

## PERSPECTIVE

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## Heterolytic cleavage of dihydrogen (HCD) in metal nanoparticle catalysis†

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The heterolytic cleavage of H<sub>2</sub> into H<sup>+</sup> and H<sup>−</sup>, and the subsequent transfer of these hydrogen species to polarized C=X groups, is a useful strategy to obtain high selectivities in catalytic hydrogenation reactions, which are one of the most important chemical processes. In the homogeneous field, this transformation is catalyzed by a wide variety of metal–ligand complexes, leading to the selective reduction of ketones, aldehydes, and imines, to name a few. In many cases, a ligand–metal cooperative mechanism promotes the activation of the hydrogen molecule; that is, a Lewis basic site of the ligand binds the proton, and simultaneously, a coordinated metal atom acts as a Lewis acid and takes the hydride. Similar to homogeneous catalysts based on organometallic complexes, in the field of metal nanoparticle catalysis the use of supports, ligands and additives can promote heterolytic H<sub>2</sub> splitting by a cooperative mechanism with the metal, and thus produce the selective reduction of thermodynamically more stable unsaturated groups. This successful approach has allowed very high selectivities to be obtained, with both supported and non-supported metal nanoparticles (MNPs). This Perspective aims to carry out a critical review of recent examples on heterolytic cleavage of dihydrogen mediated by MNPs, making a selection of the most representative works in search for general features and new steps to be followed. As will be shown, promising advances have been made through the use of supported MNPs functionalized by organic ligands, which combine the advantages of homogeneous and heterogeneous catalysts (activity, selectivity, stability and recyclability), and thus represent a new research area with tremendous industrial interest.

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## 1. Introduction

## 1.1 Homogeneous systems

Catalytic hydrogenation of unsaturated organic substrates is one of the most abundant chemical transformations. It uses both homogeneous and heterogeneous catalysts, and for the latter it is one of the oldest reactions marking the start of the discipline with the well-known studies of Sabatier. Large-scale applications involving methanation of carbon oxides, ammonia synthesis, and hydrogenation of fats are over a century old. Homogeneous metal complex catalytic systems were discovered only halfway through the 20th century; several simple metal salts showed hydrogenation activity, but copper salts stood out in the early days. The late start of homogeneous metal catalysis and the relative ease of molecular studies in solution led to a strong focus on

mechanistic aspects right from the start. Activation of dihydrogen by metal complexes has been a main theme in chemistry ever since the beginning of homogeneous



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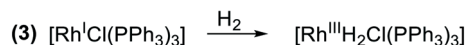
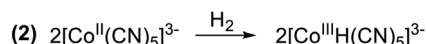
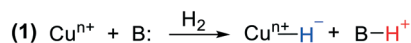
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catalysis.<sup>1,2</sup> In the copper catalysis mentioned above, heterolytic cleavage of dihydrogen (HCD) dominated the mechanistic scene, *i.e.* a copper salt and a base would react with H<sub>2</sub> producing a hydride at copper together with the conjugate acid of the base (Scheme 1, reaction (1)). We won't go into the details of the valence state of Cu, but as one can see during this reaction the valence of the metal does not change. Further reaction can reduce the metal ion, as investigated by Halpern. Three reactions for hydrogen activation were initially distinguished by Halpern, the second being homolytic cleavage, particularly studied for bimetallic complexes, *e.g.* Co carbonyls and Co cyanides, in which the dihydrogen molecule splits into two hydrogen "radicals" (Scheme 1, reaction (2)). This "radical" view was certainly stimulated by the organic radical reactions shown by the same complexes. The third reaction by which dihydrogen engages in homogeneous catalysis is oxidative addition to low valent metal complexes, which became popular with the new rhodium hydrogenation catalysts, *e.g.* the Wilkinson catalyst,<sup>3</sup> soon expanded by enantioselective versions, notably by Kagan,<sup>4</sup> again with important mechanistic insight contributed by Halpern.<sup>5</sup> In an oxidative addition reaction, both hydrogen atoms formally accept an electron from the transition metal and become hydride anions and the metal increases its valence state by two (Scheme 1, reaction (3)). In particular for 16-electron square-planar group 9 and 10 complexes, this has been a popular model in the classroom, as it explains well the changes in the complexes that take place during the reaction. Reaction (2), mechanistically the homolytic pathway, can also be regarded as an oxidative addition to a bimetallic complex, the valence of each metal being raised by one, and this would be in line with the organometallic formalism of hydrides.



**Scheme 1** (1) Heterolytic H<sub>2</sub> cleavage by a copper salt and a base. (2) Homolytic H<sub>2</sub> cleavage by a Co cyanide complex. (3) Homolytic H<sub>2</sub> cleavage with an oxidative addition reaction by the Wilkinson catalyst.

The oxidative addition model has dominated the mechanistic work on hydrogenation for decades, perhaps until the late 1980s. However, as was already pointed out by Pauling,<sup>6</sup> the charge distribution in a complex does not change as much as the formal description would suggest upon oxidative addition, and the bond between hydrogen and the "electronegative" noble metal should be regarded as covalent and the hydride formed does not have a "hydridic" character in middle and late transition metal hydrides.<sup>7</sup> This can be illustrated by many examples of group 9 and 10 metals of which the hydrides are stable in the presence of strong acids, not quite a property of main group or early transition metal (ETM) hydrides. For example, the stable species (diphosphine)PdH<sup>+</sup> (abundant in palladium carbonylation chemistry), various metal hydrides used in heterolytic hydrogenation of alkenes, and (triphenylphosphine)PdH<sup>+</sup> (for electrochemical reduction of CO<sub>2</sub>) do not react with acid.<sup>8–10</sup> "Oxidative addition" of hydrogen to electropositive main group metals such as elemental Ca and Mg leads to hydride species (because of their reversibility considered as hydrogen storage materials),<sup>11,12</sup> but these hydrides are usually not active in alkene hydrogenation and less so as catalysts. Hydrides of electropositive metals such as LiAlH<sub>4</sub>



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can be used for the stoichiometric hydrogenation of C=O and C=N bonds, but under more forcing conditions they may react with alkenes, as is well known for aluminium, used on a large industrial scale for oligomerization of ethene, although chain transfer is stoichiometric.

Hydrides of immobilized metal complexes were reviewed by Copéret, who noted that the hydrides of group 4–6 early-transition-metals are “hydridic” in character. Numerous reports exist about their application in alkene conversions.<sup>13</sup> Their preparation usually involves hydrogenolysis of organometallic precursors and rarely oxidative addition or heterolytic cleavage of H<sub>2</sub>; in view of their “very hydridic character they react fast with protic solvents, *etc.*”.

For the synthetic chemist and the electrochemist, the “thermodynamic hydricity” of hydrides is a useful yardstick that can tell about the reactivity of the hydride in important reactions such as reduction of aldehydes and CO<sub>2</sub> or water splitting; the hydricity tells us whether a hydride can be transferred from the metal to CO<sub>2</sub>, or the reverse.<sup>14,15</sup> Thermodynamic hydricity, or hydride donating ability, is the free energy required to heterolytically cleave a metal hydride bond (it is a measurable and/or calculable quantity, expressed in kJ mol<sup>−1</sup>; we will use “hydridic” (character) in the usual loose, qualitative way). For instance, Kubiak *et al.*<sup>15</sup> described very nicely the linear relationship between hydricity and reversible reduction potential of M(diphosphine)<sub>2</sub><sup>n+</sup> complexes and their corresponding hydrides, and indicated which ones are thermodynamically capable of transferring a hydride to CO<sub>2</sub> (overall thermodynamics requires a base to complete the reaction to formate rather than formic acid). In simple catalyst complexes of this type, hydricity parallels catalyst reactivity, as reported by Wiedner, Linehan *et al.*<sup>16</sup> In many practical hydrogenation catalysts of hetero atom–carbon bonds, however, this is not the case, as the catalytic process involves multiple interactions with Lewis acid sites, bases, and protic sites.

HCD on a transition metal with the formal formation of a proton and a hydride anion does not normally produce a hydridic hydride as one can imagine from the reaction already studied by Halpern in which H<sub>2</sub> reacts with Ag<sup>+</sup> to give “AgH”, Ag being only slightly less electronegative than hydrogen (Pauling values 1.9 (Ag) and 2.1 (H), respectively; Pd in the example above has 2.2). The unstable “AgH” may decompose into Ag and H<sup>+</sup> and this way the silver salt is reduced.

After two decades of a dominating position of oxidative addition in mechanistic studies, heterolytic cleavage came to the forefront again in the 1990s with the revival of the “other” Wilkinson catalyst,<sup>17</sup> the one based on Ru instead of Rh, *via* the work of Noyori and Shvo on hydrogenation and transfer hydrogenation but also in relation to the work on hydricity cited above.<sup>18–20</sup> The development of the chemistry of dihydrogen complexes contributed to new insights as, for instance, dihydrogen complexes of Ru were found to react with amines giving a ruthenium hydride and an ammonium salt, as was shown in 1990 by Chinn and Heinekey.<sup>21</sup> An early example in which a basic ligand functionality fulfils the role of the base was reported by Fryzuk, who described the heterolytic cleavage

of H<sub>2</sub> across an Ir–amide bond of a P–N–P ligand.<sup>22</sup> Later, however, they reported that H<sub>2</sub> activation occurred in this particular instance by oxidative addition–reductive transfer rather than as a direct heterolytic cleavage of H<sub>2</sub>.<sup>23</sup> Ever since, a wide variety of ligand involvements in metal-catalysed reactions have been reported and reviewed (pincer catalysis,<sup>24–26</sup> outer-sphere hydrogenation catalysis,<sup>27</sup> functional ligands,<sup>28</sup> models for hydrogenase,<sup>29</sup> cooperating ligands,<sup>30</sup> ligand-assisted proton transfer,<sup>31–34</sup> outer-sphere hydrogen transfer,<sup>35</sup> *etc.*). In particular for the heterolytic hydrogenation of CO<sub>2</sub>, ligand-assisted pathways are of crucial importance.<sup>36</sup> HCD in such systems was reviewed by Ito and Ikariya.<sup>37</sup>

## 1.2 Heterogeneous systems

In heterogeneous catalysis, the earliest hydrogenation reaction known is that of the platinum catalysed combination of hydrogen and oxygen in Döbereiner's lamp with the use of the catalyst discovered by the Davy nephews,<sup>38</sup> commercialized in 1823.<sup>39</sup> The French chemist Paul Sabatier is considered the father of the hydrogenation process (Nobel prize 1912, shared with Victor Grignard). In 1897, he discovered that small amounts of nickel catalysed the addition of hydrogen to molecules such as carbon dioxide in what is now known as the Sabatier process. A few years later, Wilhelm Normann disclosed in patents the hydrogenation of liquid oils, the beginning of fat hardening which became a huge industry. Other reactions comprise the hydrogenation of nitrogen (Haber–Bosch process, 1905) and the hydrogenation of coal-derived carbon monoxide (Fischer–Tropsch process, 1922) which converts coal to liquid fuels. Thus, the early heterogeneous hydrogenations have been game changers in the world's energy, agriculture, and food manufacture. The understanding of heterogeneous hydrogenation on a molecular level is usually stated to begin with the publication of Horiuti and Polanyi,<sup>40</sup> while taking into account various findings by others such as physisorption and chemisorption of substrates and dissociative chemisorption of dihydrogen<sup>41</sup> described by Langmuir (Nobel prize 1932). Also in 1934, the Farkas brothers in collaboration with Rideal reported that the transfer of hydrogen to ethene, hitherto described as a transfer of the H<sub>2</sub> molecule to ethene on Ni and Pt surfaces, occurs stepwise, H by H (Fig. 1a).<sup>42</sup> Based on calculations and kinetics, the preferred mechanism for H/D exchange and hydrogenation of ethene by Horiuti and Polanyi involves the formation of ethylmetal intermediates, taking place in their view at as many as 3 or 4 surface metal atoms without particular placement mentioned of the four metal atoms (Fig. 1b). A discarded mechanism was actually what we would call today a  $\sigma$ -bond metathesis at two adjacent metal atoms (thus a five-membered ring  $\cdots\text{M}-\text{H}\cdots\text{C}-\text{H}\cdots\text{M}'\cdots$ , Fig. 1c), instead of one metal atom as we know from organometallic complexes (four-membered ring, Fig. 1d).

Polanyi's “insertion” mechanism has stood the test of time with flying colours, as has the dissociative chemisorption of dihydrogen into hydrogen atoms on the metal surface. One of the key contributions of transition-metal surfaces to



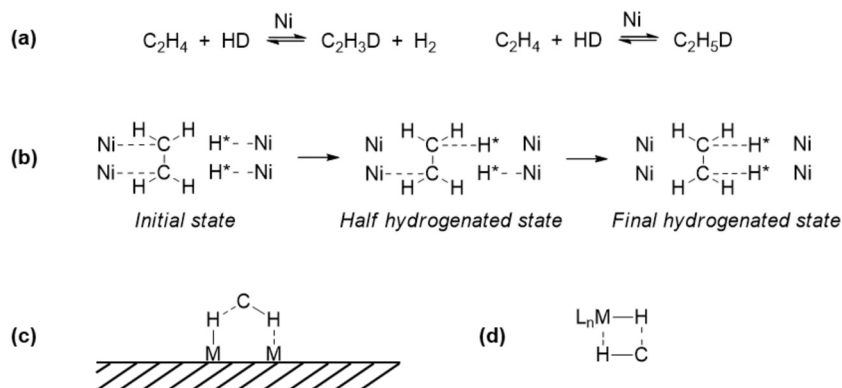


Fig. 1 (a) Exchange and addition reactions of ethylene and HD mediated by Ni reported by the Farkas brothers and Rideal, which led them to propose that transfer of  $\text{H}_2$  to ethylene on Ni and Pt surfaces occurs stepwise. (b) Mechanism for hydrogenation of ethene by Horiuti and Polanyi. (c) Five-membered ring  $\sigma$ -bond metathesis. (d) Four-membered ring  $\sigma$ -bond metathesis.

catalysis is their ability to facilitate the dissociation of the stable  $\text{H}_2$  molecules into H atoms adsorbed on the surface together with multiple C–H activation/isomerization mechanisms.<sup>43,44</sup> As an illustration of the latter, we mention (1) the two adsorption modes of ethene on Pt,  $\eta^2$ -Pt and  $\eta^2$ - $\mu^2$ -Pt<sub>2</sub>, named (CC) and di- $\sigma$ (CC) in heterogeneous catalysis, (2) ethyl-Pt and vinyl-Pt in several coordination modes, (3)  $\mu^2$ -ethylidene-Pt<sub>2</sub> and  $\mu^3$ -ethylidyne-Pt<sub>3</sub>, *etc.* (Fig. 2).<sup>45</sup> A seminal paper discussing similarities between the reaction of C–H and H–H on surfaces and complexes was published by Saillard and Hoffmann.<sup>46</sup> A major difference between solid metal hydrides and metal complex hydrides is the issue of hydridic character, which is completely missing for hydrides on metals. While one might hope that for metal complexes a single parameter that gives an indication of this property can be found, the hydrides on (and below) metal surfaces cannot be treated like this, because the charge, or better its reactivity, is the property of a complicated ensemble of atoms (*e.g.* metal face, shape, size, coverage, binding mode, mobility of metal atoms and hydrides, other ligands, *etc.*).<sup>47</sup> The huge number of reports about homolytic cleavage on many metals did not lead to simple comparisons of reactivities with those of metal complex hydrides.

However, in addition to the overwhelming number of publications using this homolytic paradigm, since the 1960s there has also been ample support for ionic mechanisms, although, to the best of our knowledge, not for purely metallic systems. Two lines of research will be discussed, heterolytic cleavage onto metal oxide catalysts and spill-over from metal catalysts to oxidic supports. HCD on solid oxide catalysts was reported in the 1950s and in hindsight its relation to the homogeneous, heterolytic mechanism has

been noted. Trapnell *et al.* reported that a variety of metal oxides catalysed the exchange of  $\text{H}_2$  and  $\text{D}_2$  at mild temperatures.<sup>48</sup> One mechanism proposed concerned heterolytic cleavage on non-reducible metal oxides (Fig. 3a, the same as for homogeneous systems, Scheme 1, reaction (1)); another one for reducible oxides, coined “homolytical” or “radicalar”, concerned the transfer of a proton to oxygen and an electron to the metal (Fig. 3b). Chromium oxide gel catalysed the deuteration of 1-hexene giving mostly dideuteriohexane as found by Burwell, while metallic catalysts give quite different patterns characterized by extensive multiple exchange due to reversible reactions and the various bonding modes of alkenes,<sup>43</sup> which is indicative of different mechanisms for metal and metal oxide catalysts.<sup>49,50</sup> In this early example from 1960, the cleavage of  $\text{H}_2$  on  $\alpha$ -Cr<sub>2</sub>O<sub>3</sub> as a step in the hydrogenation of 1-hexene was named “heterolytic dissociative adsorption at pair sites”, and although the valence state of Cr might have been (II) or (III),<sup>51</sup> in Burwell's view it did not change during the hydrogen activation, as is typical of both the heterogeneous and the homogeneous mechanism for HCD.<sup>52</sup> Interestingly, the early hydrogenation work involved apolar substrates such

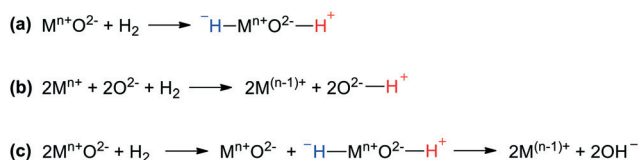


Fig. 3 (a) HCD on non-reducible metal oxide catalysts. (b) Homolytic cleavage of  $\text{H}_2$  on reducible metal oxide catalysts. (c) HCD followed by formation of a homolytic product on reducible metal oxide catalysts.

