


 Cite this: *RSC Adv.*, 2020, 10, 10411

# pH-Dependent transfer hydrogenation or dihydrogen release catalyzed by a $[(\eta^6\text{-arene})\text{RuCl}(\kappa^2\text{-}N,N\text{-dmobpy})]^+$ complex: a DFT mechanistic understanding†

 Chenguang Luo,<sup>a</sup> Longfei Li,<sup>b</sup> Xin Yue,<sup>a</sup> Pengjie Li,<sup>a</sup> Lin Zhang,<sup>a</sup> Zuoyin Yang,<sup>a</sup> Min Pu,<sup>ib</sup> Zexing Cao<sup>ib</sup>\*<sup>c</sup> and Ming Lei<sup>ib</sup>\*<sup>ac</sup>

The reaction mechanism of the pH-dependent transfer hydrogenation of a ketone or the dehydrogenation of formic acid catalyzed by a  $[(\eta^6\text{-arene})\text{RuCl}(\kappa^2\text{-}N,N\text{-dmobpy})]^+$  complex in aqueous media has been investigated using the density functional theory (DFT) method. The TM-catalyzed TH of ketones with formic acid as the hydrogen source proceeds *via* two steps: the formation of a metal hydride and the transfer of the hydride to the substrate ketone. The calculated results show that ruthenium hydride formation is the rate-determining step. This proceeds *via* an ion-pair mechanism with an energy barrier of 14.1 kcal mol<sup>-1</sup>. Interestingly, the dihydrogen release process of formic acid and the hydride transfer process that produces alcohols are competitive under different pH environments. The investigation explores the feasibility of the two pathways under different pH environments. Under acidic conditions (pH = 4), the free energy barrier of the dihydrogen release pathway is 4.5 kcal mol<sup>-1</sup> that is higher than that of the hydride transfer pathway, suggesting that the hydride transfer pathway is more favorable than the dihydrogen release pathway. However, under strongly acidic conditions, the dihydrogen release pathway is more favorable compared to the hydride transfer pathway. In addition, the ruthenium hydride formation pathway is less favorable than the ruthenium hydroxo complex formation pathway under basic conditions.

 Received 18th December 2019  
 Accepted 15th February 2020

DOI: 10.1039/c9ra10651k

[rsc.li/rsc-advances](http://rsc.li/rsc-advances)

## Introduction

The hydrogenation of polar double bonds, such as C=O and C=N, catalyzed by transition-metal (TM) complexes is a fundamentally important method to produce value-added alcohol- and amine-containing chemicals in organic syntheses and chemical industries.<sup>1,2</sup> The transfer hydrogenation (TH) of ketones using organic hydrogen sources as well as H<sub>2</sub> hydrogenation (HH) using molecular hydrogen resources have gained more attention.<sup>3</sup> The TH of ketones can be performed either in water or in organic solvents.<sup>4–8</sup> However, due to environmental and ecological advantages and reaction-specific pH selectivity, using water as the solvent for the TH of ketones is very convenient and attractive compared to traditional organic solvents.<sup>9–12</sup> Two hydrogen sources have primarily been used in TM-catalyzed TH,

namely isopropanol and formic acid. The conception of metal–ligand bifunctional cooperation is often used to explain the reaction mechanism of TH and the HH of polar double bonds catalyzed by TM complexes with different ligand environments such as diphosphine-diamine ruthenium catalyst **A** (see Scheme 1)<sup>13–19</sup> as well as those of TH using formic acid hydrogen sources.<sup>20–23</sup> For the mode of hydrogen transfer, inner-sphere and outer-sphere mechanisms are proposed based on the direct/indirect interactions between the metal center and the atoms of the ketones/imines, excluding the hydrogen atoms.

A number of TH reactions of polar double bonds catalyzed by cyclometallated TM complexes have been developed in the past decades.<sup>11,24–27</sup> Most of the TM catalysts for ketone/imine hydrogenation have a Lewis acidic site in the TM center and a Lewis basic site in the ligand moiety of the structure (TM-LB catalyst) or a Lewis basic site in the TM center and a Lewis acidic site in the ligand moiety (TM-LA catalyst).<sup>2,28</sup> In 1986, Shvo *et al.* reported a useful TH catalyst (**B** in Scheme 2) for ketones, which is an example of a ligand–metal bifunctional catalyst wherein the redox activity is distributed between the metal center and a cyclopentadienone ligand.<sup>29</sup> In 1995, Noyori *et al.* synthesized  $\eta^6\text{-arene-Ru}$  complexes bearing monotosylated 1,2-diamine moieties (**C** in Scheme 2), which are

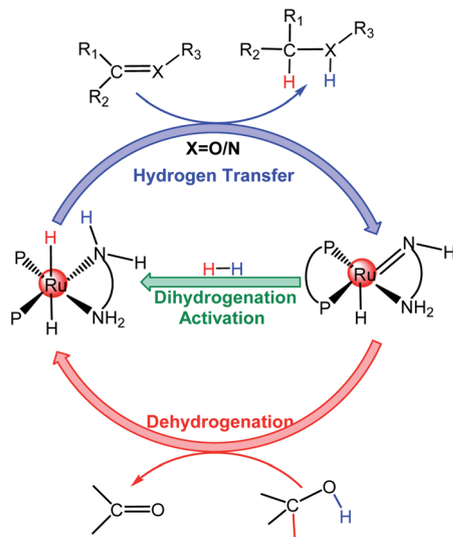
<sup>a</sup>State Key Laboratory of Chemical Resource Engineering, Institute of Computational Chemistry, College of Chemistry, Beijing University of Chemical Technology, Beijing, 100029, China. E-mail: leim@mail.buct.edu.cn

<sup>b</sup>College of Pharmaceutical Science, Hebei University, Baoding, 071002, China

<sup>c</sup>State Key Laboratory of Physical Chemistry of Solid Surfaces, Xiamen University, Xiamen, 361005, China. E-mail: zxcao@xmu.edu.cn

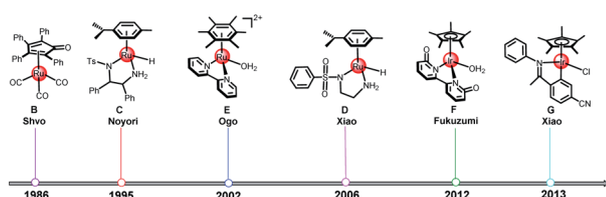
† Electronic supplementary information (ESI) available. See DOI: 10.1039/c9ra10651k





Scheme 1 The metal–ligand bifunctional mechanism for ketone/imine hydrogenation catalyzed by Ru complexes.

