

Highly Robust Hydrogen Generation by Bio- Inspired Ir Complexes for Dehydrogenation of Formic Acid in Water: Experimental and Theoretical Mechanistic Investigations at Different pH

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ABSTRACT: Hydrogen generation from formic acid (FA), one of the most promising hydrogen storage materials, has attracted much attention due to the demand for the development of renewable energy carriers. Catalytic dehydrogenation of FA in an efficient and green manner remains challenging. Here, we report a series of bio-inspired Ir complexes for highly robust and selective hydrogen production from FA in aqueous solutions without organic solvents or additives. One of these complexes bearing an imidazoline moiety (complex **6**) achieved a turnover frequency (TOF) of $322,000 \text{ h}^{-1}$ at $100 \text{ }^\circ\text{C}$, which is higher than ever reported. The novel catalysts are very stable and applicable in highly concentrated FA. For instance, complex **3** ($1 \text{ } \mu\text{mol}$) affords an unprecedented turnover number (TON) of 2,050,000 at $60 \text{ }^\circ\text{C}$. Deuterium kinetic isotope effect experiments and density functional theory (DFT) calculations employing a “speciation” approach demonstrated a change in the rate determining step with increasing solution pH. This study provides not only more insight into the mechanism of dehydrogenation of FA, but also offers a new principle for the design of effective homogeneous organometallic catalysts for H_2 generation from FA.

KEYWORDS: Formic acid dehydrogenation, Ir complexes, Mechanism, Kinetic isotope effect, pH-dependence

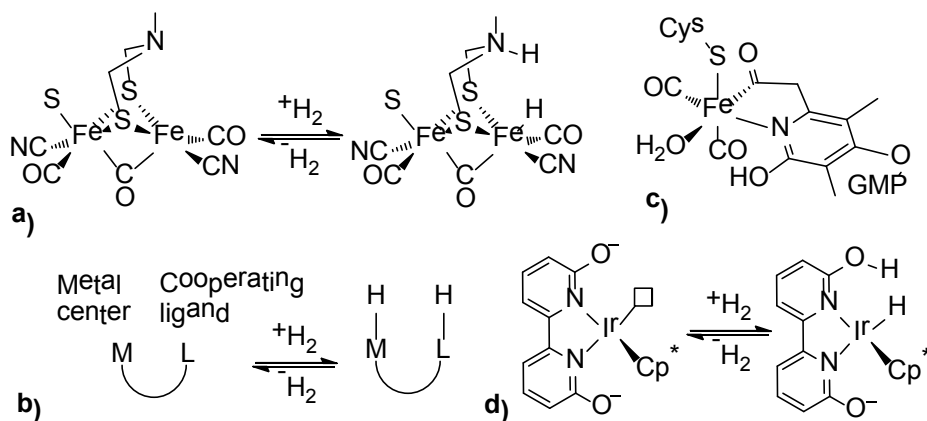
Introduction

A hydrogen economy is a promising alternative to the hydrocarbon economy for solving increasingly severe energy and environmental problems.^{1,2} Efficient hydrogen storage is one of the bottlenecks for developing a hydrogen economy.³⁻⁵ Recently, hydrogen storage using formic acid (FA) as a carrier has attracted much attention because of its favorable properties.⁶⁻⁹ While

FA has the relatively low hydrogen content of 4.4 wt%, it is non-toxic, bio-degradable, and environmental friendly. More importantly, it is a liquid under ambient conditions, which makes it easy to store, transport and handle. FA can be selectively generated by catalytic hydrogenation of CO₂ with a suitable catalyst under mild conditions and can also be used in fuel cells directly. Moreover, FA dehydrogenation can selectively release pressurized H₂/CO₂ with an appropriate catalyst. In some cases, the hydrogen storage capability (i.e., reversible interconversion between CO₂ and FA) can be achieved using the same catalyst by changing reaction conditions such as pressure, temperature, and pH.¹⁰⁻¹⁷

Various heterogeneous and homogeneous catalysts for FA dehydrogenation have been reported.^{6-8,18} Despite recent important progress,¹⁹⁻²² heterogeneous catalysts often suffer from low activity and selectivity (including the generation of CO that poisons fuel cell catalysts).^{8,19} Homogeneous catalysts are generally more effective and attractive,^{10,23-29} but are far from practical use owing to their limited catalytic efficiency and robustness. The design of highly efficient and robust homogenous catalysts remains a significant challenge. Nature uses hydrogenase enzymes to accomplish reversible H₂ oxidation (i.e., H₂ splitting and generation, Scheme 1a). An important synergistic effect between the metal center and cooperating ligands in hydrogenases has inspired many chemists to explore such a catalyst design principle (Scheme 1b).³⁰⁻³³ DuBois and Bullock et al. have developed a series of complexes with a pendent amine mimicking the structure of [FeFe]-hydrogenase to catalyze various transformations.³⁴⁻³⁶ Pincer complexes with non-innocent ligands have also shown outstanding catalytic activity in FA dehydrogenation in organic solvents.^{10,28,37} Inspired by the active center of [Fe]-hydrogenase (Scheme 1c),^{38,39} we have designed and synthesized a series of proton-responsive complexes (Scheme 1d) for efficient catalytic interconversion of formic acid/formate and H₂/CO₂ in

water.^{14,40-42} The bipyridine or bipyrimidine ligands bearing pendent OH groups are redox non-innocent, proton-responsive, and act as cooperating ligands which have been demonstrated to be largely responsible for the high activity of the complexes.⁴³ Moreover, the proton-responsive complexes bearing functional OH groups showed tunable activity and can control the equilibrium between formic acid/formate and H₂/CO₂ under different pH conditions. Under acidic conditions, they decompose FA to release H₂/CO₂, while under basic conditions they catalyze CO₂ hydrogenation to provide formate. This unique property is mainly attributed to the pH switchable OH substituent which deprotonates to give the strong electron-donating oxyanion and improves the activity. Therefore, determination of the exact species in aqueous solution under different pH is highly desirable.



Scheme 1. Bio-inspired catalyst design for H₂ splitting and generation: (a) The H₂ splitting and generation processes catalyzed by [FeFe]-hydrogenase; (b) Catalyst design with cooperating ligand bearing functional groups for H₂ activation and generation by mimicking the enzyme; (c) Structure of the guanylyl pyridinol (FeGP) cofactor in [Fe]-hydrogenase (GMP: guanine monophosphate); (d) H₂ splitting and generation with our complexes via a synergistic effect between the metal center and cooperating ligands.

Although various homogeneous catalysts with precious metals or earth-abundant metals have been developed for FA dehydrogenation, high performance (i.e., high TON and TOF) catalysts usually require the presence of organic solvents or additives.^{10,27,28,44} Catalysts based on earth-abundant metals typically suffer from low selectivity, and CO contamination is usually observed.^{26,28,45} The presence of the amine in a FA/amine system considerably decreases the hydrogen capacity and results in loss of the volatile amine. In contrast, FA dehydrogenation in water without organic additives is eco-friendly, and a highly concentrated FA solution can also provide comparable hydrogen capacity. However, reports of highly active and stable catalysts in aqueous solution are scarce.^{16,46-48} We recently reported that a five-membered azole moiety is an efficient electron donor, and its incorporation into a catalyst complex is effective for FA dehydrogenation in water.⁴⁹ This is supposedly due to the high electron donating ability of the azole moieties. Taking into account that pyridine and pyrimidine ligands bearing pendent OH groups are effective for H₂ activation, we focused on designing new catalysts that combine pyridine or pyrimidine moieties bearing pendent OH groups with electron-donating azole and azoline moieties including pyrazole, imidazole and imidazoline. Herein, we report unprecedentedly durable bio-inspired complexes for catalyzing FA dehydrogenation in water (TON of 2,050,000 at 60 °C). Furthermore, an initial TOF as high as 332,000 h⁻¹ could be achieved in 4 M HCO₂H/HCO₂Na (3/2) using a complex based on an imidazoline ligand.

The deuterium kinetic isotope effect (KIE) is an effective method for exploring reaction mechanisms.^{10,50-52} We have previously investigated the mechanism of formic acid dehydrogenation with KIE studies,⁵⁰ and demonstrated a significant effect depending on the specific ligands with or without pendent OHs in the complexes. Complexes with 6,6'-dihydroxy-

2,2'-bipyridine (OH substituents in *ortho* positions) exhibit a different pH dependence than those with 4,4'-dihydroxy-2,2'-bipyridine (OH substituents in *para* positions) owing to the two classes of complexes having different rate determining steps (RDS) in the catalytic cycle.⁴¹ Dehydrogenation of formic acid is known to exhibit a pH dependence, which suggests that the solution pH has an important influence on the reaction rate,^{14,52-54} however, to the best of our knowledge, the effect of solution pH on the mechanism has not yet been comprehensively studied. Here we demonstrate for the first time that the rate determining step for a catalyst that has a 6,6'-dihydroxy-2,2'-bipyridine-like ligand (OH substituent in an *ortho* position) changes upon altering the solution pH, and rationalize the bell-shaped pH dependence of the activity of such catalysts through a combined experimental and computational effort. In the calculations, we employed a “speciation” approach which helps to determine the actual species (i.e., fully deprotonated or partially deprotonated) in the reaction solution.

Results and discussion

Catalyst screening

The water-soluble complexes **1-6** (Chart 1) were synthesized using methods similar to those previously reported (see SI). With these complexes in hand, we carried out the catalytic dehydrogenation of formic acid in aqueous solutions at low FA concentration (1 M) and temperature (60 °C, Table 1). Complex **1** bearing a MeO group showed moderate activity (TOF: 1010 h⁻¹, entry 1) in 1 M HCO₂H solution (pH 1.7) and apparent decreased activity (TOF: 160 h⁻¹, entry 2) in 1 M HCO₂H/HCO₂Na (1/1, pH 3.5). In contrast, complex **2** with a pendent OH group showed significant improvement in activity and stability (entries 3 and 4). The TOF increased to 2740 h⁻¹ in 1 M HCO₂H, and further increased to 3620 h⁻¹ in 1 M HCO₂H/HCO₂Na

(1/1, pH 3.5). These pH dependences are consistent with our previous report that complexes with pendent OH groups in the second coordination sphere exhibit a bell-shaped pH vs. rate profile.⁴¹ In 1 M FA solution, complex **3** gave an initial TOF of 7050 h⁻¹ at 60 °C (entry 5). The initial TOF in 1 M HCO₂H/HCO₂Na (1/1, pH 3.5) reached 11,400 h⁻¹, which is triple that of complex **2** (entry 6). Similar to all the complexes with pendent OH groups, its pH vs. rate profile also was bell-shaped (Figure 1), and the TOF peaked at pH 2.8 (TOF: 18,000 h⁻¹, entry 7). Changing the position of the N in the pyrimidine ring led to a considerable decrease in the activity of the complex as shown in the pH vs. rate profile (Figure 1, **3** vs. **4**). Complex **4** gave a TOF of 5150 h⁻¹ at pH 1.7 and 60 °C (entry 8). The TOF peaked at pH 3.0 (TOF: 10,800 h⁻¹, entry 9). Replacing the pyrazole ring with an imidazole ring led to both low stability and activity at low solution pH. Complex **5** was found to have partially precipitated from the solution at some point during the reaction at pH below 2.6, leading to decreased activity. The TOF at pH 1.7 was 4290 h⁻¹, which is even lower than that of complex **4** (entry 10). Interestingly, the reaction rate increased significantly with increasing solution pH. The maximum TOF value (19,900 h⁻¹, entry 11) at pH 3.0 is even higher than that of complex **3** (Figure 1, **3** vs. **5**). When complex **6** bearing an imidazoline ring was used, we observed no apparent catalyst precipitation, but similar to complex **5** the reaction rate increased significantly with increasing solution pH (Figure 1, **6**). It exhibited the highest initial TOF of 32,500 h⁻¹ at pH 3.0 (entry 13). The catalytic efficiency of complex **6** is even higher than the most efficient Ir dinuclear THBPM complex under similar conditions.¹⁴

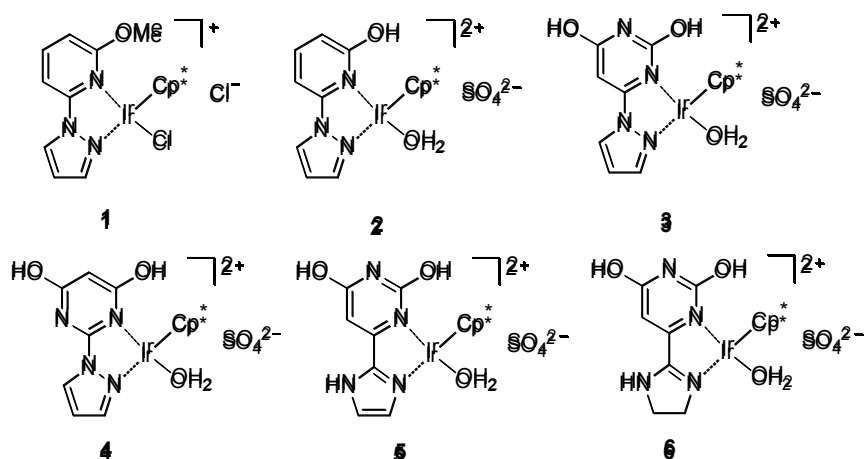


Chart 1. Novel biomimetic complexes for FA dehydrogenation in aqueous solution.

Table 1. Dehydrogenation of FA with Ir complexes **1-6**.^a

Entry	Catalyst / μM	Solution pH	Time / h	TON	TOF ^b / h^{-1}
1	1 / 100	1.7	5	2570	1010
2	1 / 100	3.5	5	430	160
3	2 / 100	1.7	7	10,000	2740
4	2 / 100	3.5	7	5000	3620
5	3 / 100	1.7	7	10,000	7050
6	3 / 100	3.5	1	5500	11,400
7	3 / 100	2.8	1.5	8700	18,000
8	4 / 100	1.7	3.5	10,000	5150
9	4 / 100	3.0	1.5	7640	10,800
10	5 / 100	1.7	5	10,000	4290
11	5 / 100	3.0	1	7850	19,900
12	6 / 100	1.7	3	10,000	5520
13	6 / 100	3.0	0.5	7850	32,500

^a Reaction conditions: 1 M HCO_2H or $\text{HCO}_2\text{H}/\text{HCO}_2\text{Na}$ solution (10 mL) with Ir complexes 1 μmol at 60 °C. All numbers are an average of two runs. FA dehydrogenation reactions with complexes **2-6** yielded complete FA decomposition. ^b Average TOF over initial 10 min.

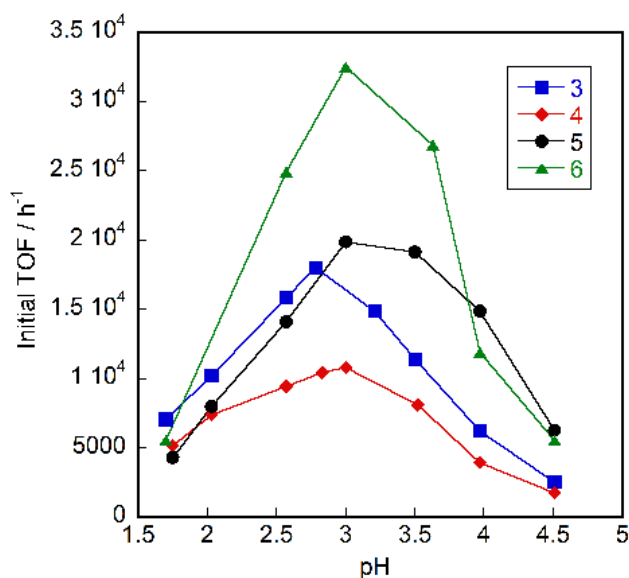


Figure 1. pH dependence of FA dehydrogenation using complexes **3-6** (100 μM) in 1 M $\text{HCO}_2\text{H}/\text{HCO}_2\text{Na}$ (10 mL) solution at 60 $^\circ\text{C}$.

Durability tests

For practical use in FA dehydrogenation, a high concentration of FA and a high temperature are required. Therefore, we carried out FA dehydrogenation under harsher conditions with complexes **3-6** that showed remarkable activity in the preliminary experiments. The experiments examining the temperature dependence with complex **3** suggested that the reaction rate increased significantly upon increasing the reaction temperature from 45 to 70 $^\circ\text{C}$. (SI, Figure S1). The FA concentration was also found to be an important factor for the increase in the reaction rate. The reaction rate reached a maximum in 4 M FA solution using either complex **3** or **4** (SI, Figures S2 and S4). In addition, the capability of generating high pressure gases is desirable for the convenience of gas delivery. We carried out the FA dehydrogenation with complex **3** in a closed

system. The inner pressure of the autoclave reached a final pressure of 4 MPa after the reaction completed (SI, Figure S5). According to the HPLC analysis, 99.4 % of the FA was converted. This is consistent with our previous reports^{14,49} and suggests that high pressure does not inhibit the catalytic activity of complex **3**. Therefore, under acidic conditions, the equilibrium between CO₂/H₂ and HCO₂H of this catalytic system is strongly biased toward gas generation, and pressure only weakly affects the equilibrium. Moreover, no CO was detected by GC (detection limit 1 ppm) from the released gas. This indicates that the catalytic reaction is highly selective and goes through no dehydration, which generates CO and H₂O. These properties are important for providing high pressure H₂ which facilitates gas delivery and utilization in fuel cells because CO is a poison for the catalyst in fuel cell electrodes.

Complexes **3** and **4** were found to be stable in highly concentrated FA solutions at both low (60 °C, SI, Figure S6) and high (100 °C, SI, Figures S9-S11) temperature. The reaction results in highly concentrated FA solution and high temperature are listed in Table 2. All the FA was completely decomposed at 100 °C (110 °C bath temperature; the actual reaction temperatures measured were close to 100 °C) with only 20 μM catalyst concentration (Table 2, entries 1-4). In contrast, complexes **5** and **6** were relatively less stable under the same conditions. Higher catalyst concentration (100 μM) was required for complete conversion of the FA (entries 5 and 6). Complex **3** showed excellent stability. This complex is quite stable in highly concentrated FA solutions (even 80 wt% FA), and all the FA could be dehydrogenated (TON: 200,000) despite the fact that the TOF decreased considerably (SI, Figure S7). Interestingly, we observed an apparent increase of reaction rate with consumption of FA and decrease of FA concentration. The rate around 9000 min is higher than that at the initial stage or around 1500 min. This is in agreement with our study of the effect of FA concentration as shown in Figures S2 and S4.

Complex **3** gave the high TOF of 178,000 h⁻¹ at 100 °C in 4 M FA (entry 1). The initial reaction rate decreased slightly upon increasing the FA concentration (entries 1-3). The reaction in 8 M FA was finished in 4 h and exhibited the high TON of 400,000 (entry 3). Using complex **4** in 4 M FA at 100 °C, the TOF of 120,000 h⁻¹ was slightly lower than that of complex **3** (entry 4 vs. entry 1). We obtained relatively lower TOF values with complexes **5** and **6** in 4 M FA (entries 5 and 6). As the pH vs. rate profile suggested, addition of formate can significantly improve the reaction rate. Thus we tried a 4 M HCO₂H/HCO₂Na (98/2) solution at 100 °C (entry 7). Accordingly, complex **3** achieved a significantly elevated initial TOF of 269,000 h⁻¹. With the same strategy, we optimized the reaction conditions and obtained the unprecedented initial TOF of 322,000 h⁻¹ with complex **6** (entry 9), which is higher than for any catalyst previously reported.^{10,25,28,55} These results indicate that both complexes **3** and **6** show much higher activity than the most effective dinuclear Ir complex previously reported.¹⁴ These water soluble complexes in aqueous media are even more efficient than the very recently reported PNP-Ru in DMF/Et₃N for FA dehydrogenation.¹⁰

Based on the results obtained in these preliminary experiments with highly stable and efficient complex **3**, we sought to maximize its TON for FA dehydrogenation. Unprecedentedly high TONs (more than 2 million) were reproducibly obtained in highly concentrated aqueous FA solutions for long reaction times (Figure 2). Sodium formate was not added to increase reaction rate because sodium formate can't be decomposed and decreases the total TON. Considering that the practical use would probably be carried out under milder conditions to achieve a lower energy cost, we carried out the reaction at the relatively lower temperature of 60 °C. The reaction proceeded smoothly using only 1 μmol of catalyst (catalyst concentration: 5 μM) in 200 mL 6 M FA aqueous solution. After 150 h, the rate of dehydrogenation became slower due to the

decrease of FA concentration. After 55 L of gas was released (167 h), a degassed 50 wt% solution of FA (74 g, approximately 0.8 mol FA) was added (see SI). The addition of FA restarted the gas generation and indicated that the catalyst was still active after the first run. Finally, 100.2 L of gas was evolved and afforded the extraordinary TON of 2,050,000 which, to the best of our knowledge, is higher than ever reported for either an aqueous or organic solvent system.

Table 2. H₂ generation from dehydrogenation of FA at high concentration and high temperature with Ir complexes **3-6**.^a

Entry	Catalyst / μ M	Conc. Substrate	& Time / h	TON	TOF ^b / h ⁻¹
1	3 / 20	4 M FA	2	200,000	178,000
2	3 / 20	6 M FA	3.5	300,000	176,000
3	3 / 20	8 M FA	4	400,000	173,000
4	4 / 20	4 M FA	3.5	200,000	120,000
5	5 / 100	4 M FA	1	40,000	89,400
6	6 / 100	4 M FA	1.5	40,000	68,600
7	3 / 20	4 M 98/2 ^d	6	196,000	269,000
8 ^c	6 / 40	4 M 80/20 ^d	5	80,000	258,000
9 ^c	6 / 40	4 M 68/32 ^d	0.5	68,000	322,000

^a Reaction conditions: 0.2-1 μ mol Ir complexes in 10 mL HCO₂H or HCO₂H/HCO₂Na solution at 100 °C (110° C bath temperature; the actual reaction temperatures measured were close to 100 °C). All numbers are an average of two runs. ^b Average TOF over initial 10 min. ^c 50 mL HCO₂H/HCO₂Na solution. ^d The ratio of HCO₂H to HCO₂Na.

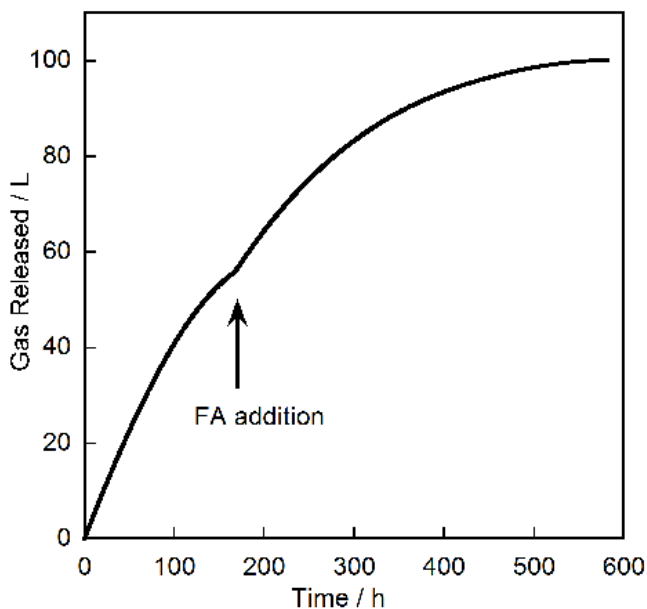
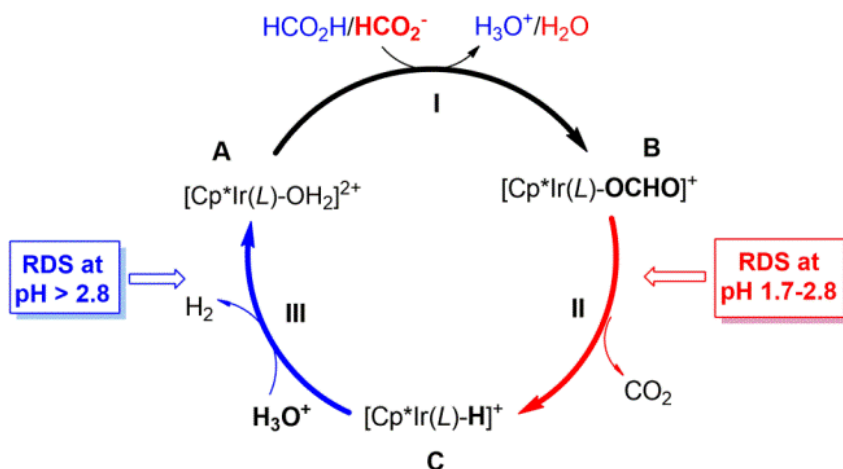


Figure 2. Time course of formic acid dehydrogenation using complex **3** (1 μmol) in a 6 M aqueous FA solution (200 mL, 1.2 mol). Additional FA (50 wt% solution 74 g, ~ 0.8 mol) at 60 $^{\circ}\text{C}$ was added after 167 h. A total of 100.2 L gas was generated (see SI).

KIE studies

The proposed mechanism for FA dehydrogenation (Scheme 2) starts with the binding of HCO_2H or formate resulting in a formate-bound intermediate **B** (Step I). Then, β -hydride elimination with concomitant CO_2 evolution leads to the formation of an Ir-hydride intermediate **C** (Step II). Following this step, H_2 is formed via proton transfer from a hydroxy group of the pyrimidine ring of **3** to an Ir-hydride moiety (Step III) assisted by a water molecule,⁴³ completing the catalytic cycle (Schemes 3 and 4).



Scheme 2. Change in rate determining step with altered solution pH for complex **3** in FA dehydrogenation.

We have reported different rate determining steps for complexes with and without pendent OH groups based on deuterium kinetic isotope effect studies.⁴¹ However, the KIE studies were limited to the regime in which the reaction rates were increasing (see Figure 1). The KIE results can explain the rate increase with increasing solution pH, but cannot explain the reaction rate decrease with further increase of solution pH. The high catalytic activity of these complexes at low pH and high pH above and below the peak motivated further studies of the KIE values at pH values on both sides of the peak of the bell-shaped activity curve. Accordingly, we performed a KIE study with complex **3**, which peaked at pH 2.8 as shown in the pH vs. rate profile of Figure 1. The reactions were carried out in 1 M formic acid solution (pH 1.7) and 1 M $\text{HCO}_2\text{H}\text{-HCO}_2\text{Na}$ (1/1) solution (pH 3.5), and the results are shown in Table 3. The KIE study at pH 1.7 suggests that DCO_2D (KIE: 2.0, entry 3) is more influential than D_2O (KIE: 1.5, entry 2) on the reaction rate. Therefore, the β -hydride elimination (Scheme 2, Step II), which involves Ir–D bond formation when DCO_2D is used, was designated as the rate-determining step. This result is

consistent with our previous report⁴¹ and other reports.^{26,51,52} The examination of FA concentration as shown in Figure S2 and S4 also supports step II as being rate-limiting around pH 1.7. The rate improvement from 1 M to 4 M FA could be explained by considering a pre-equilibrium associated with the formation of a formate bound Ir complex (**B** in Scheme 2) prior to the rate determining β -hydride elimination step (see Figure S3 and SI for details). Since formation of **B** requires deprotonation, decreasing the pH with increased FA concentration will slow down the reaction. On the other hand, increasing concentration of one of the reactants (e.g., HCO₂H) will favor formation of **B**, hence the net outcome of increasing FA concentration will be determined by these two counter-effects. Using a simple rate expression for the β -hydride elimination step including the pre-equilibrium for the formation of **B** indicates that the rate will be enhanced by 1.6 times by changing the FA concentration from 1 M to 4M which is in good agreement with the experimental observation (Figure S2). Further increase in FA concentration will slow down the deprotonation step for the formation of **B** to a greater extent and will result in a decrease in the overall reaction rate.

When a KIE study was carried out at pH 3.5 using HCO₂H-HCO₂Na in the ratio of 1/1, the results were surprising. The KIE value (2.7) using D₂O is higher than that (1.5) of using DCO₂D-DCO₂Na (entry 2 vs. entry 3), suggesting that H₂O or H₃O⁺ is involved in the rate-determining step. Therefore, as shown in Scheme 2, the RDS should be H₂ formation (Step III) when pH > 2.8. Based on these results, we propose that the rate determining step is β -hydride elimination (Scheme 2, Step II) at pH 1.7-2.8 and changes to H₂ formation (Scheme 2, Step III) when pH > 2.8 (see Computational Studies below). We carried out a similar KIE study with complex **6** (Table 4). Similarly, deuterated solvent showed much more influence than the deuterated substrate upon changing solution pH from 1.7 to 3.5. These results are consistent with those of

complex **3**, and indicate the altering of the RDS. The interesting RDS change is attributed to the loss or gain of one or more protons by the intermediate along the reaction pathway,⁵⁶ therefore detailed computational investigation of the actual forms of the intermediates of the proton responsive catalysts was performed using DFT calculations.

Table 3. Kinetic isotope effect in the dehydrogenation of FA using Ir complex **3**.^a

Entry	Substrate/ Solvent (pH 1.7)	TOF ^b / KIE ^c h ⁻¹		Substrate/ Solvent (pH 3.5)	TOF ^b / KIE ^c h ⁻¹	
	1	HCO ₂ H / H ₂ O	14,000	-	HCO ₂ H-HCO ₂ Na / H ₂ O	16,600
2	HCO ₂ H / D ₂ O	9570	1.5	HCO ₂ H-HCO ₂ Na / D ₂ O	6130	2.7
3	DCO ₂ D / H ₂ O	6870	2.0	DCO ₂ D-DCO ₂ Na / H ₂ O	11,200	1.5
4	DCO ₂ D / D ₂ O	4430	3.2	DCO ₂ D-DCO ₂ Na / D ₂ O	5090	3.3

^a Reaction conditions: 10 mL 1 M FA solution or FA/formate (1/1), 1 μmol catalyst, 70 °C.

^b Average TOF over the initial 10 min. The errors are less than 2 %.

^c KIE = TOF(entry 1)/TOF(entry n), (n = 2, 3, and 4). The exchange of all labile protons was taken into account.

Table 4. Kinetic isotope effect in the dehydrogenation of FA using Ir complex **6**.^a

Entry	Substrate/ Solvent (pH 1.7)	TOF ^b / KIE ^c h ⁻¹		Substrate/ Solvent (pH 3.5)	TOF ^b / KIE ^c h ⁻¹	
	1	HCO ₂ H / H ₂ O	13,300	-	HCO ₂ H-HCO ₂ Na / H ₂ O	28,800
2	HCO ₂ H / D ₂ O	9080	1.5	HCO ₂ H-HCO ₂ Na / D ₂ O	23,200	1.2
3	DCO ₂ D / H ₂ O	5770	2.3	DCO ₂ D-DCO ₂ Na / H ₂ O	25,300	1.1
4	DCO ₂ D / D ₂ O	3440	3.9	DCO ₂ D-DCO ₂ Na / D ₂ O	18,200	1.6

^a Reaction conditions: 10 mL 1 M FA solution or FA/formate (1/1), 1 μmol catalyst, 70 °C.

^b Average TOF over the initial 10 min. The errors are less than 2 %.

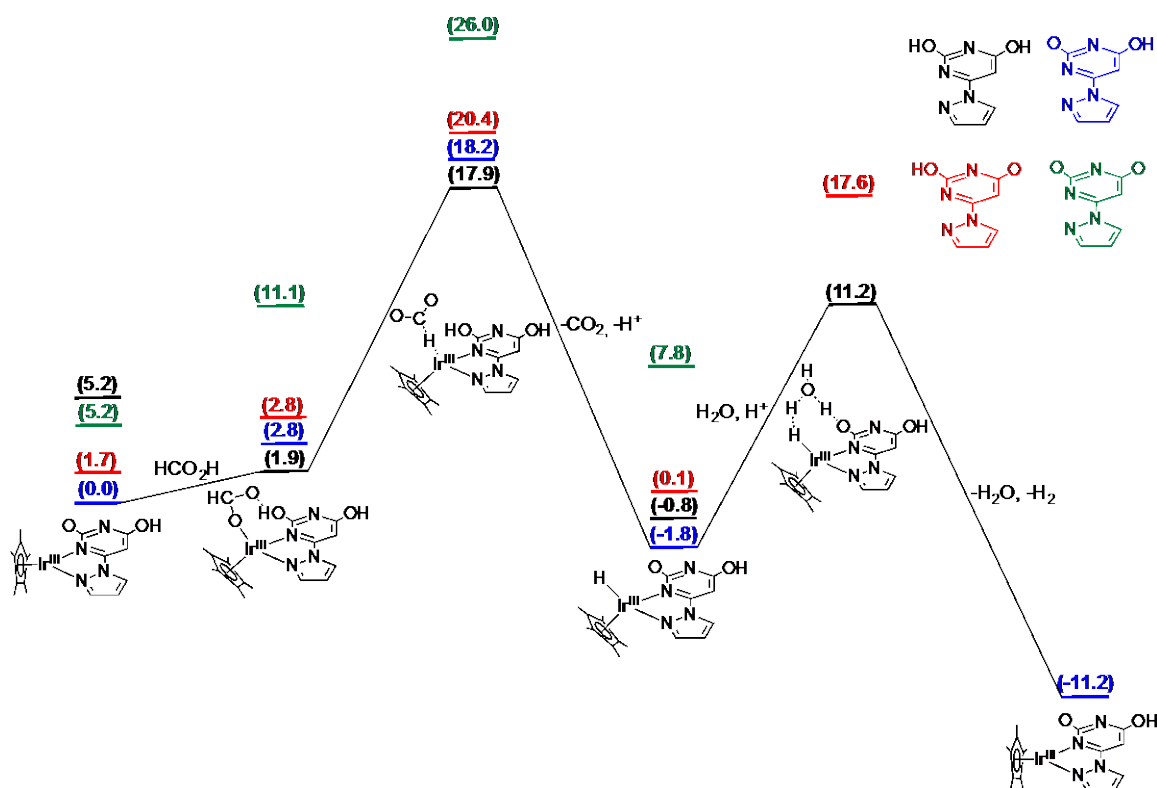
^c KIE = TOF(entry 1)/TOF(entry n), (n = 2, 3, and 4). The exchange of all labile protons was taken into account.