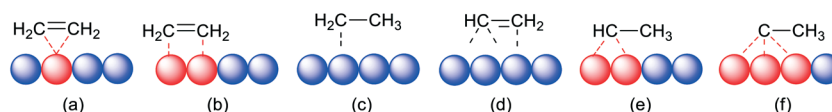


Fig. 2 Adsorption modes of ethene on Pt: (a)  $\eta^2$ -Pt (CC); (b)  $\eta^2$ - $\mu^2$ -Pt<sub>2</sub> (di- $\sigma$ (CC)); (c) ethyl and (d) vinyl in different coordination modes; (e)  $\mu^2$ -ethylidene-Pt<sub>2</sub>; (f)  $\mu^3$ -ethylidyne-Pt<sub>3</sub>.





as alkenes, and not the typical substrates one would expect for an otherwise heterolytic mechanism.<sup>53</sup>

In an HCD on a metal oxide, a metal hydride and a hydroxide are formed, and the energy gain should largely compensate for the loss of the binding energy of H<sub>2</sub> (436 kJ mol<sup>-1</sup>; upper values for Pt-H 260 kJ mol<sup>-1</sup> and O-H 460 kJ mol<sup>-1</sup>). For most transition metals, the balance will allow the addition of dihydrogen to their metal oxide. Liu reported a combined DFT and experimental study of metal ions on silica and found that the species with the highest heat of metal-hydride formation were the most active ones for propene hydrogenation under mild conditions.<sup>54</sup> Late, high-valent transition metals and post-transition metals were much more active than early transition metals, although the formation of the M(H)-O(H) intermediate is not always thermodynamically favorable. Amongst the active metals were also indium and zinc, in accord with the DFT calculations on hydride formation. Clearly, for electropositive main group metals this will mostly not be the case and hydride formation does not occur. For such reactions in bulk materials, thermodynamics prohibit the formation of metal hydrides, *e.g.* aluminium and zinc. However, for zinc atoms located at certain edges of particles or nanoparticles the balance may be favorable or close to thermodynamically feasible, or as an accessible transition state, sometimes en route to reduction of Zn.<sup>55</sup> In a catalytic process, the required energy will add to the overall activation barrier, but catalysis *via* metal hydrides may still occur.<sup>56</sup> For instance, Pydko and Van Santen calculated that ethane will cleave heterolytically on certain Zn ions in a zeolite to give ethyl-“zinc” and hydroxide, and ethene and hydrogen *via*  $\beta$ -elimination.<sup>57</sup> Hydrogen activation on non-reducible oxides has long been under debate, but evidence in favour of it has accumulated. For instance, Cop  ret and co-workers have recently shown that  $\gamma$ -Al<sub>2</sub>O<sub>3</sub>, when treated at appropriately high temperatures, can activate H<sub>2</sub> and catalyse the hydrogenation of simple alkenes. The thermal treatment generates defect sites on the alumina surface, which effectively function as frustrated Lewis pairs (FLPs, *vide infra*).<sup>58</sup>

Above we mentioned already the addition of dihydrogen to reducible metal oxides, a process that leads to metal reduction by the electrons and formation of hydroxy groups. Like the addition to noble metals, the process is called homolytic, but radicalar has also been used. Garc  a-Melchor and L  pez studied as an example of such a process the addition of H<sub>2</sub> to CeO<sub>2</sub>, a reducible oxide, not very active in catalytic hydrogenation.<sup>59</sup> The analysis pointed out that dissociative H<sub>2</sub> adsorption takes place through a heterolytic pathway followed by the transfer of a hydrogen atom that finally yields the homolytic product (Fig. 3c). From the numerous examples of ETM oxides, we mention the work by Calatayud *et al.*,<sup>60</sup> who studied H<sub>2</sub> dissociation on various rutile TiO<sub>2</sub> facets by DFT. The topology of the surface had a moderate effect on kinetics and thermodynamics as all four surfaces initially gave heterolytic dissociation to hydride-hydroxyl surface pairs (as in the study by L  pez), which then rearranged by proton and electron transfer to the thermodynamically more favourable reduction product.

We now turn briefly to the phenomenon called spill-over, which is relevant to nanoparticle catalysed hydrogenations. Prins reviewed spill-over in detail a few years ago.<sup>61</sup> The name spill-over was coined by Boudart to explain a phenomenon first described by Khoobiar a few years earlier.<sup>62,63</sup> Khoobiar described in 1964 that Pt particles on a yellow WO<sub>3</sub> support under a hydrogen atmosphere apparently catalysed the reduction of WO<sub>3</sub> to bluish, reduced oxides, while hydrogen alone did not react with WO<sub>3</sub> under the same conditions. It was proposed that dihydrogen undergoes dissociative chemisorption on Pt, and that the H-atoms migrate (spill over) to the support and convert to a proton and an electron, see Fig. 4. The proton binds to an oxygen anion and the electron reduces W(vi) to W(v). Both species are mobile in/on the support and further spill-over can occur. Water and alcohols enhance the process and mobility of the protons by providing a more favourable binding of the protons. Huizinga and Prins provided EPR evidence for the reduction of the TiO<sub>2</sub> support by H<sub>2</sub> dissociated on Pt particles (Fig. 4).<sup>64</sup>

Clearly, for this process to occur one needs a reducible support, but n-type semiconductors such as ZnO may also undergo spill-over in such a way, the electrons being stored in the conduction band.<sup>61</sup> Many publications report on the participation of spilled-over hydrogen in catalysis on non-reducible oxides (alumina, silica); other scientists remain skeptical and many different explanations for the influence of nearby supports have been brought forward. For spill-over to silica and alumina the debate continues, as it is difficult to exclude other explanations. However, Van Bokhoven *et al.* have shown that, under highly controlled laboratory experiments in a nanofabricated system, spill-over from Pt particles to iron oxide particles was much more effective on reducible titania than on “non-reducible” alumina.<sup>65,66</sup>

A further effect of hydrogen spill-over to reducible oxides is the creation of oxygen deficiencies at the surface due to formation of water and reduction of the metal. The open metal sites are now available for coordination to a Lewis-donor substrate, *e.g.* the aldehyde of an unsaturated enal. Somorjai and Baker have brought this forward as an explanation for the high activity and high selectivity for alcohol formation in the hydrogenation of unsaturated aldehydes by Pt NPs on reducible oxides compared to non-reducible oxides.<sup>67,68</sup> Calculations showed that hydrogen atom transfer to the support-coordinated aldehyde is a favoured process, while in the absence of Lewis base-acid interaction it is not (a H atom and an electron are transferred

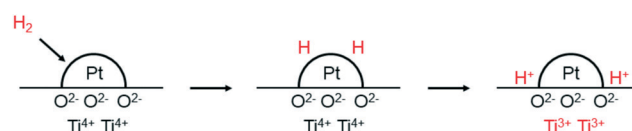


Fig. 4 Adsorption of H<sub>2</sub> on Pt and spill-over of H atoms to the TiO<sub>2</sub> support, producing protons and Ti<sup>3+</sup> cations. Adapted with permission from ref. 64. Copyright 1981. American Chemical Society.



to the organic moiety). It was proposed that the alkoxy/alkoxide formed migrates to the Pt surface to react with a second H atom, although in the general view of heterolytic mechanisms a proton transfer from the support to alkoxide seems a more logical step. Note that the preferred reaction of unmodified Pt and Pd with hydrogen and unsaturated aldehydes is hydrogenation of the C=C bond.<sup>69</sup> Unlike homogeneous systems, the presence of nearby protons on a support was usually not taken into account in the mechanistic considerations and calculations in the hydrogenation of unsaturated ketones and aldehydes with Pt, Pd, Ru, and Au catalysts.<sup>70</sup> As in homogeneous systems the game is open to several possibilities, with initiation by proton transfer, atom transfer, hydride transfer or migration, or simultaneous (outer-sphere) transfer of the hydride and proton.<sup>19,71,72</sup> For the present discussion the conclusion that the participation of hydride ions in spill-over systems is the least probable amongst the mechanistic pathways suffices.

The spill-over mechanism received a lot of interest,<sup>61,73</sup> but less attention has been paid to the interaction of dihydrogen at the edges of metal particles and support materials. In the last decade, however, mechanistic interpretations involving the bifunctional behaviour of the metal and support, in particular for NPs, have gained importance. At the edges of MNPs on supports one can imagine an HCD reaction similar to that on metal oxides or in metallic complexes with the metal as the hydride acceptor and the oxide as the base (Fig. 5), as was suggested by Fujitani *et al.* to explain the high activity of small, supported Au NPs in HD exchange reactions; the larger the perimeter interface, the higher the catalytic activity.<sup>74</sup> Proof was provided by García *et al.*, who described the HCD by Au NPs on ceria NPs, leading to the concomitant formation of Au-H and positively charged bridged OH groups, besides the generation of water from framework oxygen atoms, as studied by vibration spectroscopy.<sup>75</sup> Interestingly, the reaction of the hydrogen loaded catalyst with CO<sub>2</sub> and O<sub>2</sub> gave formic acid and hydrogen peroxide, respectively. Following this discovery, heterolytic cleavage has slowly re-established its position in the reports on heterogeneous hydrogenation pathways and evidence is also growing.<sup>59,76–78</sup>

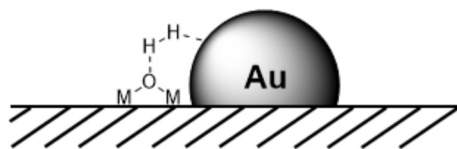


Fig. 5 Heterolytic H<sub>2</sub> cleavage at the edges of MNPs (for example AuNPs) supported on metal oxide surfaces.

It should be borne in mind that all processes described above for supported metal particles can take place simultaneously: homolytic cleavage on the metal only, spill-over to reducible and non-reducible oxides, heterolytic cleavage at the perimeters (which might also be succeeded by spill-over of hydrogen atoms in either direction), and heterolytic cleavage on the support. The species formed will be related to different catalytic actions and control of the catalyst performance will require control over the formation of all species.

Potentially, a cleaner method to achieve heterolytic dihydrogen activation on MNPs could be the use of ligands on the NPs or stabilizing agents surrounding the NPs. For practical purposes, MNPs in solution have the disadvantage that they are difficult to separate from the reaction mixture after reaction, but nanofiltration or immobilization on passive supports might solve this. Fig. 6a illustrates the basic idea; a nearby base on the metal surface aids the HCD, the same way as in homogeneous systems (Scheme 1, reaction (3)) and on metal oxides (Fig. 3a).<sup>79</sup> In the last decade, this technique has found applications at increasing speed and, for instance, Rossi *et al.* reported highly active Au/SiO<sub>2</sub> alkyne semi-hydrogenation catalysts in which piperazine was used as a ligand and base for a highly efficient HCD (*vide infra* for more details).<sup>80</sup> One of us introduced secondary phosphine oxides (SPOs) on MNPs to this end, as it was thought that the P-atom would bind to the metal and the O-atom would be free as a proton acceptor, while a neighbouring metal atom can bind a hydride (Fig. 6b).<sup>81</sup> Both for piperazine and SPO, the heterolytic action was supported by DFT calculations; for both heterolytic pathways the activation barrier was considerably lowered compared to the homolytic ones.<sup>80,82</sup> When the reaction occurs across unbound M-B (base) sites, the system is called a frustrated Lewis pair, FLP,<sup>83</sup> but for bound M-B sites the term FLP is also used, as in the area of FLP chemistry and catalysis.<sup>84</sup>

### 1.3 Enantioselective hydrogenation

The best proof for ligand or metal-ligand complex involvement in catalysis is the use of chiral ligands and the induction of chirality in the product. As far as we could trace this back, asymmetric metal catalysis probably started in the area of heterogeneous catalysis when Yoshiharu Izumi *et al.* used Pd on silk fibres for the hydrogenation of oxime derivatives to produce chiral amino acid precursors with modest ees.<sup>85</sup> The authors used several metals and subsequently they switched to metal catalysts, especially RANEY® Ni, modified with chiral hydroxyacids, as reviewed in 1971.<sup>86</sup> Cinchona molecules have been widely employed as

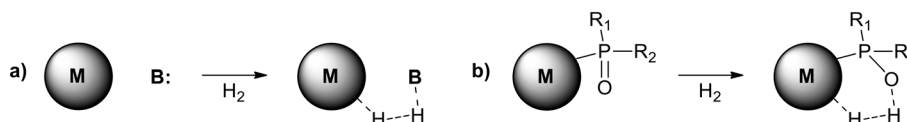


Fig. 6 Heterolytic splitting of H<sub>2</sub> by (a) MNPs and a nearby base; (b) SPO-ligated MNPs.



chiral modifiers and accelerators on metal catalysts since their introduction by Orito *et al.*<sup>87</sup> They obtained 79% ee for the hydrogenation of keto-esters with Pt/C. The scope has since then been extended, the mechanism has been studied in detail, many ligand variations have been added, industrial applications were investigated, *etc.*<sup>88</sup> Notably, the substrates are activated ketones. For the chiral catalysts mentioned here, one usually invokes a hydrogen bond interaction between the ligand and the substrate to enforce a certain orientation of the latter with respect to the catalyst surface. The cinchona molecules contain a tertiary amine that functions as a proton receptor/donor and often it was thought that for the creation of this ammonium donor a protic solvent was needed. However, it was shown that in aprotic solvents and the gas-phase, the proton can also be delivered by the Pt-surface (as a spill-over, one might say),<sup>89,90</sup> although Hahn and Baiker described the process consistently as H atom transfer.<sup>91</sup> Mechanistically, the chemistry of these modifying chiral ligands on the surface could be closely related to heterolytic cleavage. However, in the vast literature on ligand-induced, heterogeneous, asymmetric hydrogenations, heterolytic cleavage does not seem to play a role in the discussions and we mention this catalysis only here in the introduction.

#### 1.4 Frustrated Lewis pairs

About 15 years ago, Erker and Stephan introduced a new concept and route for the HCD, *viz.* the use of FLPs.<sup>92</sup> Typically, the initial Lewis base and acid contained main group elements such as phosphorus and boron as the central atoms. Even more exciting was the finding that the reagent can be used for the hydrogenation of alkenes.<sup>93</sup> The boron Lewis acids used contain fluoro-substituted aryl groups and thus the resulting hydrides are not very hydridic. Transition metal based FLPs (TMFLPs) have also been developed with borane and alumane as the Lewis acids. In the reactions of TMFLPs, the proton is transferred to the transition metal and in several examples the hydride may well possess a hydridic character, but no data on hydricity have been reported.<sup>84,94</sup> Apart from the two ligand-modified Au NPs mentioned above that were categorized as FLPs, there are only a few uses of FLPs on the surface of MNPs so far. To facilitate a practical use of FLP hydrogenation, several ways for immobilization have been studied. Solid state FLP catalysts were reviewed by Qu *et al.*<sup>95</sup> The review features FLPs taken from the solution work of which one partner of the pair is immobilized on a solid. This work closely resembles the solution work, and indeed MNPs are not involved. Intramolecular FLPs have been immobilized on the usual range of supports, including polymers. Several of the now FLP classified examples coincide with the metal oxide examples included above, but we want to single out one particular support here, *viz.* N-doped carbon materials in which the N atoms are assigned the role of the Lewis base and Pd NPs the role of the Lewis acid (see section 3.1).<sup>96</sup>

In this perspective we focus on hydrogenation catalysis by metal nanoparticles that are mechanistically based on heterolytic cleavage of dihydrogen. HCD into  $H^+$  and  $H^-$ , and their subsequent transfer of these hydrogen species to polarized  $C=X$  groups is a useful stratagem for obtaining high selectivities in hydrogenation catalysis.<sup>97</sup> Only reports that explicitly comment on this mechanism will be considered and preferably the mechanistic suggestions will be supported by experiments or theoretical calculations. Firstly, we will discuss MNPs in solution stabilized by ligands and polymers, followed by MNPs on various supports, including a particular section on supported MNPs functionalized with organic ligands. Finally, HCD by “isolated” metal oxide NPs will be examined.