efficient TH catalysts for ketones.<sup>30</sup> In 2006, Xiao *et al.* made a breakthrough by improving catalyst efficiency using iridium catalysts with *N*-sulfonyl ethylenediamine as the ligand (**D** in Scheme 2) in water with HCOONa as the hydrogen source.<sup>31</sup> Some cyclometallated TM complexes are single-site catalysts. They can be modified in the ligand moiety to provide a Lewis basic or acidic site and then can become bifunctional catalysts like **B**, **C**, and **D**. In 2002, Ogo *et al.* reported the TH of ketones using HCOONa or HCOOH in water with cyclometalated single-site Ru complexes (**E** in Scheme 2).<sup>32</sup> In 2012, Fukuzumi *et al.* reported a single-site cyclometalated Ir complex bearing a bpyO ligand (**F** in Scheme 2), which could catalyze aliphatic alcohol dehydrogenation at room temperature in a basic aqueous solution.<sup>33</sup> In 2013, Xiao *et al.* reported another cyclometalated single-site Ir complex (**G** in Scheme 2), which was shown to be an excellent catalyst for the TH of carbonyl compounds in water using formate as the hydrogen source.<sup>27</sup> As discussed above, it is obvious that the mechanism of TH reactions catalyzed by cyclometallated single-site TM complexes is different from that of bifunctional TM catalysts. However, reports concerning the mechanism of TH with formic acid as the hydrogen source and catalyzed by cyclometallated single-site TM complexes have been sporadic.<sup>22</sup> The TM-catalyzed TH of ketones with formic acid as the hydrogen source proceeds *via* two sequential steps. First, hydride transfer from the formate moiety to the metal

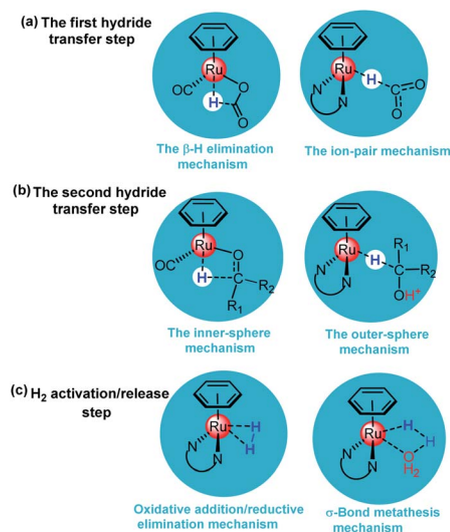


Scheme 2 The development of cyclometallated TM complexes.

center (formation of the metal hydride) occurs and then a hydride transfer step from the metal to the ketone substrate follows. There are two modes for metal hydride formation: the  $\beta$ -hydrogen elimination mechanism and the ion-pair mechanism (see Scheme 3a).<sup>22</sup>

For the  $\beta$ -hydrogen elimination mechanism, it has been stated that the formate anion must occupy two coordination sites before cleavage of the C–H bond of the formate can occur.<sup>34</sup> In 2013, Yang *et al.*<sup>35</sup> investigated the mechanism of formic acid dehydrogenation catalyzed by an iron complex *via* the DFT method and their study indicated that  $\beta$ -hydrogen elimination was involved in the transition state (TS) of metal hydride formation. In the ion-pair mechanism, the metal hydride is formed by the direct hydride transfer from the formate moiety to the metal center in the form of an ion pair and the Ru–O bond is broken before metal hydride formation.<sup>36</sup> In 2015, Xiao *et al.*<sup>22</sup> investigated the mechanism of imine reduction with formic acid catalyzed by a single-site cyclometallated Ir catalyst, the calculated results showed that metal hydride formation proceeds *via* an ion-pair mechanism.<sup>37</sup> If the TM center of the metal hydride species is coordinatively unsaturated, the substrates prefer to coordinate with the TM center according to the  $\beta$ -hydrogen elimination mechanism along an inner-sphere pathway.<sup>38</sup> If it is coordinatively saturated, the substrate will coordinate with the TM center according to the ion-pair mechanism along an outer-sphere pathway, which is supported by experimental observations.<sup>22,27,32,39</sup> The reaction pathways of the second hydride transfer step from the metal to the ketone substrate are similar to the reversible processes of the first hydride transfer step mentioned above (see Scheme 3b).

The non-cooperation mechanism of H<sub>2</sub> activation/release for single-site TM catalysts can be divided into the classical oxidative addition/reductive elimination mechanism and the  $\sigma$ -bond



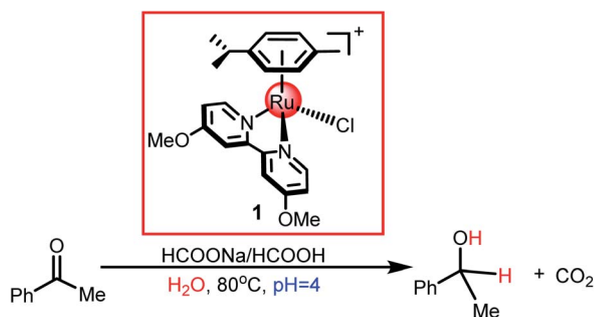
Scheme 3 (a) Representative reaction modes of formate decarboxylation to produce the metal hydride (b) Representative reaction modes of hydride transfer from the metal to the ketone substrate. (c) Representative reaction modes of H<sub>2</sub> activation/release.



metathesis (hydrogenolysis) mechanism (see Scheme 3c).<sup>28</sup> In the classical oxidative addition/reductive elimination mechanism, the coordination of H<sub>2</sub> affords a typical dihydrogen complex [M–H<sub>2</sub>], which quickly undergoes oxidative addition to form a dihydride intermediate [H–M–H] along a homolytic splitting pathway. The oxidation state of the metal center increases as the dihydride complex is formed in the oxidative addition of H<sub>2</sub>. In the  $\sigma$ -bond metathesis mechanism, the metal hydride [M–H] generally prefers to activate/release H<sub>2</sub> *via* a heterolytic splitting pathway. During the dehydrogenation, solvents, including water or formate, can assist H<sub>2</sub> release.

Previous studies on the TH of ketones in aqueous media have revealed that water can accelerate ketone reduction and that pH has a dramatic effect on the catalytic activities of cyclometallated Ru, Rh, and Ir complexes.<sup>12,24,25,31,32,40–42</sup> This effect was also observed in the dehydrogenation reactions of formic acid.<sup>43–48</sup> In 2003, Ogo *et al.* reported pH-dependent TH of a variety of carbonyl compounds catalyzed using a cyclometallated Ir catalyst in water.<sup>25</sup> They proposed that the dehydrogenation of formic acid is achieved under strongly acidic conditions, which was confirmed *via* a GC analysis. The TH of ketones and formic acid dehydrogenation catalyzed by the same cyclometallated catalyst in water are two competitive pathways at different pH values.