Computational Studies

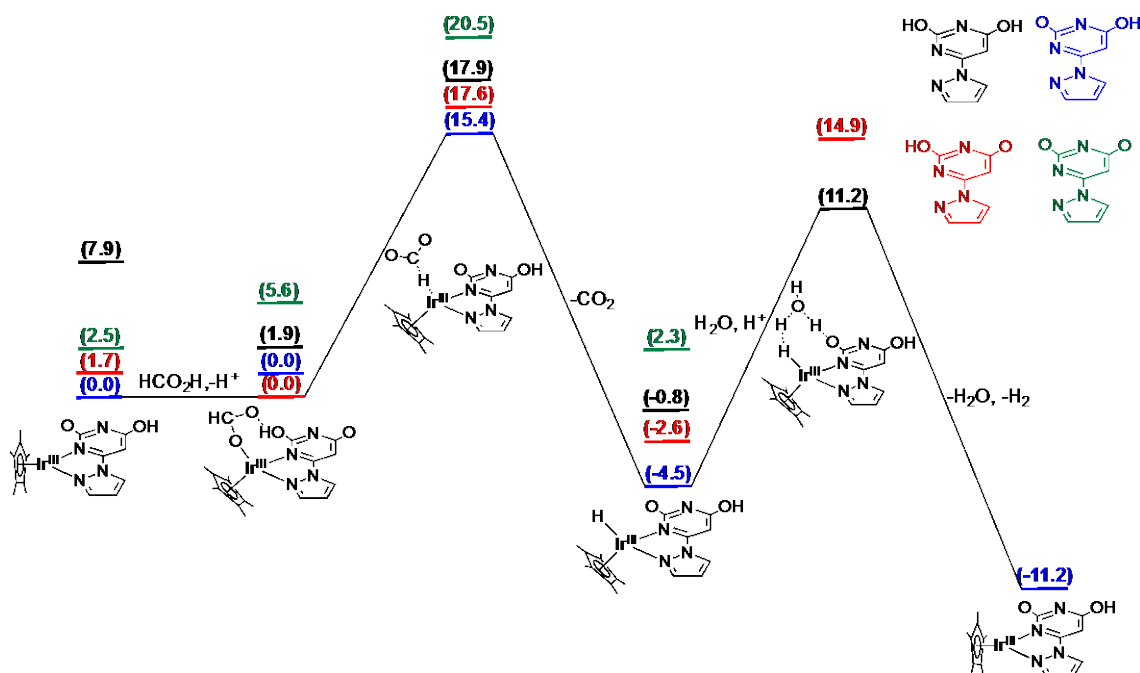
We performed density functional theory (DFT) calculations at the M06 level of theory⁵⁷ with the SMD aqueous continuum solvation model⁵⁸ (see Computational Methods in the SI for details) to investigate the mechanistic details associated with FA dehydrogenation by complexes **2**, **3**, **5**, and **6**. We have demonstrated that the proton-responsive complexes containing a proximal -O^- group can facilitate the hydrogen splitting through hydrogen bonding (Scheme 1d).⁴³ Thus, in the reverse reaction, the proximal -O^- can also assist the hydrogen formation. A sulfonamide ligand-assisted FA dehydrogenation via hydrogen bonding has been reported.⁵⁹ Further considering the different hydrogen bonding model and different electron donating ability between -OH and -O^- , it is highly desirable to determine the real forms of the proton-responsive complexes in the solution at different pH, i.e., the protonated form or a fully or partially deprotonated form. Therefore we applied a speciation approach in which we calculated the relative free energies of the intermediates and transition-state structures for all possible protonation states of the complexes and followed the changes in relative free energies with changes in pH. For instance, for FA dehydrogenation by **3**, we generated free-energy profiles at pH 1.7 and 3.5 following the relative free energies of intermediates and transition state structures differing in protonation states as depicted in Schemes 3 and 4. Similar free-energy plots for complexes **2**, **5**, and **6** are presented in the SI (Schemes S1-S19).

For the following energetic analysis of the proposed catalytic cycle, we have used the free energies associated with the most stable species for each reaction intermediate as shown in Schemes 3 and 4 connected by a black solid line. The reasoning behind this convention (or “speciation” approach) is that deprotonation of an OH group will be more facile than traversing a transition state for β -hydride elimination or H_2 formation so that the most stable protonation state

of each species will be predominant in the solution. At pH 1.7, the binding of formic acid is uphill by 1.9 kcal/mol and the following β -hydride elimination step features a free energy of activation (ΔG^\ddagger) of 16.0 kcal/mol (Scheme 3). H_2 formation from the generated Ir-hydride species occurs with $\Delta G^\ddagger = 13.0$ kcal/mol, releasing $\text{H}_{2(\text{g})}$ and regenerating the catalyst (Scheme 3). At pH 3.5, the formation of the formate complex requires no energy input ($\Delta G = 0.0$ kcal/mol) and the ΔG^\ddagger for the following β -hydride elimination step is 15.4 kcal/mol (Scheme 4). The following H_2 formation step involves a ΔG^\ddagger of 15.7 kcal/mol at pH 3.5 (Scheme 4). The calculated ΔG^\ddagger s are 16.0 kcal/mol and 13.0 kcal/mol at pH 1.7 corresponding to the β -hydride elimination and H_2 generation steps, respectively, indicating that the former is the rate-determining step (Scheme 3). The calculated overall energy barrier for the β -hydride elimination step (17.9 kcal/mol) including the free-energy cost for HCO_2H binding is consistent with the experimental observation of an activation energy of 18.7 kcal/mol (SI, Figure S1).



Scheme 3. Proposed mechanism for FA dehydrogenation by complex **3** at pH 1.7 and 333.15 K. The relative free energies are reported in units of kcal/mol. The speciation of the protonation state of each intermediate and transition state is indicated by the color code of the structures in the upper right of the scheme.



Scheme 4. Proposed mechanism for FA dehydrogenation by complex **3** at pH 3.5 and 333.15 K. The relative free energies are reported in units of kcal/mol. The speciation of the protonation state of each intermediate and transition state is indicated by the color code of the structures in the upper right of the scheme.

On the other hand, at pH 3.5, $\Delta G^\ddagger = 15.4$ kcal/mol for β -hydride elimination and $\Delta G^\ddagger = 15.7$ kcal/mol for H_2 generation, indicating that the latter is becoming the rate-determining step as the pH is increased (Scheme 4). Based on these data, the β -hydride elimination is the rate-determining step at low pH, but the free-energy cost for H_2 generation increases with increasing

pH. The catalyst displays the highest rate at moderate pH when the ΔG^\ddagger s of the two steps are very close, consistent with the experimental observation of the bell shaped pH vs. rate profile (Figure 1) and the H₂ generation step becoming rate-determining with further increase in pH as suggested by the KIE experiments. It should be kept in mind that the difference in the ΔG^\ddagger s at pH 3.5 is probably within the error in both the calculated and experimental values, but the trend from pH 1.7 should be more reliable.

Table 5. Calculated kinetic isotope effects for β -hydride elimination and H₂ generation steps for complex **3** at T = 343.15 K.

Substrate/Solvent	β -hydride elimination	H ₂ generation*
HCO ₂ H/D ₂ O	1.0	4.7
DCO ₂ D/H ₂ O	2.4	1.0
DCO ₂ D/D ₂ O	2.4	4.5

* The reactant complex, similar to the transition state, involves additional explicit water for KIE calculations. The exchange of all labile protons was taken into account.

We also calculated KIEs for the β -hydride elimination and H₂ generation steps for complex **3** associated with deuterium substitution either on the substrate or solvent (Table 5). It is clear from the calculated KIEs that β -hydride elimination is mostly affected by deuteration of formic acid and the H₂ generation step is more affected by the deuteration of water. The experimentally observed dependence of KIEs on pH, taken together with the calculated KIEs, further support the notion of a change in rate determining step from β -hydride elimination to H₂ generation with increasing pH, and at pH 3.5 is approaching the value at which that change will occur.

The calculated ΔG^\ddagger s for the β -hydride elimination and H_2 generation steps at different pH for complexes **2**, **3**, **5** and **6** provide some insight for the design of improved catalysts for FA dehydrogenation (Table 6). Comparison of ΔG^\ddagger s of the rate determining steps at pH 1.7 and 3.5 for complexes **2** and **3** indicates that the substitution of the hydroxy pyridine ligand in **2** with the dihydroxy pyrimidine ring in **3** decreases the free energies of activation, consistent with the experimentally observed enhanced activity of **3**. For complexes **2** and **3**, the ΔG^\ddagger s clearly indicate the change of RDS from β -hydride elimination to H_2 generation with increasing pH from 1.7 to 3.5. Although the ΔG^\ddagger s of complexes **5** and **6** showed no switching of RDS at 333.15 K, the ΔG^\ddagger s for the H_2 generation step are getting larger and the difference between the two steps is becoming smaller. From this viewpoint, this tendency also supports the change of RDS with increasing solution pH. From the results of our calculations, we expect the switchover point for complexes **5** and **6** to be larger than the experimental observation (pH 3.1), which could be attributed to the associated errors in the DFT calculation. We also calculated the free energies of activation at 373.15 K for complexes **3** and **6**. At this temperature, the H_2 formation step becomes the rate-determining step for both complex **3** ($\Delta G^\ddagger = 17.6$ kcal/mol) and complex **6** ($\Delta G^\ddagger = 20.0$ kcal/mol) with the former showing the highest activity at pH 2.8 whereas the latter showing enhanced activity at the slightly elevated pH of 3.1 (SI, Schemes S1-S19).

Table 6. Calculated free energies of activation (ΔG^\ddagger) associated with relevant TSs for β -hydride elimination and H_2 generation steps for complexes **2**, **3**, **5**, and **6** at $T = 333.15$ K for different pH values.

Free Energies of Activation (kcal/mol) Associated with Relevant TSs					
		Complex 2	Complex 3	Complex 5	Complex 6
pH 1.7	β -hydride elimination	17.2	16.0	18.1	18.4
	H ₂ generation	15.7	13.0	12.9	13.6
pH 3.5	β -hydride elimination	15.9	15.4	16.7	17.6
	H ₂ generation	16.1	15.7	14.8	16.4

Conclusions

We have designed and synthesized a series of biomimetic Ir complexes with hydroxyl substituted pyridine-azole and pyrimidine-azole/azoline ligands. These water-soluble complexes are demonstrated to be highly efficient and stable in catalytic FA dehydrogenation at high temperature in highly concentrated FA aqueous solutions. We have achieved the unprecedented TOF of 322,000 h⁻¹ and TON of 2,050,000 with the most effective complexes **6** and **3**, respectively. These complexes are capable of providing high pressure CO-free H₂. The extraordinary catalytic performance indicates that these complexes should be highly useful in the practical supply of H₂ from FA and contributes significantly to the development of a hydrogen economy. Investigation of the mechanism using deuterium KIE studies and DFT calculations suggests that the rate-determining steps are altered upon changing the solution pH. The DFT calculations employing a “speciation” approach provide insight into the mechanism and support the RDS change. This mechanistic insight rationalizes the bell-shaped pH vs. rate profile, and helps to determine the optimal conditions of FA dehydrogenation by addition of formate and adjustment of the solution pH.

ACKNOWLEDGMENT

Y. H. and Y. M. thank the Japan Science and Technology Agency (JST), CREST for financial support. W.-H. W. thanks the financial support from Dalian University of Technology (the Fundamental Research Funds for the Central Universities, Grant No. DUT14RC(3)082; Grant No. 844401) and National Natural Science Foundation of China (Grant No. 21402019). The work at BNL was carried out under contract DE-SC00112704 with the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, and utilized resources at the BNL Center for Functional Nanomaterials.

Supporting Information. Experimental section containing general procedures and synthesis of ligands and Ir complexes, a plot of TOF vs. temperature for FA dehydrogenation, plots of FA dehydrogenation, TOF vs. [HCO₂H], plots of time courses of FA decomposition, computational methods, free-energy profiles derived from DFT calculations for FA dehydrogenation for complexes **2-6** at pH 1.7 and 3.5, tables of optimized coordinates and energies, and ¹H NMR spectra of ligands. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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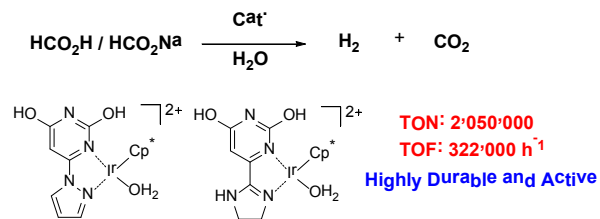
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Insert Table of Contents Graphic and Synopsis Here



Newly designed water soluble complexes exhibited unprecedented activity and durability for hydrogen generation from catalytic dehydrogenation of formic acid in aqueous solutions.

Supporting Information

Highly Robust Hydrogen Generation by Bio-Inspired Ir Complexes for Dehydrogenation of Formic Acid in Water: Experimental and Theoretical Mechanistic Investigations at Different pH

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Experimental section

General. Unless otherwise noted, materials were purchased from commercial suppliers and used without further purification. All manipulations were carried out under an argon atmosphere using standard Schlenk techniques or in a glovebox, and all aqueous solutions were degassed prior to use. ^1H NMR and ^{13}C NMR spectra were recorded on Bruker Avance 400 and 500 spectrometers using tetramethylsilane or sodium 3-(trimethylsilyl)-1-propanesulfonate as an internal standard. Elemental analyses were performed by a CE Instruments EA1110 elemental analyzer. pH values were measured on an Orion 3-Star pH meter with a glass electrode after calibration with standard buffer solutions. The amount of evolved gas was measured by a wet gas meter (Shinagawa Corp., W-NK-05; test limit: 1 mL) which is of the volumetric positive displacement type, using a chamber sealed with silicon oil. The pressure of the gases generated from the dehydrogenation of FA in a closed system was monitored by a Swagelok pressure transducer (S model, PTI-S-MG16-22AQ). ESI-MS data were collected on a Waters Micromass ZQ Mass Spectrometer. The content of the generated gas was analyzed by a GL Science GC-390 gas chromatograph. H_2 was detected by a TCD (thermal conductivity detector) using an activated 60/80 carbon column and CO_2 and CO were detected using an FID equipped with a methanizer using a Porapak Q 80/100 column at 50 °C. Formate concentrations were monitored by an HPLC on an anion-exclusion column (Tosoh TSK gel SCX(H^+)) using aqueous H_3PO_4 solution (20 mM) as eluent and a UV detector ($\lambda = 210$ nm). Water used in the reactions was obtained from a Simplicity water purification system.

General procedure for catalytic dehydrogenation of FA/formate: A freshly prepared 5 mM solution of catalyst (200 μL) was added to a deaerated 1 M aqueous HCO_2H - HCO_2Na solution (10 mL), and the mixture was stirred at the desired temperature. The reaction equipment is shown in Figure S8. The volume of evolved gas was determined by a wet gas meter. The TOF was determined according to the gas released. Since the reaction rates were fast and the reactions were usually completed in about 1 h, the average TOF of the initial 10 min was adopted for the initial TOF. After the reaction was complete, the residual FA in the solution was quantified with HPLC. The TON was calculated based on the catalyst loading and concentration of residual formate. The reactions were generally carried out more than twice and average values of the TOF are presented. Their errors are less than 5% under the same conditions of temperature and atmospheric pressure. The KIE experiments are carried out in the same method only using deuterated substrates and/or solvent.

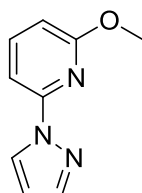
Procedure for durability test of complex 3: A freshly prepared 10 mM solution of catalyst **3** (100 μL , 1 μmol) was added to a deaerated 6 M aqueous HCO_2H solution (200 mL, 1.2 mol FA),

and the mixture was stirred at 60 °C. After 167 h, when the volume of evolved gas exceeded 55 L, the reaction solution was cooled in an ice bath to suspend the dehydrogenation. A degassed approximately 50 wt% solution of FA (74 g, ~0.8 mol FA) was added to the reaction solution. The mixture solution was warmed to 60 °C, and the gas release was restarted. After 580 h, HPLC detection indicated FA was almost completely decomposed and only 1.8 mM FA was remained in the final 190 mL reaction solution. Finally, 100.2 L of gas was evolved and afforded a total TON of 2,050,000.

Procedure for catalytic dehydrogenation of FA/formate in a closed system: A high-pressure glass cylinder (Hiper Glass Cylinder, Taiatsu Techno Co., 10 mL, Max. 5 MPa) equipped with a pressure transducer was charged with complex **3** (1 μmol) and a freshly degassed 2 M FA solution (6.5 mL). The apparatus was flushed with Ar and sealed. After it was transferred into a water bath of 70 °C, the reaction was started by vigorously stirring. The internal pressure was recorded automatically. The induction period is due to the slowly increasing temperature of the reaction solution. After the reaction was complete, the residual FA in the solution was quantified with HPLC.

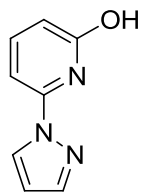
1. Synthesis of ligands

1.1. 2-methoxy-6-(1H-pyrazol-1-yl)pyridine.



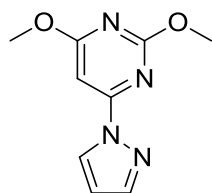
The ligands were synthesized according to the reported processes.¹ To a mixture of 2-bromo-6-methoxypyridine (0.94 g, 5 mmol), pyrazole (0.51 g, 7.5 mmol), Cu₂O (0.14 g, 1 mmol), ninhydrin (0.36 g, 2 mmol), and KOH (0.56 g, 10 mmol) was added dry DMSO (5 mL) and stirred at 120 °C under N₂ atmosphere for 24 h. After cooling to room temperature, 5 mL water was added. The resulting mixture was filtered through celite and washed with water (5 mL×2). The filtrate was extracted by dichloromethane (DCM, 10 mL×3). The combined organic phase was washed with brine (10 mL×3), dried over anhydrous Na₂SO₄, and concentrated to give the crude product as a black solid. It was then purified by column chromatography on alumina using DCM as eluent to give the product (0.76 g, 4.3 mmol, yield: 87 %) as a yellow solid. ¹H NMR (CDCl₃, 400 MHz): δ = 8.52 (d, *J* = 2.0, 1H), 7.72 (s, 1H), 7.69 (t, *J* = 8.0 Hz, 1H), 7.52 (d, *J* = 8.0 Hz, 1H), 6.63 (d, *J* = 8.0 Hz, 1H), 6.45 (s, 1H), 3.99 (s, 3H). ¹³C NMR (CDCl₃, 125 MHz): δ = 163.47, 149.83, 142.23, 141.25, 127.27, 108.05, 107.70, 104.13, 53.79.

1.2. Synthesis of 6-(1H-pyrazol-1-yl)pyridin-2-ol



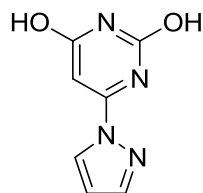
A mixture of 2-methoxy-6-(1H-pyrazol-1-yl)pyridine (0.41 g, 2.36 mmol), 4 mL HBr aq. (47 %), and 5 mL acetic acid was refluxed for 24 h. Solvent was removed under reduced pressure. To the residue, 5 mL water was added. The solution pH was adjusted to 7 with 1 M NaOH. DCM was added to extract the product. The combined organic phase was dried over anhydrous Na_2SO_4 and concentrated to give the product (0.38 g, 2.30 mmol, yield: 98%) as a white solid. ^1H NMR (CDCl_3 , 400 MHz): δ = 8.23 (d, J = 2.4 Hz, 1H), 7.71 (s, 1H), 7.65 (dd, J = 8.4, 7.6 Hz, 1H), 7.08 (d, J = 7.6 Hz, 1H), 6.60 (d, J = 8.4 Hz, 1H), 6.46 (d, J = 2.4 Hz, 1H). ^{13}C NMR (CDCl_3 , 125 MHz): δ = 162.86, 146.83, 142.55, 142.47, 127.45, 110.69, 108.84, 100.59.

1.3. 2,4-dimethoxy-6-(1H-pyrazol-1-yl)pyrimidine



2,4-Dimethoxy-6-(1H-pyrazol-1-yl)pyrimidine was prepared with the same method as described in procedure 1 using 4-bromo-2,6-dimethoxyl-pyrimidine (1.09 g, 5 mmol) as a starting material. The crude product was purified with column chromatography on alumina with DCM/n-hexane (1/1) as an eluent to afford product (0.43 g, yield: 42 %) as a white solid. ^1H NMR (CDCl_3 , 400 MHz): δ = 8.53 (d, J = 2.4 Hz, 1H), 7.76 (s, 1H), 6.95 (s, 1H), 6.46 (t, J = 2.4 Hz, 1H), 4.04 (d, J = 12 Hz, 6H). ^{13}C NMR (CDCl_3 , 125 MHz): δ = 173.74, 165.46, 159.76, 143.65, 128.24, 108.63, 87.80, 55.38, 54.78.

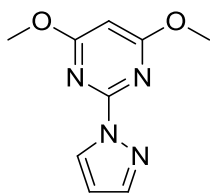
1.4. 6-(1H-pyrazol-1-yl)pyrimidine-2,4-diol



A mixture of 2,4-dimethoxy-6-(1H-pyrazol-1-yl)pyrimidine (0.37 g, 1.77 mmol) and trimethylsilyl iodide (1 mL, 8.87 mmol) in 15 mL dry acetonitrile was refluxed under nitrogen

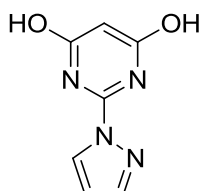
atmosphere for 24 h. After cooling to room temperature, 5 mL MeOH was added and stirred for 30 min. The resulting precipitant was collected by filtration and washed with IPA (5 mL) and ether (5 mL) respectively to give the product (0.11 g, 35 %) as a pale solid. ^1H NMR ($\text{d}^6\text{-DMSO}$, 400 MHz): δ = 11.56 (s, 1H), 11.18 (s, 1H), 8.58 (d, J = 2.8 Hz, 1H), 7.92 (d, J = 1.8 Hz, 1H), 6.67 (dd, J = 2.8, 1.8 Hz, 1H), 5.96 (d, J = 1.8 Hz, 1H). ^{13}C NMR ($\text{d}^6\text{-DMSO}$, 125 MHz): δ = 163.79, 150.35, 146.67, 143.46, 129.46, 109.80, 85.83.

1.5. 4,6-dimethoxy-2-(1H-pyrazol-1-yl)pyrimidine



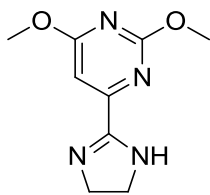
4,6-Dimethoxy-2-(1H-pyrazol-1-yl)pyrimidine was synthesized with the same method as described in procedure 1 using 2-chloro-4,6-dimethoxypyrimidine (0.7 g, 4 mmol) as a starting material. The crude product was purified with column chromatography on alumina with DCM/*n*-hexane (1/1) as an eluent to afford product (0.19 g, yield: 23 %) as a yellow solid. ^1H NMR (CDCl_3 , 400 MHz): δ = 8.62 (s, 1H), 7.89 (s, 1H), 6.53 (s, 1H), 6.00 (s, 1H), 4.07 (s, 6H). ^{13}C NMR (CDCl_3 , 125 MHz): δ = 172.55, 152.35, 144.33, 142.68, 130.06, 54.85. ESI-MS(+): m/z 207.1 $[\text{M}+\text{H}]^+$; 229.1 $[\text{M}+\text{Na}]^+$.

1.6. 2-(1H-Pyrazol-1-yl)pyrimidine-4,6-diol



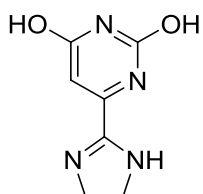
2-(1H-Pyrazol-1-yl)pyrimidine-4,6-diol was synthesized from 4,6-dimethoxy-2-(1H-pyrazol-1-yl)pyrimidine (0.19 g, 0.91 mmol) using the method as described in procedure 1.4. After the reaction, 5 mL MeOH was added and stirred for 30 min at room temperature. The solvent was removed to give black oil which was recrystallized with IPA to give the product as a white solid (0.12 g, 74 %). ^1H NMR ($\text{d}^6\text{-DMSO}$, 400 MHz): δ = 12.03 (br, 2H), 8.46 (d, J = 2.4 Hz, 1H), 7.81 (d, J = 1.6 Hz, 1H), 6.57 (dd, J = 2.4, 1.6 Hz, 1H), 5.61 (s, 1H). ^{13}C NMR ($\text{d}^6\text{-DMSO}$, 125 MHz): δ = 163.94, 150.47, 146.77, 143.58, 129.59, 109.94, 85.97.

1.7. 4-(4,5-dihydro-1H-imidazol-2-yl)-2,6-dimethoxypyrimidine



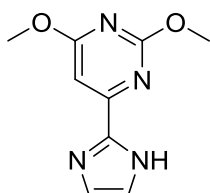
The ligands were synthesized according to the reported processes.² A mixture of 4-bromo-2,6-dimethoxyl-pyrimidine (1.09 g, 5 mmol), ethylenediamine (1.7 mL, 25 mmol), tert-butyl isocyanide (850 μ L, 7.5 mmol), cesium carbonate (2.12 g, 6.5 mmol), PdCl₂ (53.2 mg, 0.3 mmol), dppp (0.25 g, 0.6 mmol) in 25 mL dry toluene was stirred under N₂ atmosphere at reflux for 24 h. After the reaction, the mixture was filtered through a pad of celite which was washed with ethyl acetate. The filtrate was concentrated and the residue was purified with column chromatography on alumina (DCM/n-hexane: 2/1) to give the product (0.35 g, 34 %) as a white solid. ¹H NMR (CDCl₃, 400 MHz): δ = 7.19 (s, 1H), 4.01 (d, J = 8.8 Hz, 6H), 3.84 (s, 4H). ¹³C NMR (CDCl₃, 125 MHz): δ = 171.39, 164.18, 161.78, 156.35, 98.30, 53.91, 53.23.

1.8. 6-(4,5-Dihydro-1H-imidazol-2-yl)pyrimidine-2,4-diol



6-(4,5-Dihydro-1H-imidazol-2-yl)pyrimidine-2,4-diol was synthesized from 4-(4,5-dihydro-1H-imidazol-2-yl)-2,6-dimethoxypyrimidine (0.104 g, 0.5 mmol) according to the procedure 1.4. The crude product was recrystallized with IPA to give the product as a white solid (78 mg, 86 %). ¹H NMR (d⁶-DMSO, 400 MHz): δ = 11.69 (s, 1H), 10.91 (s, 2H), 6.21 (d, 1H), 4.00 (s, 4H). ¹³C NMR (d⁶-DMSO, 125 MHz): δ = 163.13, 159.44, 150.84, 136.59, 105.08, 45.44. ESI-MS(+): m/z 180.9 [M+H]⁺.

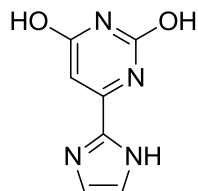
1.9. 4-(1H-imidazol-2-yl)-2,6-dimethoxypyrimidine



The ligands were synthesized according to the reported processes.³ 4-(4,5-Dihydro-1H-imidazol-2-yl)-2,6-dimethoxypyrimidine (0.2 g, 0.9 mmol) was added to a suspension of barium permanganate (1.2 g) in dry DCM and the mixture was heated to reflux for 30 h. The mixture was filtered through celite and washed with DCM. The filtrate was evaporated to give the crude product which was recrystallized with DCM/n-hexane (1/1) to

afford the product (0.12 g, 65 %) as a white solid. ^1H NMR ($\text{d}^6\text{-DMSO}$, 400 MHz): δ = 12.89 (s, 1H), 7.33 (s, 1H), 7.15 (s, 1H), 6.96 (s, 1H), 4.00 (s, 3H), 3.94 (s, 3H). ^{13}C NMR ($\text{d}^6\text{-DMSO}$, 125 MHz): δ = 172.24, 165.58, 157.81, 143.75, 130.75, 120.13, 95.11, 55.07, 54.30. ESI-MS(+): m/z 207.1 $[\text{M}+\text{H}]^+$; 229.0 $[\text{M}+\text{Na}]^+$.

1.10. 6-(1H-Imidazol-2-yl)pyrimidine-2,4-diol



6-(1H-Imidazol-2-yl)pyrimidine-2,4-diol was synthesized from 4-(1H-imidazol-2-yl)-2,6-dimethoxypyrimidine (70 mg, 0.34 mmol) using the method as described in procedure 1.4. The crude product was recrystallized with IPA to give the product as a white solid (58 mg, 96 %). ^1H NMR ($\text{d}^6\text{-DMSO}$, 400 MHz): δ = 11.14 (s, 1H), 10.68 (s, 1H), 7.38 (d, J = 6.4 Hz, 2H), 6.08 (s, 1H). ^{13}C NMR ($\text{d}^6\text{-DMSO}$, 125 MHz): δ = 164.72, 151.36, 141.98, 138.02, 125.87, 95.96.

2. Synthesis of complexes

2.1. Complex $[\text{Cp}^*\text{Ir}(6\text{-MeO-py-pz})\text{Cl}]\text{Cl}$ (1)

$[\text{Cp}^*\text{IrCl}_2]_2$ (119.5 mg, 0.15 mmol) and 2-methoxy-6-(1H-pyrazol-1-yl)pyridine (52.6 mg, 0.3 mmol) was stirred in 10 mL MeOH under Ar atmosphere at 40 °C for 6 h. The solution was filtered to remove the insoluble solid. The filtrate was concentrated and dried at 50 °C under vacuum for 5 h to give the product as a yellow solid (162.5 mg, 94 %). ^1H NMR (D_2O , 400 MHz): δ = 8.68 (d, J = 3.2 Hz, 1H), 8.24 (d, J = 3.2 Hz, 1H), 8.20 (t, J = 4.4 Hz, 1H), 7.66 (d, J = 8 Hz, 1H), 7.20 (d, J = 8 Hz, 1H), 6.98 (t, J = 3.2 Hz, 1H), 4.23 (s, 3H), 1.69 (s, 15H). ^{13}C NMR (D_2O , 125 MHz): δ = 163.80, 146.94, 145.79, 143.50, 132.09, 112.55, 106.51, 104.39, 90.54, 58.41, 8.91. ESI-MS(+): m/z 502.1 $[\text{M}-\text{Cl}-\text{H}]^+$. Anal. Calc. for $\text{C}_{19}\text{H}_{24}\text{Cl}_2\text{IrN}_3\text{O}$: C 39.79, H 4.22, N 7.33. Found: C 39.72, H 4.33, N 7.10.

2.2. Complex $[\text{Cp}^*\text{Ir}(6\text{-HO-py-pz})(\text{OH}_2)]\text{SO}_4$ (2)

$[\text{Cp}^*\text{Ir}(\text{OH}_2)_3]\text{SO}_4$ (143.7 mg, 0.3 mmol) and 6-(1H-pyrazol-1-yl)pyridin-2-ol (48.3 mg, 0.3 mmol) in 15 mL H_2O was stirred at 40 °C under Ar atmosphere for 5 h. The solution was filtered to remove the insoluble solid. The filtrate was concentrated and dried at 50 °C under vacuum for 5 h to give the product as a yellow solid (174.2 mg, 96 %). ^1H NMR (D_2O , 400 MHz): δ = 8.68 (d, J = 3.2 Hz, 1H), 8.37 (d, J = 3.2 Hz, 1H), 8.00 (t, J = 4.0 Hz, 1H), 7.42 (d, J = 8 Hz, 1H), 6.98 (t, J = 3.2 Hz, 2H), 1.69 (s, 15H). ^{13}C NMR (D_2O , 125 MHz): δ = 164.09, 146.86, 145.24, 143.88, 132.22, 112.45, 110.61, 101.99, 89.77, 8.97. ESI-MS(+): m/z 488.1

[M-H₂O-H]⁺. Anal. Calc. for C₁₈H₂₄IrN₃O₆S: C 35.87, H 4.01, N 6.97. Found: C 35.69, H 3.91, N 6.60.

2.3. Complex [Cp*Ir(2,4-(HO)₂-pm-pz)(OH₂)]SO₄ (3)

[Cp*Ir(OH₂)₃]SO₄ (95.8 mg, 0.2 mmol) and 6-(1H-pyrazol-1-yl)pyrimidine-2,4-diol (35.6 mg, 0.2 mmol) in 10 mL H₂O was stirred at 40 °C under Ar atmosphere for 5 h. The solution was filtered to remove the insoluble solid. The filtrate was concentrated and dried at 50 °C under vacuum for 5 h to give the product as a yellow solid (116.7 mg, 94 %). ¹H NMR (D₂O, 400 MHz): δ = 8.55 (d, *J* = 2.4 Hz, 1H), 8.38 (s, 1H), 6.95 (s, 1H), 6.12 (s, 1H), 1.71 (s, 15H). ¹³C NMR (d⁶-DMSO, 125 MHz): δ = 164.52, 154.67, 151.32, 144.25, 132.79, 112.49, 95.66, 84.62, 8.27. ESI-MS(+): *m/z* 505.1 [M-H₂O-H]⁺. Anal. Calc. for C₁₇H₂₃IrN₄O₇S·2H₂O: C 31.14, H 4.15, N 8.54. Found: C 31.52, H 3.95, N 8.18.

2.4. Complex [Cp*Ir(4,6-(HO)₂-pm-pz)(OH₂)]SO₄ (4)

[Cp*Ir(OH₂)₃]SO₄ (42.5 mg, 89 μmol) and 2-(1H-pyrazol-1-yl)pyrimidine-4,6-diol (15.9 mg, 89 μmol) in 10 mL H₂O was stirred at 40 °C under Ar atmosphere for 5 h. The solution was filtered to remove the insoluble solid. The filtrate was concentrated and dried at 50 °C under vacuum for 5 h to give the product as a yellow solid (50.8 mg, 92 %). ¹H NMR (d⁶-DMSO, 400 MHz): δ = 11.38 (s, 1H), 8.65 (s, 1H), 8.45 (s, 1H), 7.03 (s, 1H), 5.29 (s, 1H), 1.73 (s, 15H). ¹³C NMR (d⁶-DMSO, 125 MHz): δ = 169.56, 168.59, 152.58, 145.18, 133.44, 111.97, 96.26, 87.89, 8.83. ESI-MS(+): *m/z* 505.1 [M-H₂O-H]⁺. Anal. Calc. for C₁₇H₂₃IrN₄O₇S·0.5H₂O: C 32.48, H 3.85, N 8.91. Found: C 32.62, H 3.63, N 8.66.

