, and other phenolics can undergo hydrogenation of their aromatic ring on a metal catalyst, followed by subsequent decomposition into C\(_1\) and C\(_2\) gaseous molecules (e.g., methane, CO, CO\(_2\), and ethane) via C=C bond cleavage. Previous studies have shown that the selectivity of guaiacol HDO is sensitive to the nature of the metal catalyst\(^24\). In particular, Fe is highly selective for the C-C bond cleavage\(^25\)--\(^27\), while those C=C hydrogenation
and C–C hydrogenolysis reactions are favored on noble metals (e.g., Pt and Pd). 1Pt@SiO₂ showed a low guaiacol conversion of 15.8% with catechol and PAC as the predominant products, the selectivities of which were 70.2% and 17.9%, respectively (Entry 1, Table 1). The main activity of 1Pt@SiO₂ is likely contributed by the residual active sites on the SiO₂ support, because the guaiacol conversion (11.2%) and the combined selectivity of catechol and PAC (84.3%) obtained on the parent SiO₂ support (Entry 2, Table 1) were close to those of 1Pt@SiO₂. In contrast, a referenced
non-encapsulated Pt/C catalyst (3 wt% Pt) exhibited a nearly complete guaiacol conversion (97.4%, Entry 3 in Table 1), in which the C₀ and C₂ gaseous products (71.9% selectivity) were formed from the successive C–C–O hydrogenolysis reactions prevailing among the products. The above data clearly indicate that most of the Pt nanoparticles on 1Pt@SiO₂ are not accessible to the bulky guaiacol reactants due to their successful encapsulation within the micropores of SiO₂.

Different from the above catalysts, a guaiacol conversion of 94.8% with selectivities to PAC and BTX of 46.4% and 48.5%, respectively, was achieved on 10Fe@SiO₂ (Entry 4, Table 1). As expected, guaiacol can readily adsorb on the Fe active sites because of the preferential deposition of the Fe nanoparticles in the macropores of SiO₂ and proceed the chemical transformations expected on Fe surface to form fully deoxygenated BTX products.²⁶,²⁷ The 1Pt@10Fe@SiO₂ sample showed a guaiacol conversion and HDO selectivities nearly identical to those obtained on 10Fe@SiO₂ (Entry 5, Table 1), suggesting that the coexistence of Pt with Fe, but with spatial isolation achieved here, does not interfere with the catalytic properties of Fe in the hierarchically- porous SiO₂ framework. It has been reported that a close contact between Pt and Fe particles under the HDO reaction condition can improve the catalytic activity of Fe particles via hydrogen spillover from the Pt sites to the Fe ones,²⁵,²⁶ because the activity of Fe catalysts is generally limited by strongly bound O atoms on the Fe surfaces that hinder the activation of H₂ molecules.²⁸ This similarity of catalytic performance between 1Pt@10Fe@SiO₂ and 10Fe@SiO₂ in guaiacol HDO (verified within 10–60 kPa H₂ partial pressures, Supplementary Fig. 8) thus provides compelling evidence for the unlikely hydrogen spillover between isolated Pt and Fe nanoparticles on the nonreducible SiO₂ support under the reaction conditions studied, which is consistent with previous reports.²⁵,¹⁸

Figure 4 shows that the formation rate of BTX (rBTX) on 1Pt–10Fe@SiO₂, a descriptor of the catalytic HDO activity, is sensitive to the H₂ partial pressure and the source of the H₂ feed (details of conversion and selectivity shown in Supplementary Table 1). As the H₂ partial pressure changed from 10 to 50 kPa, the rBTX value increased accordingly from 0.44 to 0.84 mol mol⁻¹ Fe⁻¹ h⁻¹ consistent with the fact that H₂ is required for the oxygen removal from guaiacol. On the other hand, in lieu of feeding H₂ directly, H₂ is expected to be generated in situ via methanol steam reforming (CH₃OH + H₂O → CO₂ + 3H₂) on Pt under the conditions studied,²⁹,³⁰ if CH₃OH and H₂O are co-fed into the reactor. We found that an equimolar mixture of CH₃OH and H₂O (4.8 kPa each) generated an effective H₂ partial pressure of 11 kPa (corresponding to 88% methanol conversion), which was measured from the effluent of the reactor (Supplementary Table 1). At this reaction condition, rBTX reached 1.28 mol mol⁻¹ Fe⁻¹ h⁻¹, about three times the rate when H₂ was directly fed with a similar partial pressure (0.44 mol mol⁻¹ Fe⁻¹ h⁻¹). This rBTX value achieved with co-feeding CH₃OH and H₂O was even higher than that obtained with co-feeding 50 kPa H₂ (0.84 mol mol⁻¹ Fe⁻¹ h⁻¹, Fig. 4). Such a promoting effect of the CH₃OH–H₂O mixture was further confirmed for guaiacol HDO at lower conversions (i.e., ~30%, Supplementary Fig. 9), in which PAC existed as the main products. Moreover, an activity enhancement of the guaiacol conversion was also observed for the 1Pt@SiO₂ catalyst (Supplementary Fig. 10), despite 1Pt@SiO₂ was much less active than 1Pt–10Fe@SiO₂ and showed a high selectivity to PAC (Table 1). It is thus suggested that this promoting effect brought forth by the co-feeding of CH₃OH and H₂O appears to be general for the catalysts with spatially-separated H₂-activation and hydrodeoxygenation sites.