Recently, Espino *et al.*<sup>49</sup> developed a pH-dependent catalytic system for the TH of acetophenone using functional cyclometallated  $[(\eta^6\text{-arene})\text{RuCl}(\kappa^2\text{-}N,N\text{-dmobpy})]^+$  as the catalyst precursor, which showed only one open coordination site upon chloride dissociation (see Scheme 4). They proposed a tentative reaction mechanism for cyclometallated  $[(\eta^6\text{-arene})\text{RuCl}(\kappa^2\text{-}N,N\text{-dmobpy})]^+$  (**1**) catalyzed acetophenone hydrogenation, which included two steps: hydride transfer from the formate to the metal center *via* the  $\beta$ -hydrogen elimination mechanism and hydride transfer to substrates *via* the inner-sphere mechanism. In the past decades, a lot of experimental and theoretical studies have been performed to unveil the nature of the TH and HH of ketones catalyzed by TM complexes and to understand the preference for different hydrogen resources in different catalytic systems. However, the reaction mechanism of pH-dependent transfer hydrogenation or dihydrogen release catalyzed by a  $[(\eta^6\text{-arene})\text{RuCl}(\kappa^2\text{-}N,N\text{-dmobpy})]^+$  complex is still unclear. Herein, a DFT study was performed to investigate the nature of this system, which may provide insights to develop an



Scheme 4 Reduction of ketones catalyzed by the cyclometallated ruthenium single-site complex.

understanding on the TH and HH of polar double bonds catalyzed by cyclometallated single-site TM complexes.

## Computational methods

In accordance with our previous computational studies,<sup>3,19,50–55</sup> all calculations in this study were carried out using the DFT method with  $\omega$ B97X-D<sup>56</sup> using the Gaussian 09 program.<sup>57</sup> The SMD polarizable continuum model in water as the solvent was employed in the calculations.<sup>58</sup> The effective core potential (ECP) of Ru with a double- $\zeta$  valence basis set (LANL2DZ) was chosen to describe Ru and the 6-31G(d,p) basis set for other atoms.<sup>59,60</sup> All the transition states were confirmed *via* vibrational analysis and characterized by only one imaginary frequency. Intrinsic reaction coordinate (IRC) calculations were performed to confirm all transition states connecting two desired minima. All relative energies of the stationary points along the reaction pathway are relative to complex **2**. Unless otherwise stated, the energy values in the following parts are free energies. Gibbs free energies were calculated at 298.15 K.

## Results and discussion

The reaction mechanism of ketone transfer hydrogenation or dehydrogenation of formic acid catalyzed by the  $[(\eta^6\text{-arene})\text{RuCl}(\kappa^2\text{-}N,N\text{-dmobpy})]^+$  complex is shown in Fig. 1. Two steps exist in this process: the first step is the formation of a metal hydride and the second step is hydride transfer from the metal to the ketone substrate to complete TH or the dihydrogen release. The mechanisms of TH using ketones and the dehydrogenation of formic acid have the same first step. Initially, the catalytic precursor **1** undergoes an aquation step to give complex **2**, which is the active catalytic species in the catalytic cycle.<sup>49</sup> Then, the H<sub>2</sub>O of complex **2** is substituted with the formate anion to form **3**. The next step is the metal hydride

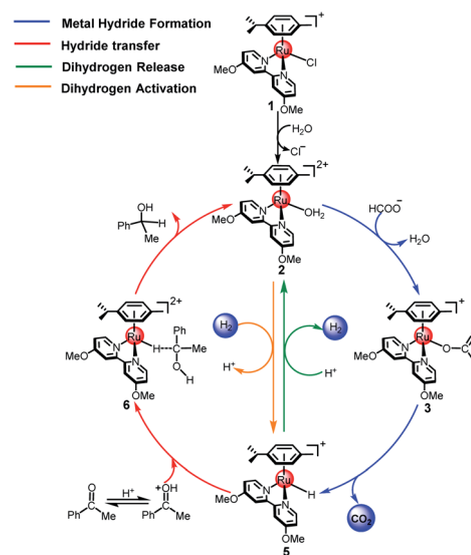


Fig. 1 The catalytic cycle for the TH of ketones catalyzed by cyclometallated ruthenium single-site complexes.



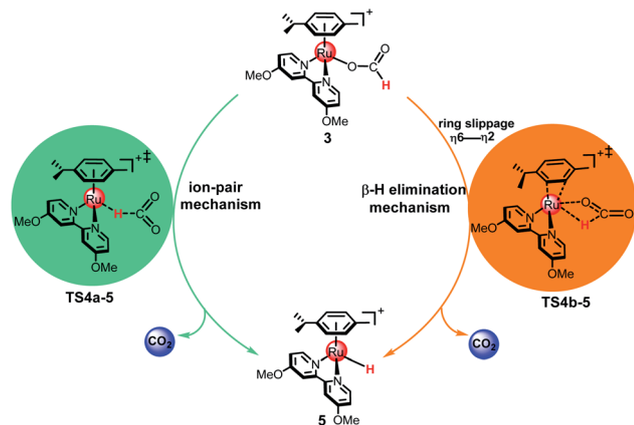


Fig. 2 The ion-pair mechanism and  $\beta$ -hydrogen elimination mechanism for the formation of metal hydride 5.

formation from 3 to 5. Subsequently, the dihydrogen release pathway and the hydride transfer pathway compete to release dihydrogen or complete the TH of the ketone. The hydride of 5 coordinates to the protonated ketone (in acidic aqueous media) to form intermediate 6 or reacts with hydrated protons to release dihydrogen. The reverse process of dihydrogen release, the dihydrogen activation, is also considered. Finally, the active catalytic species 2 is regenerated.

### Metal hydride formation

As shown in Fig. 2, two possible mechanisms for the formation of metal hydride 5 from 3 exist: the ion-pair mechanism and the  $\beta$ -hydrogen elimination mechanism. In the ion-pair

mechanism, the intermediate 5 is formed *via* TS4a-5, which forms an ion pair between the dissociated formate and the cationic 16e Ru(II) complex. The  $\beta$ -hydrogen elimination mechanism involves a change in the arene ring from  $\eta^6$ - to  $\eta^2$ -coordination by ring slippage from 3 to TS4b-5. Then, the metal hydride 5 is formed *via* TS4b-5 with a four-membered ring structure.

**The ion-pair mechanism.** In the ion-pair mechanism, the H<sub>2</sub>O ligand in the catalytic species 2 is substituted with HCOO<sup>-</sup> to form intermediate 3 in the presence of the formate anion. This process is exergonic by 12.1 kcal mol<sup>-1</sup> (see Fig. 3). Then, 3 undergoes conformational conversion to produce intermediate 4a. This process is endergonic by 12.7 kcal mol<sup>-1</sup>. Subsequently, the metal hydride intermediate 5 is formed by a hydride transfer step from the formate to the Ru center *via* TS4a-5, with an energy barrier of 1.4 kcal mol<sup>-1</sup>. Finally, the C–H bond cleavage affords the metal hydride 5 and releases one molecule CO<sub>2</sub>. The Ru–H distance is 2.953 Å in 3, indicating that there is little interaction between the hydrogen atom of the formate and the Ru(II) center. From 4a to 5, the Ru–H distance decreases from 1.840 Å to 1.598 Å, showing that a covalent Ru–H bond is formed in 5. Notably, the Ru–O bond cleavage is accompanied by charge separation and is more difficult than the C–H bond activation of the formate moiety in 3. The energy barrier for this ion-pair pathway is 14.1 kcal mol<sup>-1</sup> from 3 to TS4a-5.