2.5. Complex [Cp*Ir(2,4-(HO)₂-pm-im)(OH₂)]SO₄ (5)

[Cp*Ir(OH₂)₃]SO₄ (142.5 mg, 0.29 mmol) and 6-(1H-imidazol-2-yl)pyrimidine-2,4-diol (53 mg, 0.29 mmol) in 15 mL H₂O was stirred at 40 °C under Ar atmosphere for 9 h. The solution was filtered to remove the insoluble solid. The filtrate was concentrated to ~2 mL and hold at 5 °C overnight. The precipitated yellow solid was collected by filtration and washed with cold water. The yellow solid was dried at 50 °C under vacuum for 5 h (140 mg, 76 %). ¹H NMR (D₂O, 400 MHz): δ = 7.38 (s, 1H), 7.19 (s, 1H), 6.01 (s, 1H), 1.63 (s, 15H). ¹H NMR (d⁶-DMSO, 400 MHz): 10.89 (s, 1H), 7.86 (s, 1H), 7.57 (s, 1H), 6.12 (s, 1H), 1.69 (s, 15H). ¹³C NMR (d⁶-DMSO, 125 MHz): δ = 168.71, 157.91, 127.47, 123.18, 95.95, 87.73, 84.72, 55.90, 8.99. ESI-MS(+): *m/z* 505.2 [M-H₂O-H]⁺. Anal. Calc. for C₁₇H₂₃IrN₄O₇S·1/2H₂O: C 32.43, H 3.85, N 8.91. Found: C 32.42, H 3.70, N 8.75.

2.6. Complex [Cp*Ir(2,4-(HO)₂-pm-imidazoline)(OH₂)]SO₄ (6)

[Cp*Ir(OH₂)₃]SO₄ (142.5 mg, 0.29 mmol) and 6-(4,5-Dihydro-1H-imidazol-2-yl)pyrimidine-2,4-diol (54 mg, 0.29 mmol) in 15 mL H₂O was stirred at 40 °C under Ar

atmosphere for 15 h. The solution was filtered to remove the insoluble solid. The filtrate was concentrated to ~2 mL and hold at 5 °C overnight. The precipitated yellow solid was collected by filtration and washed with cold water. The yellow solid was dried at 50 °C under vacuum for 5 h (125 mg, 70 %). ¹H NMR (D₂O, 400 MHz): 6.07 (s, 1H), 4.24 (t, *J* = 10.4 Hz, 2H), 3.94 (t, *J* = 10.4 Hz, 2H), 1.69 (s, 15H). ¹³C NMR (D₂O, 125 MHz): δ = 174.96, 171.04, 160.32, 155.24, 103.86, 89.93, 55.63, 48.64, 11.71. ESI-MS(+): *m/z* 506.9 [M-H₂O-H]⁺. Anal. Calc. for C₁₇H₂₅IrN₄O₇S·CH₃OH: C 32.28, H 4.36, N 8.36. Found: C 32.24, H 4.00, N 7.98.

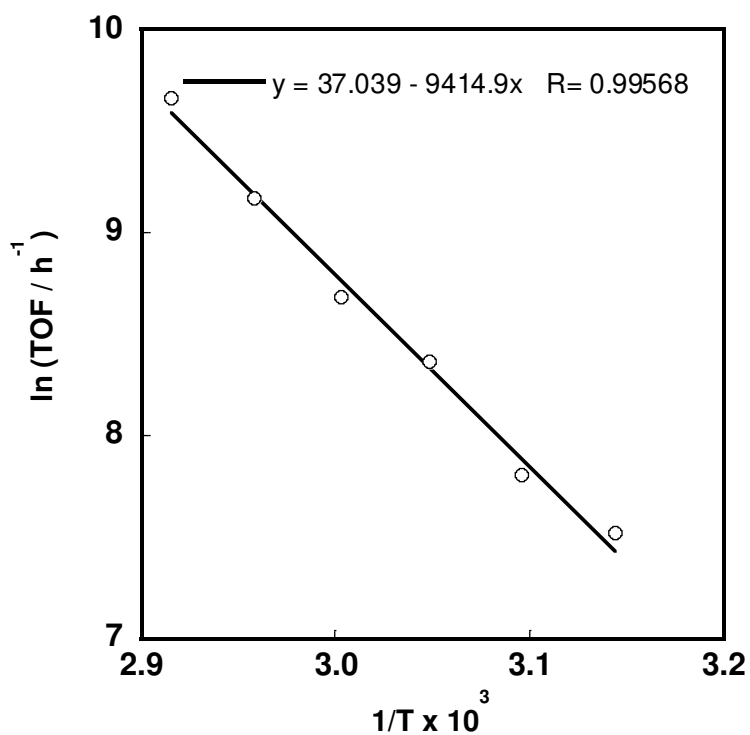


Figure S1. Arrhenius plot for FA dehydrogenation using complex **3** (1 μmol) in 10 mL FA solution (1 M). The activation energy is calculated to be 18.7 kcal mol⁻¹.

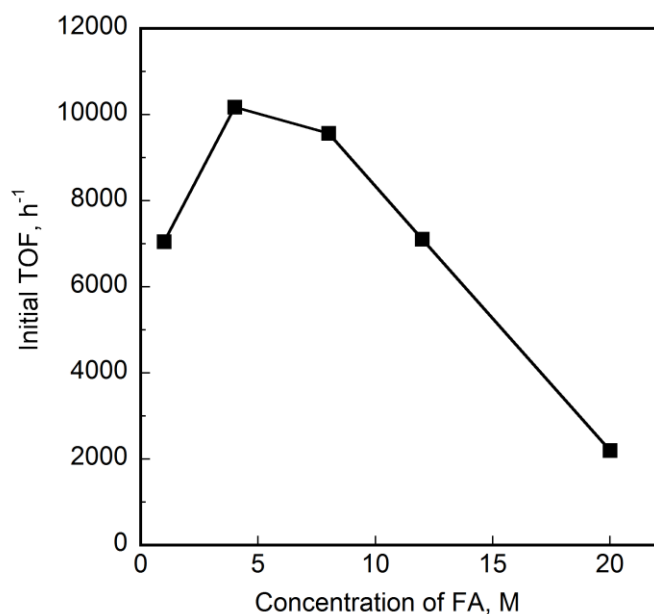
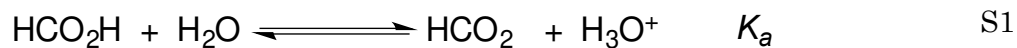


Figure S2. Plot of initial TOF vs concentration of formic acid in the dehydrogenation of formic acid solutions (10 mL) using complex **3** (1 μmol) at 60 $^{\circ}\text{C}$.

The effect of the change in FA concentration on the overall reaction rate can be explained assuming a pre-equilibrium for the formation of a formate bound Ir complex prior to the rate-determining β -hydride elimination step.



$$K_a = 10^{-\text{p}K_a} \quad \text{S2}$$

$$[\text{H}_3\text{O}^+] = \frac{1}{2} \left(K_a + [\text{HCO}_2]_0 \right) + \sqrt{\left(K_a + [\text{HCO}_2]_0 \right)^2 + 4K_a[\text{HCOOH}]_0} \quad \text{S3}$$

$$\text{pH} = -\log([\text{H}_3\text{O}^+]) \quad \text{S4}$$

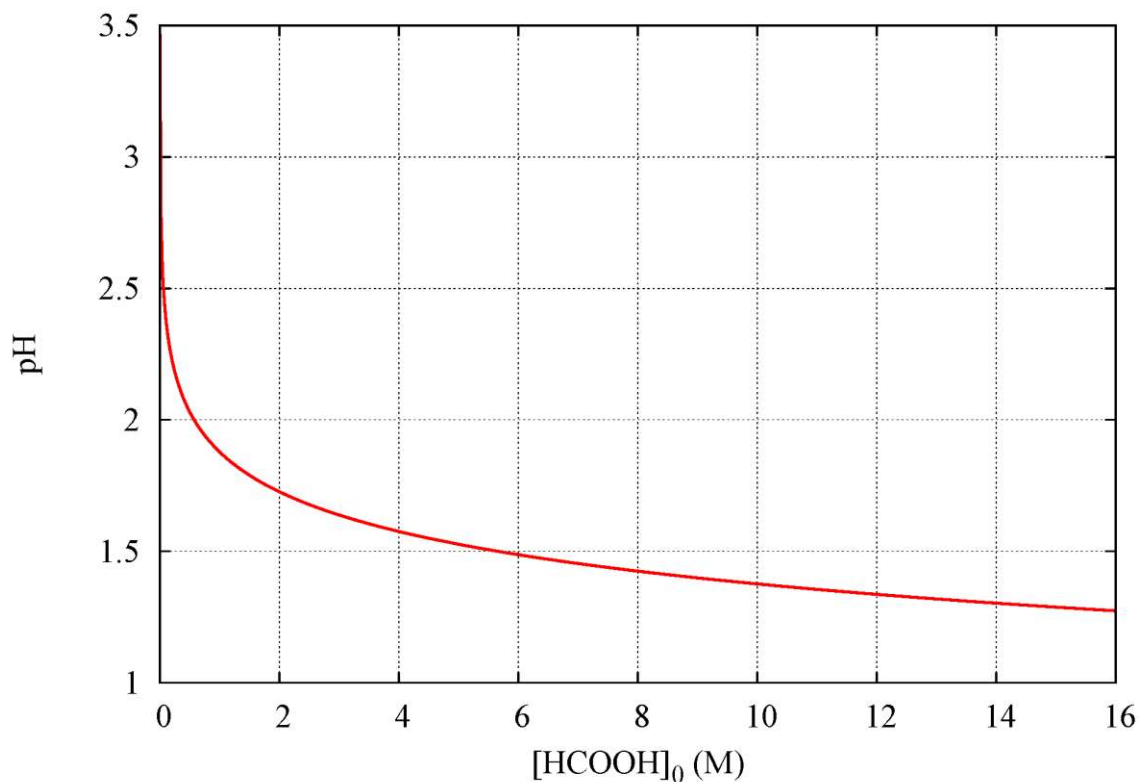
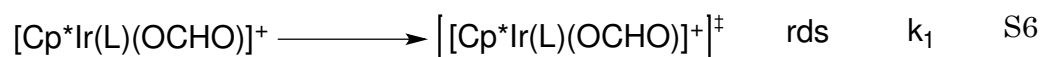


Figure S3. Dependence of the pH on $[\text{HCOOH}]_0$ for the case of $[\text{HCO}_2^-]_0 = 0$.



$$\text{Rate} = k_1 \times [\text{Cp}^*\text{Ir}(\text{L})(\text{OCHO})^+] \quad \text{S7}$$

$$[\text{Cp}^*\text{Ir}(\text{L})(\text{OCHO})^+] = K_{\text{eq}} \times \frac{[\text{HCO}_2\text{H}] \times [\text{Cp}^*\text{Ir}(\text{L})(\text{H}_2\text{O})^{2+}]}{[\text{H}_3\text{O}^+]} \quad \text{S8}$$

$$\text{Rate} = k_1 \times K_{\text{eq}} \times \frac{[\text{HCO}_2\text{H}] \times [\text{Cp}^*\text{Ir}(\text{L})(\text{H}_2\text{O})^{2+}]}{[\text{H}_3\text{O}^+]} \quad \text{S9}$$

Based on these rate expressions, a change in concentration of FA (HCO_2H) from 1 M to 4 M would be expected to enhance the rate by a factor of 4 whereas the consequent change in pH from 1.7 (1 M) to 1.3 (4 M), which was measured by a pH meter, will decrease the rate by a factor of 2.5 so that the overall reaction rate is predicted to increase by about a factor of 1.6. According to Eqs. S1-S4, the concentration of H_3O^+ would be expected to increase by a factor of 2.01 upon changing the concentration of FA from 1 M to 4 M.

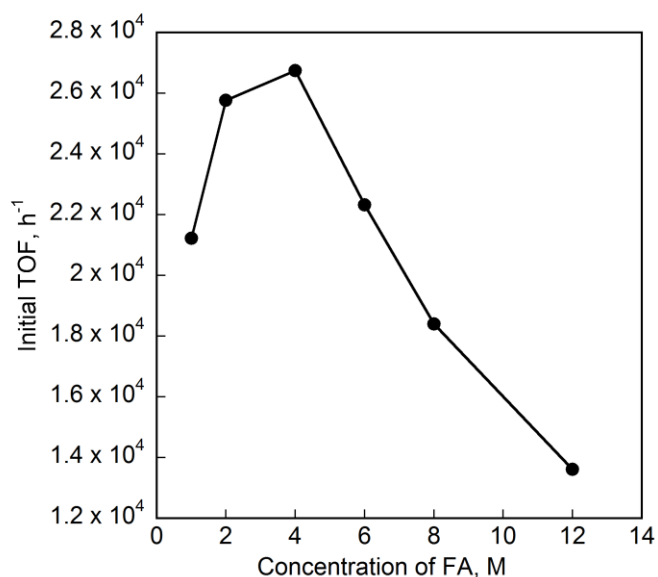


Figure S4. Plot of initial TOF vs concentration of formic acid in the dehydrogenation of formic acid solutions (10 mL) using complex **4** ($1 \mu\text{mol}$) at $80 \text{ }^\circ\text{C}$.

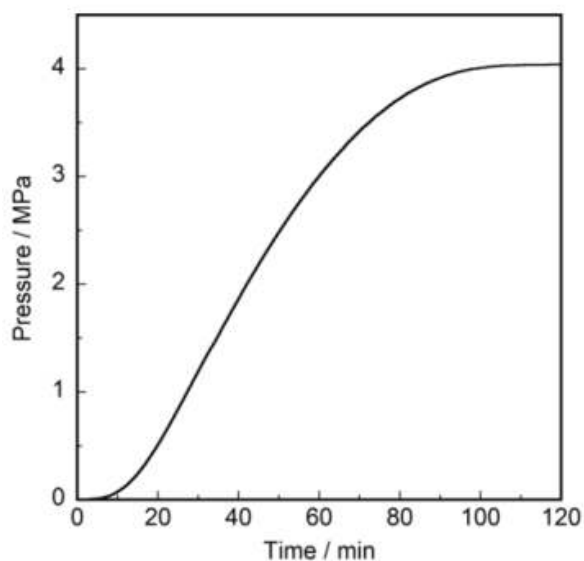


Figure S5. Time course of the pressure of an autoclave for formic acid dehydrogenation using complex **3** ($1 \mu\text{mol}$) in 2 M formic acid solution (6.5 mL) at $70 \text{ }^\circ\text{C}$. Conversion of formic acid:

99.4 %; TON: ~12,900. The induction period is due to the increasing of system temperature.

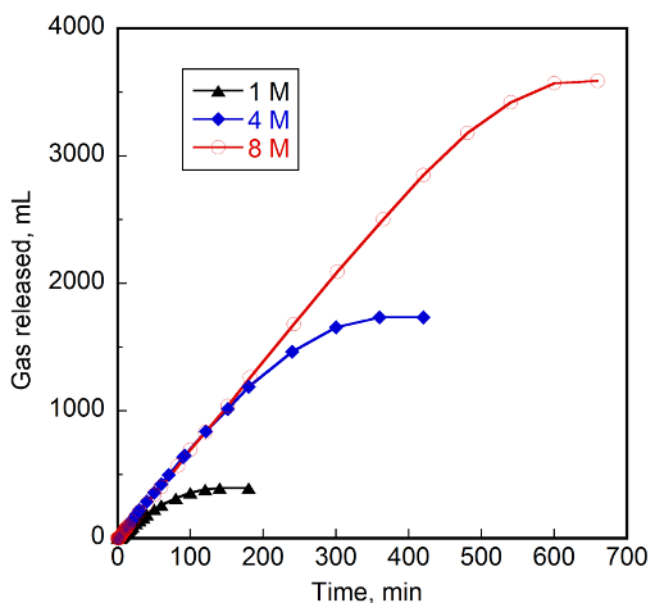


Figure S6. Time course of FA dehydrogenation using complex **3** (1 μmol) in 10 ml FA solutions (a. 1 M; b. 4 M; c. 8M) at 60 °C.

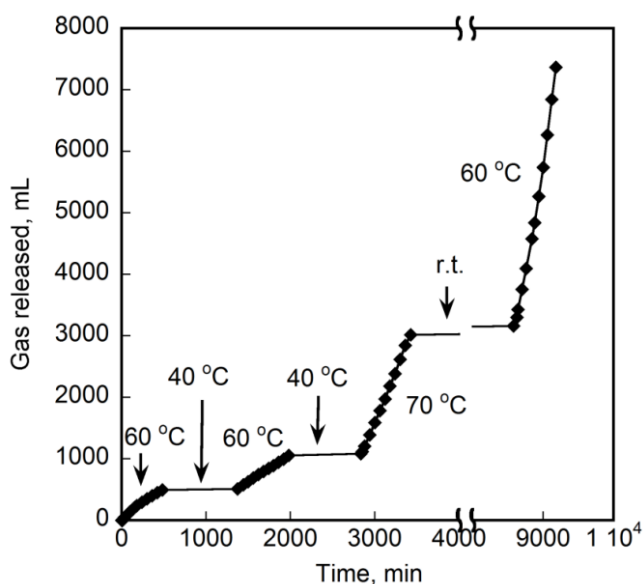


Figure S7. Time course of FA dehydrogenation using complex **3** (1 μmol) in 10 ml FA solution (80 wt%). In this experiment, the durability of complex **3** was tested with highly concentrated FA at different temperatures. The rate increase at 60 °C around 9000 min is due to the decrease of FA concentration. FA was completely decomposed (TON: 200,000).

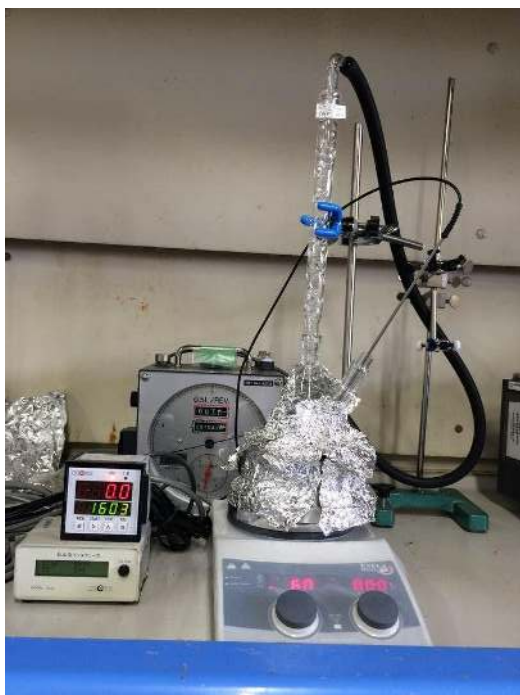


Figure S8. FA dehydrogenation equipment. The volume of evolved gases and the reaction temperature were recorded automatically by a wet gas meter equipped with a recorder (Shinagawa Corp., W-NK-05; Test limit: 1 mL).

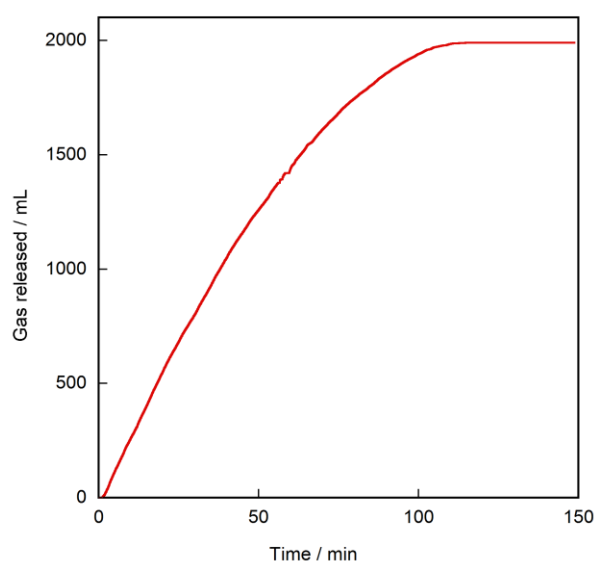


Figure S9. Time course of FA dehydrogenation using complex **3** ($0.4 \mu\text{mol}$) in 10 ml FA solutions (4 M) at 100°C corresponding to Entry 1 in Table 2.

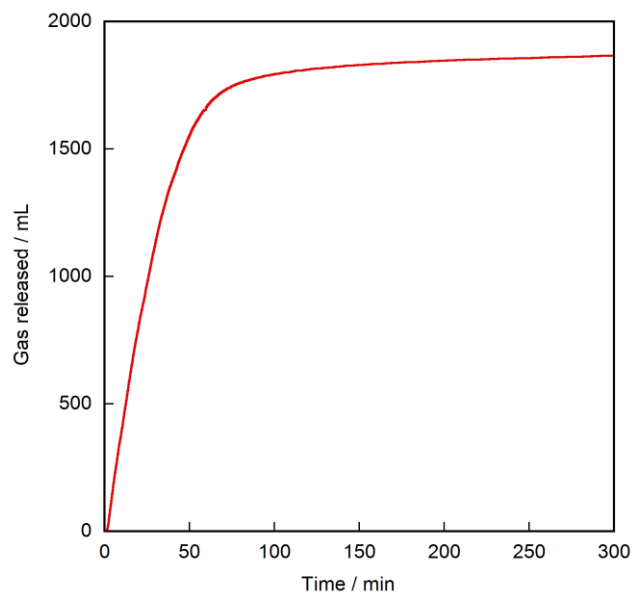


Figure S10. Time course of FA dehydrogenation using complex **3** (0.2 μmol) in 10 ml $\text{HCO}_2\text{H}/\text{HCO}_2\text{Na}$ solutions (98/2, 4 M) at 100 $^\circ\text{C}$ corresponding to Entry 7 in Table 2.

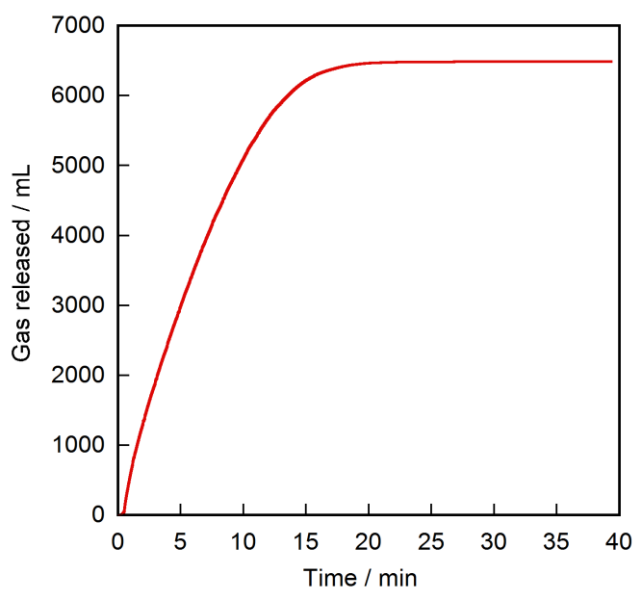


Figure S11. Time course of FA dehydrogenation using complex **6** (2 μmol) in 50 ml $\text{HCO}_2\text{H}/\text{HCO}_2\text{Na}$ solutions (68/32, 4 M) at 100 $^\circ\text{C}$ corresponding to Entry 9 in Table 2.

Computational Methods

All geometries were fully optimized at the M06 level of density functional theory⁴ with the SMD aqueous continuum solvation model⁵ using the Stuttgart [8s7p6d2f | 6s5p3d2f] ECP60MWB contracted pseudopotential basis set⁶ on Ir and the 6-31G(d,p) basis set⁷ on all other atoms. Non-analytical integrals were evaluated using the *integral=grid=ultrafine* option as implemented in the Gaussian 09 software package.⁸ The nature of all stationary points was verified by analytic computation of vibrational frequencies, which were also used for the computation of zero-point vibrational energies, molecular partition functions (with all frequencies below 50 cm⁻¹ replaced by 50 cm⁻¹ when computing free energies), and for determining the reactants and products associated with each transition-state structure (by following the normal modes associated with imaginary frequencies). Partition functions were used in the computation of 298 K, 313 K, and 383 K thermal contributions to free energy employing the usual ideal-gas, rigid-rotator, harmonic oscillator approximation.⁹ Free-energy contributions were added to single-point M06 electronic energies computed with the SDD basis set on Ir and the 6-311++G(2d,p) basis set on all other atoms to arrive at final, composite free energies.

Solvation effects associated with water as solvent were accounted for using the SMD aqueous continuum solvation model. A 1 M standard state was used for all species in solution, thus, an adjustment for the 1 atm to 1 M standard-state concentration change of $RT \ln(24.5)$, (1.89 kcal/mol at 298 K) was added to the computed gas-phase free energies. In the case of water, the 1 atm gas-phase free energy is adjusted by the sum of a 1 atm to 55.6 M standard-state concentration change, or 4.27 kcal/mol, and the experimental 1 M to 1 M self-solvation free energy, -6.32 kcal/mol, yielding an overall correction of -2.05 kcal/mol to the gas-phase free energy. The 1 M to 1 M solvation free energy of the proton was taken from experiment as -264.0 kcal/mol.¹⁰⁻¹³

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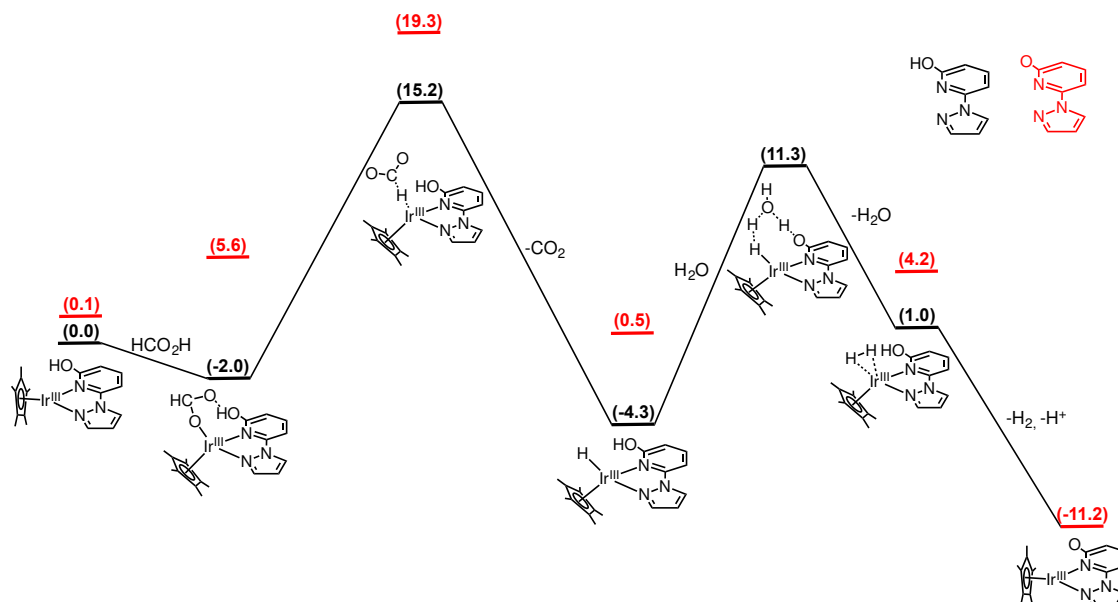
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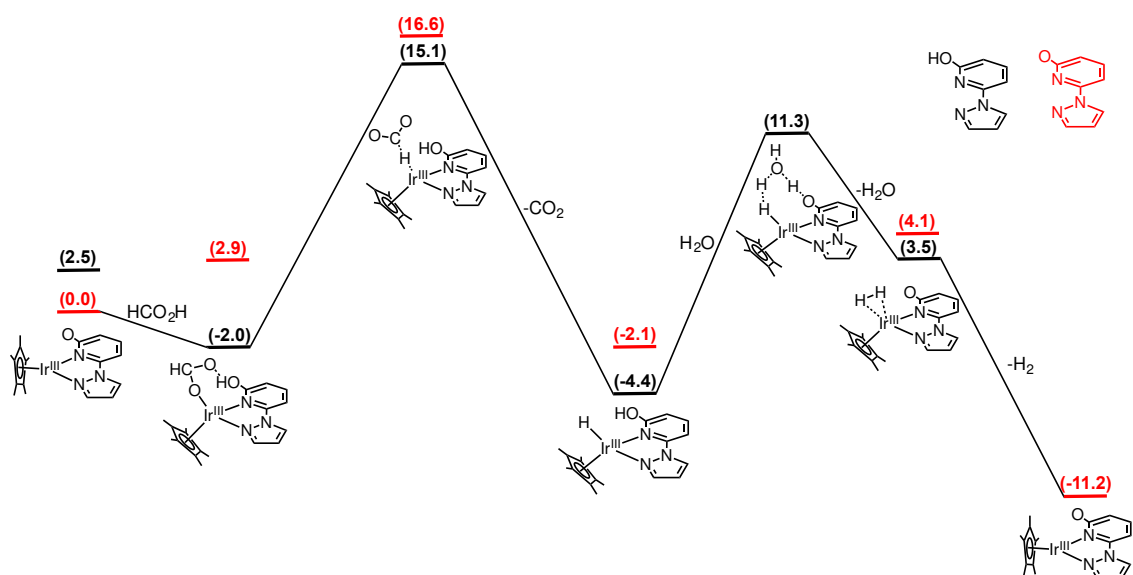
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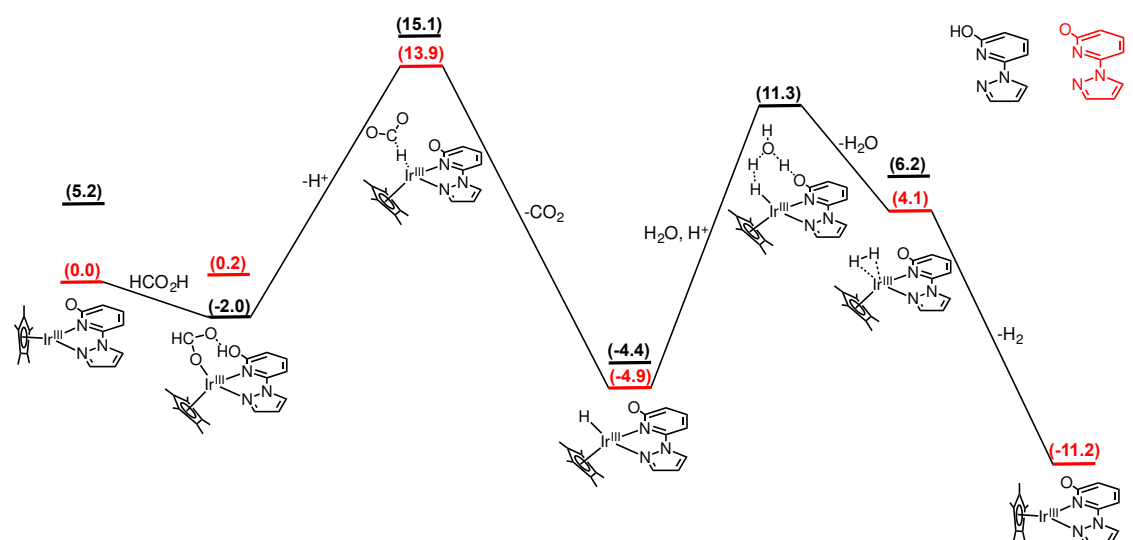
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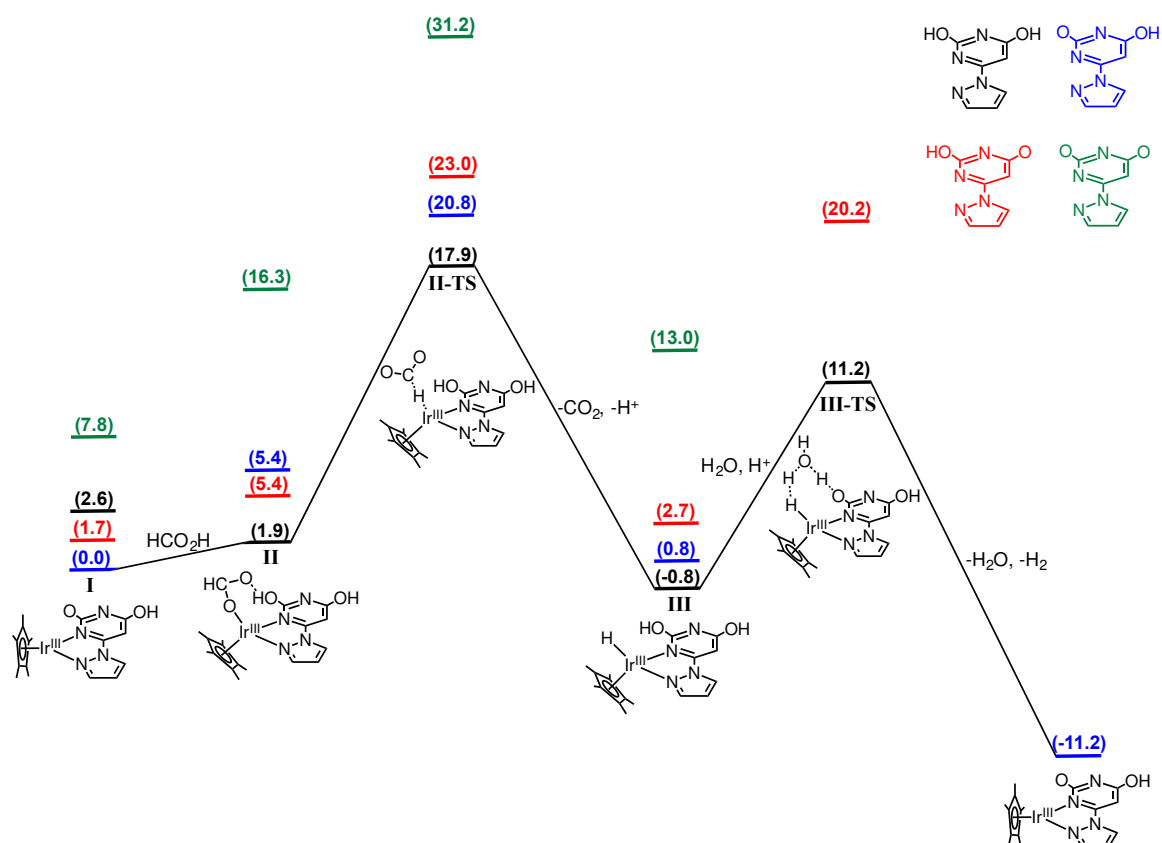
Scheme S1. Proposed mechanism for HCO₂H dehydrogenation by complex **2** at pH 0.0 and 333.15 K. The relative free energies are reported in units of kcal/mol.