To better quantify the effects of H₂ generated in situ via methanol steam reforming, we further examined how low the CH₃OH and H₂O partial pressures could be to reach a rBTX value similar to that when directly feeding with 50 kPa H₂. As shown in Fig. 4, a comparable rBTX of 0.72 mol mol⁻¹ Fe⁻¹ h⁻¹ was achieved when 0.5 kPa CH₃OH and 0.5 kPa H₂O were co-fed (vs 0.84 mol mol⁻¹ Fe⁻¹ h⁻¹ for directly feeding with 50 kPa H₂), representing a much lower effective H₂ partial pressure of mere 1.3 kPa. It is worth noting that rBTX on 10Fe@SiO₂ was only about a quarter of that for 1Pt–10Fe@SiO₂ when the same mixture of CH₃OH and H₂O was fed (Supplementary Fig. 11), which excludes possible roles of CH₃OH and H₂O directly involved in promoting guaiacol HDO on the accessible Fe particles (e.g., transfer hydrogenation

Table 1 Catalytic performance of gas-phase guaiacol hydrodeoxygenation.³⁰

| Entry | Catalyst | Guaiacol conversion (%) | Carbon selectivity (%) | Formation rate of BTX (mol mol Fe⁻¹ h⁻¹) |
|-------|----------|--------------------------|------------------------|----------------------------------------|
|       |          |                          | Catechol | PAC | BTX | C₈−C₉ |
| 1     | 1Pt@SiO₂ | 15.8                     | 17.9     | 70.2 | 3.5    | <0.1 |<0.1 | 0.01 |
| 2     | SiO₂     | 11.2                     | 33.1     | 51.2 | <0.1  | <0.1 | 0.09 |
| 3     | 3 wt% Pt/C | 97.4                  | 0.5      | 18.9 | 5.3    | 71.9 | 0.83 |
| 4     | 10Fe@SiO₂ | 94.8                     | 5.1      | 46.4 | 48.5  | <0.1 |<0.1 | 0.83 |
| 5     | 1Pt–10Fe@SiO₂ | 96.3 | 2.2               | 48.1 | 48.8 | <0.1 |<0.1 | 0.84 |

*Reaction condition: 450 °C, 0.5 kPa guaiacol, 50 kPa H₂, balanced by N₂, 0.25 g of SiO₂ 1 h⁻¹ space velocity.

**Denoted for phenol, anisole, and cresol.

†Denoted for benzene, toluene, and xylene.

²⁴Mainly including methane, CO, CO₂, and ethane.
Driven by methanol and thus confirms that the Pt sites within the micropores of 1Pt@-10Fe@SiO₂ are responsible for the H₂ production from methanol steam reforming. Moreover, negligible formation of C₁-C₂ products in the presence of co-fed CH₃OH and H₂O (Supplementary Table 1 of the supporting information) indicates that no Pt species migrated out from the small pores, otherwise significant C=C cleavage reactions would occur as observed for the Pt/C catalyst (71.9% selectivity, Table 1 of the manuscript). Here, we speculate that CH₃OH or H₂O may only not act as a hydrogen source during guaiacol HDO, but also transport active H species produced on the Pt sites to the Fe particles, accounting for the increased r_BTX.