**The  $\beta$ -hydrogen elimination mechanism.** A feature of the  $\beta$ -hydrogen elimination mechanism is the ring slippage of the arene ring from  $\eta^6$ -coordination to  $\eta^2$ -coordination, *i.e.*, from intermediate 3 to intermediate 4b. This process is endergonic by 8.2 kcal mol<sup>-1</sup>. Then, 5 is formed after hydride transfer from the formate to the Ru center *via* TS4b-5, which has a four-membered ring structure with an energy barrier of 24.0 kcal mol<sup>-1</sup>. In comparison

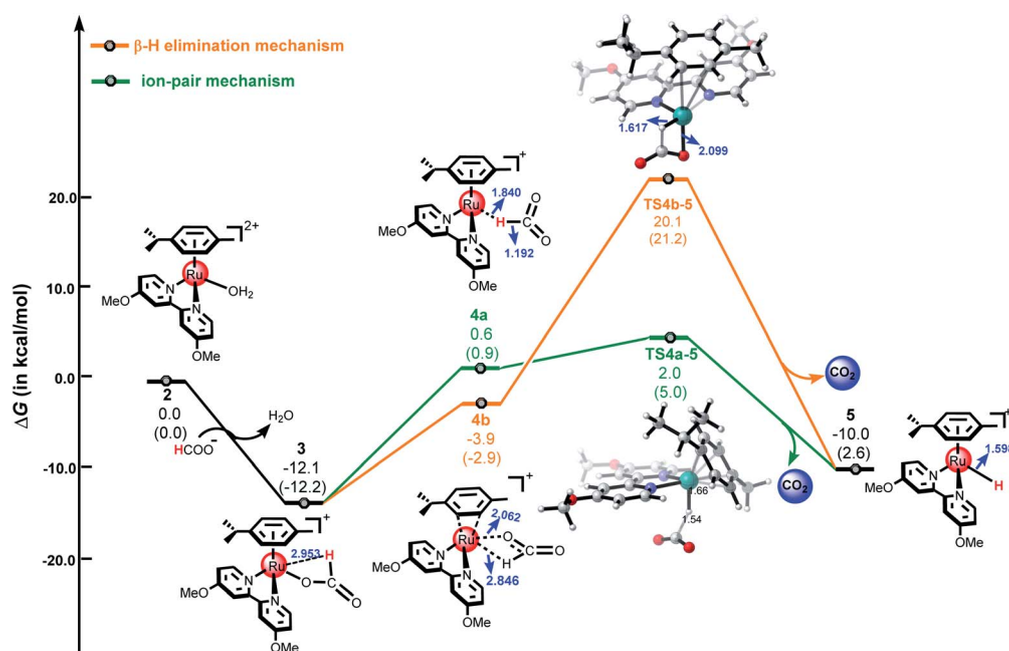


Fig. 3 The free energy profiles for the formation of metal hydride 5. All energies are denoted in kcal mol<sup>-1</sup> and interatomic distances are shown in Å, the values in parentheses are electronic energies.



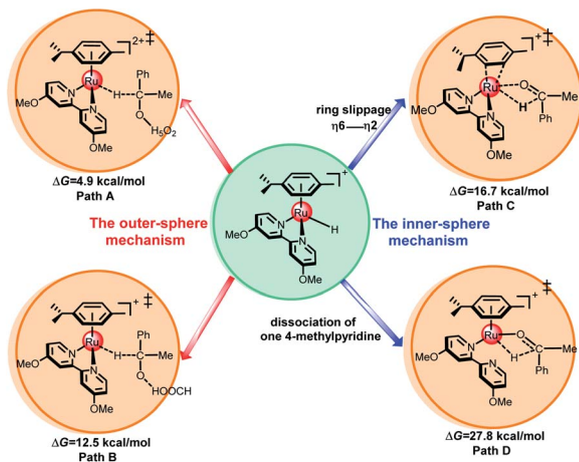


Fig. 4 Four possible pathways of hydride transfer from the metal to the ketone substrate. The values of  $\Delta G$  (in  $\text{kcal mol}^{-1}$ ) indicate the free energy barriers of the four possible pathways.

to **TS4a-5** from the ion-pair mechanism, the most notable feature here is that the oxygen of the formate moiety coordinates with the Ru center while the Ru–O bond distance changes from 2.062 Å in **4b** to 2.099 Å in **TS4b-5**. The energy barrier for this  $\beta$ -hydrogen elimination pathway is 32.2  $\text{kcal mol}^{-1}$  from **3** to **TS4b-5**, which is 18.1  $\text{kcal mol}^{-1}$  higher than that of the ion-pair pathway (14.1  $\text{kcal mol}^{-1}$ ), indicating that the  $\beta$ -hydrogen elimination mechanism is much less favorable than the ion-pair mechanism.

### Ketone hydrogenation or dihydrogen release

Fig. 4 shows four possible pathways for the second hydride transfer to complete the TH of ketones. There are two outer-sphere pathways and two inner-sphere pathways for hydride transfer from the metal center to the carbon atom of the ketone substrate, which belong to the ion-pair mechanism and  $\beta$ -hydrogen elimination mechanism, respectively.

As shown in Fig. 5, the ketone substrate can be stabilized with a hydrated proton ( $\text{H}_5\text{O}_2^+$ ) through hydrogen bonding under acidic conditions in path A, this process is endergonic by 3.6  $\text{kcal mol}^{-1}$  (see Fig. S8<sup>†</sup>). The use of the hydrated proton ( $\text{H}_5\text{O}_2^+$ ) here corresponds to an acidic aqueous medium.<sup>12</sup> First, the protonated ketone coordinates with metal hydride **5** to form intermediate **6**. Then, the hydride is transferred from the Ru center to the carbon atom of the protonated ketone *via* **TS6-7** with an energy barrier of 4.9  $\text{kcal mol}^{-1}$ . Subsequently, intermediate **7** is formed, the free energy of complex **7** is  $-14.1 \text{ kcal mol}^{-1}$ . From **6** to **TS6-7**, the Ru–H bond length increases from 1.578 Å to 1.639 Å and the C–H distance decreases from 2.431 Å to 1.697 Å. The calculated energy barriers of path B, path C, and path D are 12.5  $\text{kcal mol}^{-1}$ , 16.7  $\text{kcal mol}^{-1}$ , and 27.8  $\text{kcal mol}^{-1}$ , respectively (see Fig. 4 above and Fig. S1–S3 in ESI<sup>†</sup>). The calculated results demonstrate that hydride transfer from the metal to the ketone substrate *via* the ion-pair mechanism (path A) is more favorable than the three other possible pathways.