Scheme S2. Proposed mechanism for HCO_2H dehydrogenation by complex **2** at pH 1.7 and 333.15 K. The relative free energies are reported in units of kcal/mol.



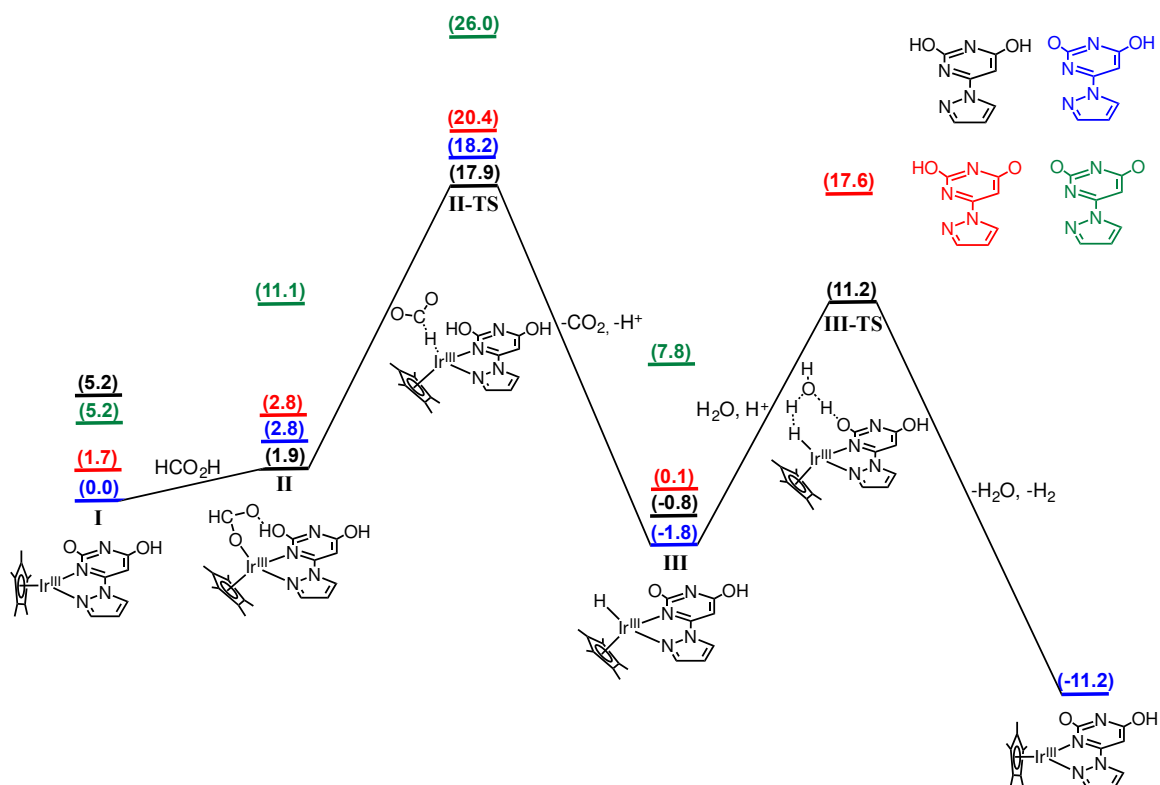
Scheme S3. Proposed mechanism for HCO_2H dehydrogenation by complex **2** at pH 3.5 and 333.15 K. The relative free energies are reported in units of kcal/mol.



The calculated distribution of the protonation states at each intermediate and relative free energies (ΔG^*) obtained as a Boltzmann weighted average

	I	II	II-TS	III	III-TS
	1.85%	98.99%	98.73%	91.77%	100.00%
	6.85%	0.51%	0.04%	0.42%	0.00%
	91.30%	0.50%	1.22%	7.81%	-
	0.00%	0.00%	0.00%	0.00%	-
ΔG^*	0.0	1.7	17.7	-0.9	11.0

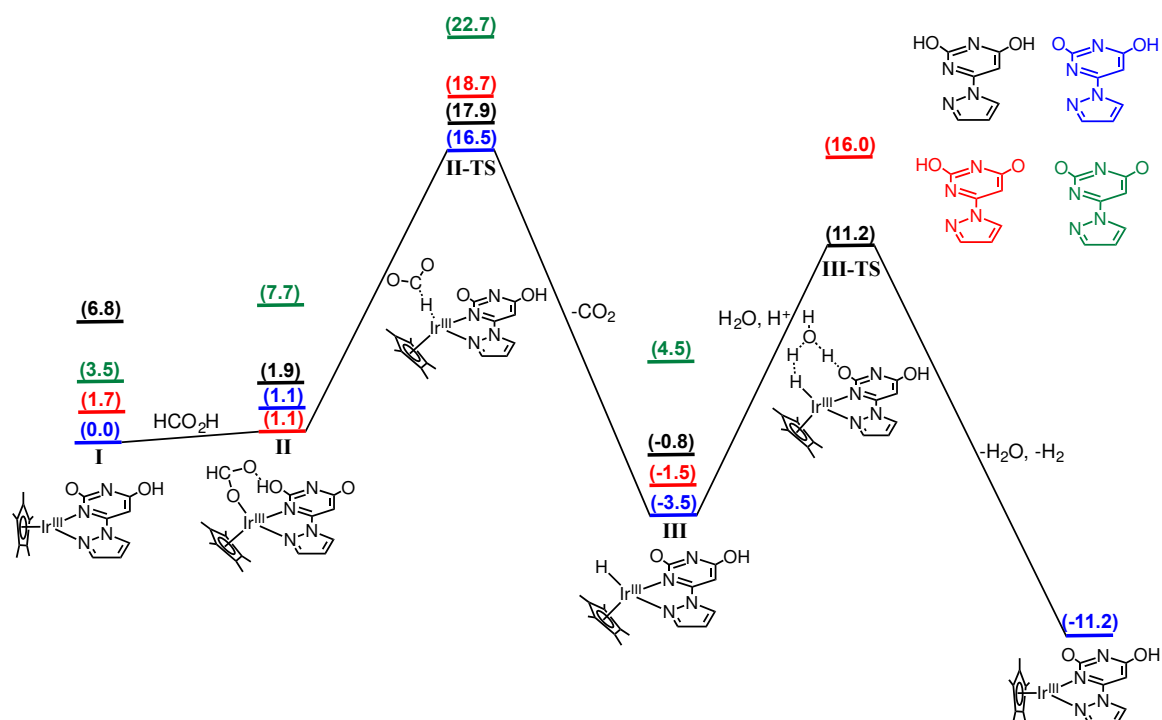
Scheme S4. Proposed mechanism for HCO₂H dehydrogenation by complex **3** at pH 0.0 and 333.15 K (the relative free energies are reported in units of kcal/mol) along with calculated distribution of the protonation states at each intermediate and relative free energies (ΔG^*) obtained as a Boltzmann weighted average..



The calculated distribution of the protonation states at each intermediate and relative free energies (ΔG^*) obtained as a Boltzmann weighted average

	I	II	II-TS	III	III-TS
	0.04%	66.26%	60.82%	18.19%	99.99%
	6.97%	16.95%	1.37%	4.22%	0.01%
	92.96%	16.79%	37.81%	77.59%	-
	0.04%	0.00%	0.00%	0.00%	-
ΔG^*	0.0	2.1	17.9	-1.7	11.1

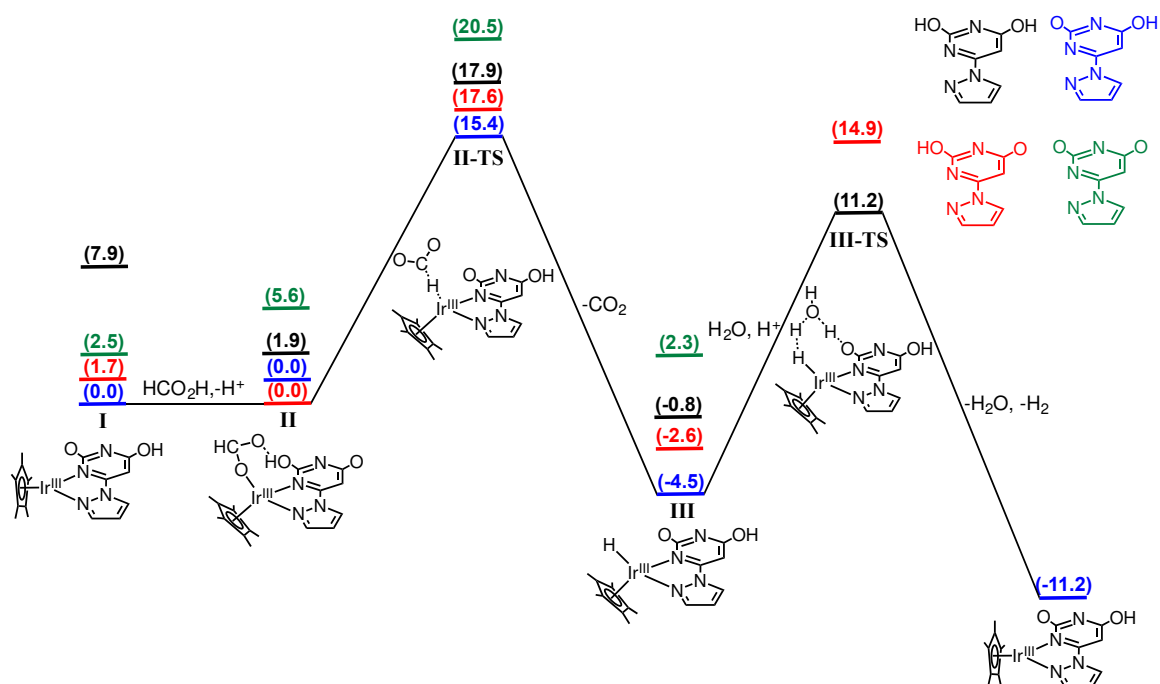
Scheme S5. Proposed mechanism for HCO₂H dehydrogenation by complex **3** at pH 1.7 and 333.15 K (the relative free energies are reported in units of kcal/mol) along with calculated distribution of the protonation states at each intermediate and relative free energies (ΔG^*) obtained as a Boltzmann weighted average



The calculated distribution of the protonation states at each intermediate and relative free energies (ΔG*) obtained as a Boltzmann weighted average

	I	II	II-TS	III	III-TS
	0.00%	13.49%	10.98%	1.74%	99.93%
	6.94%	43.45%	3.11%	5.07%	0.07%
	92.61%	43.05%	85.91%	93.19%	-
	0.45%	0.00%	0.01%	0.00%	-
ΔG*	0.0	1.1	16.6	-3.5	11.1

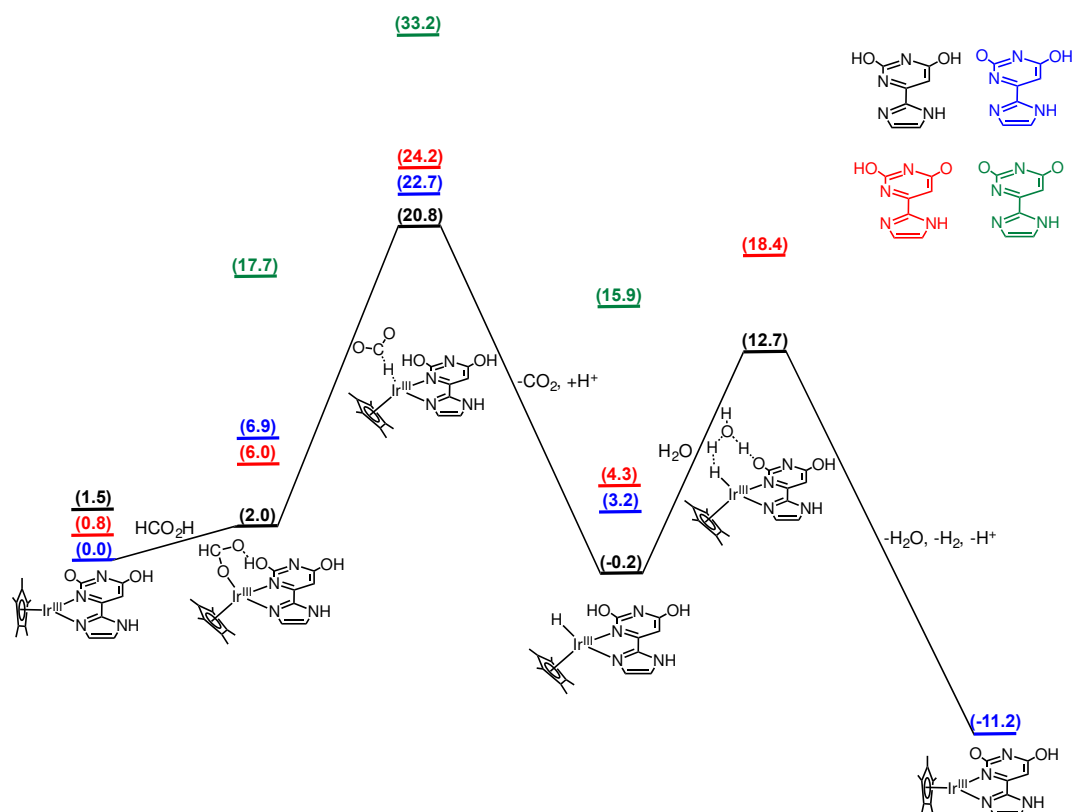
Scheme S6. Proposed mechanism for HCO₂H dehydrogenation by complex **3** at pH 2.8 and 333.15 K (the relative free energies are reported in units of kcal/mol) along with calculated distribution of the protonation states at each intermediate and relative free energies (ΔG*) obtained as a Boltzmann weighted average.



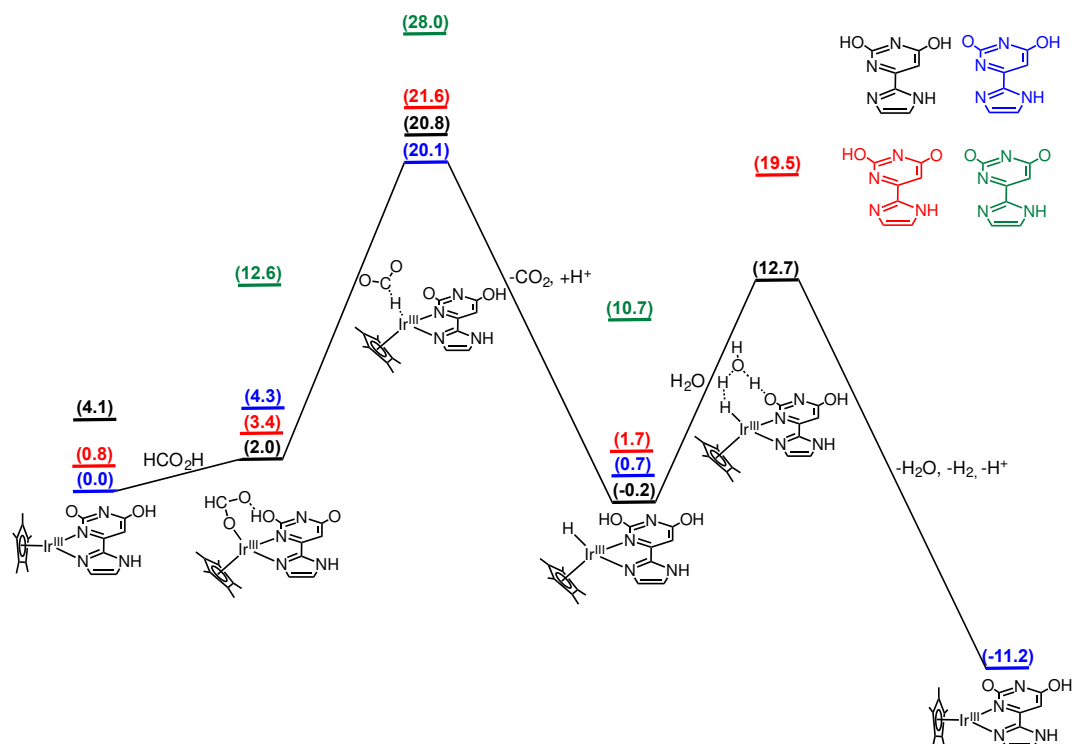
The calculated distribution of the protonation states at each intermediate and relative free energies (ΔG*) obtained as a Boltzmann weighted average

	I	II	II-TS	III	III-TS
	0.00%	3.02%	2.40%	0.35%	99.63%
	6.82%	48.71%	3.41%	5.14%	0.37%
	90.97%	48.26%	94.15%	94.50%	-
	2.20%	0.01%	0.04%	0.00%	-
ΔG*	0.0	-0.1	15.4	-4.6	11.0

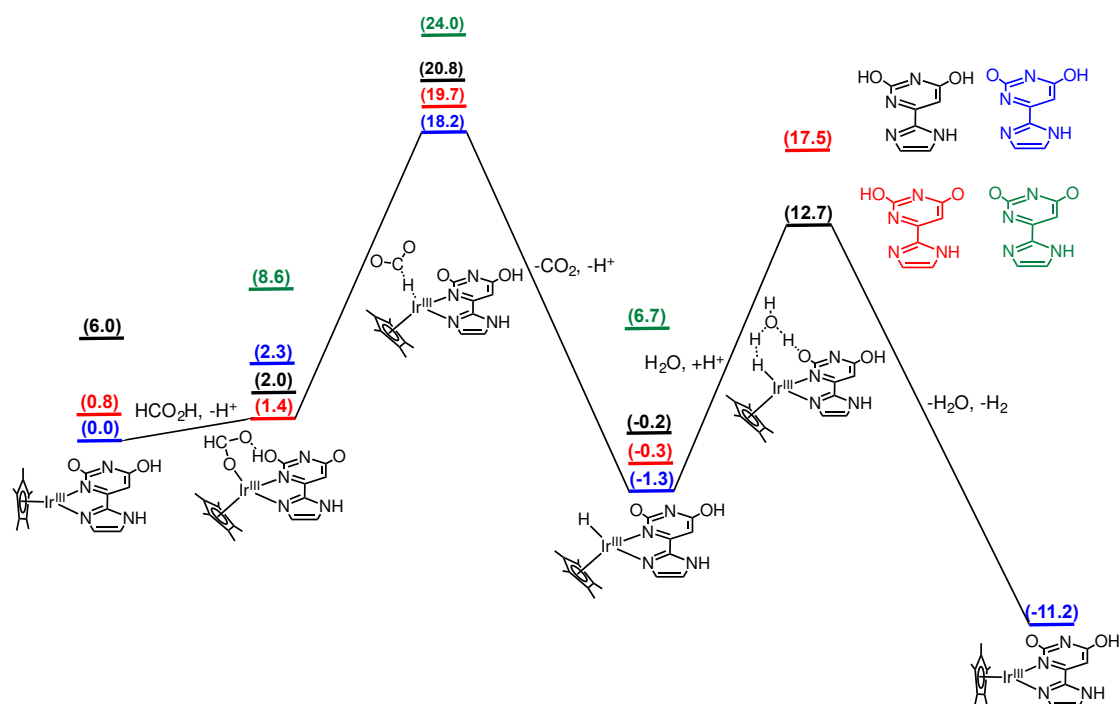
Scheme S7. Proposed mechanism for HCO₂H dehydrogenation by complex **3** at pH 3.5 and 333.15 K (the relative free energies are reported in units of kcal/mol) along with calculated distribution of the protonation states at each intermediate and relative free energies (ΔG*) obtained as a Boltzmann weighted average.



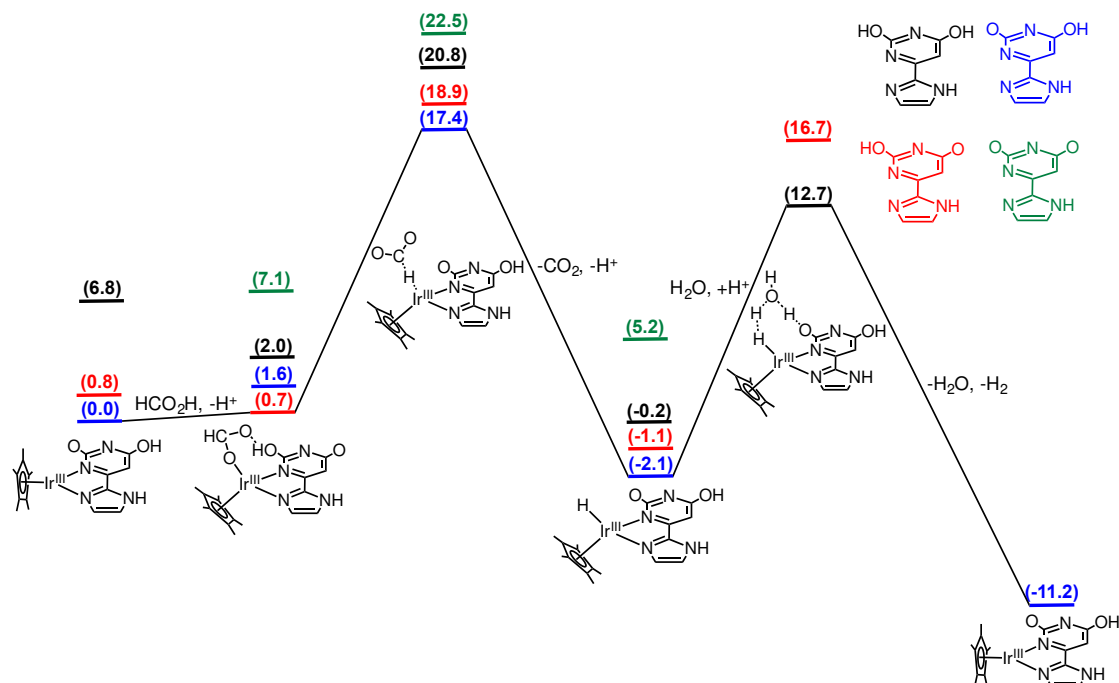
Scheme S8. Proposed mechanism for HCO₂H dehydrogenation by complex **5** at pH 0.0 and 333.15 K. The relative free energies are reported in units of kcal/mol.



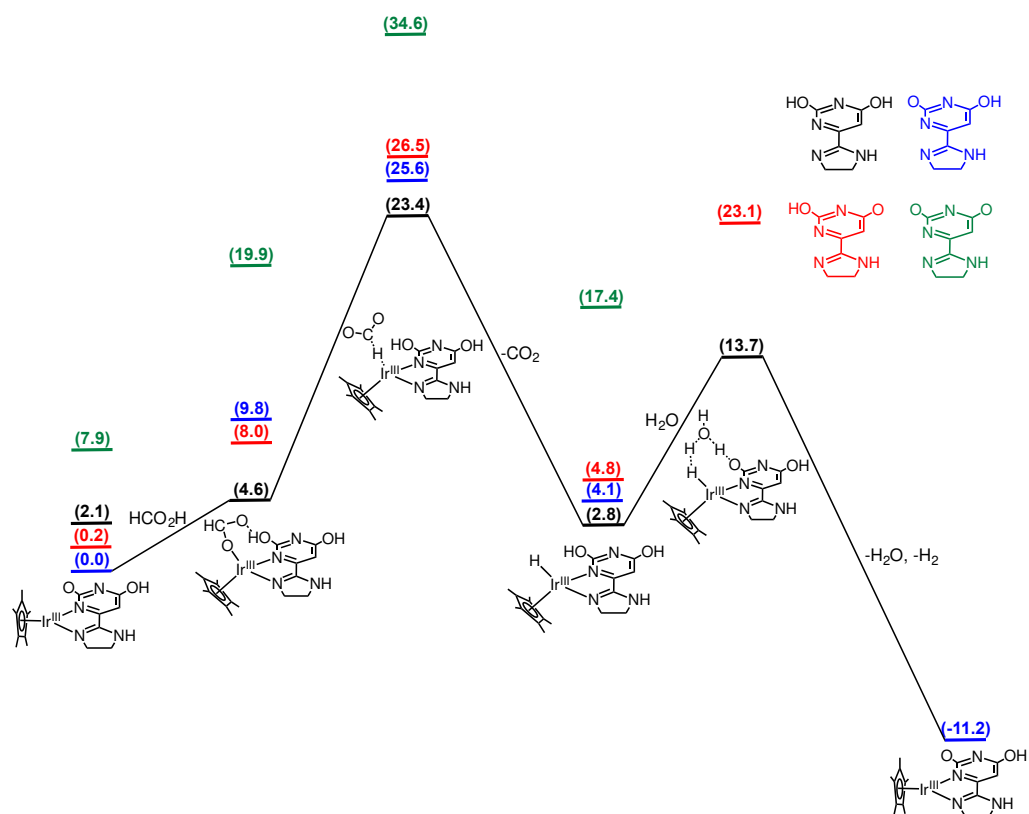
Scheme S9. Proposed mechanism for HCO₂H dehydrogenation by complex **5** at pH 1.7 and 333.15 K. The relative free energies are reported in units of kcal/mol.



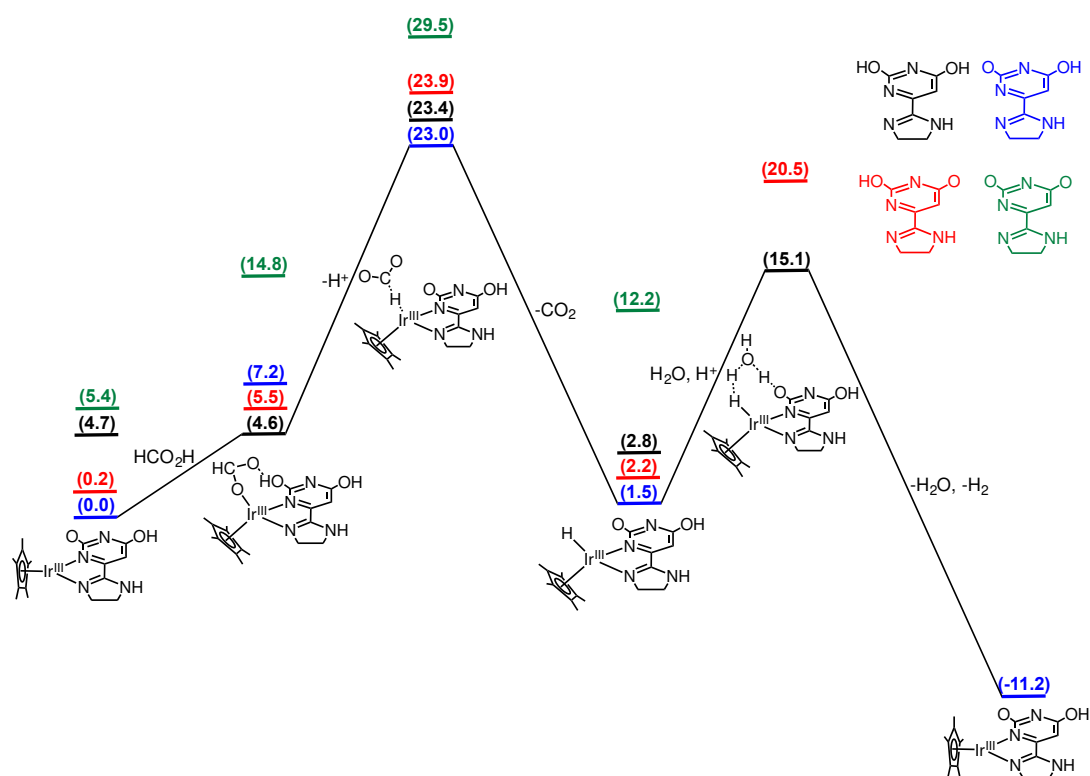
Scheme S10. Proposed mechanism for HCO_2H dehydrogenation by complex **5** at pH 3.0 and 333.15 K. The relative free energies are reported in units of kcal/mol.



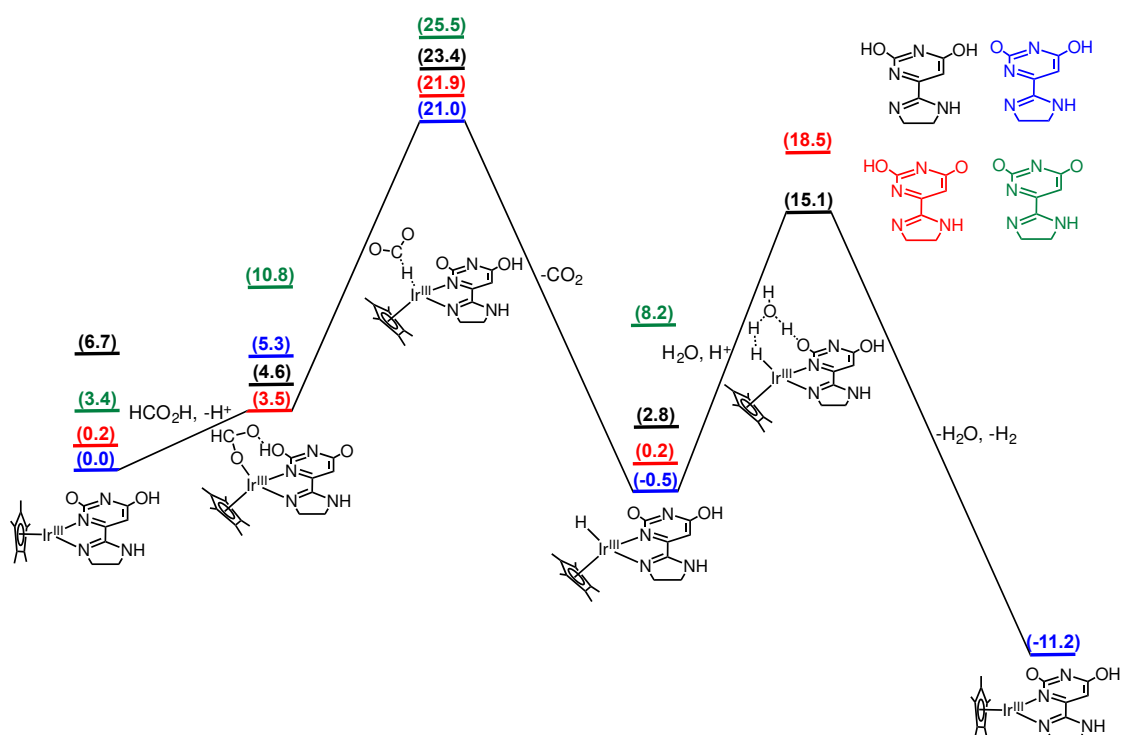
Scheme S11. Proposed mechanism for HCO_2H dehydrogenation by complex **5** at pH 3.5 and 333.15 K. The relative free energies are reported in units of kcal/mol.



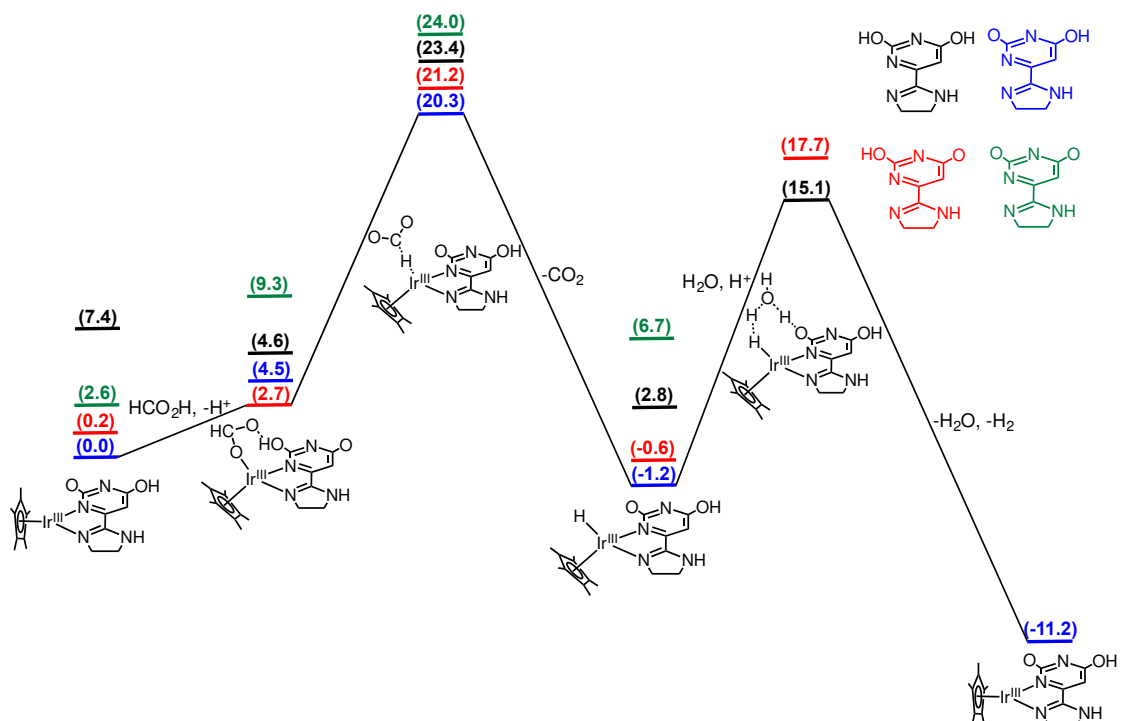
Scheme S12. Proposed mechanism for HCO₂H dehydrogenation by complex **6** at pH 0.0 and 333.15 K. The relative free energies are reported in units of kcal/mol.