Hydrogen spillover enabled by molecular carriers over SiO₂. Previous studies have reported that hydrogen spillover between two different metal sites on a nonreducible support is limited within the sub-nanometer scale and decays sharply as the distance increases⁵,¹⁷, due to the unstable nature of atomic H species on the nonreducible support. To confirm our hypothesis on the promoted hydrogen spillover over Pt@Fe@SiO₂ in the presence of the CH₃OH/H₂O mixture, we changed the distance between the Pt and Fe particles in the Pt@-10Fe@SiO₂ catalysts by varying the Pt loading and physically mixing the 1Pt@SiO₂ and 10Fe@SiO₂ catalysts (mass ratio 1:1; denoted as 1Pt@SiO₂ + 10Fe@SiO₂) with different degrees of separation. As described above, the xPt@10Fe@SiO₂ catalysts (x = 0.5, 1.0, 2.0 wt%) possessed Pt particles of similar sizes (~1.6 nm, Fig. 2 and Supplementary Fig. 7), which ensures that the increase of the Pt loading merely increased the areal density of the Pt particles and did not change their intrinsic activity. By assuming homogeneous distributions for both the Pt particles within the micropores and the Fe particles within the macropores, the average distances between the nearest Pt and Fe particles (<d_Pt,Fe>) in the xPt@10Fe@SiO₂ catalysts (x = 0.5, 1.0, 2.0 wt%) are estimated to be 10, 57, and 3.4 nm, respectively. We further increased <d_Pt,Fe> to the magnitude of 10³ nm for the well-mixed 1Pt@SiO₂ + 10Fe@SiO₂ sample and 10⁷ nm for the case with a dual-bed configuration (10Fe@SiO₂ as the downstream bed, Fig. 5). Guaiacol HDO on these encapsulated Pt-Fe catalysts with co-feeding CH₃OH and H₂O as the H₂ source showed that, even though the apparent H₂ production rate (r_BTX) increased exponentially from 0.17 to 0.92 mol mol⁻¹Fe⁻¹ h⁻¹ as <d_Pt,Fe> decreased from 10⁻⁷ to 3.4 nm (Fig. 5), further confirming the existence of effective hydrogen spillover between spatially isolated Pt and Fe particles over SiO₂ brought forth by the CH₃OH or H₂O additive, especially in the length scale of several nm. It is also worth pointing out that the similar HDO rates obtained for the well-mixed and layered 1Pt@SiO₂ + 10Fe@SiO₂ catalysts (Fig. 5) exclude the possible role of gas-phase diffusion in the H transport process, since gaseous radicals can diffuse at a millimeter scale under similar conditions³¹,⁳². In other words, it is likely that the oxygenate-bound H radicals mainly transport over the SiO₂ surface (instead of diffusing in gas phase), benefiting from the stabilization of these radicals by the oxide surface via H-bonding and van der Waals interactions.

We postulate that the transfer of active H atoms generated on the Pt sites to the Fe surface requires a molecular carrier, which should be capable of accepting the atomic H species to form a stable enough intermediate for long-distance transport and eventual H release. Intuitively, the most possible carrier candidates include not only CH₃OH and H₂O that were co-fed to generate H₂ for the HDO reaction, but also formaldehyde (HCHO) that could form from dehydrogenation of CH₃OH on the Pt sites, although only a trace of HCHO was detected from the effluent because of the thermodynamic limit at the HDO reaction condition. Here, H-atom addition energy (HAE) was calculated via density functional theory (DFT) for these three candidates and used to assess their ability to carry a H atom. These HAE values were defined as the energy change for the addition of a H atom to a molecule (M) in gas phase, in which E_M, E_H, and E_MH are the energies for gaseous M, H, and the H-added MH product, respectively. According to DFT calculations (Fig. 6), the H-addition of H₂O is endothermic (HAE: 58 kJ mol⁻¹), even more unfavorable than that for H₂ (HAE: 16 kJ mol⁻¹). Although the H-addition of CH₃OH is exothermic (HAE: −125 kJ mol⁻¹, Fig. 6), CH₃OH is unstable by the attack of a H atom, leading to the formation of a methyl radical and a H₂O molecule. In contrast, HCHO is the most likely carrier for the hydrogen spillover on Pt@Fe@SiO₂, because of the highly favorable nature of this H-addition process (HAE: −154 kJ mol⁻¹, Fig. 6) and the relatively high stability of the corresponding H-added product (i.e., CH₂OH⁻). The radical nature of such oxygenate transporting species during guaiacol HDO on 1Pt@-10Fe@SiO₂ was further confirmed by the observation of a

![Figure 5](https://example.com/fig5.png)  
**Fig. 5** Effects of Pt-Fe distance on BTX formation rates. r_BTX is shown here as a function of <d_Pt,Fe> for guaiacol HDO with H₂ generated in situ from methanol stream reforming (450 °C, 0.5 kPa guaiacol, 0.5 kPa methanol +0.5 kPa H₂O, balanced by N₂, 0.5 g·s·ml⁻¹ space velocity).