## 2. Heterolytic cleavage of $H_2$ on MNPs in solution

### 2.1 MNPs stabilized by polymers

First, we will discuss MNPs synthesized by reaction of metal precursors and reducing agents in the presence of polymers that prohibit the formation of bulk metal by stabilizing the MNPs formed initially. Among the different types of nanoparticle-based systems employed to carry out catalytic transformations that involve a heterolytic cleavage of  $H_2$ , MNPs stabilized by polymers have shown a limited efficacy. This is probably due to the fact that heterolytic  $H_2$  activation requires an intimate contact between the metal surface atoms and the stabiliser to promote the desired cooperative effect. There are only two reports on HCD mediated by MNPs stabilised with polymers. Sánchez-Delgado and co-workers described the synthesis of ruthenium nanoparticles (RuNPs) of 1–2 nm in size immobilised on cross-linked poly(4-vinylpyridine) (PVPy).<sup>98</sup> This catalytic system was employed for the hydrogenation of quinoline (substrate/Ru = 84, methanol, 30 bar  $H_2$ , 120 °C; TOF = 66  $h^{-1}$ ), leading to the formation of 1,2,3,4-tetrahydroquinoline with high selectivity. Catalytic experiments by adding an external base or acid and with solvents of different polarities suggested an ionic mechanism through a heterolytic activation of dihydrogen. It was proposed that RuNPs are stabilised in the vicinity of the pyridine groups of the polymer (Fig. 7a), in such a way that  $H_2$  is heterolytically cleaved through the cooperation between RuNPs and basic N-pyridine atoms of the polymer (Fig. 7b). Subsequently, an outer sphere concerted transfer of proton and hydride species to the substrate would take place (Scheme 1, reaction (1)), in a way similar to homogeneous ruthenium systems.<sup>99</sup> Highly polar solvents provide higher activities, which supports a heterolytic activation of dihydrogen. Similarly, the addition of small amounts of a base such as triethylamine ( $Et_3N$ ) promotes heterolytic  $H_2$  splitting and, as a result, a faster hydrogenation is observed (TOF = 120  $h^{-1}$ ). On the other hand, an increase of activity was also achieved by the use of acetic acid as an additive (TOF = 180  $h^{-1}$ ). The authors suggested that the acid may protonate the N-pyridyl atoms, which would facilitate the interaction



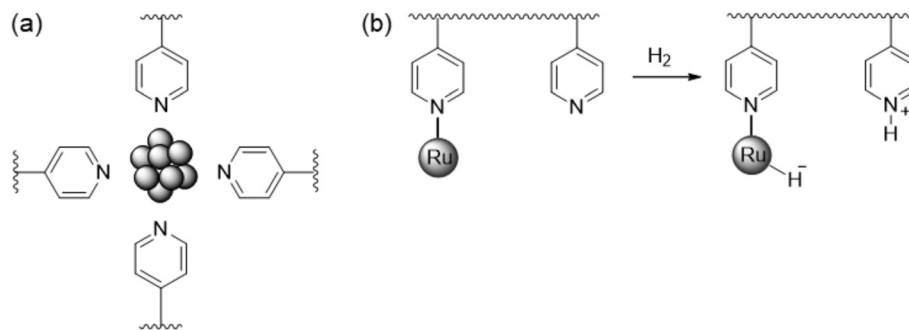


Fig. 7 (a) Representative scheme of RuNPs stabilised by poly(4-vinylpyridine). (b) Possible heterolytic H<sub>2</sub> cleavage on PVPy-stabilised RuNPs.<sup>98</sup>

with the substrate molecules by hydrogen bonding. However, given that the well-known ability of Ru to activate H<sub>2</sub> by itself and that RuNPs prepared by reduction with NaBH<sub>4</sub> or H<sub>2</sub> usually have a layer of hydrides,<sup>100</sup> one might wonder if another hydrogenation mechanism might be operative.

In a following study,<sup>101</sup> PVPy-stabilised RuNPs of 3.1 nm were applied as a hydrogenation catalyst of arenes (TOF = 82 h<sup>-1</sup> for toluene at 10 bar H<sub>2</sub>, 120 °C, THF, substrate/Ru = 100) and N-heteroaromatic compounds (TOF = 66 h<sup>-1</sup> for quinoline at 30 bar H<sub>2</sub>, 120 °C, methanol, substrate/Ru = 84). The system is highly active and maximal TOFs of 2000 h<sup>-1</sup> and 150 h<sup>-1</sup> for toluene and quinoline, respectively, were achieved at 50 bar H<sub>2</sub> and 150 °C. Powder XRD and XPS analyses showed that the RuNPs are mainly in the Ru<sup>0</sup> state, while a series of experiments provided insights into the catalytic reaction mechanism. Interestingly, two types of active sites that produce two different hydrogenation pathways were identified: one associated with a polar mechanism for C=N bond hydrogenation (A), and the other associated with a non-polar mechanism for the common hydrogenation of the aromatic ring (B). An increase in the catalytic activity in the reduction of quinoline was observed with an increase in the solvent polarity, which points to an HCD for the hydrogenation of N-heteroaromatics. This was supported by the results obtained after adding 10 eq. of an external base (Et<sub>3</sub>N) or acid (HBF<sub>4</sub>) to the reaction mixture, which led to an enhancement of the activity from 66 h<sup>-1</sup> up to 120 h<sup>-1</sup> and 180 h<sup>-1</sup>, respectively. The heterolytic H<sub>2</sub> activation would occur through a cooperative effect in which the N atoms of the polymer operate as a Lewis base and neighbouring ruthenium atoms act as a Lewis acid (Fig. 8).

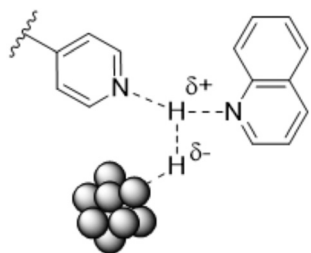
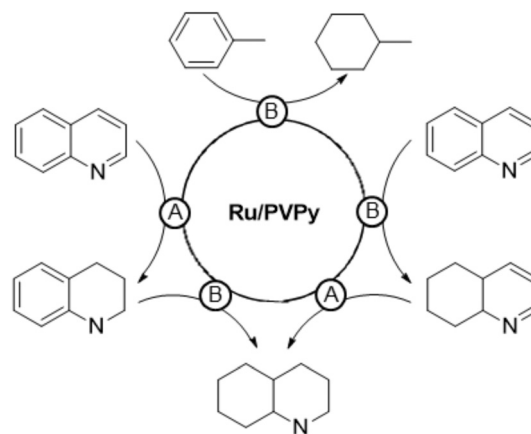


Fig. 8 Heterolytic cleavage of H<sub>2</sub> mediated by a RuNPs/PVPy system.

In addition, hydrogen bonding between N atoms of the substrate and pyridine groups of the support (or the N atom of Et<sub>3</sub>N) would favour the protonation of the former. As in the previous case,<sup>98</sup> the acid would promote the protonation of N-pyridyl atoms of the polymer or substrate, thus leading to the same hydrogen bonded intermediate (Fig. 8). On the other hand, no effect of solvent polarity or addition of an external base was observed in the hydrogenation of toluene, which suggests a homolytic cleavage of dihydrogen for the reduction of arenes.

Substrate competition experiments with quinoline and toluene supported the existence of these two hydrogenation pathways taking place on different active sites (Scheme 2). Toluene is hydrogenated only after quinoline is fully reduced to decahydroquinoline, which indicates that the heterocyclic ring is hydrogenated at one type of active site (type A), while the carbocyclic ring of quinoline is reduced at another site (type B). Thus, toluene is hydrogenated once reduction of quinoline finishes and type B sites are unblocked. Finally, selective poisoning experiments with thiophene provided further evidence of this dual hydrogenation mechanism. This compound inhibits the hydrogenation of both toluene and the carbocyclic ring of quinoline, suggesting that the thiophene molecule selectively blocks type B active sites.



Scheme 2 Proposed hydrogenation pathways through two different types of active sites.<sup>101</sup>

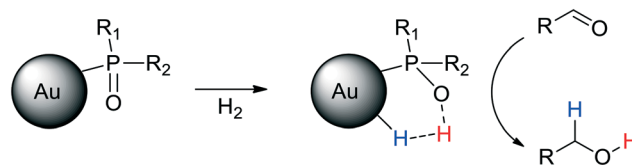




## 2.2 MNPs stabilized by ligands

Different strategies have been developed to achieve a heterolytic splitting of dihydrogen in MNP catalysis. However, in comparison with supported MNPs, there is little precedent for HCD with *homogeneous* ligand-stabilized nanoparticles dispersed in solution. A successful approach developed by van Leeuwen and coworkers entails the use of ligands that act both as a stabilising agent for MNPs and a heterolytic activator for H<sub>2</sub> through a cooperative effect with a neighbouring metal atom. Secondary phosphine oxides (SPOs) were employed to stabilise nanoparticles with applications as hydrogenation catalysts, in such a way that SPO ligands and surface metal atoms act cooperatively in HCD, which favours the selective reduction of polarized unsaturated groups. Indeed, RuNPs stabilized by SPOs showed an increase in selectivity towards the carbonyl group in the hydrogenation of acetophenone in comparison with a classic tertiary monophosphine (from 26 to 47%). The participation of P=O moieties in H<sub>2</sub>/D<sub>2</sub> exchange reactions supported the heterolytic H<sub>2</sub> splitting mediated by surface ruthenium atoms and P=O groups.<sup>81</sup> Next, these authors described the synthesis of gold nanoparticles (AuNPs) ligated by a specific SPO, *tert*-butyl(naphthalen-1-yl)phosphine oxide.<sup>102</sup> The SPO-ligated AuNPs were exceptionally selective towards the carbonyl functionality in the hydrogenation of a wide range of substituted aldehydes (Scheme 3).

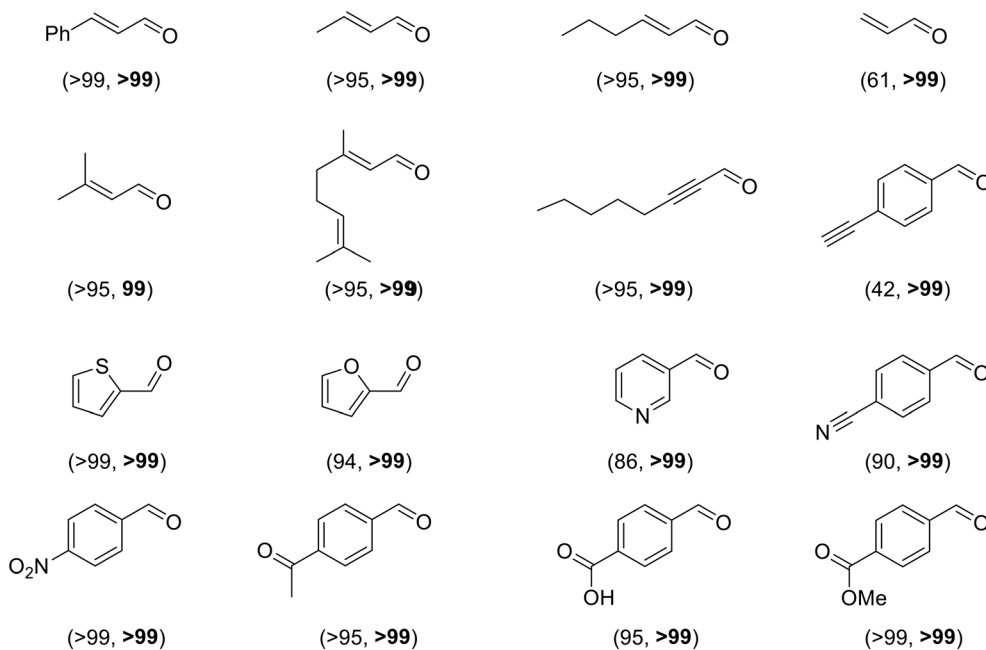
Several control experiments demonstrated that the SPO ligand plays a crucial role in the catalytic process, thus supporting a hydrogenation mechanism through an HCD in which the oxygen atom of the SPO acts as a Lewis base and binds the proton, H<sup>+</sup>, whereas a neighbouring gold atom operates as a Lewis acid and takes the hydride, H<sup>−</sup> (Scheme 4).



**Scheme 4** Heterolytic splitting of H<sub>2</sub> by SPO-ligated AuNPs and the following hydrogenation of aldehydes.

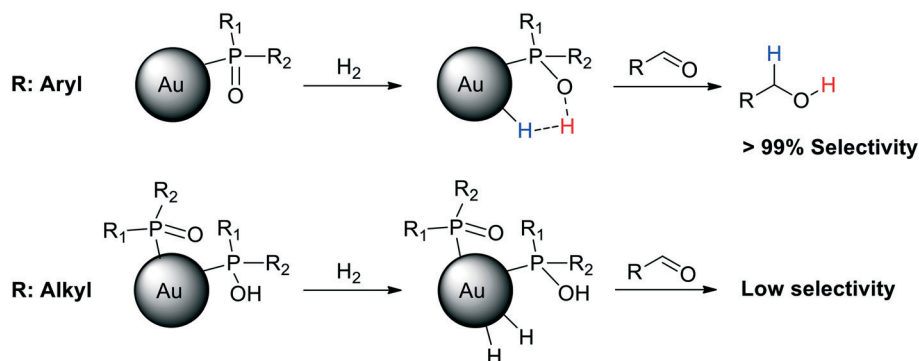
A subsequent study revealed that the degree of polarization of the P=O bond is the key for the HCD, as was proved by cross polarization magic angle spinning (CP-MAS) NMR spectroscopy.<sup>103</sup> To this end, a series of AuNPs ligated by different SPOs were synthesized and thoroughly characterized, showing important differences in the morphology and catalytic properties depending on the substituents in the SPO. CP-MAS NMR demonstrated that AuNPs stabilised with aryl SPOs exhibit a strong polarity of the P=O bond, thus favouring the HCD. These nanoparticles present Au(I) atoms and SPO anions at the surface and show very high selectivity in aldehyde hydrogenation (Scheme 3). On the other hand, AuNPs ligated by aliphatic phosphine oxides display low polarity in the P=O bond, which hampers the heterolytic H<sub>2</sub> activation. These alkyl-stabilised AuNPs have both Au(I) and Au(0) atoms at the NP surface, and also contain POH species. Consequently, they display active sites of different nature, leading to lower activity and selectivity in the hydrogenation of substituted aldehydes (Scheme 5).

This heterolytic hydrogenation mechanism by SPO-stabilised AuNPs was further supported through theoretical calculations.<sup>82</sup> A cooperative effect was identified at the AuNP-



**Scheme 3** Selective hydrogenation of substituted aldehydes catalysed by SPO-ligated AuNPs (conversion, selectivity). Reaction conditions: AuNPs (0.01 mmol Au), substrate (0.5 mmol), THF/hexane (5 mL), 18 hours, 40–60 °C, 40 bar H<sub>2</sub>.<sup>102</sup>





**Scheme 5** AuNPs ligated by aryl SPOs exhibit a strong polarity of the P=O bond, which favours the HCD and thus the selective hydrogenation of aldehydes. AuNPs ligated by alkyl SPOs show a low polarity of the P=O bond, which hinders the heterolytic H<sub>2</sub> activation. These AuNPs display low selectivity in aldehyde hydrogenation.<sup>103</sup>

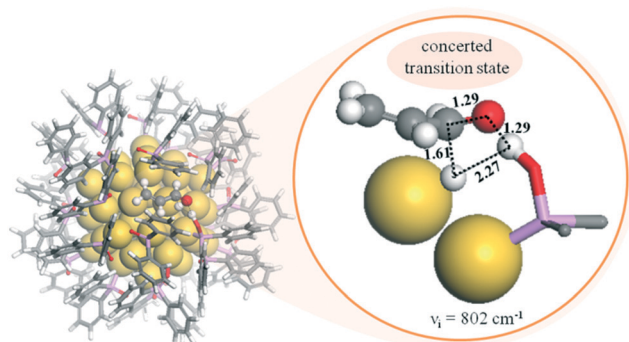
SPO interface by DFT studies, which acts as a frustrated Lewis pair (FLP). This FLP favours the heterolytic H<sub>2</sub> splitting and its addition to the C=O bond of acrolein by a concerted mechanism, leading to the selective formation of the unsaturated alcohol (Fig. 9). Crucially, the active sites available at the NP surface and the basicity difference between substrate and SPO ligand determine the activity of the AuNPs.