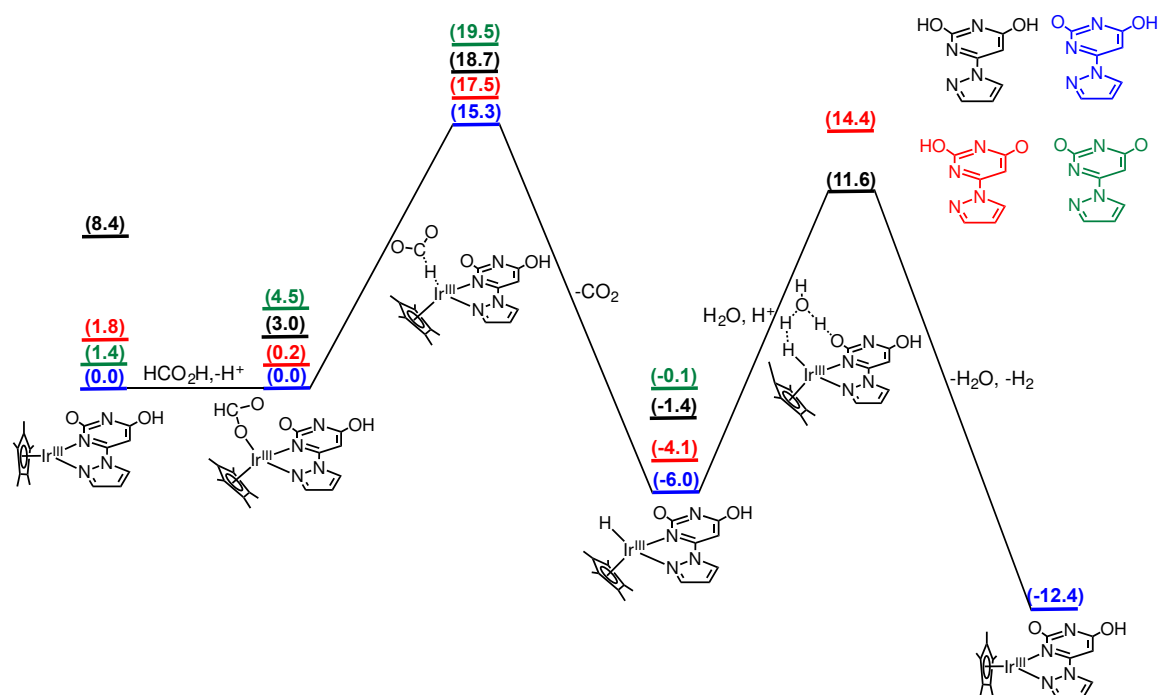
Scheme S13. Proposed mechanism for HCO₂H dehydrogenation by complex **6** at pH 1.7 and 333.15 K. The relative free energies are reported in units of kcal/mol.



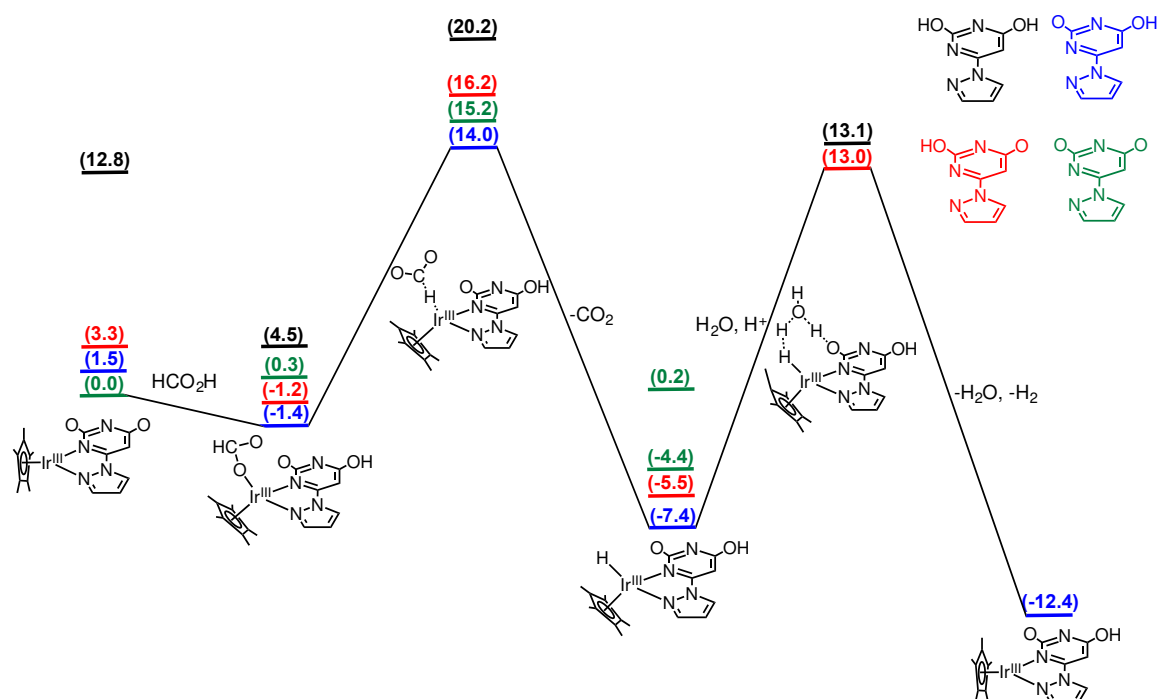
Scheme S14. Proposed mechanism for HCO₂H dehydrogenation by complex 6 at pH 3.0 and 333.15 K. The relative free energies are reported in units of kcal/mol.



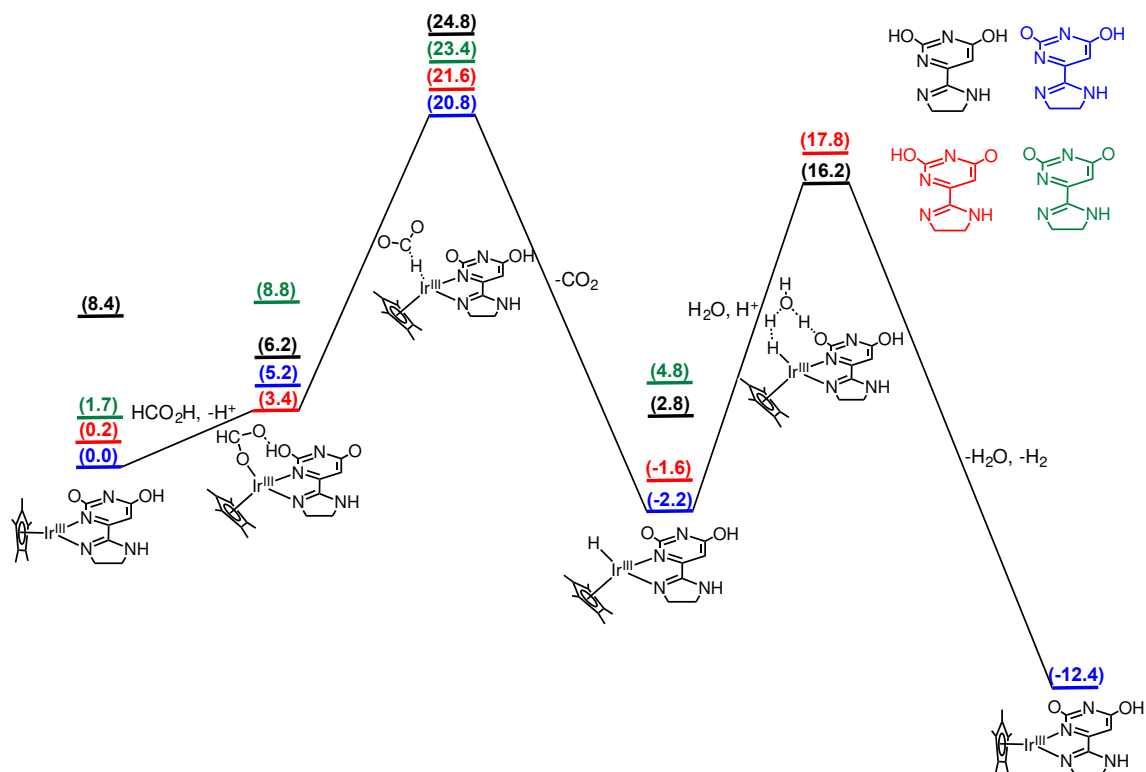
Scheme S15. Proposed mechanism for HCO₂H dehydrogenation by complex 6 at pH 3.5 and 333.15 K. The relative free energies are reported in units of kcal/mol.



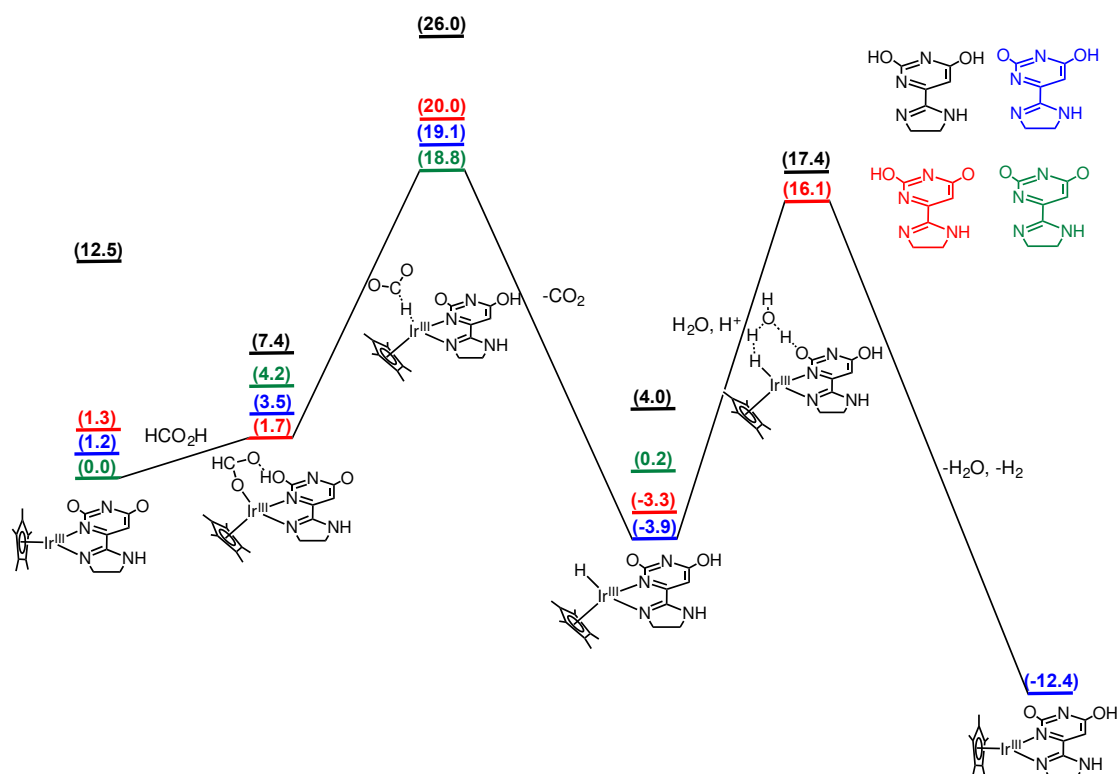
Scheme S16. Proposed mechanism for HCO₂H dehydrogenation by complex **3** at pH 1.8 and 373.15 K. The relative free energies are reported in units of kcal/mol.



Scheme S17. Proposed mechanism for HCO₂H dehydrogenation by complex **3** at pH 3.5 and 373.15 K. The relative free energies are reported in units of kcal/mol.



Scheme S18. Proposed mechanism for HCO_2H dehydrogenation by complex **6** at pH 1.8 and 373.15 K. The relative free energies are reported in units of kcal/mol.



Scheme S19. Proposed mechanism for HCO_2H dehydrogenation by complex **6** at pH 3.5 and 373.15 K. The relative free energies are reported in units of kcal/mol.

Optimized Coordinates and Energies

H₂

E = -1.16958423009 a.u.

1	0.000000	0.000000	0.371071
1	0.000000	0.000000	-0.371071

H₂O

E = -76.3842736863 a.u.

8	0.000000	0.000000	0.117679
1	0.000000	0.758416	-0.470718
1	0.000000	-0.758416	-0.470718

CO₂

E = -188.498068316 a.u.

6	0.000000	0.000000	0.000000
8	0.000000	0.000000	1.164889
8	0.000000	0.000000	-1.164889

HCO₂H

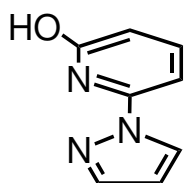
E = -189.673964877 a.u.

6	0.118981	0.368882	-0.000060
1	0.024662	1.467320	0.000153
8	1.174162	-0.218996	-0.000050
8	-1.043929	-0.284497	0.000039
1	-1.780414	0.347330	0.000295

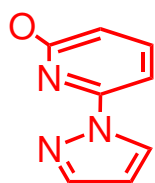
HCO₂

E = -189.204614480 a.u.

6	0.000000	0.000000	0.331383
1	0.000000	0.000000	1.458848
8	0.000000	1.128378	-0.215447
8	0.000000	-1.128378	-0.215447



(2)



(2-H⁺)

(2)

E = -1042.14791953 a.u.

77	0.405976	0.069578	-0.069929
7	-0.865693	1.729678	-0.124500
7	-1.669849	-0.738993	-0.047183
6	-2.660523	0.177557	-0.004485
6	-3.998562	-0.138377	0.044633
1	-3.958556	2.711528	0.098467
1	-4.763665	0.628912	0.078588
6	-1.951850	3.691714	-0.057776
6	-4.328267	-1.494026	0.040469
6	-0.711491	3.054862	-0.144999
6	-2.009176	-2.038097	-0.076122
6	2.343278	0.152943	-0.986090
6	2.060316	-1.239921	-0.670069
6	2.365030	0.901921	0.232397
6	2.037794	-0.010230	1.299408
6	1.893423	-1.340375	0.736288
6	2.582286	0.674687	-2.351303
1	2.395969	1.751359	-2.409154
1	1.945324	0.168366	-3.084032
1	3.626360	0.493141	-2.638693

6	1.983555	-2.319436	-1.685723
6	2.691980	2.340857	0.394908
1	1.307149	-2.041560	-2.502527
1	2.487907	2.910188	-0.517559
1	1.637892	-3.262049	-1.255165
1	2.127769	2.787470	1.220475
1	2.975641	-2.483203	-2.125133
1	3.759207	2.450919	0.623703
6	1.943887	0.313155	2.740516
6	1.626155	-2.554798	1.548203
1	1.208062	-0.325082	3.240038
1	0.749679	-2.417593	2.191459
1	1.671792	1.360524	2.901892
1	1.464885	-3.436785	0.924719
1	2.917353	0.138550	3.218500
1	2.486237	-2.747498	2.201739
1	-2.146738	4.753874	-0.051271
8	-0.986745	-2.877014	-0.166476
1	-1.294580	-3.797519	-0.171758
1	0.279886	3.482778	-0.223335
6	-2.880725	2.678782	0.017002
7	-2.204822	1.504206	-0.024357
6	-3.340720	-2.455179	-0.022398
1	-3.567659	-3.516879	-0.040920
1	-5.371894	-1.791341	0.079438

(2)-HCO₂

E = -1231.41016669 a.u.

77	-0.482356	0.124858	-0.156696
7	0.947989	1.646846	-0.645574
7	1.526482	-0.816727	-0.161084
6	2.557572	0.035050	-0.004110
6	3.809963	-0.348954	0.422017
1	4.083159	2.431942	-0.267713
1	4.611736	0.370556	0.545358
6	2.204535	3.492510	-0.872349

6	3.996013	-1.706806	0.691132
6	0.919591	2.932549	-0.972900
6	1.734383	-2.127207	0.048830
6	-1.044009	0.917312	1.752031
6	-1.056051	-0.513841	1.821246
6	-2.030065	1.317693	0.761609
6	-2.634837	0.130223	0.240644
6	-2.032180	-1.014436	0.877141
6	-0.186436	1.827406	2.557498
1	0.071223	2.735795	2.001822
1	0.746552	1.331351	2.847506
1	-0.704093	2.132563	3.475428
6	-0.203668	-1.339074	2.717484
6	-2.383079	2.715576	0.400080
1	0.795911	-0.904088	2.835321
1	-1.560637	3.405447	0.613986
1	-0.092326	-2.360313	2.340610
1	-2.640274	2.804574	-0.660691
1	-0.661051	-1.395668	3.713070
1	-3.252563	3.044316	0.983172
6	-3.671847	0.071958	-0.821108
6	-2.477754	-2.425667	0.729957
1	-3.523376	-0.796650	-1.471790
1	-2.859941	-2.624552	-0.275894
1	-3.662989	0.973603	-1.441522
1	-1.666689	-3.130882	0.933753
1	-4.667782	-0.017364	-0.369079
1	-3.288440	-2.634655	1.440051
1	2.496346	4.511069	-1.082584
8	0.744690	-2.979542	-0.161776
1	0.059465	-2.597759	-0.783758
1	-0.015292	3.396917	-1.263509
6	-0.685122	-1.285922	-2.809947
1	-0.766082	-1.277645	-3.913931
8	-0.824629	-2.362637	-2.206554
8	-0.447183	-0.140574	-2.302527
6	3.019392	2.466867	-0.459616
7	2.236578	1.362565	-0.327458

1	4.964164	-2.055430	1.038691
6	2.968298	-2.607173	0.497977
1	3.085852	-3.672361	0.667272

(2)-HCO₂-TS

E = -1231.37126145 a.u.

77	-0.491130	0.084241	0.107845
7	0.741097	1.780430	0.402867
7	1.549434	-0.542751	-0.398733
6	2.508722	0.401275	-0.348696
6	3.839612	0.168212	-0.628085
1	3.744195	2.955152	-0.043055
1	4.571981	0.965803	-0.576701
6	1.774144	3.756066	0.656070
6	4.197849	-1.133953	-0.963857
6	0.568958	3.043207	0.773532
6	1.926357	-1.802079	-0.667106
6	-2.611129	-0.316844	0.450859
6	-2.093053	-1.293406	-0.469883
6	-2.445527	0.980994	-0.127634
6	-1.873294	0.812548	-1.453428
6	-1.662840	-0.583456	-1.656699
6	-3.218415	-0.617854	1.773436
1	-3.095876	0.215570	2.472361
1	-2.777958	-1.513789	2.222635
1	-4.294265	-0.801618	1.654503
6	-2.195382	-2.769115	-0.296059
6	-2.905306	2.264287	0.468072
1	-1.880675	-3.079123	0.706639
1	-2.740194	2.287805	1.550874
1	-1.581535	-3.304818	-1.024923
1	-2.394539	3.123117	0.021076
1	-3.235665	-3.093002	-0.431077
1	-3.981061	2.394307	0.292864
6	-1.592970	1.916264	-2.409722
6	-1.081880	-1.209306	-2.873527

1	-0.869347	1.612188	-3.172455
1	-0.405252	-0.522756	-3.392898
1	-1.193792	2.798038	-1.895165
1	-0.525307	-2.119430	-2.630064
1	-2.512796	2.225103	-2.922863
1	-1.882026	-1.482948	-3.572892
1	1.944489	4.798091	0.884176
8	0.950806	-2.703740	-0.587033
1	1.290505	-3.591740	-0.779989
1	-0.402437	3.381467	1.113098
1	0.000733	-0.275424	1.704751
6	0.756477	-1.380002	2.495201
8	-0.094000	-2.156618	2.813929
8	1.891546	-1.007781	2.523979
7	2.044329	1.662254	0.041437
6	2.693412	2.850427	0.189845
6	3.244438	-2.134142	-0.979878
1	3.493090	-3.166154	-1.208594
1	5.232881	-1.365515	-1.196971

(2)-Hydride

E = -1042.88101969 a.u.

77	0.417381	0.041001	-0.329833
7	-0.858898	1.709574	-0.228220
7	-1.645064	-0.759230	-0.152933
6	-2.634083	0.151752	-0.061655
6	-3.973630	-0.165226	0.050421
1	-3.947369	2.678310	0.135497
1	-4.732858	0.606170	0.114185
6	-1.949299	3.664762	-0.059609
6	-4.305914	-1.515264	0.077343
6	-0.705180	3.028000	-0.211058
6	-1.992206	-2.054701	-0.144108
6	2.529677	-0.310708	-0.685598
6	2.025021	-1.409282	0.081797
6	2.350459	0.883894	0.087655

6	1.831387	0.500396	1.397154
6	1.627587	-0.894060	1.385597
6	3.171525	-0.402456	-2.025339
1	3.053781	0.527120	-2.591623
1	2.746885	-1.220073	-2.616927
1	4.247035	-0.593359	-1.913883
6	2.162583	-2.847780	-0.282879
6	2.829200	2.241983	-0.293499
1	1.936377	-3.019994	-1.340851
1	2.630273	2.456737	-1.349439
1	1.499440	-3.480758	0.313154
1	2.354693	3.022310	0.310724
1	3.192694	-3.185192	-0.105363
1	3.913029	2.321949	-0.136344
6	1.553274	1.458716	2.500776
6	1.069903	-1.734508	2.480254
1	0.928623	1.009559	3.279110
1	0.597552	-1.126260	3.257710
1	1.039334	2.355119	2.130859
1	0.322082	-2.439721	2.098367
1	2.487719	1.793070	2.969763
1	1.864083	-2.327094	2.952552
1	-2.140718	4.726853	-0.011480
8	-0.977013	-2.907878	-0.277932
1	-1.302588	-3.821562	-0.263418
1	0.281309	3.461267	-0.319969
1	0.044707	0.054488	-1.912790
6	-2.873558	2.654536	0.014218
7	-2.190200	1.478941	-0.090562
1	-5.345827	-1.813737	0.170396
6	-3.315859	-2.474527	-0.018257
1	-3.538658	-3.537432	-0.006916

(2)-Hydride-H₂O-TS

E = -1119.26766982 a.u.

77	0.491509	0.081163	-0.314195
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7	-0.834707	1.687950	-0.674912
7	-1.530692	-0.764558	-0.264772
6	-2.504384	0.139481	-0.025292
6	-3.762782	-0.162875	0.433062
1	-3.883186	2.647152	-0.092799
1	-4.502057	0.608956	0.615514
6	-1.967266	3.620495	-0.729110
6	-4.032355	-1.522249	0.656534
6	-0.730240	2.988113	-0.927748
6	-1.799427	-2.107821	-0.116674
6	2.656936	-0.175997	0.021299
6	1.938079	-1.211546	0.713101
6	2.224302	1.092472	0.522679
6	1.257120	0.841667	1.582167
6	1.087895	-0.568903	1.697369
6	3.655614	-0.398683	-1.056886
1	3.717938	0.461963	-1.730092
1	3.411290	-1.285260	-1.651459
1	4.649063	-0.556274	-0.617964
6	2.199516	-2.671251	0.585714
6	2.747807	2.424440	0.116303
1	2.403103	-2.954938	-0.452365
1	2.997952	2.445998	-0.949820
1	1.355379	-3.267269	0.942199
1	2.021935	3.219820	0.312642
1	3.080104	-2.947106	1.180551
1	3.659984	2.663399	0.677878
6	0.574096	1.882974	2.395105
6	0.180003	-1.261049	2.649507
1	-0.380187	1.515847	2.788673
1	-0.730289	-0.677905	2.830799
1	0.374742	2.786164	1.807298
1	-0.112757	-2.249120	2.280980
1	1.199524	2.174751	3.247987
1	0.683509	-1.400369	3.614366
1	-2.200030	4.666422	-0.866601
8	-0.919399	-2.969318	-0.418904
1	-0.137683	-2.569284	-1.770990

1	0.220064	3.403359	-1.241243
1	0.512491	0.018530	-2.054763
8	0.293434	-2.177982	-2.596754
1	1.178421	-2.570527	-2.646317
1	0.448440	-0.945377	-2.168979
6	-2.829271	2.624443	-0.333727
7	-2.119213	1.466818	-0.304333
1	-5.007076	-1.816347	1.037934
6	-3.083981	-2.474099	0.383616
1	-3.276800	-3.532877	0.530819

(2)-H₂

E = -1043.31634261 a.u.

77	0.458800	0.018873	-0.348875
7	-0.817953	1.707685	-0.403880
7	-1.587071	-0.749136	-0.197039
6	-2.569830	0.157382	-0.033007
6	-3.897874	-0.174848	0.127189
1	-3.845454	2.685752	0.278818
1	-4.660415	0.586184	0.249017
6	-1.892407	3.666393	-0.210137
6	-4.217152	-1.531232	0.099871
6	-0.672831	3.025280	-0.478443
6	-1.917237	-2.047064	-0.264821
6	2.640796	-0.052763	-0.416805
6	2.153261	-1.324966	0.083704
6	2.235967	0.974540	0.485454
6	1.431819	0.354209	1.526496
6	1.450796	-1.072129	1.302075
6	3.450757	0.122234	-1.647239
1	3.356984	1.134735	-2.050847
1	3.152559	-0.591986	-2.421065
1	4.509459	-0.055583	-1.418795
6	2.460834	-2.644032	-0.530291
6	2.596534	2.411504	0.395791
1	2.309147	-2.622943	-1.614744

1	2.651781	2.751511	-0.643508
1	1.841138	-3.441195	-0.111917
1	1.889153	3.045674	0.937836
1	3.511588	-2.901461	-0.347024
1	3.585797	2.561119	0.846564
6	0.797782	1.038903	2.683712
6	0.806629	-2.071848	2.187306
1	-0.135928	0.539960	2.967007
1	-0.157368	-1.713635	2.564137
1	0.568308	2.084034	2.452590
1	0.655212	-3.029520	1.684062
1	1.465431	1.022930	3.554112
1	1.454034	-2.240077	3.057438
1	-2.085857	4.728927	-0.200935
1	0.290567	3.452486	-0.727434
1	0.050613	0.169541	-2.200652
1	0.102242	-0.634747	-2.109996
8	-0.903904	-2.871383	-0.512464
1	-1.217541	-3.789244	-0.549457
7	-2.120311	1.482749	-0.079420
6	-2.791722	2.659192	0.037596
1	-5.250700	-1.841669	0.220342
6	-3.233658	-2.479242	-0.103971
1	-3.454597	-3.541121	-0.156949

(2-H⁺)

E = -1041.69661678 a.u.

77	0.378612	0.058165	-0.054846
7	-0.889679	1.713404	-0.114516
7	-1.606278	-0.766600	-0.040732
6	-2.638382	0.115117	0.002407
6	-3.960553	-0.225966	0.051562
1	-3.997435	2.631519	0.095266
1	-4.743527	0.523350	0.083051
6	-2.008913	3.654188	-0.061265
6	-4.254651	-1.606553	0.051970

6	-0.756868	3.042537	-0.142428
6	-1.873970	-2.134884	-0.089230
6	2.297024	0.198765	-1.000214
6	2.043268	-1.204495	-0.699488
6	2.340810	0.926967	0.227282
6	2.038305	-0.001659	1.283858
6	1.901019	-1.327516	0.707585
6	2.502159	0.742302	-2.364268
1	2.298463	1.816994	-2.403803
1	1.854204	0.238715	-3.089540
1	3.541352	0.581031	-2.679711
6	1.988688	-2.273576	-1.727755
6	2.636396	2.371950	0.406604
1	1.299004	-2.006487	-2.536944
1	2.428302	2.945653	-0.502338
1	1.673429	-3.230886	-1.306712
1	2.054591	2.799773	1.230359
1	2.982276	-2.403158	-2.175712
1	3.698289	2.506067	0.646848
6	1.953855	0.305755	2.730255
6	1.684821	-2.558554	1.511289
1	1.237455	-0.353152	3.231354
1	0.814883	-2.455978	2.169834
1	1.657899	1.344921	2.903883
1	1.537168	-3.437792	0.880733
1	2.935376	0.151041	3.198322
1	2.561534	-2.733203	2.147799
1	-2.225558	4.712278	-0.061077
8	-0.941639	-2.953356	-0.201185
1	0.226353	3.488256	-0.221492
6	-2.918983	2.621184	0.017217
7	-2.221613	1.462608	-0.017248
6	-3.253959	-2.531022	-0.012195
1	-3.458675	-3.597683	-0.026005
1	-5.292856	-1.926228	0.096871

(2-H⁺)-HCO₂

E = -1230.94387299 a.u.

77	0.432924	0.092383	0.061070
7	-0.897699	1.730296	0.125894
7	-1.577109	-0.750662	-0.157054
6	-2.609809	0.116036	-0.133125
6	-3.933912	-0.218640	-0.282812
1	-4.030965	2.574375	0.232190
1	-4.715690	0.532326	-0.265840
6	-2.061059	3.618942	0.442217
6	-4.212962	-1.583243	-0.464326
6	-0.788154	3.033722	0.361408
6	-1.829192	-2.107290	-0.313827
6	1.622925	0.781692	-1.572451
6	1.437829	-0.636799	-1.693633
6	2.333410	1.033088	-0.330613
6	2.603035	-0.234156	0.274342
6	2.049856	-1.273828	-0.542647
6	1.186527	1.817109	-2.546160
1	0.926427	2.755027	-2.043223
1	0.312003	1.485107	-3.115199
1	1.992753	2.032698	-3.258354
6	0.785651	-1.338103	-2.831307
6	2.796959	2.349691	0.187266
1	0.024822	-0.707552	-3.303444
1	2.302856	3.181742	-0.324345
1	0.303990	-2.263017	-2.500793
1	2.606668	2.446038	1.262871
1	1.530500	-1.594409	-3.595278
1	3.877545	2.461705	0.030768
6	3.324562	-0.422874	1.559049
6	2.269466	-2.732350	-0.338275
1	3.106526	-1.395348	2.009131
1	2.068124	-3.032781	0.694888
1	3.068093	0.361740	2.279509
1	1.633525	-3.331662	-0.994573
1	4.406687	-0.365786	1.382619
1	3.315218	-2.982324	-0.559914

1	-2.297257	4.658223	0.618907
8	-0.887245	-2.933208	-0.305359
1	0.188990	3.488310	0.470675
6	0.003564	-0.806895	2.958045
1	-0.258177	-0.586163	4.015235
8	0.367040	-1.944761	2.647195
8	-0.127397	0.212631	2.190196
6	-2.949985	2.587787	0.247617
7	-2.220072	1.457852	0.060628
1	-5.244126	-1.904629	-0.592106
6	-3.198455	-2.501295	-0.479832
1	-3.390253	-3.562412	-0.614268

(2-H⁺)-HCO₂-TS

E = -1230.91393172 a.u.

77	-0.475626	0.075789	0.103821
7	0.754151	1.765478	0.406648
7	1.525196	-0.549850	-0.437264
6	2.504184	0.372460	-0.365906
6	3.833138	0.141570	-0.629328
1	3.755613	2.926046	-0.046038
1	4.572702	0.930386	-0.552448
6	1.792892	3.734924	0.670780
6	4.175362	-1.172614	-0.985462
6	0.586385	3.027969	0.788549
6	1.851056	-1.871905	-0.704825
6	-2.597811	-0.298980	0.478493
6	-2.103506	-1.284017	-0.450395
6	-2.436123	0.993193	-0.105862
6	-1.875582	0.818662	-1.435797
6	-1.681826	-0.580241	-1.642255
6	-3.194863	-0.595955	1.807849
1	-3.058636	0.236390	2.505883
1	-2.754577	-1.494227	2.252920
1	-4.273316	-0.773764	1.703439
6	-2.235345	-2.758397	-0.279504

6	-2.880003	2.282008	0.491887
1	-1.904652	-3.080413	0.714388
1	-2.704685	2.306719	1.573268
1	-1.648121	-3.304748	-1.022589
1	-2.366019	3.136123	0.039185
1	-3.285231	-3.059696	-0.392528
1	-3.956104	2.423242	0.327086
6	-1.596272	1.922128	-2.393925
6	-1.131668	-1.218367	-2.868231
1	-0.880844	1.614701	-3.163243
1	-0.497081	-0.526091	-3.431359
1	-1.183742	2.799359	-1.881627
1	-0.535645	-2.102823	-2.621766
1	-2.516849	2.241559	-2.899451
1	-1.948714	-1.535518	-3.528784
1	1.970499	4.774549	0.904875
8	0.968329	-2.761442	-0.650192
1	-0.382108	3.367113	1.135195
1	0.032038	-0.305215	1.684382
6	0.801796	-1.441296	2.472443
8	-0.053156	-2.216343	2.777133
8	1.931665	-1.059533	2.503108
7	2.052188	1.643626	0.039543
6	2.705798	2.825069	0.192548
6	3.218543	-2.151903	-1.026170
1	3.464639	-3.181182	-1.272764
1	5.211680	-1.410380	-1.214480

(2-H⁺)-Hydride

E = -1042.42193685 a.u.

77	0.428329	0.040035	-0.365743
7	-0.862675	1.701089	-0.329772
7	-1.587102	-0.776966	-0.249003
6	-2.590279	0.100882	-0.058055
6	-3.906891	-0.232742	0.163774
1	-3.946113	2.608637	0.164497

1	-4.669389	0.523228	0.313754
6	-1.984343	3.638152	-0.164734
6	-4.202383	-1.604427	0.187342
6	-0.737103	3.024260	-0.363540
6	-1.865006	-2.136179	-0.290064
6	2.582486	-0.189155	-0.535013
6	2.038676	-1.375808	0.069620
6	2.289797	0.916922	0.323818
6	1.637401	0.403008	1.521398
6	1.480342	-0.992651	1.357100
6	3.352295	-0.138158	-1.809125
1	3.253364	0.835902	-2.299670
1	3.012692	-0.907060	-2.511126
1	4.419788	-0.310145	-1.617726
6	2.284663	-2.768395	-0.403557
6	2.737297	2.322817	0.116906
1	2.195892	-2.843799	-1.493068
1	2.706931	2.603533	-0.941976
1	1.580488	-3.477657	0.040303
1	2.118668	3.031398	0.677864
1	3.299260	-3.090515	-0.131989
1	3.771631	2.448804	0.463572
6	1.210699	1.241085	2.675683
6	0.838134	-1.940447	2.308395
1	0.471367	0.725991	3.297367
1	0.218261	-1.418610	3.044577
1	0.766311	2.186761	2.341114
1	0.206005	-2.661961	1.778772
1	2.066493	1.496145	3.314577
1	1.600365	-2.510335	2.855588
1	-2.199580	4.696643	-0.142198
8	-0.952611	-2.959694	-0.546085
1	0.233099	3.473217	-0.537585
1	0.146717	0.039121	-1.975363
6	-2.878543	2.607144	-0.006252
7	-2.176076	1.447273	-0.109660
1	-5.224299	-1.926744	0.374149
6	-3.217315	-2.531287	-0.032423

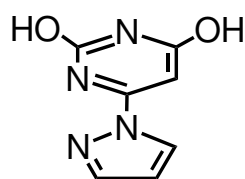
1	-3.426990	-3.597592	-0.033125
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(2-H⁺)-H₂

E = -1042.86400682 a.u.