![Figure 6](https://example.com/fig6.png)  
**Fig. 6** DFT-derived H-atom addition energies (HAE). The calculations were carried out at the B3LYP/6-311 + G(d,p) level of theory.
significant inhibiting effect of NO, a radical chain inhibitor, on the BTX formation rate no matter whether H2 was directly fed or generated in situ from a mixture of CH3OH and H2O (Supplementary Figs. 12 and 13). It is also worth noting that the possible role of H2O in promoting the H transfer over SiO29 was excluded based on the negligible effect of H2O addition on the HDO activity of 1Pt@-10Fe@SiO2 at 10 kPa H2 (Supplementary Fig. 14). On the other hand, a slight deactivation of the 1Pt@-10Fe@SiO2 catalyst was observed with time-on-stream when cofeeding CH3OH and H2O (Supplementary Fig. 13), which is mainly due to the oxidation of metallic Fe by H2O or active site blocking by adsorbed oxygenate species27,28. This suggests that a sufficient H2 partial pressure, which is either cofed or formed in situ, is requisite for promoting the Fe-based catalysts from deactivation during HDO.

The ability of carrying H atoms is hypothesized to be general for oxygenate molecules containing a carbonyl functional group (e.g., acetaldehyde, propanal, acetone, and methyl formate) based on their negative HAE values (DFT data shown in Supplementary Fig. 15). Therefore, we further examined the effects of these oxygenate additives on the rates of guaiacol HDO over 1Pt@-10Fe@SiO2, and these experiments were conducted without cofeeding H2O in order to avoid the possible formation of surface OH groups on the SiO2 support that may promote the hydrogen spillover.9 It is also worth noting that the cofed C2-C3 oxygenates are stable and showed negligible conversions during the guaiacol HDO reaction, while the conversion of CH3OH is around 30%, mainly leading to the formations of H2, HCHO and CO2. As expected, all of them were able to promote the HDO rates (Fig. 7a), and the enhanced BTX formation rates approximately reached maximum values \( r_{BTX,max} \) as either the H2 or the oxygenate additive partial pressure became sufficiently high (Fig. 7a and Supplementary Fig. 16, respectively), reflecting the intrinsic ability of these oxygenate additives in carrying H atoms under the HDO condition. As shown in Fig. 7b, \( r_{BTX,max} \) increased with the oxygenate additives in the order of methyl formate < acetone < acetaldehyde < propanal < formaldehyde, consistent with the trend of their HAE values. This excellent correlation strongly supports that these oxygenate additives containing a carbonyl functional group act as an effective H-carrier to promote guaiacol HDO on Fe surface (illustrated in Fig. 7c).

**Discussion**

In summary, bimetallic Pt@-Fe@SiO2 catalysts with Pt and Fe nanoparticles encapsulated in the respective micropores and macropores of the hierarchically structured SiO2 support were prepared successfully via a dual template method and used here to examine the H transfer between these isolated metal sites during guaiacol hydrodeoxygenation. The distinct C–O/C=O hydrogenolysis selectivities of the Pt and Fe nanoparticles and the inaccessibility of the bulky guaiacol reactant to the micropores allowed us to experimentally confirm that H spillover over the SiO2 surface is hard to occur at conditions relevant to catalysis, due to the nonreducible nature of SiO2. More importantly, we found that the H migration from Pt to Fe sites can be dramatically enhanced when a small oxygenate additive (e.g., C1-C3 carbonyl compounds and esters) is present, resulting in much improved hydrodeoxygenation activity than the monometallic Fe and Pt catalysts. The stabilization of active H atoms conferred by the cofed oxygenate additive that underpins the achieved H spillover over nonreducible oxides also reflects from the decrease of activation barrier for the kinetically-relevant hydrogenating steps involved in guaiacol hydrodeoxygenation on Fe surface. It is noteworthy that these carbonyl H-carriers can be generated in situ from dehydrogenation or steam reforming of their alcohol precursors, which act concomitantly as a source of H2 for the hydrodeoxygenation reaction. The above findings demonstrate a molecule-assisted strategy for establishing an efficient transfer of active chemical species between catalytic sites that are spatially isolated on an inert support, and may bring an alternative
approach to optimize or design novel systems of multifunctional catalysis.