Following this approach, the same authors proposed a heterolytic H<sub>2</sub> activation in the hydrogenation of several unsaturated aldehydes catalysed by SPO-ligated IrNPs.<sup>104,105</sup> Indeed, very high chemoselectivities towards the carbonyl group were observed in the hydrogenation of cinnamaldehyde (99%), *p*-cyanobenzaldehyde (>99%) and 2-octynal (96% selectivity), suggesting a heterolytic splitting of H<sub>2</sub> mediated by the oxygen atom of the SPO and a neighbouring iridium atom. Interestingly, the catalytic activity of these NPs was compared with those of an analogous Ir-SPO complex and several supported IrNPs

previously described.<sup>105</sup> The SPO-stabilised IrNPs showed lower catalytic activity but higher robustness than the complex, providing high selectivities for aldehydes that poison the molecular catalyst, *e.g.* cyano- and alkyne-substituted aldehydes, whereas the supported IrNPs were generally less selective.

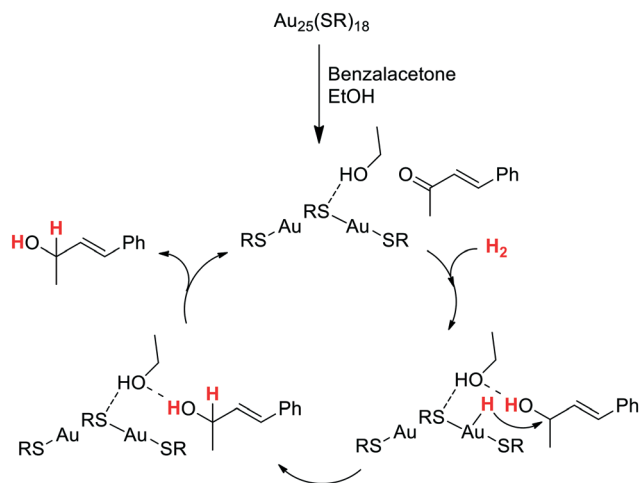
There are other reports describing a HCD carried out by ligand-stabilized MNPs *via* other mechanisms rather than a ligand-metal cooperation. In this context, very high selectivities were obtained by Jin *et al.* in the hydrogenation of  $\alpha,\beta$ -unsaturated carbonyl compounds catalyzed by the atomically precise Au<sub>25</sub>(SR)<sub>18</sub> cluster (SR = thiolate).<sup>106</sup> The authors suggested that the hydrogenation process occurs by a homolytic mechanism involving the adsorption of H<sub>2</sub> at low coordination surface gold atoms and activation of the C=O group on electron-rich gold sites located in the core. By contrast, Jiang and coworkers proposed that the H<sub>2</sub> activation by Au<sub>25</sub>(SR)<sub>18</sub> clusters takes place through a heterolytic splitting mechanism in which H<sub>2</sub> is cleaved by the C=O group of the substrate and a surface gold atom.<sup>107</sup> DFT studies analysed the possible pathways for the hydrogenation of benzalacetone catalysed by Au<sub>25</sub>(SR)<sub>18</sub>, discarding a direct dihydrogen activation on the cluster as that proposed by Jin. Indeed, the transition state corresponding to this path displays a barrier of 2.14 eV, which indicates that this route is very unlikely at the temperature at which the catalytic reaction was performed (0 °C).<sup>106</sup> Instead, the theoretical calculations revealed a first coadsorption of both benzalacetone and dihydrogen on the Au<sub>25</sub>(SR)<sub>18</sub> cluster, and a subsequent HCD through the cooperation of a Au atom of the cluster and the carbonyl oxygen atom of benzalacetone (Scheme 6). The study also shows that the process is assisted by one molecule of ethanol (solvent), which forms a hydrogen bond with the OH group of the partially hydrogenated reactant. Then, the hydride species generated at a Au atom is transferred to this intermediate, leading to the selective formation of the unsaturated alcohol product.

DFT calculations are in good agreement with the 100% selectivity observed at 0 °C. Indeed, the energy profile shows



**Fig. 9** Left: DFT adsorption configuration of 27 Ph<sub>2</sub>PO (SPO) ligands on a Au<sub>27</sub> NP. Right: Structure, distances (Å), and imaginary frequency (cm<sup>-1</sup>) determined for the transition state corresponding to the concerted mechanism in the selective hydrogenation of the C=O bond of acrolein on a SPO-ligated AuNP. The inset displays the formation of the four-membered ring and the distances. Yellow spheres: gold atoms. Stick models: adsorbed Ph<sub>2</sub>PO ligands. Color codes: C, grey; O, red; H, white; P, pink. Reprinted with permission from ref. 82. Copyright 2017. American Chemical Society.

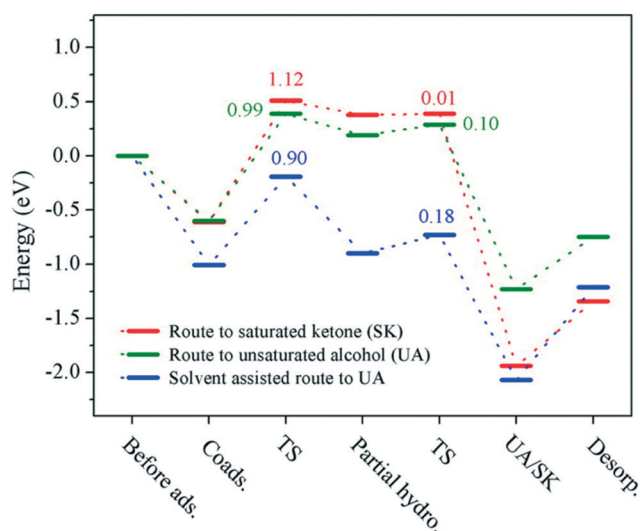




**Scheme 6** Proposed catalytic cycle for the chemoselective hydrogenation of benzalacetone catalyzed by the  $\text{Au}_{25}(\text{SR})_{18}$  cluster. The cluster is represented as  $\text{RS-Au-SR-Au-SR}$ .<sup>107</sup>

an activation energy of 0.90 eV for the ethanol-assisted process, while the barriers in the solvent-free route are 0.99 and 1.12 eV for the hydrogenation of  $\text{C}=\text{O}$  and  $\text{C}=\text{C}$  bonds, respectively (Fig. 10).

The use of a base as an additive is a very successful strategy to achieve a heterolytic  $\text{H}_2$  splitting in MNP catalysis.

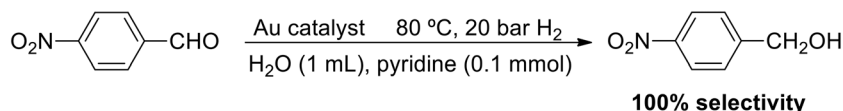


**Fig. 10** Energy profiles for the chemoselective hydrogenation of benzalacetone to the corresponding saturated ketone (red) and unsaturated alcohol (UA) through solvent-free (green) and solvent-assisted (blue) pathways. Reprinted with permission from ref. 107. Copyright 2015. American Chemical Society.

In line with this, Jin and coworkers proposed a heterolytic mechanism in the selective hydrogenation of 4-nitrobenzaldehyde catalysed by gold nanoparticles and nanorods stabilised with citrate and CTABr, respectively (Scheme 7).<sup>108</sup> Both Au catalysts are very selective, but the nanorods are more active than the NPs and allow an efficient recycling by centrifugation. 100% selectivity towards 4-nitrobenzyl alcohol was observed at 80 °C and 20 bar  $\text{H}_2$  in a transformation that requires the additional use of pyridine. In fact, no reaction was observed in the absence of a base. It was suggested that pyridine binds the proton of  $\text{H}_2$  and forms a  $\text{PyH}^+$  pyridinium species, whereas the gold catalyst abstracts the corresponding hydride,  $\text{H}^-$ . These  $\text{PyH}^+$  and  $\text{H}^-$  would also favour the adsorption of the substrate through the aldehyde and nitro groups ( $-\text{CHO}$  and  $-\text{NO}_2$ ).

Previously, Jin described the same process under analogous conditions ( $\text{H}_2\text{O}$  as a solvent, 80 °C, 20 bar  $\text{H}_2$ , 0.1 mmol pyridine) catalysed by water-soluble thiolate-stabilized gold clusters ligated by glutathione ( $\text{H-SG}$ ) and captopril (Capt), such as  $\text{Au}_{15}(\text{SG})_{13}$ ,  $\text{Au}_{18}(\text{SG})_{14}$ ,  $\text{Au}_{25}(\text{SG})_{18}$ ,  $\text{Au}_{25}(\text{Capt})_{18}$ , and  $\text{Au}_{38}(\text{SG})_{24}$ .<sup>109</sup> These atomically precise clusters provided complete selectivity towards the nitrobenzyl alcohol product, for which DFT studies revealed that both  $-\text{NO}_2$  and  $-\text{CHO}$  functionalities interact with the  $\text{S-Au-S}$  surface of the cluster. In parallel, the cluster would activate the  $\text{H}_2$  molecule with the aid of pyridine. There is no mention of heterolytic  $\text{H}_2$  splitting but the prerequisite of pyridine points toward a base-assisted heterolytic process.

In a further step,  $\text{CeO}_2$ -supported  $\text{Au}_{25}(\text{SR})_{18}$  clusters were employed as a catalyst for aldehyde hydrogenation.<sup>110</sup> Interestingly, the presence of a Lewis acid in the reaction medium such as  $\text{Cu}^+$ ,  $\text{Cu}^{2+}$ ,  $\text{Ni}^{2+}$  and  $\text{Co}^{2+}$  enhances the catalytic activity, while the use of a base ( $\text{NH}_3$  or pyridine) is also required. Once more, 100% selectivity towards the alcohol product was obtained in the reduction of 4-nitrobenzaldehyde, among other substrates. UV-vis spectroscopy, MALDI and ESI-MS showed the generation of new cluster  $\text{Au}_{25-n}(\text{SR})_{18-n}$  ( $n = 1-4$ ) species, and DFT calculations elucidated the structure and mechanism for the speciation of  $\text{Au}_{24}(\text{SR})_{17}$ , which involves the abstraction of a  $\text{Au-SR}$  fragment from the  $\text{Au}_{25}(\text{SR})_{18}$  cluster with the participation of a Lewis acid. As a result, the  $\text{Au}_{24}(\text{SR})_{17}$  species may present different open metal sites for adsorption of substrates: a gold atom of the cluster core ( $\text{Au1}$ ), a gold atom on a staple motif ( $\text{Au2}$ ), and the metal ion employed as a Lewis acid ( $\text{M1}$ ). The DFT studies allowed the authors to propose a hydrogenation mechanism for each of these active sites, all of them involving a base-assisted heterolytic  $\text{H}_2$  cleavage (Fig. 11). In this way, higher conversions are



**Scheme 7** Chemoselective hydrogenation of 4-nitrobenzaldehyde catalysed by AuNPs and Au nanorods.



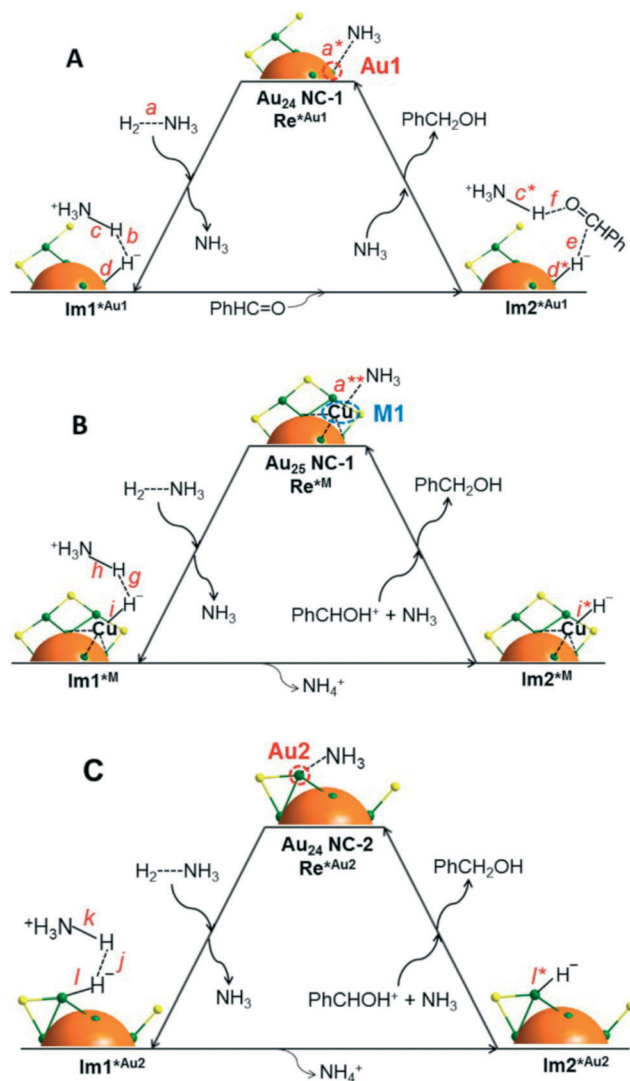


Fig. 11 Proposed catalytic mechanism for benzaldehyde hydrogenation in the presence of  $\text{NH}_3$  on (A) a gold atom of the cluster core (Au1), (B) the metal ion employed as a Lewis acid (M1), and (C) a gold atom on a staple motif (Au2). Colours: Au, green; S, yellow. Reprinted with permission from ref. 110. Copyright 2015. American Chemical Society.

observed with stronger bases such as  $\text{NH}_3$ , which activates the H–H bond and abstracts a  $\text{H}^+$  to generate  $\text{NH}_4^+$ .

Recently, the same authors described the heterolytic hydrogenation of nitriles mediated by thiolate-stabilised nickel  $\text{Ni}_6(\text{SR})_{12}$  clusters. The clusters consist of a double crown in which the Ni atoms form a hexagonal ring, while the twelve S atoms are disposed as bridging Ni–S–Ni below and above the ring plane (Fig. 12).<sup>111</sup> Consequently, the Ni sites are shielded by these sulphur atoms, which limits the access of substrates. Indeed, very low conversions and complete selectivities towards the primary amine were obtained in the hydrogenation of 4-cyanotetrahydropyran. Interestingly, the use of  $\text{NH}_3$  dramatically enhances the catalytic activity. DFT calculations indicate that the  $\text{Ni}_6(\text{SR})_{12}$  cluster is activated through the cleavage and subsequent

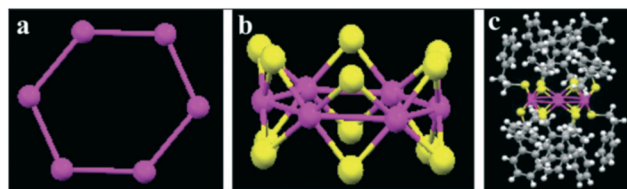


Fig. 12 Structure of the  $\text{Ni}_6(\text{SR})_{12}$  cluster (R:  $\text{SC}_2\text{H}_4\text{Ph}$ ): (a)  $\text{Ni}_6$  structure. (b) Framework of  $\text{Ni}_6$  plus 12 S atoms. (c) Crystal structure. Colors: Ni, magenta; S, yellow; C, gray; H, white. Reproduced from ref. 111 with permission from The Royal Society of Chemistry.

insertion of  $\text{H}_2$  into a Ni–S–Ni unit, leading to the formation of a hetero-bridge Ni–H–H–S chain, which undergoes heterolytic  $\text{H}_2$  cleavage. In this case,  $\text{NH}_3$  does not act as a heterolytic activator for dihydrogen. Instead, this base would play a crucial role by preventing a recombination reaction that regenerates the cluster structure and leads to the deactivation of the catalyst. The theoretical studies reveal that  $\text{NH}_3$  is adsorbed on the double crown and partially cleaves the Ni–S–Ni bridge, thus maintaining two Ni sites (Ni– $\text{NH}_3$  and Ni–S) open.

Finally, the work of Häkkinen and Zheng is worth noting, in which atomically precise copper-hydride nanoclusters ( $\text{CuNCs}$ ) ligated by thiolates were employed as a catalyst for ketone hydrogenation.<sup>112</sup> The  $[\text{Cu}_{25}\text{H}_{10}(\text{SR})_{18}]^{3-}$   $\text{CuNC}$  contains 10 hydrides that serve as the hydrogen source for the catalytic process. DFT calculations using formaldehyde as a model substrate reveal that this molecule takes a hydride from the cluster to generate a  $[\text{Cu}_{25}\text{H}_9\text{OCH}_3(\text{SR})_{18}]^{3-}$  intermediate containing an alkoxy group, which then reacts with dihydrogen through two possible catalytic pathways (Fig. 13). In the first route,  $\text{H}_2$  is heterolytically cleaved in the proximity of the Cu–O bond to form a proton that binds to the alkoxy group and affords the alcohol product, whereas the corresponding hydride species restores the cluster. For the second pathway, it is proposed that the alkoxy intermediate could bind a second hydrogen atom and form the product. Subsequently, the dissociation of  $\text{H}_2$  leads to the replenishing of these two hydride vacancies and thus the regeneration of the catalyst. Theoretical studies indicate that both routes are kinetically and thermodynamically feasible, and that there is no preference for a particular one. However, additional experimental proofs and the study of the solvent-accessible surface area suggest that the second route is the preferred pathway.