Recently, several studies have shown that metal hydride species ( $[\text{M}-\text{H}]^+$ ) can react with hydrated protons to release  $\text{H}_2$  in an acidic aqueous medium.<sup>43,44,47,48,61</sup> As mentioned above, the dihydrogen release pathway and the second hydride transfer pathway (path A) could be two competitive processes driven by the cyclometallated ruthenium single-site species **5**. As shown in Fig. 5, the energy barrier of the dihydrogen release pathway is 9.4  $\text{kcal mol}^{-1}$ , which is 4.5  $\text{kcal mol}^{-1}$  higher than that of path A (4.9  $\text{kcal mol}^{-1}$ ). Obviously, the hydride transfer pathway *via* an outer-sphere ion-pair mechanism (path A) is more favorable than the dihydrogen release pathway, which is in agreement with experimental results.<sup>49</sup>

### The influence of pH on reactivity

Espino *et al.* found that this  $[(\eta^6\text{-arene})\text{RuCl}(\kappa^2\text{-}N,N\text{-dmobpy})]^+$  system is pH-dependent in experiments.<sup>49</sup> A pH value of 4 was found to be the most favorable and the reduction becomes slow or stagnant outside of a reasonable pH value. Furthermore, the

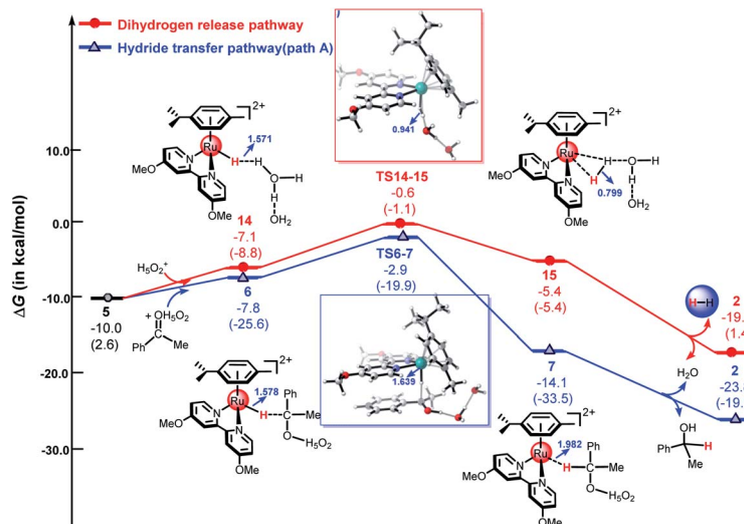


Fig. 5 The free energy profiles of the hydride transfer pathway (path A) and dihydrogen release pathway. All energies are denoted in  $\text{kcal mol}^{-1}$  and interatomic distances are shown in Å, the values in brackets are electronic energies.



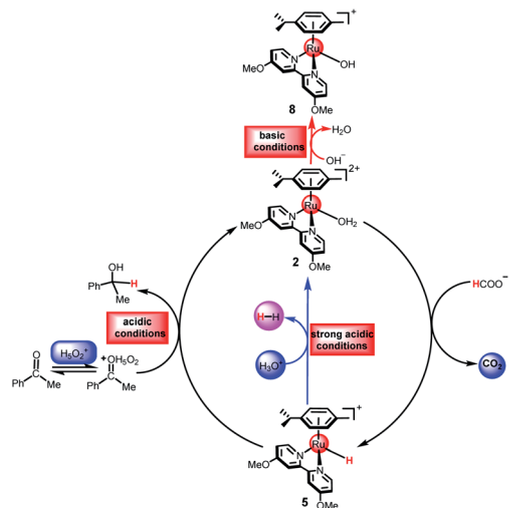
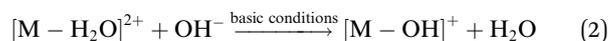
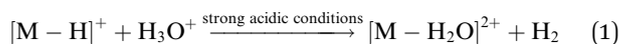


Fig. 6 Proposed mechanism for TH of ketones catalyzed by cyclometallated ruthenium single-site complex at different pH values.

experimental results have shown that the reversible formation of unreactive  $[\text{Ru}-\text{OH}]^+$  at high pH levels (basic condition) leads to inactivity. Several cases of pH-dependent selective TH of ketones have been reported by several groups.<sup>24,32,62</sup> For those active cyclometallated TM single-site complexes  $[\text{M}-\text{H}_2\text{O}]^{2+}$  and metal hydrides  $[\text{M}-\text{H}]^+$ , Ogo *et al.* proposed that (1) under strongly acidic conditions, the protonation of the  $[\text{M}-\text{H}]^+$  leads to the release of  $\text{H}_2$ , which was confirmed by GC analysis (see eqn (1)); (2) under basic conditions,  $[\text{M}-\text{H}_2\text{O}]^{2+}$  is predominantly deprotonated to form a hydroxo complex  $[\text{M}-\text{OH}]^+$ , which easily leads to the termination of the reaction (see eqn (2)). In 2018, Xu *et al.*<sup>11</sup> pointed out that even though a metal

hydride is generated, reduction did not occur under neutral or basic conditions. Thus, the pH-dependence is also related to the proton-mediated activation of the ketones.



That is to say, different pH values may result in different reaction reactivities. As shown in Fig. 6, the active species 5 can react with  $\text{H}_3\text{O}^+$  to release  $\text{H}_2$  under strongly acidic conditions or with the protonated ketone to undergo the second hydride transfer pathway. The active intermediate 2 can form the hydroxo intermediate 8 leading to the termination of the reaction under basic conditions or the formation of the metal hydride 5 under acidic conditions. The metal hydride formation pathway and the metal hydroxo formation pathway are the two processes open to the catalytic species 2 and the chosen pathway is dependent on the pH values.

Herein, we used the DFT method to investigate the reaction mechanism for the TH of ketones catalyzed by a cyclometallated ruthenium single-site complex at different pH values. We used different hydrated protons including  $\text{H}_3\text{O}^+$ ,  $\text{H}_5\text{O}_2^+$ , and  $\text{H}_7\text{O}_3^+$  to indicate the different pH values of the solution. Fig. 7 shows the free energy profiles of the dihydrogen release pathway and the hydride transfer pathway mediated by  $\text{H}_3\text{O}^+$ . The energy barrier of the dihydrogen release pathway is  $1.1 \text{ kcal mol}^{-1}$ . Interestingly, the reaction energy barrier along the hydride transfer pathway (Path A) is  $3.5 \text{ kcal mol}^{-1}$ . In general, the dihydrogen release pathway, driven by the cyclometallated TM single-site system, is more favorable than the hydride transfer pathway under strongly acidic conditions, which is in agreement with