Methods

Catalyst preparation. The Pt-Fe/SiO2 catalysts with Pt and Fe particles deposited on the respective micropores and macropores of the hierarchically structured SiO2 support were prepared via a dual-template step-impregnation method as described in our previous study19. In a typical preparation procedure, surfactant Tergitol 15-S-5 (used as the micropore template, Aldrich), concentrated HCl, and tetraethoxysilicate (TEOS, Sinopharm Chemical Reagent Co. 99.0%) were sequentially added into a mixed solvent of de-ionized water, ethanol (Sinopharm Chemical Reagent Co. 99.5%) with the molar ratio of TEOS: water: HCl, and surfactant Tergitol 15-S-5 (used as the micropore template, Aldrich) at liquid nitrogen temperature (−196 °C), in which each sample was degassed at 150 °C for 4 h before measurements. Images of transmission electron microscopy (TEM) were collected with a Tecnai G2 F20 field emission microscope operating at 200 kV, and the size distributions of the Pt and Fe particles were obtained by counting ca. 200 particles in each image. Hydrogen–temperature programmed reduction (H2–TPR) experiments were carried out in a homemade fixed-bed reactor. 100 mg of sample was in situ pre-treated in flowing Ar (30 mL min−1) at 200 °C for 1 h before test. After cooling down to 30 °C, the sample was heated to 800 °C (ramping rate 10 °C min−1) in a 5 vol% H2/Ar flow (30 mL min−1) in a 5 vol% H2/Ar flow (30 mL min−1), in which the H2 uptake was measured and quantified using a thermal conductivity detector (TCD).

Catalytic tests. Guaiacol hydrodeoxygenation was conducted under atmospheric pressure in a vertical fixed-bed quartz tube reactor with an internal diameter of 8 mm and a length of 750 mm. 200 mg of catalyst diluted with 800 mg of quartz particles (0.18–0.25 mm) was placed between two layers of quartz wool in the center of the reactor. The temperature of the catalyst was monitored using a K-type thermocouple placed in the middle of the catalyst bed. Prior to catalytic measurements, the catalyst was treated in flowing 50 vol% H2/N2 (50 mL min−1) at 450 °C (ramping rate 5 °C min−1) for 2 h. Guaiacol and oxygenate additives with set partial pressures were introduced separately using syringe pumps and transferred into the reactor by a N2 flow. All transfer lines were heated to around 210 °C to avoid condensation of liquid reactants or products. The components of the effluent from the reactor were analyzed quantitatively using an on-line gas chromatograph with a DB-WAXETR column (50 m × 0.320 mm) connected to a flame ionization detector (FID) for separating oxygenates and hydrocarbons and a TDX-01 column connected to a thermal conductivity detector (TCD) for separating small molecules (CH4, CO, CO2, and H2). N2 was used as an internal standard, and response factors for guaiacol and the hydrodeoxygenation products were determined using standard chemicals with known concentrations. The mass balance for each experiment, unless otherwise noted, was at least 95%. Guaiacol conversions and product selectivities were reported on a carbon basis as described elsewhere27.

Theoretical calculation. The values of hydrogen affinity (Eq. (1)) for various oxygenates, H2O, and H2 were calculated via the Gaussian software package at the hybrid B3LYP functional level of theory. The standard 6–311+G(d,p) basis set was used for all atoms, and the Grimme’s D3BJ dispersion correction was taken into account in the electronic energy calculations. Geometry optimizations were performed using the Berny geometry algorithm with convergence criteria of 1.0 × 10−6 Ha for energy and 1.5 × 10−5 Ha Bohr−1 for the maximum residual forces on each atom.

Data availability

The authors declare that all the relevant data within this paper and its Supplementary Information file are available from the corresponding authors upon a reasonable request.

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**Author contributions**

Y.W., Shu.W., and M.T. conceived the idea for the project. M.T., Ya.Y., Yi.Y. and J.C. conducted the catalyst synthesis, structural characterization, and catalytic experiments. Shu.W. conducted the theoretical calculations. M.T. drafted the manuscript under the guidance of Y.W and Shu.W. Z.Z., G.F., J.L., Sha.W. and the authors mentioned above all discussed and commented on the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

**Supplementary information**

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**Correspondence** and requests for materials should be addressed to Shuai Wang or Yong Wang.

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