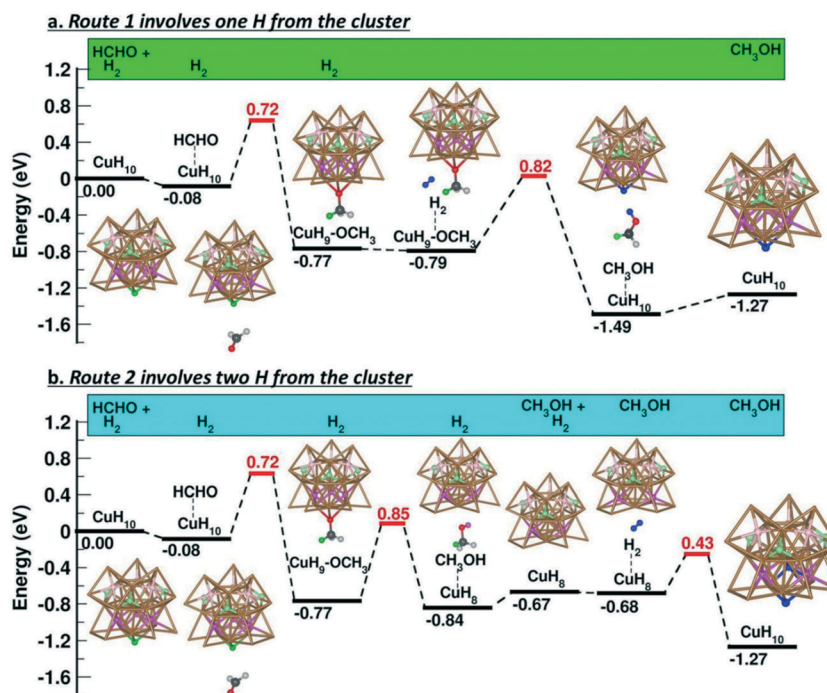
### 3. Supported MNPs

#### 3.1 Ligand-free MNPs

Supported transition metal catalysts are of great importance for the industrial synthesis of numerous chemicals.<sup>113,114</sup> These industrial catalysts are often based on expensive metals well-dispersed on cheap, high-surface area, porous supports. A high dispersion is convenient since most of the existing metals on the surface are accessible for the substrates and ready for catalysis.<sup>115,116</sup> In this respect, supported metal







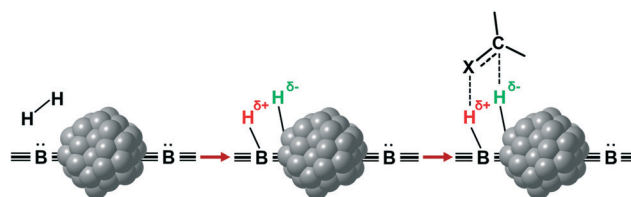
**Fig. 13** Energy profiles of pathways 1 and 2 for the hydrogenation of formaldehyde catalysed by  $[\text{Cu}_{25}\text{H}_{10}(\text{SR})_{18}]^{3-}$ . Colours: frame of the copper cluster, brown; hydrides, green, pale green, pink and pale pink; C (HCHO), dark grey; H (HCHO), light grey; O (HCHO), red; H ( $\text{H}_2$ ), blue. Reprinted with permission from ref. 112. Copyright 2019. American Chemical Society. <https://pubs.acs.org/doi/10.1021/acsnano.9b02052>. Further permissions related to the material excerpted should be directed to the ACS.

nanoparticles (MNPs) have been demonstrated to be a promising type of green catalyst. Their large surface area, together with their higher stability and recyclability compared to MNPs in solution, makes supported MNPs ideal catalysts for industrial applications.<sup>117</sup> In fact, supports not only increase the stability and recyclability of MNPs, but they can also be used to modify the MNPs' electronic/steric properties, and therefore alter the catalytic properties.

As was mentioned in the Introduction, selective hydrogenation of polar functional groups ( $\text{C}=\text{N}$ ,  $\text{C}=\text{S}$  or  $\text{C}=\text{O}$ ) with homogeneous catalysts is in several cases due to a metal ligand cooperation, which dissociates  $\text{H}_2$  in a heterolytic way (forming heteroatom- $\text{H}^+$ /metal- $\text{H}^-$  pairs). On the other hand, it has been established that supported-MNPs activate hydrogen in a homolytic way, forming sometimes  $\text{O}-\text{H}^+$  on the support through hydrogen spillover.<sup>61</sup> In addition to this, a cooperative activation of  $\text{H}_2$  can be performed by metal-basic site interfaces. Indeed, well-defined nanostructured materials based on MNPs immobilized on basic supports have been demonstrated to be promising catalysts for selective hydrogenation reactions. Basic functional groups of the support in close proximity to active metal sites promote the heterolytic  $\text{H}_2$  cleavage and thereby boost the hydrogenation of some substrates, such as heteroaromatics, aldehydes, ketones or carboxylic acids. Here, MNPs and the basic centres of the supports work together in the dissociation of  $\text{H}_2$  through a heterolytic mechanism (Fig. 5). In addition, the hydrogen transfer occurs through an outer sphere mechanism in which there is no

direct interaction between the MNP surface and the substrate (Fig. 14), and prevents poisoning of the catalyst.

The function of supports is generally to avoid MNP aggregation and to facilitate the recyclability of the catalysts. However, the number of reports in which the support plays an active role in the hydrogenation mechanism is growing fast. Indeed, there is a great variety of examples in the literature about heterolytic  $\text{H}_2$  activation by supported MNPs. Following previous work with polymer-stabilized MNPs (see section 2.1),<sup>98,101</sup> Sanchez-Delgado *et al.* reported one of the first examples in 2012 by using Pd nanoparticles supported on MgO (Pd/MgO) for the hydrogenation of alkenes, quinolines and biodiesel.<sup>118</sup> The high activity of Pd/MgO in comparison with commercial Pd/SiO<sub>2</sub> and Pd/Al<sub>2</sub>O<sub>3</sub> was explained by an ionic hydrogenation mechanism promoted by the combination of MNPs and the basic sites of the support as shown in Fig. 5. Shortly thereafter, the same research group reported the analogous catalyst containing Ru NPs (Ru/MgO), which is



**Fig. 14** MNP and basic support working in synergy for heterolytic  $\text{H}_2$  dissociation and the subsequent hydrogenation of  $\text{R}_2\text{C}=\text{X}$  (where  $\text{X} = \text{N}, \text{S}, \text{O}$ ) through an outer sphere mechanism.



capable of hydrogenating heteroaromatics without catalyst poisoning, normally observed in the presence of nitrogen- and sulfur-containing species.<sup>119</sup> This tolerance to poisoning is probably due to the surface ionic hydrogenation pathways shown in Fig. 14. Indeed, after mechanistic studies, they proposed a dual homolytic/heterolytic mechanism, in which two kinds of active sites operate simultaneously: i) the surface oxygen atoms of the support (basic centres) next to metal active sites (Ru atoms) for HCD (site A, Fig. 15); ii) Ru atoms far away from the basic support act as common metallic sites splitting  $H_2$  homolytically (site B, Fig. 15). All this explains the high activity and stability of Ru/MgO in the hydrogenation of N- and S-heteroaromatics. This novel mechanism was verified through a competition test between quinolone and toluene, where the hydrogenation of toluene was basically blocked until most of the quinolone was hydrogenated. In addition, an inhibition experiment with thiophene confirmed that after blocking the Ru surface with the heteroarene, the hydrogenation of toluene was totally suppressed, but instead the N-heterocyclic ring of quinolone was fully hydrogenated. A similar dual homolytic/heterolytic hydrogenation mechanism was proposed for Rh nanoparticles immobilized on basic supports, such as MgO, CaO or SrO.<sup>120</sup> Here, it was observed that the hydrogenation rate of N-heterocycles increases with the basicity of the support ( $Rh/MgO < Rh/CaO < Rh/SrO$ ).

Jagadeesh in collaboration with Beller reported a series of Co and Fe heterogeneous catalysts prepared by pyrolysis of molecular complexes with nitrogen ligands immobilized on activated carbon.<sup>121,122</sup> The resulting catalysts, non-noble MNPs supported on N-doped graphene, are able to activate dihydrogen by HCD.<sup>123</sup> More specifically, N-doped graphene coated Co- and Fe catalysts show a high activity/selectivity in the hydrogenation of nitroarenes under mild conditions (70 °C and 20 bar). Here, the basicity of the graphene support is essential for the HCD. Zhang *et al.* recently described another interesting study about HCD in MNPs supported on a N-doped carbon material.<sup>96b</sup> More specifically, a new catalyst based on Pd NPs supported on nitrogen-doped porous carbon (Pd/NC-BT) gave a high activity in hydrodeoxygenation of a great number of carbonyl compounds, affording the corresponding alkanes. Here, again, the nitrogen atoms promote the HCD in cooperation with the Pd NPs. Fig. 16 shows the proposed

mechanism where first the  $H_2$  molecule and the carbonyl compound adsorb on the catalyst surface. Then, HCD generates  $N-H^+$  and  $Pd-H^-$  species, which react with the polarized carbonyl group, generating the corresponding alcohol. Finally, the hydrogenolysis of the alcohol group also takes place through the heterolytic cleavage of another  $H_2$  molecule. The same group reported a similar system, nitrogen-doped carbon supported Pd NPs, for the selective hydrogenation of quinolone derivatives into the corresponding 1,2,3,4-tetrahydroquinolines.<sup>96a</sup> Here, in the same way as in the previous case, the nitrogen atoms act as a Lewis base promoting the HCD. Rossi and collaborators proposed an analogous cooperative action between gold and nitrogen atoms for Au NPs encapsulated in nitrogen-doped carbon and supported on  $TiO_2$  (Au@N-doped carbon/ $TiO_2$ ).<sup>124</sup> Here, as well as in the previous examples, N atoms facilitate the heterolytic  $H_2$  cleavage, which also explains the high activity of this catalyst in the semihydrogenation of alkynes to alkenes.

Along the same line, Martínez-Prieto *et al.* recently proposed the same dual mechanism as Fig. 15 for  $H_2$  dissociation in the selective hydrogenation of palmitic acid into the corresponding alcohol (1-hexadecanol) catalysed by Ru NPs supported on reduced-graphene oxide doped with N (Ru/ $NH_2$ -rGO).<sup>125</sup> Here, the basic centres that promote HCD are the amino and pyrrolic groups present in the N-doped graphene (Fig. 17b). To distinguish between sites A and B, they used the hydrogenation of acetophenone as a model reaction (Fig. 17a). They proposed that the ketone groups of acetophenone are hydrogenated by site A, while the aromatic rings are hydrogenated on the Ru surface, site B. Comparing the initial hydrogenation rates of Ru NPs supported on reduced graphene oxide doped (Ru/ $NH_2$ -rGO) and non-doped with N (Ru/rGO), it was observed that Ru/ $NH_2$ -rGO hydrogenates the ketone much faster than the phenyl group. On the other hand, for Ru/rGO, which mainly contains site B, the hydrogenation of the ketone is slower than that of the aromatic ring (Fig. 17b and c). Therefore, they concluded that site A in Ru/ $NH_2$ -rGO enhances the hydrogenation of carbonyl groups in both substrates, acetophenone and palmitic acid.

In 2017, Hosono *et al.* reported a new heterogeneous catalyst based on Ru NPs supported on nanocages of  $12CaO \cdot 7Al_2O_3$  (Ru/HT-C12A7), which exhibits a high chemoselectivity in the hydrogenation of heteroarenes.<sup>126</sup> Ru/HT-C12A7 hydrogenates N- and O-heteroarenes with high yields in a solvent-free system. The high activity and stability were explained by the high percentage of strong basic sites of HT-C12A7 in close proximity to the Ru atoms, which promotes HCD (Fig. 18). In our opinion, this unique nanocage with many basic sites that facilitate the formation of  $H^+$  is a promising material for heterolytic  $H_2$  activation by supported MNPs.

Heterolytic dissociation of  $H_2$  on  $CeO_2$ -supported gold nanoparticles (Au/ $CeO_2$ ) was experimentally detected for the first time in 2010 by García, Corma *et al.* combining inelastic neutron scattering (INS) and FT-IR spectroscopy. Indeed, after treating Au/ $CeO_2$  with  $H_2$  at 150 °C, they observed that HCD takes place leading to a hydride coordinated to gold and a proton bonded to a ceria oxygen.<sup>75</sup> More specifically, they

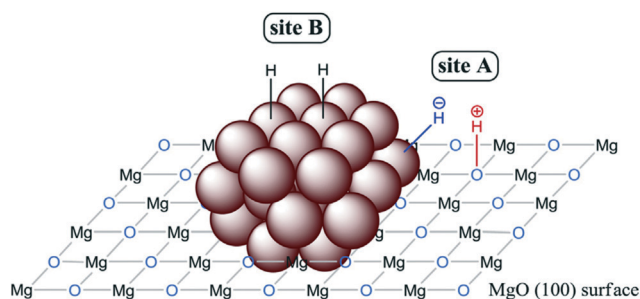


Fig. 15 Illustration of the active sites necessary for the dual homolytic/heterolytic hydrogenation mechanism proposed for Ru/MgO. Reprinted with permission from ref. 119. Copyright 2014. Elsevier.



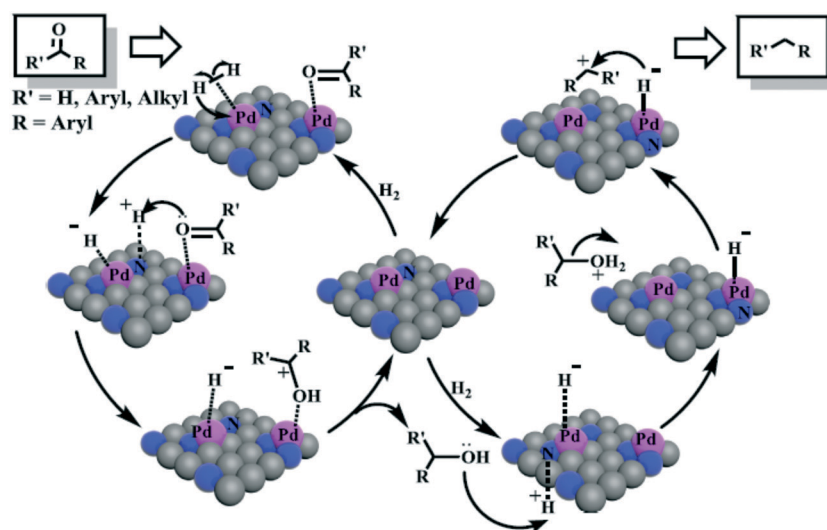


Fig. 16 Proposed mechanism for the hydrodeoxygenation of carbonyl compounds with Pd/NC-BT. Reprinted with permission from ref. 96b. Copyright 2018. Elsevier.

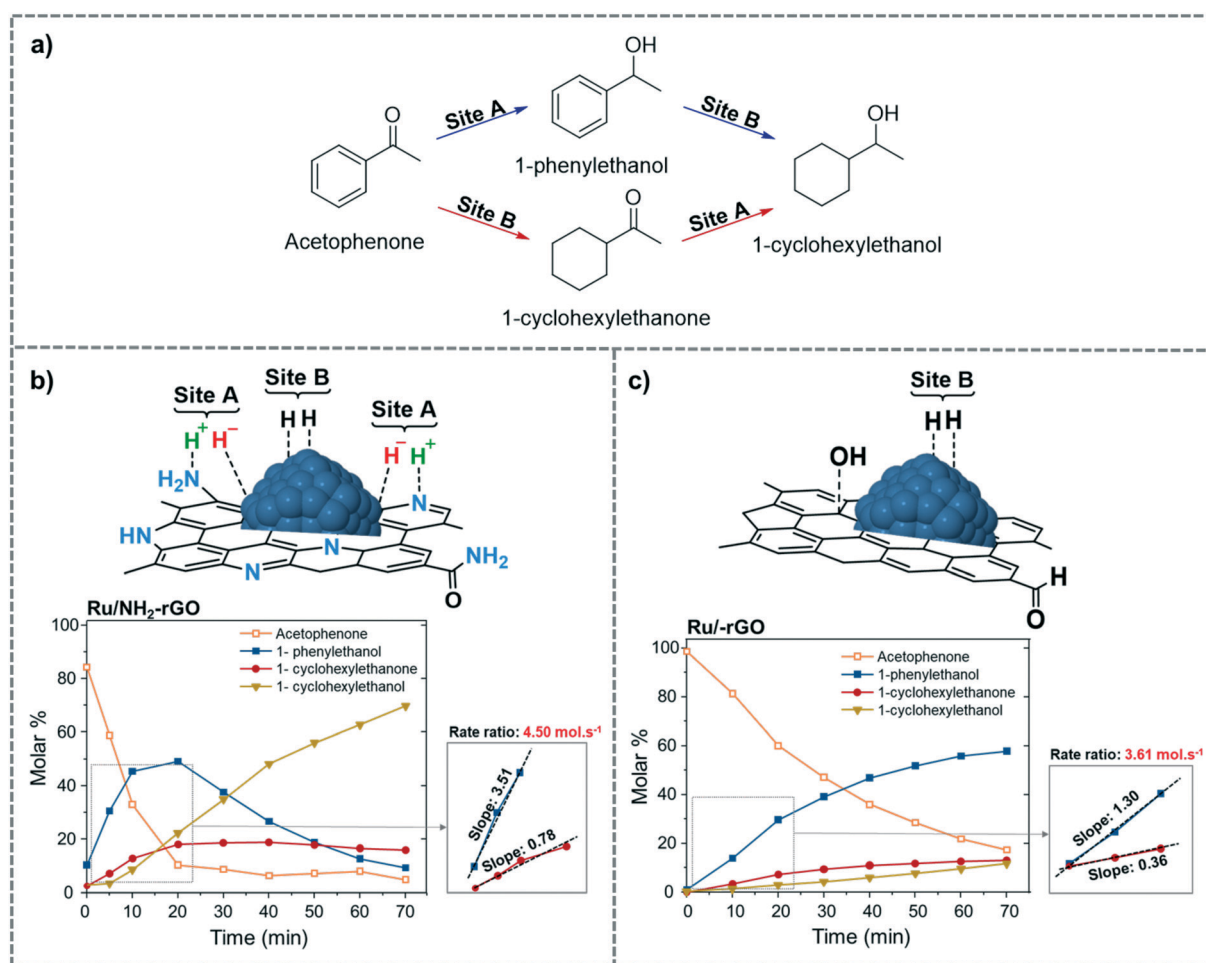


Fig. 17 Possible pathways for the hydrogenation of acetophenone to 1-cyclohexylethanol (a). Proposed active sites and time course of the product yield during the hydrogenation of acetophenone using (b) Ru/NH<sub>2</sub>-rGO and (c) Ru/rGO as catalysts. Adapted with permission from ref. 125. Copyright 2019. Elsevier.