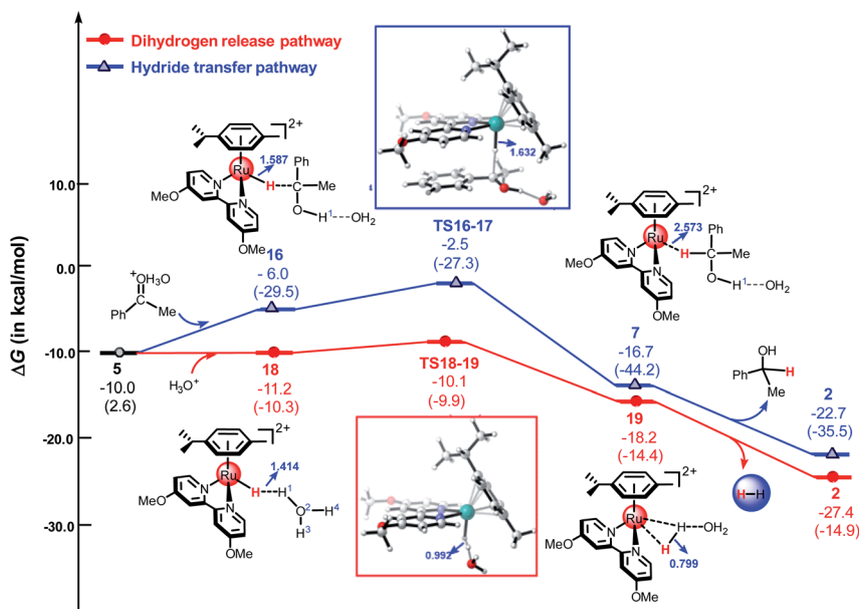


Fig. 7 The free energy profiles of the hydride transfer pathway and the dihydrogen release pathway mediated by  $\text{H}_3\text{O}^+$ . All energies are denoted in  $\text{kcal mol}^{-1}$  and interatomic distances are shown in Å, the values in parentheses are electronic energies.



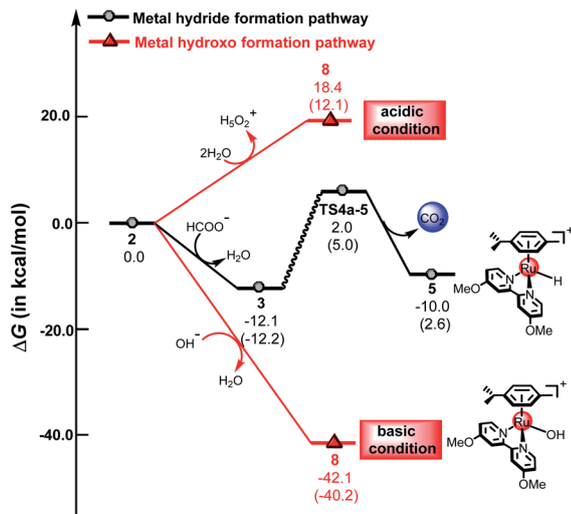


Fig. 8 The free energy profiles of the metal hydride formation pathway and the metal hydroxo formation pathway, the values in parentheses are electronic energies.

the experimental results.<sup>11,25</sup> Compared with that of the  $H^1$  atom of  $H_5O_2^+$  and  $H_7O_3^+$ , the  $H^1$  atom in  $H_3O^+$  carries a more positive charge (see Table S1 in ESI†). This means that  $H_3O^+$  is more acidity. The calculated results indicated that the dihydrogen release pathway is less favorable than the hydride transfer pathway (Path A) for the reaction mediated by  $H_5O_2^+$  or  $H_7O_3^+$  (see Fig. 5 and S4 in ESI†).

Fig. 8 demonstrates the energy profiles of the metal hydride formation pathway and the metal hydroxo formation pathway under acidic and basic conditions. The energy barrier of the metal hydride formation is  $14.1 \text{ kcal mol}^{-1}$ , which may also be formed under acidic, neutral or basic conditions. However, for the metal hydroxo formation pathway, the free energy changes dramatically in solutions of different pH values. Under acidic conditions, the metal hydride formation pathway is more favorable than the metal hydroxo formation pathway, which is endergonic by  $18.4 \text{ kcal mol}^{-1}$ . In contrast, the metal hydride formation pathway is less favorable than the metal hydroxo

formation pathway under basic conditions. The metal hydroxo formation pathway is exergonic by  $42.1 \text{ kcal mol}^{-1}$  under basic conditions, indicating that 8 is very easily formed under basic conditions. The calculated results indicate that different pH values can change the reactivity of the reaction, which may switch the preference for ketone hydrogenation or dihydrogen release in the mechanism.

### The modes of dihydrogen activation

Fig. 9 shows three typical HH or TH catalysts and their modes of dihydrogen activation. While catalyst A<sup>30</sup> is an efficient HH catalyst, catalysts C<sup>30,63</sup> and 2 (ref. 49) show good TH activities. Meanwhile, catalyst C was reported to be able to drive HH reactions under acidic conditions.<sup>17,64,65</sup> Our previous theoretical works investigated the preference for the TH or HH of ketones and concluded that  $H_2$  coordination is essential for dihydrogen activation.<sup>3</sup> Under neutral or basic conditions, the  $16e$  species  $RuN_C$ , with strong delocalized  $\pi$ -bonds, is hard to break to provide a vacant d-orbital for  $H_2$  coordination, thus the  $H_2$  activation barrier is high. On the contrary,  $RuN_A$  is a  $16e$  species, which could provide a vacant d-orbital for  $H_2$  coordination, thus  $H_2$  could be activated. Whereas under acidic conditions, dihydrogen can easily coordinate with  $RuN_C$  to form a stable  $\eta^2-H_2$  intermediate with the assistance of TfOH, which indicates that protonic acid interrupts the hyperconjugative effect of the Ru–N double bonds, resulting in a significant reduction in the energy barrier for dihydrogen activation. The single-site TM catalysts usually operate *via* non-cooperation mechanisms including the classical oxidative addition/reductive elimination mechanism and the  $\sigma$ -bond metathesis mechanism.<sup>28</sup> Interestingly, the catalytic species 2 does not have a M–H/N–H bifunctional framework and the mode of dihydrogen activation catalyzed by 2 is different from those catalyzed by A and C (metal–ligand bifunctional cooperation). It is obvious that dihydrogen heterolytic splitting, activated by 2, belongs to the ion-pair mechanism, which requires metal-solvent cooperation (see TS18-19 in Fig. 9). The  $H_2$  with assistance from the solvent,  $H_2O$ , prefer to coordinate with the TM center similar to an end-on coordination mode. The complex 2 shows only one open coordination site upon  $H_2O$  ligand