77	0.474981	0.016466	-0.411974
7	-0.809739	1.696007	-0.533165
7	-1.520553	-0.767961	-0.338905
6	-2.508340	0.109574	-0.070179
6	-3.802782	-0.240796	0.220614
1	-3.799875	2.630356	0.328615
1	-4.564685	0.503713	0.421653
6	-1.882042	3.642053	-0.241907
6	-4.084726	-1.619353	0.221915
6	-0.675120	3.018027	-0.587760
6	-1.782924	-2.127446	-0.421039
6	2.673415	-0.084200	-0.195698
6	2.093061	-1.337820	0.248271
6	2.156967	0.969648	0.607382
6	1.220891	0.381426	1.557398
6	1.240127	-1.047724	1.361590
6	3.643007	0.047648	-1.312882
1	3.647139	1.061295	-1.724557
1	3.415377	-0.655528	-2.120515
1	4.655375	-0.176033	-0.953293
6	2.452923	-2.674919	-0.295189
6	2.547313	2.401553	0.533377
1	2.448956	-2.671240	-1.390618
1	2.718381	2.720498	-0.500301
1	1.762745	-3.450582	0.046553
1	1.790356	3.054040	0.978211
1	3.463427	-2.951146	0.031781
1	3.481991	2.554093	1.087469
6	0.468547	1.107975	2.614703
6	0.470346	-2.026712	2.167855
1	-0.469095	0.594288	2.854659
1	-0.480274	-1.605435	2.511107

1	0.224836	2.129172	2.302093
1	0.262986	-2.942048	1.608008
1	1.061972	1.171415	3.535224
1	1.053679	-2.294453	3.057978
1	-2.085833	4.702203	-0.204430
1	0.273470	3.455805	-0.873045
1	0.501632	0.173741	-2.285812
1	0.320366	-0.619160	-2.167263
8	-0.878283	-2.922354	-0.766154
7	-2.089437	1.453547	-0.147863
6	-2.760032	2.618631	0.032091
1	-5.091869	-1.953180	0.460233
6	-3.116393	-2.535923	-0.090417
1	-3.325634	-3.601836	-0.113724



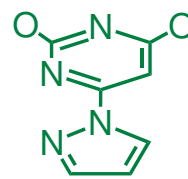
(3)



(3-H⁺)



(3-H⁺)



(3-2H⁺)

(3)

E = -1133.41235417 a.u.

77	0.599443	0.048643	-0.028115
7	-0.398500	1.902633	-0.036073
7	-1.544996	-0.415096	-0.095509
6	-2.397869	0.637751	-0.043117
6	-3.758388	0.506516	0.013452
1	-3.327824	3.327872	-0.054442
1	-4.442260	1.345442	0.058916
6	-1.193369	4.002428	-0.046165
6	-4.222325	-0.815764	0.036317
6	-0.053659	3.187893	-0.037897
6	-2.119762	-1.631517	-0.091495

6	2.457784	-0.218240	-1.061309
6	1.958256	-1.535092	-0.696800
6	2.695775	0.525310	0.142428
6	2.280386	-0.300715	1.242932
6	1.852237	-1.586039	0.719085
6	2.697769	0.239848	-2.447899
1	2.682175	1.331500	-2.518694
1	1.948411	-0.168270	-3.133971
1	3.682555	-0.108851	-2.786449
6	1.632258	-2.601899	-1.676020
6	3.262273	1.894726	0.238097
1	0.999369	-2.215056	-2.482874
1	3.081589	2.466415	-0.678185
1	1.116511	-3.442940	-1.206368
1	2.835194	2.446692	1.082392
1	2.556662	-2.974241	-2.135158
1	4.346989	1.839121	0.390399
6	2.325561	0.055990	2.678505
6	1.412196	-2.722074	1.566701
1	1.507627	-0.417897	3.230611
1	0.630580	-2.413228	2.269940
1	2.271558	1.138736	2.825561
1	1.034464	-3.555123	0.969626
1	3.270074	-0.297556	3.113581
1	2.262741	-3.081059	2.159907
1	-1.228133	5.081770	-0.050634
8	-1.280808	-2.639566	-0.171512
1	-1.776440	-3.476468	-0.160039
1	0.992788	3.467080	-0.035434
8	-5.523434	-1.008951	0.121943
1	-5.713698	-1.962933	0.138007
7	-3.416719	-1.873730	-0.018554
6	-2.262337	3.140688	-0.048475
7	-1.761904	1.876590	-0.044916

(3)-HCO₂

E = -1322.67414470 a.u.

77	-0.672907	0.051507	-0.151473
7	0.330369	1.846143	-0.749591
7	1.466866	-0.428730	-0.338984
6	2.289767	0.636423	-0.225608
6	3.610863	0.530992	0.123186
1	3.220316	3.326141	-0.449972
1	4.278158	1.377944	0.228189
6	1.120553	3.942172	-0.918619
6	4.046886	-0.779862	0.359061
6	-0.006573	3.102522	-0.999198
6	2.021821	-1.641181	-0.162435
6	-1.226538	0.727949	1.794547
6	-0.877440	-0.660791	1.863079
6	-2.371479	0.851749	0.905619
6	-2.712531	-0.458864	0.445164
6	-1.786641	-1.405218	1.017754
6	-0.549688	1.837402	2.517359
1	-0.569171	2.767533	1.938477
1	0.496536	1.588795	2.727776
1	-1.047096	2.029573	3.476209
6	0.239955	-1.232479	2.660626
6	-3.099749	2.108372	0.590626
1	1.114758	-0.570766	2.661538
1	-2.463891	2.987714	0.730708
1	0.547119	-2.212115	2.281365
1	-3.471313	2.109997	-0.439401
1	-0.078388	-1.362340	3.702321
1	-3.964215	2.213996	1.258253
6	-3.798536	-0.792894	-0.510317
6	-1.870548	-2.885988	0.902655
1	-3.511498	-1.623107	-1.164627
1	-2.333922	-3.189036	-0.041034
1	-4.060633	0.066209	-1.135846
1	-0.884653	-3.356341	0.967264
1	-4.698508	-1.097903	0.038549
1	-2.484602	-3.288033	1.718731
1	1.156695	5.008883	-1.084800

8	1.280194	-2.704282	-0.356103
1	0.459750	-2.495320	-0.898374
1	-1.037184	3.351296	-1.223171
8	5.304091	-0.946206	0.741071
1	5.478881	-1.895218	0.859563
7	3.280430	-1.853852	0.204411
6	-0.830263	-1.453038	-2.786778
1	-1.067717	-1.508954	-3.866583
8	-0.606924	-2.508723	-2.168887
8	-0.814840	-0.266969	-2.320583
6	2.168738	3.119673	-0.596476
7	1.668430	1.855416	-0.496853

(3)-HCO₂-TS

E = -1322.63530342 a.u.

77	-0.675224	0.048705	0.092702
7	0.287244	1.915609	0.390882
7	1.448272	-0.273391	-0.344106
6	2.275441	0.781887	-0.186577
6	3.641018	0.704306	-0.308825
1	3.134987	3.480887	0.100473
1	4.304172	1.550067	-0.175741
6	1.028482	4.024435	0.621075
6	4.132432	-0.570423	-0.598346
6	-0.077094	3.154908	0.685525
6	2.050991	-1.450499	-0.568280
6	-2.690089	-0.702725	0.420713
6	-2.005259	-1.568076	-0.504964
6	-2.748231	0.608751	-0.147936
6	-2.148545	0.552466	-1.470954
6	-1.700154	-0.784032	-1.686082
6	-3.240714	-1.112387	1.738600
1	-3.300171	-0.264680	2.428712
1	-2.630776	-1.894500	2.202008
1	-4.254436	-1.513220	1.608650
6	-1.858574	-3.040805	-0.334473

6	-3.420884	1.790851	0.454042
1	-1.483947	-3.293851	0.663640
1	-3.257200	1.838630	1.536226
1	-1.174967	-3.470501	-1.071097
1	-3.065978	2.725558	0.008459
1	-2.834195	-3.529557	-0.454355
1	-4.503489	1.735240	0.282388
6	-2.067032	1.691995	-2.422377
6	-1.014703	-1.284249	-2.905890
1	-1.311603	1.518741	-3.195046
1	-0.415148	-0.499669	-3.379002
1	-1.819062	2.627487	-1.907397
1	-0.357920	-2.129021	-2.679839
1	-3.032236	1.840897	-2.923433
1	-1.757261	-1.625423	-3.638399
1	1.035506	5.088464	0.806721
8	1.243334	-2.492510	-0.622114
1	1.763544	-3.292889	-0.807362
1	-1.108126	3.373296	0.935662
7	3.351075	-1.641671	-0.716742
1	-0.159480	-0.279985	1.695147
6	0.725670	-1.249723	2.512802
8	-0.018643	-2.124430	2.843364
8	1.804731	-0.735517	2.536631
8	5.438925	-0.719169	-0.746320
1	5.635815	-1.653588	-0.929753
7	1.623987	1.966292	0.134740
6	2.094472	3.240399	0.270178

(3)-Hydride

E = -1134.14598673 a.u.

77	0.621632	0.021113	-0.325622
7	-0.401036	1.864623	-0.208608
7	-1.567414	-0.442413	-0.186370
6	-2.401363	0.612540	-0.086573
6	-3.767366	0.496617	0.023004

1	-3.316185	3.301605	0.088201
1	-4.438564	1.343581	0.096726
6	-1.185805	3.965989	-0.054824
6	-4.244502	-0.814611	0.045884
6	-0.050022	3.141757	-0.182063
6	-2.156519	-1.645504	-0.179384
6	2.614634	-0.726946	-0.697559
6	1.923983	-1.693644	0.103072
6	2.677061	0.496489	0.054299
6	2.133725	0.234154	1.384802
6	1.668509	-1.093519	1.410156
6	3.199130	-0.957671	-2.047085
1	3.232473	-0.033191	-2.632685
1	2.624846	-1.699488	-2.611616
1	4.227060	-1.331580	-1.952620
6	1.761394	-3.137802	-0.224821
6	3.383894	1.740702	-0.358771
1	1.552769	-3.289821	-1.289386
1	3.258451	1.935818	-1.429545
1	0.948442	-3.593262	0.347875
1	3.017864	2.613781	0.192446
1	2.683961	-3.683701	0.014619
1	4.460354	1.656399	-0.158441
6	2.062295	1.252850	2.466326
6	0.994320	-1.802201	2.531211
1	1.421404	0.925502	3.290652
1	0.693715	-1.112199	3.325534
1	1.669117	2.206344	2.090563
1	0.099854	-2.333462	2.184392
1	3.061071	1.455554	2.873971
1	1.665226	-2.551679	2.971015
1	-1.209464	5.044864	-0.006883
8	-1.335711	-2.670702	-0.324972
1	-1.852651	-3.493947	-0.316187
1	0.993272	3.419458	-0.268697
8	-5.550344	-1.002218	0.172166
1	-5.740241	-1.955423	0.165933
7	-3.452697	-1.880247	-0.054233

1	0.281970	0.124477	-1.907366
7	-1.758815	1.844565	-0.102253
6	-2.255411	3.114518	-0.006491

(3)-Hydride-H₂O-TS

E = -1210.53544284 a.u.

77	0.650846	0.037087	-0.268735
7	-0.341567	1.882380	-0.547383
7	-1.489564	-0.403780	-0.193858
6	-2.316429	0.653569	-0.131952
6	-3.666668	0.566460	0.091390
1	-3.222868	3.361645	-0.231894
1	-4.331453	1.420142	0.134560
6	-1.113230	3.985786	-0.643939
6	-4.122728	-0.744971	0.284223
6	0.010681	3.146043	-0.740503
6	-2.046440	-1.668142	-0.096022
6	2.698801	-0.750928	-0.121702
6	1.833882	-1.549548	0.698001
6	2.656268	0.601357	0.350151
6	1.776360	0.636133	1.506768
6	1.267264	-0.680496	1.712247
6	3.505253	-1.243831	-1.267780
1	3.600965	-0.482681	-2.048746
1	3.066467	-2.143991	-1.709874
1	4.515814	-1.500059	-0.924754
6	1.710916	-3.031315	0.646464
6	3.465738	1.731234	-0.179937
1	1.718623	-3.404699	-0.382971
1	3.579635	1.662765	-1.266989
1	0.792816	-3.376699	1.128442
1	3.013206	2.699749	0.055232
1	2.558676	-3.490060	1.171861
1	4.470695	1.719535	0.261176
6	1.464425	1.839032	2.323215
6	0.321968	-1.108740	2.775755

1	0.452442	1.785773	2.739299
1	-0.274377	-0.267239	3.143161
1	1.540579	2.758291	1.732842
1	-0.362779	-1.882438	2.411304
1	2.168845	1.923606	3.159860
1	0.874469	-1.526834	3.626557
1	-1.140645	5.059292	-0.760711
8	-1.322763	-2.674704	-0.284457
1	-0.419563	-2.530507	-1.691211
1	1.045114	3.395586	-0.944461
8	-5.419920	-0.897063	0.567338
1	-5.600053	-1.846447	0.666522
7	-3.366097	-1.819882	0.198384
1	0.456446	-0.047529	-2.021192
8	0.069869	-2.234774	-2.512416
1	0.910881	-2.717737	-2.509352
1	0.350232	-1.003206	-2.130636
6	-2.171720	3.154681	-0.377647
7	-1.680327	1.885916	-0.321643

(3-H⁺)

E = -1132.95929547 a.u.

77	0.558361	0.047446	-0.001679
7	-0.443181	1.884661	-0.011957
7	-1.535457	-0.447535	-0.050026
6	-2.428577	0.588368	-0.016190
6	-3.771483	0.432154	0.018523
1	-3.380531	3.275840	-0.035576
1	-4.454671	1.274103	0.048580
6	-1.250892	3.976281	-0.029757
6	-4.297014	-0.914377	0.029650
6	-0.105475	3.175264	-0.019789
6	-2.113626	-1.674463	-0.051910
6	2.372795	-0.178604	-1.117032
6	1.938618	-1.510374	-0.725767
6	2.650234	0.576295	0.070812

6	2.307288	-0.253714	1.190154
6	1.894974	-1.556689	0.692392
6	2.542399	0.278090	-2.515644
1	2.455554	1.366060	-2.594977
1	1.797666	-0.182198	-3.173110
1	3.535985	-0.011327	-2.882968
6	1.615518	-2.591829	-1.689955
6	3.187339	1.960140	0.134059
1	0.904632	-2.245356	-2.448861
1	2.925023	2.533376	-0.761426
1	1.191522	-3.467907	-1.193921
1	2.812036	2.495383	1.013105
1	2.529268	-2.900682	-2.213244
1	4.281674	1.933196	0.202665
6	2.403455	0.111026	2.622443
6	1.531263	-2.702670	1.563925
1	1.613215	-0.371861	3.206626
1	0.789978	-2.408307	2.315444
1	2.335329	1.193460	2.766654
1	1.128502	-3.541042	0.990280
1	3.368404	-0.224665	3.025334
1	2.422927	-3.049187	2.101561
1	-1.297976	5.055222	-0.038118
8	-1.231926	-2.669304	-0.102916
1	-1.726348	-3.504858	-0.092171
1	0.938631	3.462369	-0.019937
8	-5.514441	-1.145715	0.078985
7	-3.385609	-1.954930	-0.013909
6	-2.313211	3.100029	-0.028478
7	-1.803660	1.845749	-0.018368

(3-H⁺)-HCO₂

E = -1322.21314350 a.u.

77	-0.642845	0.060698	-0.142113
7	0.391449	1.853692	-0.692156
7	1.455944	-0.464606	-0.363648

6	2.321914	0.571413	-0.202410
6	3.617718	0.422682	0.169038
1	3.327856	3.232132	-0.462696
1	4.290046	1.265296	0.292173
6	1.237823	3.921837	-0.889936
6	4.092836	-0.918892	0.414663
6	0.084931	3.120706	-0.942212
6	1.995424	-1.692646	-0.189719
6	-1.151808	0.652085	1.843238
6	-0.878432	-0.755407	1.831796
6	-2.296705	0.887301	0.976453
6	-2.703367	-0.370979	0.437851
6	-1.824419	-1.398346	0.949140
6	-0.413215	1.682143	2.622257
1	-0.386833	2.642553	2.095722
1	0.620071	1.368160	2.808063
1	-0.892634	1.847091	3.595330
6	0.202578	-1.431229	2.597181
6	-2.952541	2.200852	0.741735
1	1.110343	-0.817712	2.640689
1	-2.258118	3.031950	0.899758
1	0.461331	-2.399845	2.158905
1	-3.352007	2.275613	-0.275217
1	-0.128863	-1.606424	3.628126
1	-3.788788	2.331912	1.440145
6	-3.810532	-0.598563	-0.526088
6	-1.996095	-2.863494	0.754756
1	-3.550971	-1.381401	-1.247095
1	-2.469973	-3.087341	-0.205891
1	-4.054570	0.313385	-1.080137
1	-1.040223	-3.394678	0.796430
1	-4.713983	-0.922457	0.005991
1	-2.639046	-3.272640	1.544939
1	1.307590	4.985926	-1.062909
8	1.195536	-2.731265	-0.422586
1	0.401472	-2.454617	-0.953707
1	-0.941922	3.398682	-1.147940
8	5.254654	-1.142828	0.805384

7	3.220883	-1.964216	0.181249
6	-0.830122	-1.268235	-2.864190
1	-1.019488	-1.245202	-3.955587
8	-0.690955	-2.367671	-2.303455
8	-0.778331	-0.113181	-2.325351
6	2.266953	3.062228	-0.586934
7	1.730713	1.820085	-0.471731

(3-H⁺)-HCO₂-TS

E = -1322.17351678 a.u.

77	-0.535007	-0.031727	0.006082
7	0.418107	1.856116	-0.071565
7	1.581023	-0.425010	0.199818
6	2.435515	0.614285	0.005318
6	3.787987	0.491602	-0.017526
1	3.320446	3.298229	-0.361908
1	4.446019	1.338336	-0.179631
6	1.178181	3.952718	-0.317628
6	4.348299	-0.821218	0.177615
6	0.049792	3.126876	-0.173759
6	2.186521	-1.594785	0.477758
6	-2.664155	-0.500654	-0.145539
6	-1.882351	-1.713741	-0.270343
6	-2.297622	0.373557	-1.213978
6	-1.319729	-0.298497	-2.042823
6	-1.083791	-1.585526	-1.465335
6	-3.728550	-0.250210	0.863504
1	-3.804731	0.812033	1.118406
1	-3.552100	-0.815426	1.783649
1	-4.700520	-0.565426	0.461264
6	-2.036888	-2.917517	0.593720
6	-2.876598	1.718753	-1.471424
1	-2.060294	-2.641842	1.654587
1	-3.156179	2.221359	-0.538292
1	-1.218533	-3.628701	0.449241
1	-2.177962	2.362446	-2.016035

1	-2.976198	-3.438471	0.366528
1	-3.783542	1.624622	-2.082396
6	-0.705496	0.237041	-3.287836
6	-0.163101	-2.606253	-2.033534
1	0.273022	-0.218973	-3.474352
1	0.796290	-2.165458	-2.328203
1	-0.565586	1.322784	-3.229659
1	0.033252	-3.418314	-1.329471
1	-1.339906	0.032409	-4.160270
1	-0.612833	-3.039690	-2.935767
1	1.195712	5.027820	-0.421469
8	1.336447	-2.571105	0.816045
1	1.859914	-3.371305	0.983725
1	-1.000560	3.390689	-0.139232
8	5.572922	-1.038002	0.114739
7	3.468642	-1.849508	0.467966
1	-0.341178	0.160365	1.690989
6	-1.266050	0.626070	2.889224
8	-1.355544	1.812303	2.791182
8	-1.486165	-0.398923	3.459624
7	1.774765	1.843061	-0.149258
6	2.258364	3.105616	-0.294327

(3-H⁺)-Hydride

E = -1133.68534303 a.u.

77	0.601445	0.025964	-0.343167
7	-0.435770	1.860520	-0.246950
7	-1.549650	-0.471709	-0.211667
6	-2.416211	0.566627	-0.082435
6	-3.762010	0.428999	0.057497
1	-3.360424	3.250570	0.081961
1	-4.426562	1.280375	0.156833
6	-1.243410	3.951449	-0.113143
6	-4.305101	-0.902936	0.079592
6	-0.100112	3.145131	-0.250058
6	-2.144630	-1.678555	-0.222313

6	2.648693	-0.604838	-0.633709
6	1.943327	-1.655449	0.046377
6	2.619789	0.556364	0.207721
6	1.989662	0.185956	1.470473
6	1.575602	-1.156755	1.366809
6	3.335124	-0.725682	-1.949329
1	3.365609	0.232275	-2.478595
1	2.835108	-1.455624	-2.594449
1	4.370184	-1.062525	-1.804944
6	1.881838	-3.078471	-0.393561
6	3.305493	1.850313	-0.064154
1	1.697494	-3.161305	-1.470362
1	3.279605	2.103215	-1.129925
1	1.091915	-3.628255	0.125674
1	2.849277	2.674208	0.495407
1	2.833695	-3.583685	-0.180598
1	4.359301	1.795125	0.240059
6	1.813844	1.112364	2.621910
6	0.856753	-1.957858	2.394180
1	1.093807	0.722467	3.348288
1	0.429725	-1.323200	3.177116
1	1.460942	2.098440	2.294060
1	0.044050	-2.540767	1.946473
1	2.766521	1.269907	3.144354
1	1.540943	-2.670252	2.873220
1	-1.285834	5.030543	-0.084282
8	-1.295689	-2.695983	-0.415911
1	-1.819211	-3.513480	-0.407041
1	0.937833	3.433902	-0.362384
8	-5.522672	-1.125495	0.228495
7	-3.416537	-1.950091	-0.081301
1	0.285789	0.116828	-1.934085
6	-2.298686	3.078689	-0.028659
7	-1.785754	1.820102	-0.111051

(3-H⁺)-Hydride-H₂O-TS

E = -1210.06449402 a.u.

77	0.667540	0.038258	-0.317077
7	-0.307425	1.869412	-0.733420
7	-1.434989	-0.441474	-0.492761
6	-2.248233	0.607709	-0.219622
6	-3.525013	0.499058	0.225868
1	-3.185038	3.312425	-0.300625
1	-4.147521	1.362362	0.437681
6	-1.092740	3.967795	-0.772675
6	-4.030012	-0.838592	0.436330
6	0.031798	3.140472	-0.920145
6	-1.988039	-1.706735	-0.350476
6	2.730538	-0.515504	0.203810
6	1.816599	-1.517307	0.700954
6	2.399810	0.737250	0.803220
6	1.283113	0.516299	1.713167
6	0.946685	-0.870983	1.658681
6	3.812329	-0.762785	-0.786231
1	4.093697	0.154629	-1.312529
1	3.505159	-1.504988	-1.531167
1	4.706371	-1.149244	-0.280770
6	1.886718	-2.979295	0.426912
6	3.115292	2.026420	0.604271
1	2.355224	-3.181733	-0.542320
1	3.492869	2.121863	-0.419449
1	0.892436	-3.436639	0.424655
1	2.465262	2.883194	0.809009
1	2.488036	-3.482268	1.195350
1	3.973817	2.092666	1.284695
6	0.624804	1.554875	2.551200
6	-0.139433	-1.535014	2.426606
1	-0.404006	1.269363	2.797700
1	-1.006724	-0.876149	2.552721
1	0.592917	2.521938	2.036138
1	-0.473934	-2.452668	1.932982
1	1.169010	1.697603	3.493180
1	0.222661	-1.804535	3.426574
1	-1.136820	5.041783	-0.881002

8	-1.264779	-2.703581	-0.677400
1	-0.287381	-2.450011	-1.860127
1	1.058105	3.397906	-1.152914
8	-5.179679	-1.023030	0.905363
7	-3.231949	-1.897212	0.104491
1	0.777824	-0.019431	-2.069339
8	0.323056	-2.172849	-2.631254
1	1.142085	-2.678578	-2.518526
1	0.608901	-0.960476	-2.186550
7	-1.634876	1.859263	-0.455722
6	-2.135485	3.119814	-0.477239

(3-H⁺)

E = -1132.96613804 a.u.

77	0.569967	0.040663	-0.013441
7	-0.414496	1.893035	-0.025276
7	-1.517281	-0.435014	-0.069475
6	-2.389712	0.596135	-0.031538
6	-3.746119	0.456374	0.008286
1	-3.352326	3.287110	-0.044481
1	-4.440478	1.286917	0.043042
6	-1.222900	3.985184	-0.038627
6	-4.180328	-0.885051	0.029485
6	-0.076725	3.182008	-0.030186
6	-2.036863	-1.737036	-0.085894
6	2.414394	-0.195973	-1.084362
6	1.948544	-1.524592	-0.716365
6	2.673989	0.543350	0.114977
6	2.287402	-0.289368	1.217506
6	1.871073	-1.583331	0.699908
6	2.620275	0.270216	-2.475127
1	2.569429	1.361232	-2.543497
1	1.871120	-0.158321	-3.149026
1	3.609252	-0.046478	-2.832253
6	1.640541	-2.593389	-1.699461
6	3.222736	1.921388	0.204079

1	0.948204	-2.233620	-2.469326
1	3.006408	2.497635	-0.701796
1	1.199544	-3.472018	-1.223523
1	2.812804	2.461050	1.064894
1	2.563684	-2.901599	-2.206549
1	4.312339	1.884149	0.323661
6	2.351340	0.062860	2.654951
6	1.478910	-2.731569	1.555788
1	1.541126	-0.415476	3.215029
1	0.701347	-2.443730	2.272513
1	2.292172	1.144920	2.806314
1	1.110265	-3.574274	0.966340
1	3.301836	-0.286799	3.079783
1	2.349048	-3.068617	2.133336
1	-1.268490	5.064192	-0.045075
8	-1.260240	-2.694667	-0.164727
1	0.967224	3.469753	-0.029448
8	-5.498768	-1.075648	0.103982
1	-5.668207	-2.032510	0.116422
7	-3.390134	-1.930376	-0.011960
6	-2.285040	3.110914	-0.038344
7	-1.773924	1.855102	-0.031275

(3-H⁺)-HCO₂

E = -1322.21359143 a.u.

77	0.628099	0.051435	0.056226
7	-0.387710	1.913273	0.133151
7	-1.513234	-0.405246	-0.152073
6	-2.362025	0.628967	-0.054984
6	-3.729344	0.526336	-0.119477
1	-3.325031	3.299251	0.284938
1	-4.405893	1.368465	-0.046453
6	-1.198538	3.982338	0.428926
6	-4.184756	-0.785136	-0.313352
6	-0.048433	3.179668	0.331386
6	-2.051739	-1.681239	-0.308178

6	1.934481	0.476538	-1.579757
6	1.498036	-0.886508	-1.677125
6	2.669251	0.621429	-0.333501
6	2.680643	-0.654693	0.312221
6	1.951863	-1.593356	-0.496195
6	1.712528	1.553446	-2.580094
1	1.603080	2.530881	-2.097945
1	0.812658	1.366048	-3.174766
1	2.565854	1.613608	-3.267200
6	0.733559	-1.478138	-2.807055
6	3.359160	1.845661	0.155587
1	0.132450	-0.719436	-3.319237
1	2.993718	2.745156	-0.349801
1	0.061231	-2.268408	-2.459713
1	3.220977	1.977989	1.234837
1	1.422253	-1.913407	-3.542108
1	4.437697	1.775234	-0.034731
6	3.319801	-0.957275	1.618183
6	1.889113	-3.061640	-0.256212
1	2.784913	-1.747603	2.153714
1	1.626606	-3.293462	0.780859
1	3.363297	-0.070021	2.258433
1	1.158244	-3.545089	-0.909042
1	4.349260	-1.300835	1.452738
1	2.870853	-3.509357	-0.458646
1	-1.241931	5.050470	0.584598
8	-1.296354	-2.664526	-0.361813
1	0.996115	3.458619	0.403475
8	-5.510501	-0.953711	-0.413460
1	-5.681562	-1.899781	-0.551000
7	-3.408776	-1.841257	-0.405429
6	-0.073231	-0.820362	2.913453
1	-0.351229	-0.568145	3.959286
8	0.093643	-2.001345	2.597244
8	0.031247	0.217012	2.164597
6	-2.258441	3.122686	0.281348
7	-1.740711	1.875742	0.107887

(3-H⁺)-HCO₂-TS

E = -1322.18120315 a.u.

77	-0.542113	-0.042203	0.008516
7	0.389333	1.852458	-0.088579
7	1.567219	-0.419337	0.176884
6	2.399324	0.615281	-0.004982
6	3.768862	0.514378	-0.017095
1	3.288626	3.301250	-0.404504
1	4.438542	1.351298	-0.169332
6	1.144979	3.948506	-0.357711
6	4.234872	-0.793309	0.178779
6	0.018088	3.120404	-0.201946
6	2.115774	-1.659066	0.495107
6	-2.663838	-0.576938	-0.098725
6	-1.847492	-1.755826	-0.290146
6	-2.359258	0.348974	-1.139926
6	-1.385030	-0.259933	-2.023444
6	-1.087204	-1.557417	-1.502541
6	-3.700103	-0.404142	0.954628
1	-3.821000	0.646799	1.238240
1	-3.460265	-0.980607	1.853155
1	-4.669550	-0.760277	0.581263
6	-1.958295	-3.004925	0.514805
6	-2.992299	1.681154	-1.332814
1	-1.942344	-2.789390	1.589599
1	-3.246666	2.147392	-0.373942
1	-1.143220	-3.700852	0.298207
1	-2.338671	2.363357	-1.886326
1	-2.903699	-3.518717	0.295918
1	-3.921174	1.576140	-1.908170
6	-0.830058	0.359116	-3.258000
6	-0.152096	-2.534149	-2.121116
1	0.122433	-0.101470	-3.540600
1	0.714018	-2.035459	-2.570319
1	-0.657295	1.433356	-3.122041
1	0.213436	-3.262765	-1.393518

1	-1.521688	0.241187	-4.102350
1	-0.666749	-3.082166	-2.920554
1	1.158400	5.022566	-0.472634
8	1.366460	-2.588051	0.840640
1	-1.032226	3.384042	-0.164591
8	5.561405	-0.975695	0.118956
1	5.741734	-1.914608	0.290016
7	3.470229	-1.835001	0.425177
1	-0.329732	0.157041	1.689642
6	-1.233040	0.635814	2.903450
8	-1.335333	1.819393	2.789696
8	-1.429181	-0.382900	3.492663
7	1.745959	1.842605	-0.171595
6	2.227449	3.106151	-0.330274

(3-H⁺)-Hydride

E = -1133.69210777 a.u.

77	0.612175	0.020983	-0.343278
7	-0.390281	1.869514	-0.254055
7	-1.516461	-0.454462	-0.220975
6	-2.362708	0.576737	-0.099937
6	-3.726131	0.456842	0.035521
1	-3.316243	3.266111	0.059902
1	-4.403030	1.296462	0.131792
6	-1.196979	3.961415	-0.126262
6	-4.171689	-0.869890	0.054218
6	-0.053035	3.152677	-0.257279
6	-2.053284	-1.739778	-0.260219
6	1.938530	-1.676074	0.031251
6	1.525318	-1.199845	1.344962
6	2.670209	-0.613551	-0.604464
6	2.629941	0.526193	0.261179
6	1.951881	0.138480	1.491949
6	1.897439	-3.090475	-0.438508
1	1.716948	-3.154298	-1.517320
1	1.113627	-3.661491	0.066675

1	2.856446	-3.585455	-0.233527
6	0.750506	-2.010836	2.324126
6	3.389313	-0.713811	-1.904266
1	-0.070655	-2.543892	1.830337
1	2.893923	-1.418647	-2.580191
1	0.323964	-1.387243	3.116344
1	3.450923	0.256059	-2.408605
1	1.394175	-2.763571	2.797361
1	4.414116	-1.073254	-1.741821
6	3.327601	1.822735	0.036134
6	1.747771	1.044805	2.655164
1	2.855377	2.637554	0.595751
1	1.402559	2.036759	2.336460
1	3.337287	2.095732	-1.024968
1	1.009632	0.642833	3.356463
1	4.370745	1.758906	0.373507
1	2.686936	1.192976	3.204198
1	-1.236920	5.040686	-0.099268
8	-1.301841	-2.709102	-0.464310
1	0.985664	3.440644	-0.364928
8	-5.488872	-1.063779	0.219971
1	-5.652516	-2.020690	0.203388
7	-3.397085	-1.924350	-0.082583
1	0.309658	0.109157	-1.940963
7	-1.741456	1.831464	-0.123155
6	-2.254248	3.092193	-0.044917

(3-2H⁺)

E = -1132.49598856 a.u.

77	0.530480	0.037414	0.000341
7	-0.465082	1.875182	-0.003078
7	-1.509138	-0.469935	-0.051287
6	-2.419626	0.543783	-0.019479
6	-3.758860	0.372384	0.012659
1	-3.415565	3.228264	-0.055964
1	-4.454784	1.204383	0.042264

6	-1.293495	3.957143	-0.036305
6	-4.250269	-0.993341	0.034475
6	-0.138943	3.169807	-0.013987
6	-2.031539	-1.786307	-0.068051
6	2.358097	-0.148073	-1.111028
6	1.940182	-1.489486	-0.734457
6	2.630263	0.595349	0.082146
6	2.289536	-0.244762	1.191688
6	1.895472	-1.549633	0.682751
6	2.526561	0.324223	-2.506132
1	2.408905	1.410185	-2.577474
1	1.797695	-0.148986	-3.172288
1	3.529638	0.066927	-2.871351
6	1.657888	-2.570192	-1.712827
6	3.148326	1.986524	0.158927
1	0.936651	-2.240838	-2.469659
1	2.885548	2.563247	-0.734361
1	1.264361	-3.467143	-1.229687
1	2.759843	2.511449	1.038684
1	2.583820	-2.839503	-2.236943
1	4.242475	1.976150	0.235877
6	2.379302	0.109770	2.628452
6	1.566756	-2.711835	1.548070
1	1.595554	-0.389857	3.207578
1	0.809419	-2.447568	2.294987
1	2.290264	1.189835	2.780974
1	1.197600	-3.562292	0.970018
1	3.348685	-0.210775	3.032897
1	2.466421	-3.028658	2.091014
1	-1.353522	5.035495	-0.049808
8	-1.204971	-2.720723	-0.137492
1	0.901540	3.468922	-0.007610
8	-5.483274	-1.212693	0.098401
7	-3.355229	-2.015472	-0.009667
6	-2.346085	3.066887	-0.039506
7	-1.821544	1.820767	-0.019837

(3-2H⁺)-HCO₂

E = -1321.73642667 a.u.