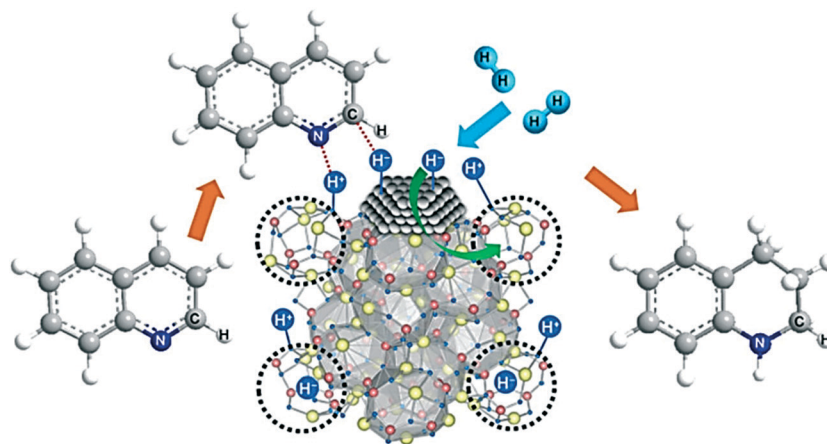


Fig. 18 Pathway proposed by Hosono *et al.* for the selective hydrogenation of heteroarenes over Ru/HT-C12A7. Reprinted with permission from ref. 126. Copyright 2017. Elsevier.

detected a new band at 400–600  $\text{cm}^{-1}$  by INS corresponding to bridging hydroxyl groups, which disappeared after exposure to  $\text{CO}_2$  or  $\text{O}_2$ . In addition, by FT-IR they observed the chemisorption of hydrogen on gold through the formation of an Au–H band, which also disappeared after treatment with  $\text{CO}_2$  or  $\text{O}_2$ . The assignments of these bands were confirmed by isotopic labelling experiments with  $\text{D}_2$  and  $^{18}\text{O}_2$ . With these results, they provided experimental evidence about the “direct” heterolytic  $\text{H}_2$  cleavage over immobilized MNPs, which is in contrast with the hitherto

commonly accepted reactivity of supported transition metal catalysts, homolytic cleavage followed by spillover.

HCD on core-gold/shell-ceria nanomaterials ( $\text{Au}@\text{CeO}_2$ ) was reported by Kaneda *et al.*<sup>127</sup> Here, the great number of interface sites between the gold core (5 nm) and the ceria shell (4 nm thickness) maximizes heterolytic  $\text{H}_2$  dissociation (Fig. 19a–c), and explains the high selectivity of this material in semi-hydrogenation reactions of alkynes, which possess a higher electrophilicity than alkenes. A couple of experiments were carried out to confirm this ionic  $\text{H}_2$  cleavage: i) the first

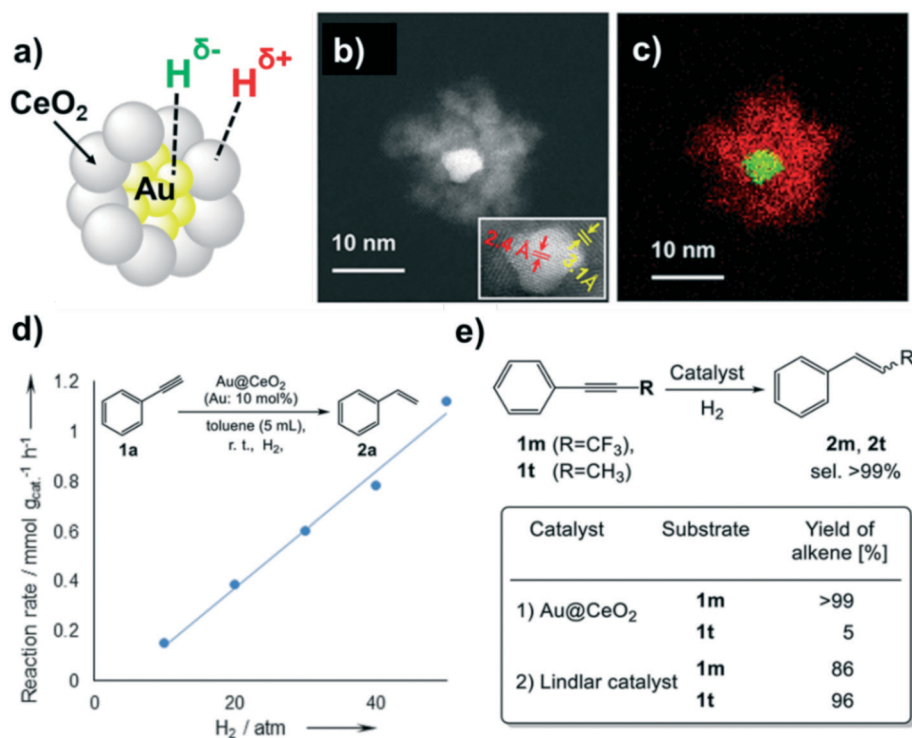


Fig. 19 (a) Illustration of  $\text{Au}@\text{CeO}_2$ . (b) HAADF-STEM image of  $\text{Au}@\text{CeO}_2$  (inset: HRTEM image showing the lattice spacing). (c) Elemental mapping of  $\text{Au}@\text{CeO}_2$  (Au: green, Ce: red). (d) Reaction rate vs.  $\text{H}_2$  pressure. (e) Hydrogenation of alkynes using  $\text{Au}@\text{CeO}_2$  or a Lindlar catalyst. Reprinted with permission from ref. 128. Copyright 2016. American Chemical Society.





order dependence of the reaction rate on  $H_2$  pressure in the semihydrogenation of phenylacetylene (Fig. 19d), which points to the participation of the polar hydrogen species in the hydrogenation mechanism; ii) the strong inductive effect on the increment of the reaction rate when using an acetylene with an electron-acceptor group ( $CF_3$ ) (Fig. 19e). The same nanomaterial but supported on hydrotalcite ( $Au@CeO_2/HT$ ) exhibits also a high selectivity in the hydrogenation of unsaturated aldehydes and ketones towards the corresponding unsaturated alcohols.<sup>128</sup> A similar selectivity due the above-mentioned  $H_2$  heterolytic cleavage was observed by the same group using the analogous silver material ( $Ag@CeO_2$ ) supported on ceria.<sup>129</sup>

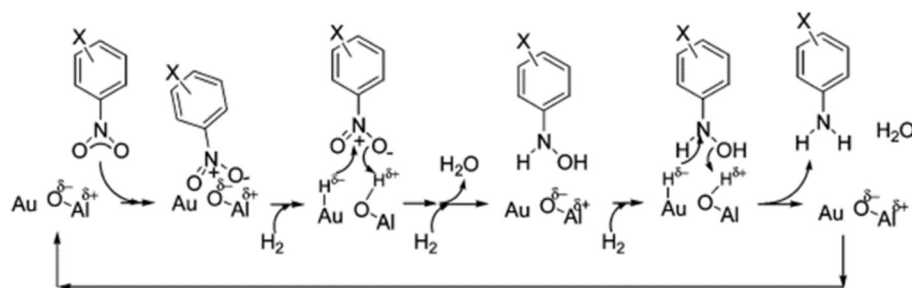
High levels of chemoselectivity in the hydrogenation of substituted nitroaromatics were also observed by Shimizu *et al.* by using  $Au/Al_2O_3$  as a catalyst.<sup>130</sup> More specifically, 2.5 nm Au NPs supported on  $Al_2O_3$  selective hydrogenate  $-NO_2$  in the presence of other reducible groups such as vinyl or carbonyl groups. The reason for this high selectivity is the cooperative work between the metal and basic sites at the nanoparticle-support interface, which heterolytically cleave  $H_2$ . The electrophilic  $H^{\delta+}$  and nucleophilic  $H^{\delta-}$  species generated at the metal-support interfaces are transferred to the polar nitro group, forming hydroxylamine, which is subsequently hydrogenated by a similar transfer mechanism to yield the final aniline (Scheme 8).

So far, we have observed that HCD by supported MNPs is highly dependent on the nature of the support. In addition, it can be also influenced by the solution surrounding the MNPs, as has been recently investigated by Dupont *et al.* More specifically, they observed that hydrogen activation proceeds, as expected, *via* heterolytic cleavage on a clean surface of Au NPs supported on  $\gamma-Al_2O_3$  ( $Au/\gamma-Al_2O_3$ ). However, after encapsulating these supported-Au nanoparticles in ionic liquids ( $IL-Au/\gamma-Al_2O_3$ ),  $H_2$  activation goes through a homolytic dissociation.<sup>131</sup> As a result, the two catalysts show totally different pathways in the selective hydrogenation of a wide scope of substrates.

Chandler *et al.*<sup>78</sup> provided experimental and theoretical evidence about HCD by supported metal nanoparticles. More specifically, based on DFT calculations with Au nanorods (Au NRs) supported on  $TiO_2$ , they suggested that  $Au-H^-$  and  $O-H^+$  species are generated. H/D isotopic exchange experiments and

kinetic studies supported the proposed mechanism. Apart from offering significant evidence for HCD on metal-support interfaces, this study also showed how water molecules block these interfaces for heterolytic  $H_2$  activation. By DFT calculations, Nie *et al.* investigated the influence of cluster size and metal-support interaction in the adsorption and dissociation of  $H_2$  over anatase  $TiO_2$  (101) and (001) surfaces.<sup>132</sup> The Au atoms that are active in HCD are neutral in charge and located on the edges and corners of the nanoparticle. In particular, HCD occurs through the transition state  $O^{2-}-H^+-H^-$ -Au on both lattices, (101) and (001) (Fig. 20a). Moreover, the existence of oxygen vacancies influences the stability of the adsorbed Au NPs and  $H_2$  molecules, but it does not affect significantly the energy barrier for  $H_2$  activation. Along the same lines are the studies of Larese *et al.* about the influence of the support on the hydrogenation of cinnamaldehyde to cinnamyl alcohol by using a series of  $Au/ZnO$  catalysts.<sup>133</sup> They prepared three different types of  $Au/ZnO$  catalysts: Au/rod-tetrapod ZnO, Au/porous ZnO, and Au/ZnO-CP synthesized by co-precipitation. Au/ZnO-CP, which presents the highest dispersion and smallest NP size, showed the best results (100% selectivity to cinnamyl alcohol at 94.9% conversion) (Fig. 20b). Therefore, it can be concluded that the Au-ZnO interaction is a key parameter for achieving a highly selective catalyst for  $C=O$  hydrogenation.

It is not always the support that provides the basic centres next to metal sites needed for heterolytic activation of  $H_2$ . In 2013, Tomishige and co-workers reported silica supported Ir NPs partially covered with  $ReO_x$  ( $Ir-ReO_x/SiO_2$ ) as a highly active catalyst for the chemoselective hydrogenation of unsaturated aldehydes into the corresponding alcohols.<sup>134</sup> Here, the basic sites are  $ReO_x$  species that work together with the Ir atoms in HCD. Later, the same group observed a similar selectivity and activity in the hydrogenation of  $\alpha,\beta$ -unsaturated ketones by using an Fe cation modified  $Ir/MgO$  catalyst [ $Ir/MgO^+Fe(NO_3)_3$ ], being one of the best examples of a heterogeneous catalyst that presents high selectivity (up to 90% at full conversion) towards the corresponding  $\alpha,\beta$ -unsaturated alcohol.<sup>135</sup> Here, the cation  $Fe^{2+}$  next to the anion  $O^{2-}$  forms the active sites for HCD. (Fig. 21a). Also, Tomishige *et al.* reported that the presence of  $Re^{n+}$  ( $Re^{3+}$  and  $Re^{4+}$ ) species on silica-supported  $RePd$  bimetallic catalysts ( $Re-Pd/SiO_2$ ) is responsible for HCD in the hydrogenation of stearic acid (Fig. 21b). On the other hand, the  $Pd(0)$  atoms



**Scheme 8** Mechanism proposed by Shimizu *et al.* for the selective hydrogenation of nitroaromatic compounds catalyzed by  $Au/Al_2O_3$ . Reprinted with permission from ref. 130. Copyright 2009. American Chemical Society.



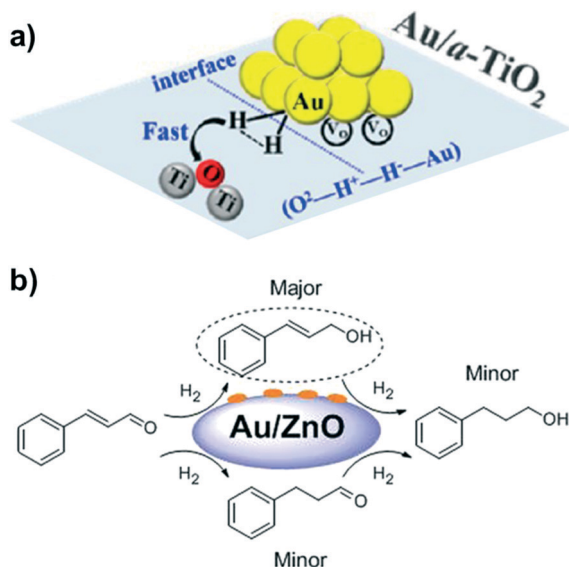


Fig. 20 (a) Adsorption and dissociation of H<sub>2</sub> at the Au/TiO<sub>2</sub> interface. (b) Selective hydrogenation of cinnamaldehyde to the corresponding unsaturated alcohol over ZnO-supported Au NPs. Reprinted with permission from ref. 132 and 133, respectively. Copyrights 2018 and 2015. American Chemical Society.

facilitate the adsorption of stearic acid at the metal surface, and promote the reduction and dispersion of Re<sup>n+</sup> species.<sup>136</sup>

In addition to metal oxide supports (MgO, CeO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, TiO<sub>2</sub> or ZnO), silica and carbon-based supports, other materials such as Fe<sub>3</sub>O<sub>4</sub> or layered double hydroxides (LDHs) have been reported for heterolytic H<sub>2</sub> activation by supported MNPs. For example, Pei, Qiao and co-workers reported gold nanoparticles on flower-like magnetite (flower-like Au/Fe<sub>3</sub>O<sub>4</sub>) as an efficient catalyst for selective hydrogenation of crotonaldehyde to the corresponding alcohol.<sup>137</sup> This catalyst (Fig. 22a), prepared by deposition of Au on hematite and a subsequent reduction, presents a stronger metal-support interaction and a higher activity/selectivity than the comparable catalyst AuNPs

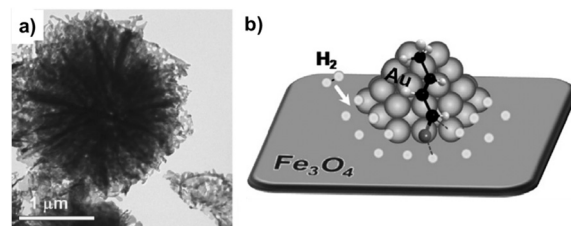


Fig. 22 (a) TEM image of flower-like Au/Fe<sub>3</sub>O<sub>4</sub>. (b) Illustration of the selective hydrogenation of crotonaldehyde to crotyl alcohol at the interface of flower-like Au/Fe<sub>3</sub>O<sub>4</sub> catalyst. O: gray, H: white, and C: black. Reprinted with permission from ref. 137. Copyright 2011. Elsevier.

supported on Fe<sub>3</sub>O<sub>4</sub>, *e.g.*, with flower-like Au/Fe<sub>3</sub>O<sub>4</sub> the selectivity is higher than 76% at full conversion. The proposed mechanism involves HCD on the interface sites Au-O-Fe and the interaction of crotonaldehyde with the hydroxyl groups on magnetite next to gold atoms. The proton of the hydroxyl group attacks the oxygen of the carbonyl, and the H-atom with the hydride character from HCD (H<sup>δ-</sup>) is added to the carbon in the α position. Finally, the other hydrogen atom, partially positively charged (H<sup>δ+</sup>), replaces the hydrogen of the hydroxyl group, closing the catalytic cycle (Fig. 22b). Due to its magnetic character, this catalyst presents exceptional recovery and recycling properties.