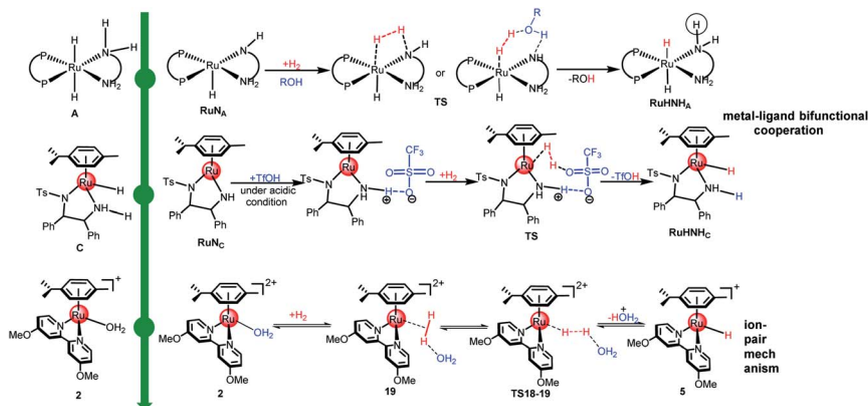


Fig. 9 Three typical TM catalysts for HH or TH of ketones and the modes of dihydrogen activation.



dissociation. The dihydrogen activation mode of the single-site complex **2** does not change the oxidation state of the Ru center of complex **2**. On the contrary, the energy barrier of H<sub>2</sub> activation is only 17.4 kcal mol<sup>-1</sup> (see Fig. 7), which is the reverse reaction of H<sub>2</sub> release. This implies that complex **2** is not only an efficient catalyst for TH but also an efficient catalyst for H<sub>2</sub>-hydrogenation in acidic aqueous media based on this calculation.

## Conclusions

In summary, we investigated the transfer hydrogenation (TH) of ketones and dihydrogen release catalyzed by  $[(\eta^6\text{-arene})\text{RuCl}(\kappa^2\text{-}N,N\text{-dmobpy})]^+$  complexes in different pH environments using the DFT method. TH of ketones proceeds *via* two sequential steps: the formation of a metal hydride and the second hydride transfer *via* an ion-pair mechanism. The calculated results show that metal hydride formation is the rate-determining step with a free energy barrier of 14.1 kcal mol<sup>-1</sup>. In addition, we analyzed the origin of pH-dependent transfer hydrogenation and dihydrogen release catalyzed by this single-site cyclometallated ruthenium complex. Under weakly acidic conditions, the favorable pathway is the metal hydride formation pathway and the hydride transfer pathway *via* an ion-pair mechanism to complete the TH of ketones. Under strongly acidic conditions, the dehydrogenation of formic acid can be achieved to release dihydrogen following the metal hydride formation pathway and the dihydrogen release pathway *via* an ion-pair mechanism. This is in agreement with experimental results. The dihydrogen activation mode of this reaction adopts an ion-pair mechanism, which is metal-solvent cooperative in nature. Such single-site cyclometallated TM catalysts are predicted to be able to drive H<sub>2</sub>-hydrogenation under acidic conditions. Meanwhile, under basic conditions, the active species  $[(\eta^6\text{-arene})\text{Ru}(\text{H}_2\text{O})(\kappa^2\text{-}N,N\text{-dmobpy})]^{2+}$  is easily deprotonated to form the hydroxo complex  $[(\eta^6\text{-arene})\text{Ru}(\text{OH})(\kappa^2\text{-}N,N\text{-dmobpy})]^+$ , which leads to the termination of the reaction.

## Conflicts of interest

The authors declare no conflict of interest.

## Acknowledgements

This work is supported by the National Natural Science Foundation of China (Grant No. 21672018), the State Key Laboratory of Physical Chemistry of Solid Surfaces (Xiamen University) (No. 201811), and the Fundamental Research Funds for the Central Universities (XK1802-6). We thank the National Supercomputing Center in Tianjin (TianHe-1) for providing part of the computational sources.

## References

- S. E. Clapham, A. Hadzovic and R. H. Morris, *Coord. Chem. Rev.*, 2004, **248**, 2201–2237.
- Y. Liu, X. Yue, C. Luo, L. Zhang and M. Lei, *Energy Environ. Mater.*, 2019, 1–21.
- M. Lei, W. Zhang, Y. Chen and Y. Tang, *Organometallics*, 2010, **29**, 543–548.
- G. Zassinovich, G. Mestroni and S. Gladioli, *Chem. Rev.*, 1992, **92**, 1051–1069.
- C. F. de Graauw, J. A. Peters, H. van Bekkum and J. Huskens, *Synthesis*, 1994, **1994**, 1007–1017.
- R. Noyori and S. Hashiguchi, *Acc. Chem. Res.*, 1997, **30**, 97–102.
- C. Wang, X. Wu and J. Xiao, *Chem.-Asian J.*, 2008, **3**, 1750–1770.
- T. Ikariya and A. J. Blacker, *Acc. Chem. Res.*, 2007, **40**, 1300–1308.
- F. Pena-Pereira, A. Kloskowski and J. Namieśnik, *Green Chem.*, 2015, **17**, 3687–3705.
- P. J. Dunn, *Chem. Soc. Rev.*, 2012, **41**, 1452–1461.
- J.-t. Liu, S. Yang, W. Tang, Z. Yang and J. Xu, *Green Chem.*, 2018, **20**, 2118–2124.
- X. Wu, X. Li, F. King and J. Xiao, *Angew. Chem., Int. Ed.*, 2005, **44**, 3407–3411.
- M. Yamakawa, H. Ito and R. Noyori, *J. Am. Chem. Soc.*, 2000, **122**, 1466–1478.
- R. Noyori, M. Yamakawa and S. Hashiguchi, *J. Org. Chem.*, 2001, **66**, 7931–7944.
- P. A. Dub and T. Ikariya, *J. Am. Chem. Soc.*, 2013, **135**, 2604–2619.
- O. Pàmies and J.-E. Bäckvall, *Chem.-Eur. J.*, 2001, **7**, 5052–5058.
- Y. Chen, Y. Tang, S. Liu, M. Lei and W. Fang, *Organometallics*, 2009, **28**, 2078–2084.
- X. Zhang, X. Guo, Y. Chen, Y. Tang, M. Lei and W. Fang, *Phys. Chem. Chem. Phys.*, 2012, **14**, 6003–6012.
- R. Feng, A. Xiao, X. Zhang, Y. Tang and M. Lei, *Dalton Trans.*, 2013, **42**, 2130–2145.
- T. Koike and T. Ikariya, *Adv. Synth. Catal.*, 2004, **346**, 37–41.
- D. G. Blackmond, M. Ropic and M. Stefinovic, *Org. Process Res. Dev.*, 2006, **10**, 457–463.
- H. Y. Chen, C. Wang, X. Wu, X. Jiang, C. R. Catlow and J. Xiao, *Chem.-Eur. J.*, 2015, **21**, 16564–16577.
- Z. Yang, Z. Zhu, R. Luo, X. Qiu, J.-t. Liu, J.-K. Yang and W. Tang, *Green Chem.*, 2017, **19**, 3296–3301.
- S. Ogo, N. Makihara and Y. Watanabe, *Organometallics*, 1999, **18**, 5470–5474.
- T. Abura, S. Ogo, Y. Watanabe and S. Fukuzumi, *J. Am. Chem. Soc.*, 2003, **125**, 4149–4154.
- J. Canivet, L. Karmazin-Brelot and G. Süß-Fink, *J. Organomet. Chem.*, 2005, **690**, 3202–3211.
- Y. Wei, D. Xue, Q. Lei, C. Wang and J. Xiao, *Green Chem.*, 2013, **15**, 629–634.
- Z. Ke, Y. Li, C. Hou and Y. Liu, *Phys. Sci. Rev.*, 2018, **3**, 20170038.
- Y. Shvo, D. Czarkie, Y. Rahamim and D. F. Chodosh, *J. Am. Chem. Soc.*, 1986, **108**, 7400–7402.
- T. Ohkuma, H. Ooka, S. Hashiguchi, T. Ikariya and R. Noyori, *J. Am. Chem. Soc.*, 1995, **117**, 2675–2676.