77	0.589208	0.057448	0.067342
7	-0.454089	1.882703	0.251880
7	-1.510364	-0.450663	-0.095304
6	-2.401689	0.561134	-0.013134
6	-3.750676	0.424819	-0.111215
1	-3.408480	3.223528	0.308302
1	-4.430359	1.268144	-0.046917
6	-1.296000	3.940212	0.523164
6	-4.253123	-0.908853	-0.334208
6	-0.134342	3.155202	0.461774
6	-2.039514	-1.738248	-0.246030
6	1.796948	0.588886	-1.613179
6	1.398341	-0.784879	-1.743923
6	2.599921	0.707765	-0.408598
6	2.686303	-0.590302	0.178898
6	1.947671	-1.523293	-0.627858
6	1.486326	1.694818	-2.556980
1	1.384726	2.651710	-2.033927
1	0.554964	1.506097	-3.100835
1	2.291974	1.802378	-3.294266
6	0.596004	-1.359671	-2.856516
6	3.287443	1.929255	0.093876
1	-0.042931	-0.600843	-3.320216
1	2.902740	2.836361	-0.382715
1	-0.044208	-2.174287	-2.504459
1	3.171018	2.036540	1.178716
1	1.259019	-1.761327	-3.633258
1	4.363029	1.875543	-0.118393
6	3.414110	-0.915116	1.432900
6	1.954610	-3.002063	-0.456595
1	2.984222	-1.787070	1.934518
1	1.854161	-3.286775	0.595593
1	3.400699	-0.071608	2.131668
1	1.143300	-3.476392	-1.014311
1	4.464105	-1.139861	1.204119

1	2.904137	-3.413941	-0.822899
1	-1.360567	5.007140	0.680666
8	-1.234425	-2.702898	-0.254111
1	0.903870	3.446707	0.566715
8	-5.487890	-1.113074	-0.481872
7	-3.366498	-1.940198	-0.378619
6	0.118656	-0.956821	2.949758
1	-0.126234	-0.761108	4.016771
8	0.359209	-2.112144	2.587724
8	0.109135	0.109157	2.237223
7	-1.802503	1.826903	0.179140
6	-2.339697	3.063155	0.337598

(3-2H⁺)-HCO₂-TS

E = -1321.70659347 a.u.

77	0.631345	0.044805	-0.108982
7	-0.343640	1.875407	-0.497045
7	-1.433213	-0.305613	0.330780
6	-2.296873	0.718115	0.175118
6	-3.645644	0.629739	0.330326
1	-3.187810	3.412701	-0.175102
1	-4.307846	1.477937	0.190733
6	-1.102842	3.965819	-0.788466
6	-4.171330	-0.663373	0.695927
6	0.005899	3.107513	-0.854836
6	-1.995637	-1.569816	0.530176
6	2.727123	-0.541577	-0.335672
6	2.046901	-1.494336	0.513799
6	2.663021	0.737112	0.290927
6	1.971352	0.589993	1.560551
6	1.611229	-0.785864	1.695255
6	3.400667	-0.866236	-1.621850
1	3.465317	0.010471	-2.274526
1	2.870183	-1.657292	-2.162805
1	4.422479	-1.222268	-1.434990
6	2.030425	-2.969128	0.298868

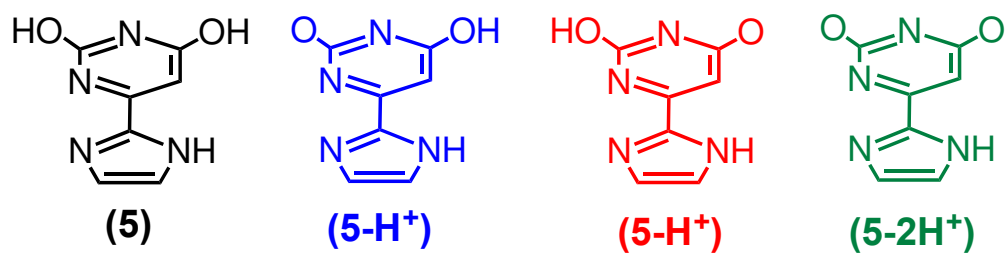
6	3.295057	1.986802	-0.214979
1	1.770796	-3.224155	-0.734903
1	3.216042	2.067284	-1.305028
1	1.312671	-3.464487	0.958808
1	2.844007	2.879265	0.230845
1	3.022896	-3.392522	0.503599
1	4.362789	1.997491	0.039712
6	1.726751	1.686530	2.536691
6	0.890488	-1.393135	2.846084
1	0.904954	1.439503	3.216786
1	0.297338	-0.647012	3.385538
1	1.468956	2.622472	2.026611
1	0.216308	-2.189250	2.514822
1	2.619913	1.879292	3.145216
1	1.606908	-1.827438	3.554991
1	-1.130604	5.021245	-1.017731
8	-1.223952	-2.562887	0.482786
1	1.026781	3.320196	-1.147594
8	-5.400627	-0.819079	0.925301
7	-3.313820	-1.719775	0.761004
1	0.144097	-0.327540	-1.701269
6	-0.682926	-1.448951	-2.480412
8	-1.767644	-0.966361	-2.591685
8	0.115061	-2.308257	-2.699457
6	-2.149492	3.179636	-0.367005
7	-1.664681	1.923370	-0.197691

(3-2H⁺)-Hydride

E = -1133.21384805 a.u.

77	0.587246	0.020792	-0.349858
7	-0.439644	1.854360	-0.272282
7	-1.516415	-0.490972	-0.224553
6	-2.394121	0.524210	-0.097871
6	-3.741093	0.380973	0.041934
1	-3.374829	3.206030	0.074771
1	-4.414065	1.226404	0.139844

6	-1.266714	3.935561	-0.134172
6	-4.249315	-0.966361	0.074351
6	-0.115573	3.143767	-0.277994
6	-2.056016	-1.779870	-0.269260
6	2.665857	-0.564657	-0.584478
6	1.946801	-1.645346	0.037501
6	2.592391	0.568993	0.284603
6	1.901063	0.164779	1.502276
6	1.503275	-1.181837	1.345119
6	3.404146	-0.646110	-1.875223
1	3.465310	0.329861	-2.368088
1	2.922337	-1.346122	-2.566070
1	4.429574	-1.000393	-1.704837
6	1.955540	-3.061354	-0.430766
6	3.268717	1.879800	0.074300
1	1.816278	-3.130865	-1.515410
1	3.281975	2.161137	-0.984702
1	1.166544	-3.649290	0.046370
1	2.777388	2.682526	0.635047
1	2.917010	-3.534957	-0.189160
1	4.310365	1.833356	0.419293
6	1.660321	1.062205	2.665916
6	0.732587	-2.012872	2.310906
1	0.916481	0.643830	3.351647
1	0.273676	-1.400485	3.093910
1	1.302929	2.049461	2.345910
1	-0.064263	-2.566973	1.800629
1	2.585902	1.224542	3.233908
1	1.386265	-2.748241	2.797704
1	-1.322791	5.014182	-0.105714
8	-1.265801	-2.735648	-0.489705
1	0.918427	3.443985	-0.396553
8	-5.480190	-1.185237	0.242159
7	-3.374344	-1.998447	-0.088798
1	0.286337	0.093644	-1.949676
6	-2.311672	3.047571	-0.042406
7	-1.785028	1.797596	-0.126911



(5)

E = -1133.45161237 a.u.

77	-0.596950	0.040312	-0.004847
7	0.352831	1.923222	-0.021160
7	1.562560	-0.426727	-0.070028
6	1.687297	1.908471	-0.042386
6	2.403362	0.651810	-0.051617
6	3.764429	0.520723	-0.023466
1	3.137234	3.432516	-0.044233
1	4.443859	1.366070	-0.006325
6	1.086676	4.019814	-0.007138
6	4.246245	-0.800744	0.011047
6	-0.035555	3.237498	0.002447
6	2.143297	-1.631340	-0.044770
6	-2.349747	-0.236565	1.170989
6	-1.873900	-1.556682	0.777668
6	-2.689006	0.497154	-0.015560
6	-2.364642	-0.340669	-1.138808
6	-1.900419	-1.624405	-0.640351
6	-2.485651	0.237018	2.567488
1	-2.382430	1.324861	2.632599
1	-1.735271	-0.225744	3.216254
1	-3.476731	-0.033664	2.955155
6	-1.454665	-2.609091	1.736319
6	-3.275982	1.859917	-0.085167
1	-0.763983	-2.205051	2.485462
1	-3.021898	2.454198	0.798481
1	-0.963848	-3.446781	1.234778

1	-2.931297	2.396037	-0.976106
1	-2.332669	-2.991489	2.271898
1	-4.369411	1.792139	-0.138984
6	-2.524499	0.002487	-2.569663
6	-1.527455	-2.762919	-1.517127
1	-1.766043	-0.495083	-3.182439
1	-0.809453	-2.452130	-2.284687
1	-2.459456	1.082318	-2.733760
1	-1.093667	-3.591101	-0.952036
1	-3.509693	-0.333703	-2.920076
1	-2.422343	-3.128979	-2.036019
1	1.213655	5.091608	0.002999
8	1.317825	-2.656504	-0.058889
1	1.824904	-3.485699	-0.038275
1	-1.075202	3.532633	0.023144
7	2.156787	3.164905	-0.033532
8	5.551406	-0.979130	0.067990
1	5.750528	-1.931013	0.105052
7	3.450085	-1.862220	-0.002693

(5)-HCO₂

E = -1322.71227144 a.u.

77	-0.672389	0.043173	-0.154151
7	0.292195	1.871842	-0.730262
7	1.484905	-0.429713	-0.322538
6	1.601824	1.891057	-0.497010
6	2.301488	0.657387	-0.220316
6	3.627776	0.548264	0.110877
1	3.036672	3.437297	-0.468537
1	4.287322	1.404307	0.203719
6	1.013836	3.969861	-0.882217
6	4.079282	-0.759381	0.348761
6	-0.091768	3.162518	-0.965241
6	2.050264	-1.631969	-0.154393
6	-1.232693	0.697611	1.794953
6	-0.877269	-0.690836	1.854541

6	-2.377550	0.820383	0.905275
6	-2.707244	-0.489240	0.432979
6	-1.779746	-1.434871	1.004117
6	-0.564092	1.805621	2.528599
1	-0.601812	2.743582	1.963319
1	0.488220	1.569538	2.722800
1	-1.052775	1.976706	3.495900
6	0.238151	-1.264697	2.653323
6	-3.120033	2.071734	0.600748
1	1.105864	-0.594496	2.676066
1	-2.502103	2.958563	0.770797
1	0.559744	-2.233748	2.258973
1	-3.470565	2.088646	-0.436635
1	-0.089919	-1.417565	3.688914
1	-4.000151	2.150104	1.251562
6	-3.787919	-0.822421	-0.529386
6	-1.857112	-2.915730	0.882004
1	-3.492112	-1.644120	-1.190669
1	-2.323077	-3.216078	-0.061346
1	-4.053317	0.040661	-1.148143
1	-0.868588	-3.381806	0.939285
1	-4.688476	-1.138957	0.011960
1	-2.465379	-3.326431	1.698169
1	1.135420	5.034399	-1.014746
8	1.322722	-2.708806	-0.345186
1	0.498313	-2.505234	-0.879912
1	-1.117934	3.427627	-1.181715
7	2.069076	3.151565	-0.583885
8	5.341577	-0.916624	0.721925
1	5.522688	-1.864106	0.842798
7	3.317036	-1.835294	0.204749
6	-0.793847	-1.470818	-2.785734
1	-1.017089	-1.528012	-3.868805
8	-0.570657	-2.525058	-2.165735
8	-0.793586	-0.284615	-2.319986

(5)-HCO₂-TS

E = -1322.67171664 a.u.

77	-0.617987	0.056568	0.033231
7	0.321144	1.935932	0.387472
7	1.551873	-0.300891	-0.200594
6	1.646378	1.975657	0.275688
6	2.375705	0.770521	-0.018058
6	3.744954	0.675324	-0.096945
1	3.062110	3.531152	0.457934
1	4.403326	1.525569	0.044221
6	1.007984	4.022141	0.738811
6	4.245768	-0.601337	-0.372111
6	-0.093175	3.207121	0.673528
6	2.155585	-1.473497	-0.416035
6	-2.725576	-0.443540	0.191623
6	-2.026474	-1.527118	-0.468851
6	-2.583200	0.729507	-0.615328
6	-1.820339	0.386773	-1.795207
6	-1.496309	-1.004492	-1.703857
6	-3.517473	-0.540113	1.445988
1	-3.457182	0.384797	2.029650
1	-3.175458	-1.365227	2.077185
1	-4.574618	-0.715494	1.206991
6	-2.009323	-2.946014	-0.014876
6	-3.191711	2.053694	-0.323706
1	-1.981138	-3.011234	1.078422
1	-3.190331	2.270189	0.750366
1	-1.140364	-3.483779	-0.405524
1	-2.671564	2.863397	-0.845539
1	-2.910603	-3.471980	-0.355866
1	-4.236339	2.060896	-0.660655
6	-1.486015	1.299532	-2.921134
6	-0.752884	-1.763374	-2.744592
1	-0.581032	0.969447	-3.442172
1	0.133797	-1.216412	-3.085839
1	-1.318978	2.323833	-2.568654
1	-0.434440	-2.745708	-2.388136
1	-2.301514	1.330683	-3.655633

1	-1.402344	-1.909617	-3.617046
1	1.104046	5.078786	0.938588
8	1.356513	-2.518648	-0.532206
1	1.892959	-3.309136	-0.711790
1	-1.134570	3.456292	0.824438
7	2.092649	3.230780	0.486788
7	3.464656	-1.666545	-0.519047
1	-0.128207	-0.267196	1.644001
6	-0.112640	-1.320564	2.778701
8	-0.953251	-0.930956	3.532712
8	0.750902	-2.092769	2.487208
8	5.557308	-0.753262	-0.483405
1	5.755841	-1.686174	-0.672357

(5)-Hydride

E = -1134.18406183 a.u.

77	0.620353	0.014345	-0.320293
7	-0.379878	1.879166	-0.201956
7	-1.585348	-0.445176	-0.170440
6	-1.705914	1.876129	-0.095332
6	-2.419865	0.627841	-0.082153
6	-3.787937	0.500344	0.008063
1	-3.141098	3.415463	0.078405
1	-4.457827	1.351201	0.069612
6	-1.092121	3.982681	-0.055407
6	-4.270921	-0.810298	0.034922
6	0.018915	3.186890	-0.177654
6	-2.176771	-1.642140	-0.167929
6	2.599825	-0.760715	-0.708049
6	1.905972	-1.718711	0.099756
6	2.682423	0.465327	0.039215
6	2.141775	0.214370	1.372162
6	1.658075	-1.108315	1.402665
6	3.173322	-1.001855	-2.060683
1	3.214325	-0.079132	-2.648667
1	2.586240	-1.737598	-2.620071

1	4.197303	-1.387996	-1.972089
6	1.732222	-3.163750	-0.219488
6	3.403901	1.698339	-0.382834
1	1.517036	-3.320665	-1.282106
1	3.269166	1.895118	-1.452238
1	0.919057	-3.610590	0.359837
1	3.057399	2.577246	0.171566
1	2.651992	-3.715293	0.018153
1	4.481286	1.598380	-0.194671
6	2.092937	1.233618	2.455179
6	0.980702	-1.802471	2.531321
1	1.424455	0.931588	3.267358
1	0.674968	-1.101093	3.313782
1	1.746327	2.203416	2.075969
1	0.088758	-2.340655	2.188873
1	3.091608	1.393573	2.882064
1	1.651096	-2.543468	2.986119
1	-1.201498	5.055430	-0.003381
8	-1.364446	-2.678577	-0.305526
1	-1.891400	-3.495103	-0.298362
1	1.057959	3.475177	-0.261220
7	-2.167229	3.141988	-0.005586
8	-5.579283	-0.997884	0.152307
1	-5.767573	-1.951349	0.151995
7	-3.477946	-1.874179	-0.051075
1	0.266650	0.117210	-1.901767

(5)-Hydride-H₂O-TS

E = -1210.57273171 a.u.

77	0.652360	0.038178	-0.262039
7	-0.314095	1.911007	-0.504377
7	-1.500485	-0.397625	-0.179388
6	-1.630081	1.921900	-0.324277
6	-2.330998	0.674606	-0.124335
6	-3.682865	0.569255	0.090880
1	-3.062891	3.469818	-0.306577

1	-4.347184	1.425548	0.131791
6	-1.024880	4.012105	-0.627950
6	-4.139502	-0.742904	0.289652
6	0.082332	3.208185	-0.690652
6	-2.056578	-1.656011	-0.102997
6	2.696641	-0.756512	-0.146658
6	1.831198	-1.594944	0.633727
6	2.651456	0.572834	0.387509
6	1.766505	0.552361	1.541515
6	1.259823	-0.774555	1.683120
6	3.500444	-1.192679	-1.317947
1	3.642430	-0.375711	-2.032392
1	3.026921	-2.030928	-1.839207
1	4.492479	-1.524720	-0.985997
6	1.706857	-3.072985	0.512310
6	3.460522	1.730632	-0.080087
1	1.833756	-3.404006	-0.524040
1	3.591626	1.712914	-1.167312
1	0.733660	-3.428287	0.862027
1	2.995996	2.684145	0.191055
1	2.483817	-3.562111	1.114239
1	4.458611	1.707963	0.375996
6	1.452694	1.716252	2.412255
6	0.314266	-1.251924	2.725902
1	0.459396	1.618641	2.863654
1	-0.301643	-0.433430	3.112705
1	1.477771	2.656376	1.850770
1	-0.352450	-2.028629	2.335744
1	2.185493	1.795513	3.224867
1	0.869228	-1.682772	3.568802
1	-1.140634	5.080460	-0.731032
8	-1.344306	-2.668297	-0.320433
1	-0.464307	-2.475483	-1.720443
1	1.115710	3.475790	-0.866163
7	-2.092337	3.186037	-0.394726
8	-5.435562	-0.899914	0.581114
1	-5.611070	-1.849802	0.683445
7	-3.377517	-1.812077	0.201654

1	0.463060	0.002909	-2.017440
8	0.016267	-2.165345	-2.544456
1	0.851906	-2.657501	-2.566218
1	0.319462	-0.953003	-2.145031

(5-H⁺)

E = -1132.99833410 a.u.

77	-0.557788	0.041465	0.000125
7	0.411993	1.902866	0.000163
7	1.549433	-0.461509	-0.001040
6	1.745515	1.870819	-0.000359
6	2.434545	0.598795	-0.000583
6	3.778859	0.436537	-0.000074
1	3.212707	3.375989	-0.000912
1	4.459003	1.283693	0.000331
6	1.166540	3.992395	-0.000196
6	4.317442	-0.909207	0.000342
6	0.037407	3.226242	0.000253
6	2.133052	-1.677280	-0.000610
6	-2.333586	-0.239586	1.154714
6	-1.903267	-1.556392	0.709825
6	-2.648998	0.552264	-0.000205
6	-2.333387	-0.240162	-1.154740
6	-1.903413	-1.556777	-0.709133
6	-2.471571	0.169558	2.571806
1	-2.391930	1.255018	2.685606
1	-1.706836	-0.305389	3.195084
1	-3.453297	-0.140610	2.954016
6	-1.549552	-2.669102	1.627403
6	-3.198220	1.932881	-0.000626
1	-0.830279	-2.341778	2.386974
1	-2.881797	2.489171	0.888322
1	-1.121273	-3.519693	1.091268
1	-2.881782	2.488616	-0.889918
1	-2.449478	-3.010837	2.154182
1	-4.294670	1.900613	-0.000657

6	-2.471144	0.168286	-2.572045
6	-1.549977	-2.669998	-1.626206
1	-1.706755	-0.307550	-3.195069
1	-0.830597	-2.343254	-2.385926
1	-2.390748	1.253624	-2.686461
1	-1.121975	-3.520481	-1.089695
1	-3.453093	-0.141436	-2.954047
1	-2.450014	-3.011688	-2.152822
1	1.307350	5.062515	-0.000289
8	1.265248	-2.688359	-0.000807
1	1.773584	-3.515335	-0.000326
1	-0.998789	3.533495	0.000576
7	2.229744	3.121434	-0.000542
8	5.538768	-1.128462	0.001149
7	3.412655	-1.951160	-0.000004

(5-H⁺)-HCO₂

E = -1322.25026126 a.u.

77	-0.640158	0.053617	-0.146856
7	0.360648	1.859014	-0.735662
7	1.471629	-0.476550	-0.344806
6	1.671096	1.840524	-0.510070
6	2.335199	0.584740	-0.225856
6	3.640905	0.433863	0.118445
1	3.147023	3.343350	-0.484649
1	4.311295	1.284279	0.208649
6	1.135190	3.939242	-0.887910
6	4.129860	-0.898416	0.392754
6	0.010128	3.164509	-0.966584
6	2.020056	-1.691374	-0.153395
6	-1.139417	0.680322	1.823458
6	-0.866369	-0.729812	1.840044
6	-2.291356	0.895431	0.960503
6	-2.695951	-0.373813	0.446750
6	-1.814721	-1.391326	0.976543
6	-0.402793	1.725415	2.584380

1	-0.375259	2.675269	2.038667
1	0.630393	1.415685	2.777465
1	-0.883362	1.909437	3.553556
6	0.213667	-1.389985	2.620169
6	-2.957148	2.200930	0.708246
1	1.118162	-0.771763	2.662085
1	-2.272055	3.039541	0.867428
1	0.479699	-2.362982	2.196298
1	-3.346816	2.264559	-0.313383
1	-0.124372	-1.553618	3.650974
1	-3.801941	2.329358	1.396866
6	-3.805426	-0.622553	-0.509635
6	-1.984835	-2.860433	0.809176
1	-3.540713	-1.409582	-1.224388
1	-2.455921	-3.103151	-0.148486
1	-4.061142	0.281245	-1.071840
1	-1.027902	-3.389154	0.861892
1	-4.703308	-0.951260	0.028892
1	-2.628701	-3.257302	1.604877
1	1.288357	4.999594	-1.020973
8	1.229295	-2.747174	-0.349340
1	0.428970	-2.492945	-0.881946
1	-1.008978	3.457452	-1.180344
7	2.171411	3.087405	-0.595871
8	5.302002	-1.105692	0.765757
7	3.256365	-1.949066	0.207567
6	-0.855498	-1.413823	-2.827442
1	-1.076846	-1.442410	-3.913329
8	-0.664809	-2.485505	-2.229062
8	-0.826522	-0.236681	-2.339415

(5-H⁺)-HCO₂-TS

E = -1322.21078896 a.u.

77	-0.536580	-0.041364	0.007857
7	0.390964	1.868124	-0.087718
7	1.597234	-0.429317	0.197629

6	1.718981	1.872540	-0.147008
6	2.446949	0.632179	0.001964
6	3.801588	0.501280	-0.008275
1	3.141352	3.413036	-0.366020
1	4.456331	1.353917	-0.166099
6	1.079323	3.966491	-0.348946
6	4.372146	-0.808575	0.187083
6	-0.024442	3.167998	-0.213451
6	2.208095	-1.589779	0.475276
6	-2.648161	-0.581126	-0.123480
6	-1.831212	-1.764563	-0.286030
6	-2.325401	0.330992	-1.175231
6	-1.338564	-0.292101	-2.033763
6	-1.050754	-1.581162	-1.486835
6	-3.704230	-0.389545	0.907084
1	-3.830755	0.666725	1.167405
1	-3.481188	-0.946945	1.821885
1	-4.666451	-0.753700	0.522907
6	-1.942220	-3.000617	0.538744
6	-2.955292	1.660539	-1.394178
1	-2.008969	-2.761135	1.606364
1	-3.215081	2.142632	-0.444608
1	-1.080220	-3.659252	0.397924
1	-2.296053	2.332784	-1.953208
1	-2.843666	-3.566929	0.269988
1	-3.880562	1.548362	-1.974027
6	-0.756525	0.287268	-3.274917
6	-0.099796	-2.556388	-2.085229
1	0.248455	-0.107844	-3.461526
1	0.836698	-2.072748	-2.386889
1	-0.682953	1.379087	-3.212094
1	0.140906	-3.368615	-1.395169
1	-1.374920	0.048183	-4.150162
1	-0.544266	-2.997192	-2.986511
1	1.181435	5.034312	-0.470318
8	1.371668	-2.582332	0.810588
1	1.909395	-3.373252	0.976032
1	-1.070871	3.442343	-0.195514

7	2.167874	3.133592	-0.305259
8	5.599471	-1.018248	0.127096
7	3.496016	-1.837961	0.470445
1	-0.343295	0.175004	1.688445
6	-1.268436	0.657670	2.897571
8	-1.367414	1.840326	2.776882
8	-1.470674	-0.360178	3.485212

(5-H⁺)-Hydride

E = -1133.72206962 a.u.

77	0.601918	0.019829	-0.340729
7	-0.421758	1.875807	-0.236408
7	-1.566901	-0.482458	-0.192821
6	-1.743892	1.842382	-0.111298
6	-2.435071	0.575035	-0.083406
6	-3.784470	0.423790	0.031833
1	-3.210013	3.347867	0.070458
1	-4.449327	1.279702	0.110350
6	-1.174208	3.966214	-0.095740
6	-4.331792	-0.907667	0.064332
6	-0.050448	3.195848	-0.227310
6	-2.162640	-1.682790	-0.200400
6	2.638225	-0.625707	-0.649042
6	1.935479	-1.674579	0.037038
6	2.623861	0.536740	0.192509
6	2.000141	0.169943	1.459334
6	1.577275	-1.171287	1.357635
6	3.313205	-0.749853	-1.970614
1	3.342295	0.207740	-2.500793
1	2.804760	-1.478024	-2.611286
1	4.348472	-1.090463	-1.836314
6	1.868778	-3.098425	-0.399995
6	3.315886	1.826116	-0.086294
1	1.673088	-3.183247	-1.474729
1	3.278790	2.079888	-1.151630
1	1.083863	-3.647059	0.128167

1	2.870946	2.652853	0.478059
1	2.822182	-3.604923	-0.196663
1	4.373068	1.765225	0.205120
6	1.844121	1.092439	2.617342
6	0.862255	-1.966762	2.392477
1	1.113343	0.714605	3.339531
1	0.441217	-1.327767	3.175252
1	1.515450	2.089068	2.296960
1	0.045212	-2.549453	1.952349
1	2.798724	1.223136	3.143855
1	1.547355	-2.678576	2.871201
1	-1.309250	5.036364	-0.051682
8	-1.321142	-2.713079	-0.376391
1	-1.855280	-3.523396	-0.363797
1	0.981349	3.505265	-0.326796
7	-2.231706	3.097557	-0.024616
8	-5.552377	-1.128576	0.201759
7	-3.440329	-1.952780	-0.069694
1	0.268843	0.112080	-1.930079

(5-H⁺)-Hydride-H₂O-TS

E = -1210.09995768 a.u.

77	0.645360	0.019094	-0.296487
7	-0.316420	1.877731	-0.636269
7	-1.478864	-0.441976	-0.298840
6	-1.617576	1.886305	-0.367834
6	-2.310448	0.633489	-0.145614
6	-3.631708	0.520897	0.159705
1	-3.040067	3.434375	-0.229993
1	-4.273605	1.391717	0.266189
6	-1.022684	3.984661	-0.667486
6	-4.164151	-0.806278	0.361139
6	0.073161	3.180774	-0.818254
6	-2.046796	-1.694509	-0.182275
6	2.718905	-0.612149	-0.002942
6	1.851048	-1.549586	0.665846

6	2.521400	0.683565	0.567168
6	1.519086	0.555126	1.617638
6	1.124601	-0.817379	1.679520
6	3.668663	-0.959104	-1.093406
1	3.879543	-0.097757	-1.734857
1	3.277201	-1.767018	-1.720994
1	4.619712	-1.302484	-0.666417
6	1.864145	-3.025952	0.474988
6	3.286129	1.914290	0.226786
1	2.007147	-3.296713	-0.576958
1	3.520568	1.959010	-0.842304
1	0.933821	-3.485242	0.820007
1	2.735183	2.821735	0.493548
1	2.689497	-3.472829	1.044637
1	4.235412	1.932964	0.777593
6	1.003212	1.658524	2.473122
6	0.113957	-1.392073	2.606571
1	-0.036465	1.473689	2.767479
1	-0.651030	-0.653840	2.871913
1	1.037492	2.619960	1.948946
1	-0.386088	-2.260920	2.166288
1	1.599872	1.754875	3.388749
1	0.598033	-1.719038	3.535375
1	-1.142696	5.054898	-0.741932
8	-1.310357	-2.708280	-0.429981
1	-0.473959	-2.476736	-1.736325
1	1.094760	3.448934	-1.052118
7	-2.077499	3.152663	-0.381885
8	-5.364644	-0.978924	0.692091
7	-3.331105	-1.871372	0.166778
1	0.625786	-0.053126	-2.041491
8	0.039920	-2.193373	-2.569918
1	0.850277	-2.724418	-2.575568
1	0.409078	-0.990626	-2.146322

(5-H⁺)

E = -1133.00324355 a.u.

77	-0.566876	0.032983	0.001848
7	0.380009	1.911869	0.006113
7	1.530527	-0.449064	-0.035797
6	1.711980	1.881086	-0.016705
6	2.397129	0.604898	-0.025976
6	3.753525	0.460121	-0.012210
1	3.179469	3.387686	-0.032181
1	4.444866	1.295737	-0.001926
6	1.134117	4.001120	0.003699
6	4.201956	-0.881383	0.005129
6	0.004609	3.232894	0.019575
6	2.051694	-1.739164	-0.015155
6	-2.348501	-0.246546	1.151436
6	-1.898571	-1.565537	0.727710
6	-2.667821	0.523697	-0.014480
6	-2.334956	-0.280969	-1.155741
6	-1.896254	-1.588140	-0.691050
6	-2.494134	0.182026	2.562359
1	-2.416299	1.269209	2.661517
1	-1.731077	-0.282848	3.195259
1	-3.476908	-0.123823	2.945221
6	-1.538728	-2.659002	1.665240
6	-3.226365	1.900646	-0.043558
1	-0.811686	-2.316481	2.410754
1	-2.943023	2.467409	0.849839
1	-1.116714	-3.521454	1.143929
1	-2.883988	2.451397	-0.926535
1	-2.434508	-2.987357	2.207577
1	-4.321979	1.861894	-0.079515
6	-2.468462	0.106397	-2.579571
6	-1.547047	-2.713359	-1.595325
1	-1.693264	-0.367446	-3.190748
1	-0.817357	-2.402088	-2.351677
1	-2.402098	1.191069	-2.708500
1	-1.134630	-3.564939	-1.048987
1	-3.443451	-0.221348	-2.964490
1	-2.447475	-3.046861	-2.126729

1	1.273816	5.071392	0.006493
8	1.288569	-2.713266	-0.009112
1	-1.031402	3.540236	0.038998
7	2.196695	3.131927	-0.017924
8	5.523880	-1.063872	0.032167
1	5.698709	-2.019851	0.051846
7	3.414750	-1.925712	0.002015

(5-H⁺)-HCO₂

E = -1322.24961170 a.u.

77	0.626210	0.045259	0.055811
7	-0.361488	1.928714	0.134781
7	-1.530715	-0.412512	-0.148403
6	-1.686863	1.902346	0.113171
6	-2.376907	0.641854	-0.054897
6	-3.744630	0.528863	-0.121466
1	-3.154453	3.408396	0.300776
1	-4.417424	1.376650	-0.053222
6	-1.108811	4.000465	0.423665
6	-4.208452	-0.782304	-0.310214
6	0.018796	3.228642	0.332218
6	-2.071031	-1.681508	-0.289484
6	1.917002	0.449758	-1.594516
6	1.476646	-0.913945	-1.675082
6	2.663190	0.603499	-0.355888
6	2.675888	-0.667946	0.300496
6	1.940981	-1.613831	-0.494731
6	1.692986	1.515289	-2.607276
1	1.610572	2.502088	-2.138885
1	0.776819	1.335085	-3.179281
1	2.531103	1.549671	-3.314684
6	0.700674	-1.511948	-2.794058
6	3.361951	1.829578	0.116519
1	0.090678	-0.757016	-3.301476
1	2.994414	2.725784	-0.393296
1	0.034980	-2.303368	-2.436879

1	3.233468	1.972290	1.195690
1	1.381557	-1.947261	-3.536403
1	4.438736	1.754900	-0.082375
6	3.327383	-0.959903	1.603196
6	1.882928	-3.080920	-0.246504
1	2.798429	-1.746656	2.149865
1	1.651228	-3.307676	0.798986
1	3.374958	-0.067628	2.236261
1	1.131007	-3.565746	-0.874250
1	4.356087	-1.303018	1.432037
1	2.856299	-3.533354	-0.477218
1	-1.241457	5.061349	0.573829
8	-1.327915	-2.677728	-0.327785
1	1.057470	3.523706	0.401349
7	-2.173630	3.148670	0.282587
8	-5.535375	-0.949259	-0.414784
1	-5.707320	-1.896108	-0.545606
7	-3.432937	-1.837589	-0.392502
6	-0.052963	-0.828505	2.919815
1	-0.323233	-0.575313	3.967667
8	0.111256	-2.010538	2.604955
8	0.045377	0.207571	2.169447

(5-H⁺)-HCO₂-TS

E = -1322.21702112 a.u.

77	-0.539951	-0.048447	0.006547
7	0.364209	1.868163	-0.096833
7	1.583255	-0.417896	0.172182
6	1.691763	1.877499	-0.160316
6	2.414887	0.635617	-0.007574
6	3.785491	0.522822	-0.007453
1	3.108983	3.424901	-0.380836
1	4.453312	1.364832	-0.152375
6	1.045964	3.968605	-0.354892
6	4.257247	-0.784802	0.182747
6	-0.055746	3.165564	-0.217327

6	2.131726	-1.649092	0.495051
6	-2.653403	-0.612044	-0.095792
6	-1.823879	-1.782282	-0.284311
6	-2.361167	0.314740	-1.140842
6	-1.379024	-0.285078	-2.022221
6	-1.064647	-1.576940	-1.494972
6	-3.692812	-0.446497	0.956226
1	-3.822212	0.603963	1.238216
1	-3.449640	-1.018646	1.856611
1	-4.659451	-0.810717	0.583273
6	-1.912239	-3.031285	0.523722
6	-3.012810	1.638332	-1.333786
1	-1.946238	-2.809141	1.596789
1	-3.262314	2.105470	-0.373982
1	-1.055749	-3.687821	0.345678
1	-2.372891	2.326989	-1.895379
1	-2.822557	-3.591573	0.272468
1	-3.946508	1.521178	-1.899010
6	-0.829075	0.325629	-3.263331
6	-0.120661	-2.544175	-2.115643
1	0.174904	-0.054921	-3.481760
1	0.735640	-2.035555	-2.572671
1	-0.762894	1.416367	-3.176833
1	0.259451	-3.265646	-1.388370
1	-1.465374	0.102348	-4.129793
1	-0.632842	-3.101609	-2.910287
1	1.143198	5.037143	-0.474436
8	1.392233	-2.585670	0.849726
1	-1.102679	3.437564	-0.193856
7	2.136942	3.140789	-0.317039
8	5.585043	-0.968538	0.121536
1	5.763917	-1.908678	0.286937
7	3.490360	-1.824725	0.422551
1	-0.325651	0.158069	1.684816
6	-1.247178	0.631129	2.905463
8	-1.357173	1.813077	2.788630
8	-1.433373	-0.390731	3.490883

(5-H⁺)-Hydride

E = -1133.72759337 a.u.