In 2011, Kaneda *et al.* described a cooperative effect between Au NPs and basic sites of layered double hydroxides (LDHs) for HCD. In particular, they selectively deoxygenate epoxides into alkenes (99% selectivity) by using a hydrotalcite-supported gold catalyst (Au/HT).<sup>138</sup> The active species, Au-H<sup>δ-</sup> and HT-H<sup>δ+</sup> (Scheme 9a), were identified by FT-IR after treatment of the catalyst with molecular hydrogen. The same catalytic system was used for the formation of terminal alkenes through the selective hydrogenolysis of allylic carbonates.<sup>139</sup> Here, the proposed mechanism also involves the formation of Au-H<sup>δ-</sup> and HT-H<sup>δ+</sup> species at the Au-HT interface. Once the active hydrogen species are formed, the allylic carbonate reacts with HT-H<sup>δ+</sup> to form the Au-allyl intermediate, which is finally attacked by the hydride (Au-H<sup>δ-</sup>) to yield the terminal alkene (Scheme 9b).

A more recent example concerning HCD at the interface of metal and basic sites of a layered double hydroxide was recently reported by Xiao and coworkers.<sup>140</sup> The formation of polar hydrogen species on this Zn-Ti LDH supported PdAu catalyst (Zn-Ti LDHs) is responsible for the selective hydrogenation of phenylacetylene to styrene. The PdAu alloy nanoparticles are placed between the layers of LDHs, and strongly interact with them. This strong interaction facilitates the heterolytic hydrogenation, and makes Zn-Ti LDH an active and selective catalyst in the semihydrogenation of phenylacetylene (90% selectivity at full conversion), suppressing the hydrogenation of styrene after full conversion. In addition, this material was presented as a promising catalyst for industrial applications due to its high stability and effective recyclability.

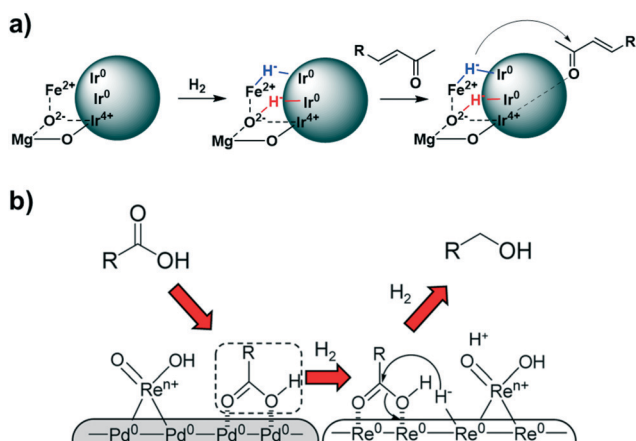
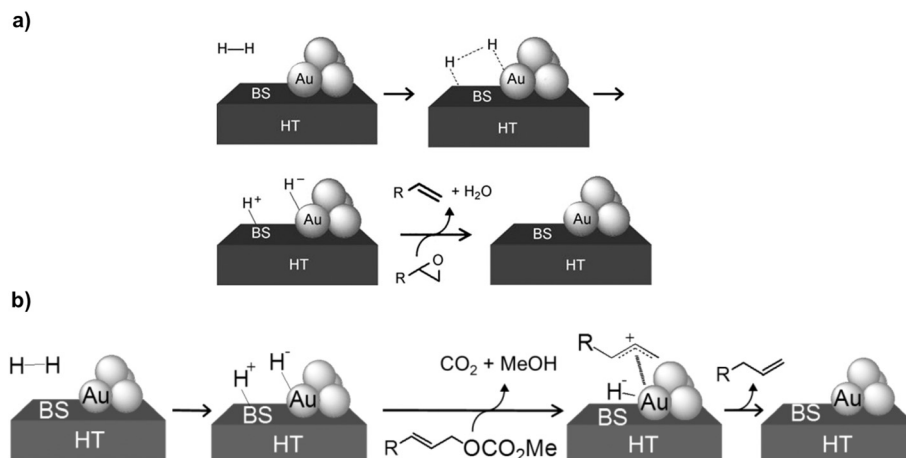


Fig. 21 Proposed mechanism for the hydrogenation of carbonyl compounds over (a) Re-Pd/SiO<sub>2</sub> and (b) Ir/MgO<sup>+</sup>Fe(NO<sub>3</sub>)<sub>3</sub>. Reprinted with permission from ref. 135 and 136, respectively. Copyrights 2017 and 2015. American Chemical Society.





**Scheme 9** Heterolytic activation of  $H_2$  at the interface between gold atoms and basic sites (BS) of HT in (a) the deoxygenation of epoxides and (b) the hydrogenolysis of allylic carbonates. Reprinted with permission from ref. 138 and 139, respectively. Copyright 2011. Wiley. Copyright 2012. Royal Society of Chemistry.

To finish this section, it is worth mentioning that atomically dispersed catalysts on oxide supports are highly efficient in HCD. Here, the presence of metal–oxygen couples only, instead of metal–metal pairs necessary for homolytic dissociation of  $H_2$ , makes the heterolytic dissociation the predominant way to cleave hydrogen. A clear example of that is the work of Zheng *et al.* concerning atomically dispersed palladium atoms on  $TiO_2$ .<sup>141</sup> Through a new photochemical procedure, stable dispersed palladium atoms on ethylene glycolate (EG)-stabilized ultrathin  $TiO_2$  nanosheets ( $Pd_1/TiO_2$ ) were prepared. The large number of  $Pd$ –O interfaces of the system essentially activates dihydrogen in a heterolytic way, which is reflected in an exceptional stability and high catalytic activity in the hydrogenation of carbonyl groups. For example, the atomically dispersed  $Pd$  catalyst exhibits a TOF much higher than commercial  $Pd$  catalysts in the hydrogenation of aldehydes, without any decrease of the activity during the catalytic process.

### 3.2 Supported MNPs functionalized with organic ligands

Unifying the advantages of ligand-stabilized and supported MNPs for heterolytic activation of dihydrogen, supported MNPs functionalized with organic ligands are presented as high potential catalysts for selective hydrogenation reactions.<sup>142</sup> Theoretically, they possess the stability and recyclability of supported MNPs together with the specific activity of metals influenced by coordinated ligands. At first, one might think that surface ligands would block the active sites of MNPs available for catalysis, but the ligands also modify the NP surface electronically and sterically, thereby changing the catalytic activity and selectivity, in the same way as they do in organometallic chemistry. Additionally, ligands can play an important role in the reaction mechanisms through metal–ligand interactions, like in the heterolytic activation of  $H_2$  in ligand-stabilized MNP catalysis discussed above (see section 2.2). In addition to the geometric and

electronic effects induced by ligands to the metal surface<sup>143–145</sup> and the metal–ligand cooperation,<sup>146</sup> there is another possible effect based on substrate–ligand interactions.<sup>147,148</sup> The possibility to control the selectivity of a given reaction by ligand–substrate interactions is reminiscent of biological catalysts, where substrates bind to enzymes through non-covalent interactions. To sum up, the functionalization of supported MNPs to improve their catalytic performance is a new strategy with high potential that has been scarcely explored.

Since this novel approach is still in its infancy, we have only found one example in the literature about HCD by supported ligand-functionalized MNPs. In 2017, Rossi *et al.* presented supported-Au NPs functionalized with basic nitrogen compounds as a frustrated Lewis pair able to dissociate dihydrogen heterolytically.<sup>80</sup> This HCD led to a high activity in the selective hydrogenation of alkynes into *cis*-alkenes. After a screening study of a series of amines with different structures and  $pK_a$  values, they observed that nitrogen-containing bases with two nitrogen atoms showed a higher activity than those with only one (Fig. 23a). Indeed, piperazine-functionalized Au NPs supported on silica ( $Au/SiO_2$ ) exhibited full conversion and 100% selectivity to styrene in the hydrogenation of phenylacetylene, giving the best result among all the N-containing bases studied. A large amount of the amine is necessary to have a higher activity and to suppress the hydrogenation of styrene to ethylbenzene. For instance, when the substrate:amine:Au ratio is 100:1:1, the conversion is very low, only 13% of the alkene is observed. However, when the ratio is 100:100:1, there was a full conversion and a selectivity to styrene of 100%. Therefore, the amount of piperazine not only plays an important role in the activation of  $Au/SiO_2$ , but also improves the selectivity of the catalyst, blocking the hydrogenation of alkenes into alkanes. Other parameters such as the solvent, temperature and support used also influence the activity, obtaining the best results in EtOH, at 100 °C with piperazine-



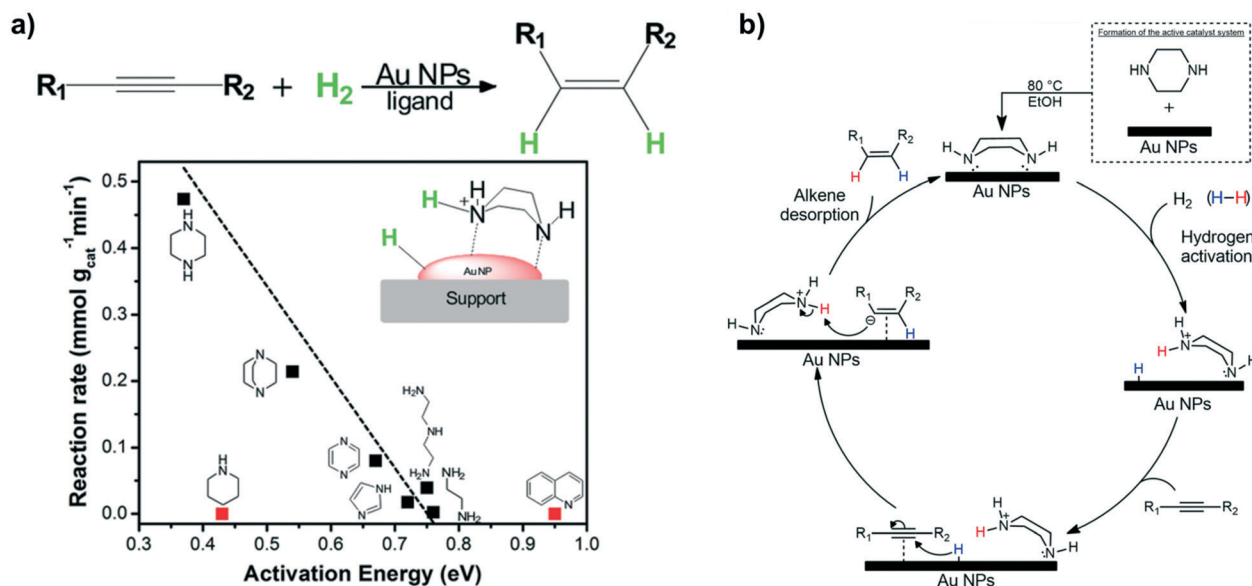


Fig. 23 (a) Top: Semihydrogenation of alkynes to alkenes using Au/SiO<sub>2</sub> functionalized with nitrogen-containing bases. Bottom: Experimental reaction rates vs. computed activation energies for heterolytic H<sub>2</sub> activation at N ligand–Au(111) interfaces. Inset: Illustration about heterolytic H<sub>2</sub> cleavage over piperazine-functionalized AuNPs supported on SiO<sub>2</sub>. (b) Proposed mechanism *via* H<sub>2</sub> cleavage. Reprinted with permission from ref. 80. Copyright 2017. American Chemical Society.

functionalized Au/SiO<sub>2</sub>. Employing these conditions, numerous terminal and internal alkynes were selectively hydrogenated to *cis*-alkenes with excellent yields. Here, as well as in homogeneous catalysis, the ligands coordinated on the metal surface promote the HCD. Au/SiO<sub>2</sub> without any amine on the surface is not active in the hydrogenation of alkynes (conversion <1%), but after functionalizing it with piperazine, it is able to activate hydrogen in a heterolytic way and hydrogenate selectively triple bonds. Fig. 23b shows the mechanism proposed by the authors, which starts with the adsorption of piperazine on the gold surface to form the active catalyst. Then, HCD takes place at the Au–N interface. Subsequently, the adsorbed alkyne reacts with the hydride of the pair Au–H<sup>δ−</sup>, forming the vinyl intermediate. Finally, this intermediate is attacked by the N–H<sup>δ+</sup> ion, affording the alkene and regenerating the catalyst. To the best of our knowledge, this is one of the few examples of HCD over ligand-functionalized supported metal nanoparticles so far, but this versatile concept can be easily adapted to other catalytic systems with other metals and basic ligands. Thanks to that, we will be able to reach catalytic reactivities characteristic of molecular complexes with tunable heterogeneous systems, having the advantages of both catalysts: (i) homogeneous: activity and selectivity; and (ii) heterogeneous: stability and recyclability.

## 4. Metal oxide NPs

The dissociation of H<sub>2</sub> on “bulk” metal oxides through a HCD mechanism has a longstanding tradition (section 1.2). This transformation has been proposed to occur on

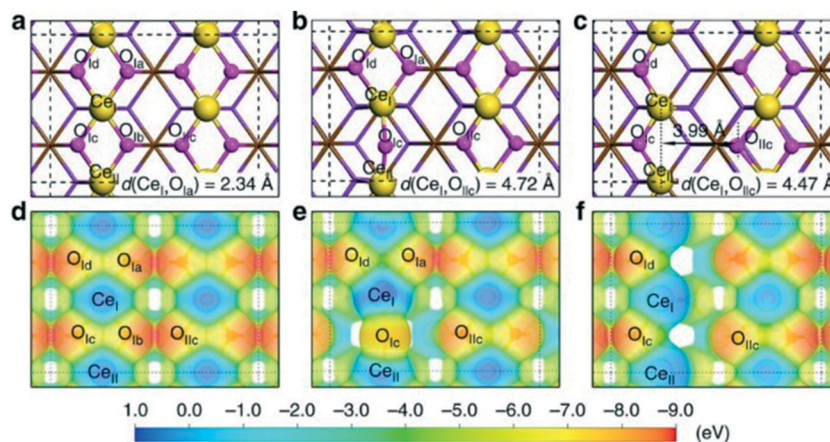
non-reducible oxides with the concomitant formation of MH<sup>−</sup>/OH<sup>+</sup> pairs. On the other hand, a homolytic cleavage that implies the generation of two O–H bonds has been suggested for reducible metal oxides (*vide supra*). Both mechanisms have long been under discussion, and indeed García-Melchor and López demonstrated that they are not excluding one another and may be interconnected (*e.g.* as in Fig. 3c).<sup>59</sup> A good example of this debate is evidenced in an excellent theoretical study of Pacchioni *et al.*<sup>149</sup> DFT calculations showed that an HCD is energetically preferred on extended surfaces (bulk) of ZrO<sub>2</sub>, for which Zr–H<sup>−</sup>/O–H<sup>+</sup> pairs are generated without reduction of zirconia. Conversely, a homolytic H<sub>2</sub> cleavage involving the formation of two O–H groups and reduction of Zr<sup>4+</sup> to Zr<sup>3+</sup> is the favoured mechanism for ZrO<sub>2</sub> NPs of 1.5 nm in size. In the present section, we focus on the use of “isolated” metal oxide nanoparticles as a tool to achieve a HCD process.

In this context, Knözinger and coworkers described a heterolytic hydrogen cleavage on MgO NPs of 5–8 nm in size.<sup>150</sup> The MgO NPs, obtained by chemical vapour deposition, were exposed to H<sub>2</sub>, leading to a HCD and the subsequent formation of Mg–H<sup>−</sup> and O–H<sup>+</sup> species on the NP surface. Under UV irradiation, the formed hydrides serve as the electron source to generate different types of colour centres at the surface and thus characterize electron-trapping sites. IR and EPR studies suggested that the surface MgO sites responsible for the heterolytic cleavage form colour centres because their respective Mg<sup>2+</sup> species operate as a trap for electrons from hydrides.