- 31 X. Wu, J. Liu, X. Li, A. Zanotti-Gerosa, F. Hancock, D. Vinci, J. Ruan and J. Xiao, *Angew. Chem., Int. Ed.*, 2006, **45**, 6718–6722.
- 32 S. Ogo, T. Abura and Y. Watanabe, *Organometallics*, 2002, **21**, 2964–2969.
- 33 Y. Maenaka, T. Suenobu and S. Fukuzumi, *J. Am. Chem. Soc.*, 2012, **134**, 9417–9427.
- 34 E. N. Yurtchenko and N. P. Anikeenko, *React. Kinet. Catal. Lett.*, 1975, **2**, 65–72.
- 35 X. Yang, *Dalton Trans.*, 2013, **42**, 11987–11991.
- 36 J. H. Merrifield and J. A. Gladysz, *Organometallics*, 1983, **2**, 782–784.
- 37 C. P. Casey, S. W. Singer and D. R. Powell, *Can. J. Chem.*, 2001, **79**, 1002–1011.
- 38 J. S. Samec, A. H. Ell, J. B. Aberg, T. Privalov, L. Eriksson and J. E. Backvall, *J. Am. Chem. Soc.*, 2006, **128**, 14293–14305.
- 39 J. E. Martins, G. J. Clarkson and M. Wills, *Org. Lett.*, 2009, **11**, 847–850.
- 40 X. Wu, D. Vinci, T. Ikariya and J. Xiao, *Chem. Commun.*, 2005, 4447–4449, DOI: 10.1039/b507276j.
- 41 J. Xiao, X. Li, J. Blacker, I. Houson and X. Wu, *Synlett*, 2006, **2006**, 1155–1160.
- 42 X. Wu and J. Xiao, *Chem. Commun.*, 2007, 2449–2466.
- 43 J. H. Barnard, C. Wang, N. G. Berry and J. Xiao, *Chem. Sci.*, 2013, **4**, 1234–1244.
- 44 E. A. Bielinski, P. O. Lagaditis, Y. Zhang, B. Q. Mercado, C. Wurtele, W. H. Bernskoetter, N. Hazari and S. Schneider, *J. Am. Chem. Soc.*, 2014, **136**, 10234–10237.
- 45 S. Fukuzumi, T. Kobayashi and T. Suenobu, *ChemSusChem*, 2008, **1**, 827–834.
- 46 C. Guan, D. D. Zhang, Y. Pan, M. Iguchi, M. J. Ajitha, J. Hu, H. Li, C. Yao, M. H. Huang, S. Min, J. Zheng, Y. Himeda, H. Kawanami and K. W. Huang, *Inorg. Chem.*, 2017, **56**, 438–445.
- 47 D. Mellmann, P. Sponholz, H. Junge and M. Beller, *Chem. Soc. Rev.*, 2016, **45**, 3954–3988.
- 48 W. H. Wang, S. Xu, Y. Manaka, Y. Suna, H. Kambayashi, J. T. Muckerman, E. Fujita and Y. Himeda, *ChemSusChem*, 2014, **7**, 1976–1983.
- 49 C. Aliende, M. Pérez-Manrique, F. A. Jalón, B. R. Manzano, A. M. Rodríguez and G. Espino, *Organometallics*, 2012, **31**, 6106–6123.
- 50 H. Li, X. Ma, B. Zhang and M. Lei, *Organometallics*, 2016, **35**, 3301–3310.
- 51 L. Li, M. Lei and S. Sakaki, *Organometallics*, 2017, **36**, 3530–3538.
- 52 L. Li, M. Lei, Y. Xie, H. F. Schaefer, 3rd, B. Chen and R. Hoffmann, *Proc. Natl. Acad. Sci. U. S. A.*, 2017, **114**, 9803–9808.
- 53 L. Li, H. Zhu, L. Liu, D. Song and M. Lei, *Inorg. Chem.*, 2018, **57**, 3054–3060.
- 54 M. Xiao, X. Yue, R. Xu, W. Tang, D. Xue, C. Li, M. Lei, J. Xiao and C. Wang, *Angew. Chem., Int. Ed.*, 2019, **58**, 10528–10536.
- 55 X. Yue, L. Li, P. Li, C. Luo, M. Pu, Z. Yang and M. Lei, *Chin. J. Chem.*, 2019, **37**, 883–886.
- 56 J. D. Chai and M. Head-Gordon, *Phys. Chem. Chem. Phys.*, 2008, **10**, 6615–6620.
- 57 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery Jr, J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, T. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, O. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, *Gaussian 09, Revision B.01*, Gaussian, Inc., Wallingford CT, 2010.
- 58 A. V. Marenich, C. J. Cramer and D. G. Truhlar, *J. Phys. Chem. B*, 2009, **113**, 6378–6396.
- 59 C. Lee, W. Yang and R. G. Parr, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1988, **37**, 785–789.
- 60 P. J. Hay and W. R. Wadt, *J. Chem. Phys.*, 1985, **82**, 299–310.
- 61 Z. Wang, S. M. Lu, J. Li, J. Wang and C. Li, *Chem.-Eur. J.*, 2015, **21**, 12592–12595.
- 62 C. A. Mebi, R. P. Nair and B. J. Frost, *Organometallics*, 2007, **26**, 429–438.
- 63 A. Fujii, S. Hashiguchi, N. Uematsu, T. Ikariya and R. Noyori, *J. Am. Chem. Soc.*, 1996, **118**, 2521–2522.
- 64 C. A. Sandoval, T. Ohkuma, N. Utsumi, K. Tsutsumi, K. Murata and R. Noyori, *Chem.-Asian J.*, 2006, **1**, 102–110.
- 65 T. Ohkuma, K. Tsutsumi, N. Utsumi, N. Arai, R. Noyori and K. Murata, *Org. Lett.*, 2007, **9**, 255–257.