HCD was also achieved by porous CeO<sub>2</sub> nanorods that exhibit a high density of surface defects.<sup>151</sup> The regulation of







**Fig. 24** Design of FLP in CeO<sub>2</sub>(110) NPs. Top: Optimized structure of (a) ideal CeO<sub>2</sub> NPs; (b) CeO<sub>2</sub> NPs with one oxygen vacancy; (c) CeO<sub>2</sub> NPs with two adjacent oxygen vacancies. Bottom: Electron-density isosurface of (d) ideal CeO<sub>2</sub> NPs; (e) CeO<sub>2</sub> NPs with one oxygen vacancy; (f) CeO<sub>2</sub> NPs with two oxygen vacancies. Electron-density isosurfaces plotted at 0.01 e bohr<sup>-3</sup>. Colour bar shows the electrostatic potential scale. Reprinted with permission from ref. 151. Copyright 2017. Springer Nature.

surface defects is key for the heterolytic splitting, since it allows the creation of oxygen vacancies. The elimination of one oxygen atom involves the reduction of two surface Ce cations (Ce<sup>4+</sup> → Ce<sup>3+</sup>) and the production of one oxygen vacancy. These vacancies avoid the electronic interaction of Ce species with linked O atoms; that is, the removal of O atoms directly bonded to Ce prevents the generation of a classic Lewis acid–base adduct (Fig. 24).

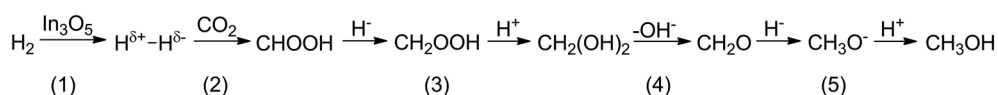
Conversely, the high concentration of surface defects leads to independent acidic and basic sites. Two neighbouring surface Ce<sup>3+</sup> cations function as a Lewis acid, while an adjacent but unbonded lattice oxygen operates as a Lewis base, in such a way that the ensemble acts as a FLP that promotes the HCD (Fig. 24). Indeed, DFT calculations showed a low activation energy (0.17 eV) for the heterolytic dissociation of the H–H bond.

The CeO<sub>2</sub> nanorods were employed as hydrogenation catalysts for alkenes and alkynes. Very high activities and excellent selectivities were observed in the hydrogenation of a series of substituted alkenes, while a decrease in selectivity was noted for alkynes (20 mg CeO<sub>2</sub>, 1 mmol substrate, toluene as a solvent, 6–30 bar, 100 °C, 12–24 h).

The HCD is a useful strategy to reduce CO<sub>2</sub> and thus obtain high-added value products. For instance, Pérez-Ramírez, López *et al.* described a heterolytic H<sub>2</sub> splitting mechanism in the hydrogenation of CO<sub>2</sub> mediated by In<sub>2</sub>O<sub>3</sub> NPs of ca. 8 nm in size.<sup>152</sup> These NPs were found to be an efficient catalyst for the synthesis of methanol with high selectivity, for which kinetic experiments showed a lower

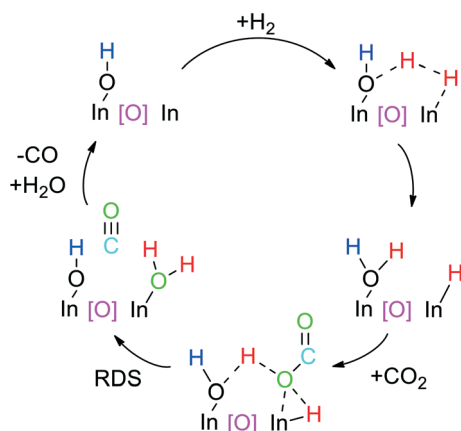
apparent activation energy in comparison with that of the reverse water-gas shift reaction (CO<sub>2</sub> + H<sub>2</sub> → CO + H<sub>2</sub>O). Indeed, a deactivation of the catalyst and therefore an inhibition process by H<sub>2</sub>O was observed. This was supported by kinetic investigations, in which an apparent reaction order of –0.9 was determined for this molecule. On the other hand, the conversion of CO<sub>2</sub> to methanol and thus the selectivity towards the latter is favored at higher H<sub>2</sub> pressures, in line with the apparent reaction order with respect to this reactant (0.5).

Experimental and DFT studies demonstrated that the In<sub>2</sub>O<sub>3</sub> (1 1 1) surface termination is the most exposed one and responsible for the process. The calculations showed that this termination is partially reduced by H<sub>2</sub> through a homolytic mechanism, leading to the generation of OH groups, the concomitant H<sub>2</sub>O release and the formation of an oxygen vacancy surrounded by 3 indium atoms. Then, the methanol synthesis process occurs in several steps (Scheme 10): (1) a heterolytic splitting of H<sub>2</sub> and CO<sub>2</sub> activation takes place at this site surrounded by an In<sub>3</sub>O<sub>5</sub> ensemble. The H<sub>2</sub> molecule is activated as a H<sup>δ+</sup>–H<sup>δ–</sup> species by the binding of H<sup>δ+</sup> to one O atom and H<sup>δ–</sup> to one indium atom, in such a way that the electron density of the latter is delocalized in the 3 indium atoms of In<sub>3</sub>O<sub>5</sub>. (2) A transfer of one H<sup>–</sup> to CO<sub>2</sub> generates a formate (CHO<sub>2</sub><sup>–</sup>) species, which forms a formic acid (CHOOH) intermediate by a subsequent transfer of H<sup>+</sup>. (3) Afterwards, a new H<sub>2</sub> molecule is heterolytically transferred to this intermediate through the formation of



**Scheme 10** Hydrogenation of CO<sub>2</sub> through a HCD catalyzed by In<sub>2</sub>O<sub>3</sub> NPs to form methanol.





**Scheme 11** Proposed catalytic cycle for the reverse water-gas shift reaction mediated by nanostructured  $\text{In}_2\text{O}_{3-x}(\text{OH})_y$ . RDS: rate-determining step.

$\text{CH}_2\text{OOH}$  by  $\text{H}^-$  insertion and its following protonation to generate methanediol ( $\text{CH}_2(\text{OH})_2$ ). (4) A third  $\text{H}_2$  molecule is heterolytically adsorbed, which implies the dissociation of an OH group from the  $\text{CH}_2(\text{OH})_2$  intermediate to fill the oxygen vacancy and the subsequent formation of formaldehyde ( $\text{CH}_2\text{O}$ ). (5) Finally, this molecule interacts with the  $\text{H}^-$  arising from this third HCD and evolves to methoxide ( $\text{CH}_3\text{O}^-$ ), which is protonated and forms methanol ( $\text{CH}_3\text{OH}$ ).

The introduction of OH groups on the surface of  $\text{In}_2\text{O}_3$  NPs leads to catalysts with a completely different behavior. The ability of nanostructured hydroxylated indium oxide ( $\text{In}_2\text{O}_{3-x}(\text{OH})_y$ ) to reduce  $\text{CO}_2$  through a process involving HCD has been studied in depth by Singh, Ozin *et al.*<sup>153</sup> In this case, we are dealing with a transformation that produces CO by the reverse water-gas shift reaction ( $\text{CO}_2 + \text{H}_2 \rightarrow \text{CO} + \text{H}_2\text{O}$ ). The  $\text{In}_2\text{O}_{3-x}(\text{OH})_y$  surface presents Lewis acid (In) and Lewis base (InOH) sites. These sites are spatially separated due to the presence of oxygen vacancies, and thus may function as a surface FLP with the ability to heterolytically cleave  $\text{H}_2$  (Scheme 11).<sup>154</sup> Then, the hydrogenated FLP adsorbs  $\text{CO}_2$ , which is converted to CO and  $\text{H}_2\text{O}$  as a co-product.

DFT calculations showed that an increase in temperature hardly affects the process. However, the energy barriers corresponding to the HCD and  $\text{CO}_2$  adsorption decrease by 0.15 and 0.19 eV at 180 °C, respectively. This enhancement of activity is associated to a larger spatial separation between the Lewis acid (In) and the O of the Lewis base (InOH) of the FLP.<sup>155</sup> Importantly, the nanocatalysts based on  $\text{In}_2\text{O}_{3-x}(\text{OH})_y$  are even more active under light, as photoexcitation reduces the activation barrier by about 20 kJ  $\text{mol}^{-1}$ .<sup>156</sup> Indeed, experimental and theoretical studies demonstrated that the photoactivation promotes the generation of electrons and holes trapped at the Lewis base (electron trapping) and Lewis acid (hole trapping) sites, respectively (Fig. 25). As a result, the Lewis acid and Lewis

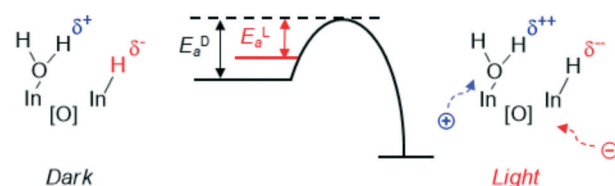
base active sites located in the vicinity of oxygen vacancies become more acidic and more basic, and thus the surface FLP shows an increased activity.<sup>156,157</sup>

In addition, Ozin and coworkers reported the photocatalytic hydrogenation of  $\text{CO}_2$  catalyzed by cuprous oxide ( $\text{Cu}_2\text{O}$ ) nanocubes of *ca.* 500 nm in size.<sup>158</sup> A process involving HCD also led to the reverse water-gas shift reaction. This inexpensive semiconductor shows great potential as a photocatalyst, but exhibits a high tendency towards irreversible disproportionation ( $\text{Cu}_2\text{O} \rightarrow \text{Cu} + \text{CuO}$ ) due to its redox instability, which hinders its use as a photocatalyst. Interestingly, the synthesis provided  $\text{Cu}_2\text{O}$  nanocubes in which the surface contains OH groups, oxygen vacancies and Cu sites with different oxidation states (0, I and II). This makes the redox disproportionation reversible, which improves the stability of  $\text{Cu}_2\text{O}$  nanocubes and thus allows the  $\text{CO}_2$  reduction through HCD.

The  $\text{Cu}_2\text{O}$  nanocubes exhibit high activity and selectivity in the reverse water-gas shift reaction under very mild conditions (25 °C, intense light of 50 suns of blue, green and red visible light,  $\text{CO}_2/\text{H}_2 = 5$ , total pressure *ca.* 4 bar). A CO production of 70.3 mmol  $\text{g}_{\text{cat}}^{-1} \text{h}^{-1}$  was obtained, which was increased up to 88 mmol  $\text{g}_{\text{cat}}^{-1} \text{h}^{-1}$  by removal of coproduced water with dehydrated zeolite LTA-3A. In addition, 139.6 mmol  $\text{g}_{\text{cat}}^{-1} \text{h}^{-1}$  was achieved by a long-term stability experiment (60 h).

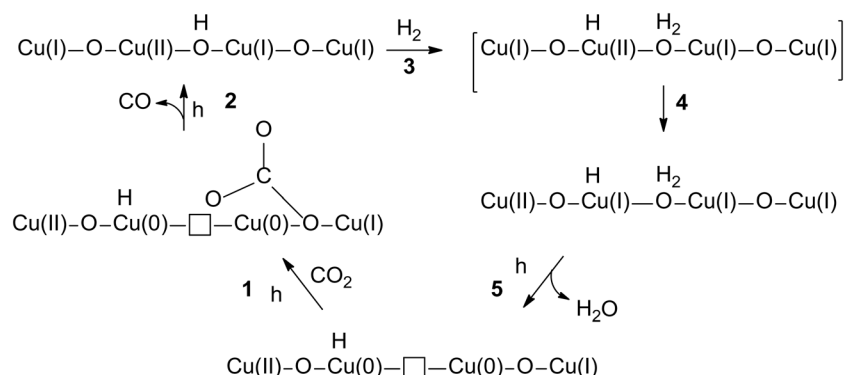
The authors suggested that  $\text{H}_2$  is heterolytically cleaved by an FLP consisting of  $\text{Cu}_2\text{O}$  and  $\text{Cu}_2(\text{OH})_3\text{Cl}$  ( $\text{Cu}(\text{I})_{2-x}\text{Cu}(\text{II})_x\text{O}_{1+x/2-y/2}(\text{OH})_y$ ). In this FLP, OH acts as a Lewis base and Cu(II) operates as a Lewis acid, leading to a HCD as follows:  $\text{Cu}(\text{II})\cdots\text{OH} + \text{H}_2 \rightarrow \text{Cu}(\text{II})\text{H}\cdots\text{OH}_2$ . In addition, the presence of Cu sites with different oxidation states favours electron delocalization ( $\text{Cu}(\text{II})\text{H}\cdots\text{OH}_2 \leftrightarrow \text{Cu}(\text{I})\text{H}\cdots\text{OH}_2$ ). Finally, a catalytic cycle was proposed on the basis of *in situ* DRIFTS studies (Scheme 12), while different control experiments demonstrated that light is a prerequisite for the process.

An HCD occurs on the surface of  $\text{Cu}_2\text{O}$  nanocubes (3 and 4), where electron delocalization between Cu(I) and Cu(II) sites also takes place (4). Then,  $\text{H}_2\text{O}$  is removed, thus generating an oxygen vacancy (5). This vacancy facilitates the adsorption of  $\text{CO}_2$  as a carbonate-like species (1), which eventually converts to CO (2).



**Fig. 25** Representation of the difference in the experimental activation energy for the reverse water-gas shift reaction in the dark (black) and in the light (red) mediated by nanostructured  $\text{In}_2\text{O}_{3-x}(\text{OH})_y$ . Adapted with permission from ref. 153. Copyright 2016. American Chemical Society.





**Scheme 12** Proposed mechanism for photocatalytic hydrogenation of CO<sub>2</sub> mediated by Cu<sub>2</sub>O nanocubes.

## 5. Conclusions and outlook

For a long time, selective hydrogenation of polar functional groups (C=O, C=N or C=S) has been mostly carried out in the laboratory using homogeneous catalysts, in which H<sub>2</sub> is heterolytically activated by a metal ligand cooperation. On the other hand, MNPs and supported metal catalysts normally cleave hydrogen in a homolytic way, showing in general lower selectivities than homogeneous catalysts. However, with the advent of new and controlled protocols for MNP synthesis, and the possibility to functionalize them with organic molecules, the use of MNPs for HCD is gaining attention from the scientific community.

This article collects recent examples of well-defined metal nanocatalysts for the heterolytic activation of H<sub>2</sub> by metal-ligand or metal-support cooperation. Here, the catalytic H<sub>2</sub> dissociation is promoted by the presence of both surface ligands or support basic sites. Even more exciting is the functionalization of supported MNPs with organic molecules to facilitate the heterolytic H<sub>2</sub> cleavage. In this way, we can enhance the selectivity of almost any supported-MNPs. These novel catalytic systems combine the advantages of ligand-stabilized MNPs (selectivity) and supported MNPs (stability and recyclability). In the majority of examples the proof for HCD is absent or only circumstantial, but above we have collected several neat examples that report spectroscopic proof and/or DFT calculations proving the feasibility of HCD. Yet, more joint theoretical-experimental studies about the ligand-metal surface interactions are needed in order to design more efficient and selective catalysts that operate through HCD.

In addition to functionalized supported MNPs, the future directions for research in this area could be oriented to the deposition of single atoms on MNPs as the simplest way to modify their reactivity. A recent example about this was reported by Pérez-Ramírez and López, where chalcogen-modified copper catalysts showed an increased selectivity in the electrochemical reduction of CO<sub>2</sub> to formic acid.<sup>159</sup> In their study, sulfur-, selenium and tellurium active centers are present on the MNP surface as adatoms, and participate in the hydrogenation mechanism playing a dual role: (i)

transferring hydrides to the CO<sub>2</sub> molecule and (ii) suppressing the formation of CO. In a similar way, chalcogen-modified MNPs could perform HCD by a cooperative work between chalcogen adatoms and surface metal atoms, just like in homogeneous systems.<sup>160</sup>

Another interesting prospect in this research area is the use of FLPs as modifying ligands, such as phosphinoboranes or aminophosphines. The main advantage of the functionalization of MNPs with FLPs is to reverse the addition of H<sub>2</sub>. In a typical HCD reaction (metal-basic centre), the hydride coordinates to the metal and it will not be “hydridic”, but in TMFLP systems the addition is inverted; this means that the transition metal accepts the proton and the Lewis acid (*i.e.* borane) the hydride, giving rise to a highly “hydridic” hydride which might be of interest for the course of the catalytic reaction.<sup>84</sup> Over the last few years, FLPs have been an important source of inspiration in chemistry, and we expect that metal nanoparticle catalysis could also take advantage of this.

## Conflicts of interest

There are no conflicts of interest to declare.

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