77	0.612136	0.014076	-0.340452
7	-0.375477	1.883410	-0.243396
7	-1.536599	-0.461002	-0.200256
6	-1.698301	1.856477	-0.123302
6	-2.385741	0.587699	-0.098409
6	-3.752249	0.453805	0.011696
1	-3.158867	3.371155	0.045758
1	-4.429918	1.297186	0.086465
6	-1.119964	3.976618	-0.112776
6	-4.200788	-0.872706	0.042941
6	0.001920	3.200398	-0.237451
6	-2.071215	-1.740034	-0.237787
6	1.933078	-1.694523	0.015836
6	1.526213	-1.221910	1.331711
6	2.661789	-0.628765	-0.618901
6	2.632366	0.506910	0.253644
6	1.958029	0.115587	1.484697
6	1.889182	-3.106837	-0.460533
1	1.699499	-3.165470	-1.538167
1	1.109557	-3.680824	0.048001
1	2.849615	-3.603686	-0.266577
6	0.755035	-2.034156	2.313159
6	3.373017	-0.724005	-1.923733
1	-0.065581	-2.570187	1.821778
1	2.870847	-1.422814	-2.601039
1	0.328115	-1.410591	3.105329
1	3.434705	0.248621	-2.422899
1	1.401714	-2.783998	2.787027
1	4.397770	-1.088035	-1.770947
6	3.335002	1.801461	0.032079
6	1.768684	1.011678	2.658883
1	2.869349	2.614762	0.599329
1	1.443659	2.014035	2.352346
1	3.339198	2.081185	-1.027391

1	1.020489	0.615794	3.353163
1	4.380080	1.732033	0.362534
1	2.708390	1.134325	3.213468
1	-1.249319	5.047683	-0.072765
8	-1.325713	-2.719268	-0.431856
1	1.035479	3.505056	-0.334013
7	-2.181431	3.114685	-0.042869
8	-5.520856	-1.069450	0.195649
1	-5.680608	-2.027072	0.188923
7	-3.420105	-1.924504	-0.067651
1	0.292841	0.102433	-1.937193

(5-2H⁺)-Hydride

E = -1133.24810891 a.u.

77	0.588922	0.014470	-0.348185
7	-0.427670	1.869776	-0.263980
7	-1.529894	-0.502966	-0.206069
6	-1.747490	1.814897	-0.129936
6	-2.411315	0.530284	-0.095603
6	-3.761144	0.368705	0.024270
1	-3.234893	3.298606	0.053318
1	-4.435780	1.217461	0.106180
6	-1.208702	3.949814	-0.128584
6	-4.270528	-0.978580	0.065517
6	-0.074997	3.196486	-0.262593
6	-2.065981	-1.784545	-0.257583
6	2.662818	-0.572953	-0.591322
6	1.945913	-1.659416	0.024968
6	2.592001	0.555866	0.285421
6	1.898465	0.145514	1.499436
6	1.500626	-1.201447	1.333034
6	3.400825	-0.645942	-1.883036
1	3.465835	0.334437	-2.366791
1	2.915711	-1.336988	-2.580688
1	4.425061	-1.006179	-1.717453
6	1.956654	-3.073319	-0.449886

6	3.268425	1.868093	0.083338
1	1.820978	-3.137710	-1.535398
1	3.282503	2.155900	-0.973978
1	1.165510	-3.663344	0.021406
1	2.774950	2.667151	0.647617
1	2.916697	-3.549672	-0.207466
1	4.309793	1.821704	0.429446
6	1.660965	1.032097	2.672715
6	0.731430	-2.036593	2.296931
1	0.904252	0.616224	3.346045
1	0.264698	-1.425866	3.076765
1	1.320498	2.028397	2.362812
1	-0.059460	-2.597478	1.784969
1	2.582685	1.173316	3.252667
1	1.388335	-2.765772	2.788802
1	-1.360356	5.017991	-0.089373
8	-1.282181	-2.749030	-0.477802
1	0.951592	3.520954	-0.369857
7	-2.253833	3.063580	-0.047285
8	-5.502893	-1.200529	0.228113
7	-3.389458	-2.005812	-0.082031
1	0.273180	0.087350	-1.948221

(5-2H⁺)-HCO₂

E = -1321.77149055 a.u.

77	0.588115	0.051635	0.063189
7	-0.431937	1.900962	0.238898
7	-1.524267	-0.457439	-0.117492
6	-1.755505	1.849370	0.174828
6	-2.413890	0.574356	-0.023010
6	-3.764691	0.421837	-0.108735
1	-3.252928	3.328040	0.320640
1	-4.444169	1.267268	-0.034550
6	-1.221396	3.960234	0.507888
6	-4.273862	-0.910321	-0.332966
6	-0.079855	3.210300	0.448516

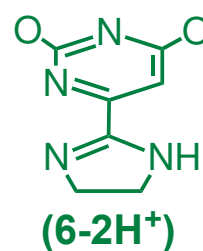
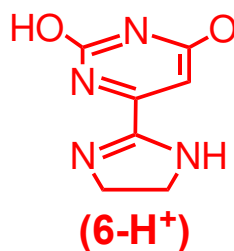
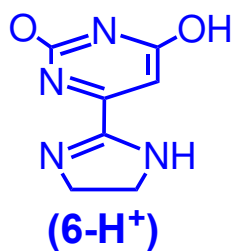
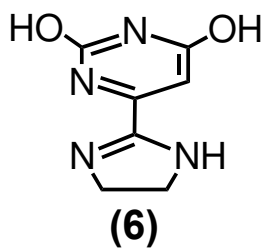
6	-2.053703	-1.737044	-0.254763
6	1.803731	0.558928	-1.614063
6	1.408686	-0.818260	-1.729392
6	2.600978	0.694126	-0.406996
6	2.680699	-0.595255	0.200424
6	1.948437	-1.541051	-0.599624
6	1.500516	1.651133	-2.576987
1	1.401102	2.616482	-2.068909
1	0.568375	1.458294	-3.118439
1	2.306660	1.744058	-3.315821
6	0.617225	-1.406304	-2.843423
6	3.285941	1.924005	0.078592
1	-0.017305	-0.652581	-3.321479
1	2.892384	2.823816	-0.404839
1	-0.026904	-2.217146	-2.489841
1	3.176407	2.043245	1.162858
1	1.287848	-1.815472	-3.609691
1	4.360331	1.874859	-0.140919
6	3.397825	-0.907446	1.464202
6	1.960521	-3.018040	-0.406301
1	2.924716	-1.735889	1.999985
1	1.809313	-3.290108	0.643485
1	3.430094	-0.038520	2.130259
1	1.181968	-3.508459	-0.996713
1	4.433627	-1.195843	1.242083
1	2.929861	-3.427514	-0.719872
1	-1.378675	5.017888	0.657016
8	-1.259627	-2.713920	-0.260813
1	0.950929	3.523487	0.549544
7	-2.267886	3.087039	0.331644
8	-5.510501	-1.113208	-0.476912
7	-3.386528	-1.938290	-0.382082
6	0.093432	-0.967419	2.943443
1	-0.155348	-0.772180	4.009913
8	0.336054	-2.123490	2.583591
8	0.084151	0.097568	2.230775

(5-2H⁺)-HCO₂-TS

E = -1321.74150867 a.u.

77	-0.632689	0.048055	0.100967
7	0.335207	1.900774	0.464053
7	1.445928	-0.326909	-0.312934
6	1.636672	1.933973	0.197077
6	2.313898	0.717672	-0.197397
6	3.660541	0.604645	-0.381663
1	3.074091	3.474013	0.253444
1	4.328174	1.456655	-0.279233
6	1.066439	3.977988	0.782094
6	4.178911	-0.697392	-0.727814
6	-0.036456	3.171438	0.828798
6	2.000761	-1.590343	-0.478935
6	-2.727527	-0.528139	0.326224
6	-2.040662	-1.515230	-0.479002
6	-2.657671	0.725026	-0.351510
6	-1.951697	0.527051	-1.605923
6	-1.589606	-0.853657	-1.679944
6	-3.416020	-0.804360	1.616438
1	-3.492412	0.096988	2.233366
1	-2.887429	-1.569887	2.195047
1	-4.433974	-1.173486	1.433582
6	-2.030548	-2.980885	-0.208418
6	-3.290440	1.996790	0.093865
1	-1.796286	-3.196034	0.840188
1	-3.285198	2.090504	1.185569
1	-1.297136	-3.501611	-0.830721
1	-2.784898	2.871346	-0.328917
1	-3.017694	-3.413280	-0.420549
1	-4.336519	2.033463	-0.237246
6	-1.708694	1.575567	-2.634343
6	-0.860120	-1.504803	-2.801392
1	-0.850491	1.324576	-3.266885
1	-0.251802	-0.782054	-3.355839
1	-1.510236	2.549983	-2.172724
1	-0.198992	-2.297462	-2.437627

1	-2.581887	1.695379	-3.289227
1	-1.571358	-1.953114	-3.506862
1	1.196283	5.028298	0.995648
8	1.239111	-2.590097	-0.375087
1	-1.051934	3.418224	1.107602
7	2.111618	3.182204	0.382893
8	5.402667	-0.861982	-0.986506
7	3.316619	-1.749841	-0.742691
1	-0.157478	-0.305474	1.696728
6	0.661643	-1.395237	2.555574
8	1.734698	-0.892327	2.676444
8	-0.138713	-2.249529	2.778723



(6)

E = -1134.65347887 a.u.

77	-0.595742	0.010416	-0.028971
7	0.354730	1.831031	-0.132078
7	1.560085	-0.476467	-0.092380
6	1.662842	1.871484	-0.014058
6	2.393453	0.600070	-0.006543
6	3.748732	0.486111	0.062177
1	3.168815	3.292082	0.025743
1	4.420264	1.334805	0.128212
6	4.244486	-0.836635	0.047134
6	2.148807	-1.675084	-0.129745
6	-2.243174	-0.397626	1.270181
6	-1.839968	-1.662882	0.690332
6	-2.679581	0.475526	0.210827

6	-2.476253	-0.225627	-1.023730
6	-1.972272	-1.556170	-0.719662
6	-2.246934	-0.086759	2.717588
1	-2.115855	0.985218	2.896706
1	-1.454510	-0.630782	3.241514
1	-3.207204	-0.387927	3.157509
6	-1.386710	-2.835129	1.482969
6	-3.256409	1.834165	0.379487
1	-0.593141	-2.557410	2.186001
1	-2.818569	2.352365	1.238992
1	-1.015792	-3.640147	0.844225
1	-3.098849	2.451478	-0.510504
1	-2.225605	-3.225428	2.072645
1	-4.337528	1.760345	0.550265
6	-2.747635	0.290544	-2.385627
6	-1.666870	-2.581896	-1.748629
1	-2.037280	-0.117974	-3.111771
1	-1.023716	-2.170070	-2.535024
1	-2.690340	1.383416	-2.416523
1	-1.170298	-3.455694	-1.320157
1	-3.756382	-0.004671	-2.703314
1	-2.596146	-2.914533	-2.228067
8	1.339051	-2.703773	-0.256849
1	1.858185	-3.526251	-0.275426
7	2.178443	3.088264	0.093354
8	5.547398	-1.002014	0.137423
1	5.761553	-1.951381	0.116115
7	3.457887	-1.896010	-0.053998
6	1.119610	4.071613	-0.178373
1	1.247118	4.475090	-1.188912
1	1.157072	4.894709	0.537023
6	-0.142119	3.213397	-0.042142
1	-0.877687	3.408672	-0.826842
1	-0.631578	3.349999	0.930380

(6)-HCO₂

E = -1323.90900112 a.u.

77	-0.675191	0.034181	-0.123080
7	0.339411	1.838203	-0.609321
7	1.491687	-0.472825	-0.347300
6	1.620174	1.869442	-0.387362
6	2.311519	0.596089	-0.165490
6	3.625853	0.464040	0.184902
1	3.159186	3.203087	-0.602216
1	4.293130	1.304923	0.338347
6	4.064143	-0.863034	0.367484
6	2.043265	-1.686417	-0.241019
6	-1.378107	0.652676	1.791761
6	-0.902914	-0.696698	1.875845
6	-2.475256	0.676856	0.836463
6	-2.683520	-0.672377	0.385513
6	-1.717025	-1.527602	1.009444
6	-0.840201	1.797238	2.574264
1	-1.159241	2.758645	2.160882
1	0.256116	1.780524	2.600604
1	-1.198094	1.744423	3.610185
6	0.225507	-1.153815	2.729879
6	-3.303197	1.854309	0.460093
1	1.032107	-0.411142	2.752628
1	-2.784573	2.794775	0.671419
1	0.641183	-2.104045	2.379206
1	-3.554606	1.837298	-0.606180
1	-0.120368	-1.300302	3.760927
1	-4.243570	1.857398	1.025800
6	-3.701910	-1.088767	-0.611991
6	-1.646656	-3.009134	0.897235
1	-3.391367	-1.983726	-1.159837
1	-2.008127	-3.358854	-0.074678
1	-3.901638	-0.290679	-1.334625
1	-0.623831	-3.375414	1.029174
1	-4.646867	-1.318490	-0.103170
1	-2.270025	-3.474099	1.671670
8	1.316534	-2.741790	-0.526689
1	0.503854	-2.483918	-1.056735

7	2.177922	3.097028	-0.367188
8	5.311181	-1.044754	0.772446
1	5.487027	-1.997748	0.854038
7	3.300007	-1.919380	0.141106
6	-0.783990	-1.239888	-2.885531
1	-0.994030	-1.222770	-3.972710
8	-0.561200	-2.336271	-2.340074
8	-0.798939	-0.091487	-2.332856
6	1.176991	4.016460	-0.935894
1	1.384297	4.181480	-2.000197
1	1.182635	4.978337	-0.420861
6	-0.118290	3.228751	-0.722938
1	-0.837917	3.336748	-1.539592
1	-0.617658	3.516088	0.213402

(6)-HCO₂-TS

E = -1323.86836917 a.u.

77	-0.618851	0.033688	0.025664
7	0.356887	1.876787	0.341282
7	1.538789	-0.349764	-0.224374
6	1.645721	1.955597	0.197261
6	2.370711	0.717450	-0.067831
6	3.734807	0.612628	-0.137880
1	3.124718	3.326526	0.575816
1	4.406467	1.455046	-0.015978
6	4.230368	-0.678110	-0.380983
6	2.131424	-1.531485	-0.416290
6	-2.712749	-0.542001	0.216762
6	-2.004462	-1.564283	-0.518208
6	-2.618862	0.683482	-0.516750
6	-1.884956	0.425140	-1.740001
6	-1.517876	-0.958088	-1.734685
6	-3.461845	-0.741303	1.484855
1	-3.443304	0.157522	2.110229
1	-3.057007	-1.574581	2.066624
1	-4.512386	-0.968207	1.259758

6	-1.956335	-3.007216	-0.148315
6	-3.270480	1.965583	-0.138451
1	-1.799006	-3.138510	0.928290
1	-3.277009	2.109178	0.947386
1	-1.151028	-3.532380	-0.669318
1	-2.774353	2.825834	-0.598609
1	-2.901068	-3.502261	-0.408588
1	-4.313834	1.964015	-0.479693
6	-1.623729	1.406572	-2.827296
6	-0.763368	-1.638165	-2.821398
1	-0.719408	1.149100	-3.388859
1	0.037590	-1.000353	-3.212467
1	-1.498356	2.420217	-2.429940
1	-0.318828	-2.578703	-2.485711
1	-2.460495	1.431477	-3.537852
1	-1.441124	-1.860582	-3.655496
8	1.324588	-2.570670	-0.520749
1	1.853312	-3.370631	-0.680859
7	2.165163	3.203485	0.271862
7	3.440963	-1.735725	-0.508989
1	-0.125998	-0.294776	1.635159
6	-0.082621	-1.320572	2.807461
8	-0.884332	-0.890255	3.580230
8	0.752389	-2.116213	2.499183
8	5.539888	-0.836629	-0.480146
1	5.739592	-1.773305	-0.649916
6	1.107671	4.046417	0.855869
1	1.109229	5.047970	0.422995
1	1.249335	4.128210	1.940798
6	-0.146684	3.246538	0.507398
1	-0.589619	3.573241	-0.444951
1	-0.919214	3.289572	1.279305

(6)-Hydride

E = -1135.38062636 a.u.

77	0.612407	0.001893	-0.300525
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7	-0.431047	1.819328	-0.130203
7	-1.558599	-0.502414	-0.138728
6	-1.728989	1.835605	-0.113606
6	-2.417226	0.552478	-0.092946
6	-3.779962	0.404499	-0.029491
1	-3.209167	3.198120	0.260298
1	-4.466357	1.243458	-0.005326
6	-4.242101	-0.917117	0.025464
6	-2.126056	-1.711182	-0.121258
6	2.620268	-0.659406	-0.755652
6	1.971622	-1.707841	-0.026897
6	2.666214	0.503486	0.088311
6	2.145001	0.127659	1.397110
6	1.703560	-1.210271	1.315835
6	3.191590	-0.773851	-2.126024
1	3.186392	0.189138	-2.646902
1	2.632829	-1.492954	-2.733982
1	4.232228	-1.119812	-2.071918
6	1.856378	-3.126117	-0.471570
6	3.335283	1.793251	-0.240468
1	1.556714	-3.198473	-1.522866
1	3.201572	2.055842	-1.295751
1	1.127395	-3.678549	0.127741
1	2.942812	2.616128	0.366411
1	2.823907	-3.635745	-0.368247
1	4.414173	1.727589	-0.046278
6	2.087910	1.036318	2.574556
6	1.058657	-2.012378	2.391447
1	1.386999	0.672931	3.332748
1	0.728667	-1.382435	3.223428
1	1.779091	2.048587	2.286065
1	0.187320	-2.560011	2.012587
1	3.076182	1.124558	3.044787
1	1.760975	-2.755471	2.791286
8	-1.294835	-2.734414	-0.227530
1	-1.805420	-3.561174	-0.209493
7	-2.300851	3.065210	-0.171524
8	-5.546857	-1.122115	0.128892

1	-5.723513	-2.077615	0.157519
7	-3.425705	-1.962668	-0.017961
1	0.231916	0.130402	-1.874573
6	0.024508	3.211210	-0.220618
1	0.862487	3.394078	0.458120
1	0.369541	3.398233	-1.247470
6	-1.223259	4.022463	0.132717
1	-1.321358	4.934407	-0.458637
1	-1.244150	4.286249	1.197454

(6)-Hydride-H₂O-TS

E = -1211.77105731 a.u.

77	0.664987	0.016338	-0.280830
7	-0.346391	1.856004	-0.555917
7	-1.466606	-0.463639	-0.261234
6	-1.621489	1.877458	-0.305569
6	-2.302594	0.595636	-0.127467
6	-3.639312	0.472474	0.139722
1	-3.151381	3.232675	-0.441393
1	-4.311803	1.315814	0.247291
6	-4.081038	-0.853972	0.303791
6	-2.002028	-1.730658	-0.191353
6	2.745839	-0.627502	-0.069708
6	1.905166	-1.585506	0.599863
6	2.559596	0.654400	0.543403
6	1.603029	0.487885	1.629991
6	1.208707	-0.883632	1.653653
6	3.653143	-0.925276	-1.209679
1	3.792555	-0.049235	-1.850713
1	3.265105	-1.742634	-1.826585
1	4.638949	-1.229402	-0.835542
6	1.889017	-3.057282	0.376333
6	3.310691	1.894581	0.207453
1	2.174033	-3.313793	-0.649140
1	3.499167	1.968584	-0.868926
1	0.899747	-3.483657	0.569387

1	2.771342	2.793944	0.520502
1	2.604017	-3.547468	1.049922
1	4.282359	1.897771	0.717732
6	1.130393	1.538600	2.573327
6	0.243611	-1.491404	2.606118
1	0.039969	1.506756	2.690893
1	-0.457842	-0.745985	2.996030
1	1.405748	2.540394	2.229656
1	-0.329007	-2.299797	2.139292
1	1.575413	1.392612	3.565545
1	0.785039	-1.918565	3.460010
8	-1.284466	-2.731155	-0.440359
1	-0.353381	-2.515849	-1.792820
7	-2.170644	3.110147	-0.212616
8	-5.363169	-1.027888	0.634942
1	-5.533689	-1.981136	0.712632
7	-3.313135	-1.908327	0.148568
1	0.579381	-0.027493	-2.031161
8	0.161255	-2.192929	-2.591236
1	0.998894	-2.681786	-2.585436
1	0.442040	-0.987523	-2.153176
6	0.117420	3.248798	-0.596537
1	0.614343	3.484224	0.355817
1	0.836526	3.403087	-1.405659
6	-1.176304	4.044474	-0.766875
1	-1.395189	4.241712	-1.823854
1	-1.170934	4.990750	-0.223606

(6-H⁺)

E = -1134.20186023 a.u.

77	-0.559599	0.013170	-0.025524
7	0.419208	1.817873	-0.108297
7	1.543529	-0.512724	-0.070645
6	1.730247	1.828761	-0.001670
6	2.424176	0.544167	0.002842
6	3.765376	0.388763	0.057353

1	3.256984	3.216736	-0.027804
1	4.445358	1.233655	0.114168
6	4.307577	-0.960025	0.049939
6	2.127317	-1.725194	-0.111808
6	-2.228948	-0.361735	1.258030
6	-1.863679	-1.635251	0.668382
6	-2.640510	0.532322	0.208154
6	-2.444399	-0.157237	-1.031292
6	-1.980336	-1.505930	-0.740751
6	-2.235195	-0.067064	2.709891
1	-2.086485	1.000540	2.901529
1	-1.453401	-0.628981	3.231119
1	-3.201593	-0.357239	3.143667
6	-1.462292	-2.831436	1.453789
6	-3.183245	1.903677	0.391497
1	-0.674540	-2.587023	2.175483
1	-2.723440	2.406297	1.249016
1	-1.102011	-3.639419	0.812503
1	-3.020819	2.523263	-0.496112
1	-2.324013	-3.203976	2.021775
1	-4.263966	1.855949	0.574006
6	-2.697575	0.381111	-2.389281
6	-1.704387	-2.528689	-1.781715
1	-2.017374	-0.062583	-3.123531
1	-1.040860	-2.130600	-2.558268
1	-2.574946	1.469002	-2.415347
1	-1.240773	-3.424644	-1.361684
1	-3.724348	0.149010	-2.701156
1	-2.641093	-2.823377	-2.271644
8	1.267408	-2.735465	-0.235540
1	1.780575	-3.559755	-0.253171
7	2.267874	3.041458	0.104223
8	5.526454	-1.174103	0.129033
7	3.407475	-1.997656	-0.051924
6	1.227167	4.036987	-0.189415
1	1.349480	4.406667	-1.213991
1	1.284884	4.882265	0.498389
6	-0.049065	3.208774	-0.016848

1	-0.799089	3.416851	-0.784516
1	-0.510282	3.365138	0.966901

(6-H⁺)-HCO₂

E = -1323.44869952 a.u.

77	-0.644937	0.048594	-0.136745
7	0.425873	1.830007	-0.654344
7	1.450770	-0.534055	-0.347298
6	1.705037	1.797514	-0.408891
6	2.335025	0.498715	-0.175022
6	3.631301	0.308001	0.175465
1	3.301595	3.064543	-0.583813
1	4.321745	1.136241	0.307378
6	4.086084	-1.048692	0.407277
6	1.969487	-1.767605	-0.211615
6	-1.178589	0.605804	1.847881
6	-0.904850	-0.803026	1.814736
6	-2.314551	0.853337	0.972356
6	-2.711872	-0.399784	0.412245
6	-1.846478	-1.435584	0.921357
6	-0.446375	1.610385	2.665917
1	-0.549973	2.619590	2.253749
1	0.622976	1.373366	2.716292
1	-0.832719	1.627921	3.692878
6	0.168408	-1.488360	2.583083
6	-2.995561	2.157501	0.749349
1	1.070729	-0.869324	2.654461
1	-2.362880	3.000515	1.043237
1	0.442208	-2.445025	2.127200
1	-3.275787	2.290063	-0.301525
1	-0.177620	-1.689386	3.604684
1	-3.914505	2.208898	1.347144
6	-3.809165	-0.601138	-0.569276
6	-2.018876	-2.898587	0.710905
1	-3.572451	-1.410156	-1.268748
1	-2.494143	-3.112200	-0.251451

1	-3.998606	0.309411	-1.147156
1	-1.063233	-3.430947	0.745124
1	-4.738117	-0.870240	-0.050472
1	-2.660706	-3.317874	1.496802
8	1.163461	-2.796804	-0.472660
1	0.373477	-2.496432	-0.996720
7	2.314162	3.002215	-0.360869
8	5.242243	-1.295088	0.801850
7	3.195800	-2.066821	0.154877
6	-0.844684	-1.242171	-2.885397
1	-1.032346	-1.205630	-3.977133
8	-0.720253	-2.351425	-2.338964
8	-0.779664	-0.095011	-2.333043
6	1.366609	3.969071	-0.937175
1	1.585590	4.122758	-2.001443
1	1.415205	4.930924	-0.423873
6	0.036476	3.243877	-0.730682
1	-0.683586	3.415144	-1.535674
1	-0.433770	3.537748	0.219327

(6-H⁺)-HCO₂-TS

E = -1323.40924424 a.u.

77	-0.537225	-0.052038	-0.002935
7	0.465601	1.804883	-0.039649
7	1.594947	-0.512256	0.138621
6	1.762177	1.815355	-0.124755
6	2.462496	0.541194	-0.006858
6	3.813053	0.397076	-0.000626
1	3.257092	3.202862	0.002999
1	4.481776	1.245531	-0.112878
6	4.367802	-0.928683	0.152998
6	2.187625	-1.693281	0.358827
6	-2.645165	-0.676734	-0.043241
6	-1.816976	-1.778972	-0.451540
6	-2.429253	0.414776	-0.943594
6	-1.512745	-0.036063	-1.980391

6	-1.135549	-1.376269	-1.665017
6	-3.611477	-0.696349	1.086991
1	-3.767915	0.302288	1.507866
1	-3.285485	-1.366693	1.888109
1	-4.584058	-1.058809	0.727923
6	-1.841244	-3.137283	0.160894
6	-3.133362	1.725713	-0.892558
1	-1.812058	-3.081096	1.255674
1	-3.277546	2.056780	0.142435
1	-0.992744	-3.744129	-0.167701
1	-2.577494	2.505384	-1.423448
1	-2.760253	-3.668607	-0.120174
1	-4.124159	1.647367	-1.358692
6	-1.070192	0.757510	-3.159169
6	-0.182940	-2.215534	-2.440628
1	-0.059032	0.477660	-3.474931
1	0.546493	-1.599055	-2.977048
1	-1.068432	1.831294	-2.940074
1	0.366411	-2.903428	-1.789721
1	-1.741668	0.597216	-4.012676
1	-0.721827	-2.816720	-3.183932
8	1.337062	-2.696701	0.615434
1	1.866858	-3.498749	0.751950
7	2.316385	3.035027	-0.337168
8	5.593661	-1.145015	0.114686
7	3.473733	-1.956977	0.361296
1	-0.305747	0.113076	1.677948
6	-1.202161	0.421265	2.957902
8	-1.438501	1.589197	2.905025
8	-1.253330	-0.642196	3.494828
6	1.259567	4.016250	-0.035377
1	1.313092	4.879159	-0.701463
1	1.353212	4.363537	1.001448
6	-0.004685	3.180260	-0.236777
1	-0.401839	3.278783	-1.258252
1	-0.805982	3.424879	0.466618

(6-H⁺)-Hydride

E = -1134.92032802 a.u.

77	0.605933	-0.000042	-0.336968
7	-0.465010	1.812589	-0.226047
7	-1.537822	-0.540323	-0.203999
6	-1.750680	1.792376	-0.052587
6	-2.417840	0.498170	-0.035275
6	-3.758941	0.326119	0.108234
1	-3.289976	3.122927	-0.193061
1	-4.435116	1.165940	0.240095
6	-4.290500	-1.015935	0.090383
6	-2.116243	-1.747511	-0.255254
6	2.666998	-0.579193	-0.637193
6	1.985749	-1.667761	0.006578
6	2.607558	0.557378	0.238425
6	1.977841	0.135968	1.483714
6	1.589541	-1.212588	1.331937
6	3.360735	-0.644513	-1.953381
1	3.368081	0.328450	-2.455493
1	2.880558	-1.367476	-2.621214
1	4.403937	-0.958786	-1.816763
6	1.959247	-3.079154	-0.474043
6	3.255512	1.878982	0.008671
1	1.801864	-3.134511	-1.556999
1	3.292499	2.127396	-1.057671
1	1.164809	-3.654851	0.010009
1	2.722292	2.683937	0.526781
1	2.912370	-3.578206	-0.251869
1	4.286746	1.872014	0.386558
6	1.798822	1.006080	2.678788
6	0.880492	-2.055961	2.332646
1	1.028559	0.616620	3.352453
1	0.416301	-1.449439	3.116905
1	1.513576	2.026590	2.395964
1	0.097055	-2.659720	1.861300
1	2.733608	1.081104	3.250327
1	1.580181	-2.751005	2.814884

8	-1.265026	-2.757239	-0.488992
1	-1.784852	-3.576914	-0.498648
7	-2.333203	3.008890	0.123444
8	-5.504802	-1.253752	0.242620
7	-3.389565	-2.038756	-0.115300
1	0.284243	0.097062	-1.930268
6	-1.331327	3.997379	-0.310859
1	-1.365732	4.898493	0.304361
1	-1.506158	4.278025	-1.357304
6	-0.029300	3.211310	-0.152001
1	0.441860	3.386449	0.826598
1	0.708284	3.433410	-0.928844

(6-H⁺)-Hydride-H₂O-TS

E = -1211.29975818 a.u.

77	0.644999	0.016529	-0.286924
7	-0.395009	1.840276	-0.579442
7	-1.461240	-0.506012	-0.280541
6	-1.666947	1.832962	-0.302958
6	-2.321263	0.542182	-0.110615
6	-3.637939	0.392176	0.188990
1	-3.220556	3.157670	-0.415319
1	-4.303754	1.241587	0.313762
6	-4.137871	-0.953638	0.369164
6	-1.996282	-1.774585	-0.189875
6	2.742755	-0.563235	-0.058348
6	1.923910	-1.547988	0.602797
6	2.518259	0.709464	0.556312
6	1.551230	0.514888	1.629015
6	1.197810	-0.869310	1.650034
6	3.674990	-0.841949	-1.183207
1	3.819634	0.040529	-1.814291
1	3.306920	-1.658305	-1.813982
1	4.656114	-1.141365	-0.792749
6	1.972671	-3.018200	0.375370
6	3.238628	1.970917	0.230018

1	2.122818	-3.261289	-0.682045
1	3.429396	2.056333	-0.845278
1	1.052837	-3.506136	0.709851
1	2.674708	2.854746	0.544660
1	2.807437	-3.459346	0.935862
1	4.207862	1.998499	0.744365
6	1.035601	1.551521	2.566148
6	0.238636	-1.503750	2.592106
1	-0.053640	1.480062	2.677522
1	-0.481209	-0.774774	2.980181
1	1.274323	2.561772	2.219568
1	-0.316205	-2.318263	2.114249
1	1.479268	1.428128	3.562109
1	0.779716	-1.926139	3.448465
8	-1.236869	-2.766540	-0.452988
1	-0.418011	-2.496748	-1.763567
7	-2.234824	3.058636	-0.197912
8	-5.332276	-1.159483	0.699618
7	-3.277177	-1.989308	0.154341
1	0.589553	-0.036334	-2.037866
8	0.077343	-2.183727	-2.597128
1	0.901693	-2.692000	-2.625262
1	0.410328	-0.979578	-2.156921
6	-1.271029	4.009400	-0.774937
1	-1.274881	4.959681	-0.238221
1	-1.509724	4.196623	-1.829811
6	0.040660	3.240943	-0.622679
1	0.543873	3.489523	0.323198
1	0.745827	3.414481	-1.440491

(6-H⁺)

E = -1134.20595169 a.u.

77	-0.570435	0.004419	-0.028150
7	0.382087	1.825502	-0.125714
7	1.528282	-0.496791	-0.079056
6	1.690019	1.842987	-0.009432

6	2.387439	0.554641	-0.003468
6	3.740412	0.421445	0.053611
1	3.213876	3.237568	-0.014195
1	4.428259	1.257543	0.111738
6	4.196532	-0.921689	0.044362
6	2.052823	-1.780976	-0.148049
6	-2.227890	-0.380538	1.267576
6	-1.843122	-1.652910	0.686858
6	-2.661590	0.496698	0.213259
6	-2.460836	-0.199777	-1.020758
6	-1.974271	-1.539313	-0.722100
6	-2.226145	-0.073244	2.716909
1	-2.093385	0.998273	2.898354
1	-1.430131	-0.617522	3.235472
1	-3.183302	-0.375550	3.162760
6	-1.417699	-2.834165	1.482418
6	-3.217879	1.864096	0.387255
1	-0.612737	-2.574446	2.179646
1	-2.760649	2.378479	1.239232
1	-1.069359	-3.651155	0.846144
1	-3.062983	2.478093	-0.505674
1	-2.263062	-3.200372	2.078537
1	-4.297716	1.808761	0.572680
6	-2.726632	0.320456	-2.383471
6	-1.701704	-2.569310	-1.756692
1	-2.013706	-0.088085	-3.107331
1	-1.043725	-2.176059	-2.540428
1	-2.664031	1.413254	-2.412403
1	-1.234784	-3.461689	-1.333414
1	-3.734583	0.030474	-2.708357
1	-2.641518	-2.867560	-2.238687
8	1.301192	-2.751861	-0.300560
7	2.223882	3.056085	0.102029
8	5.515431	-1.096049	0.138505
1	5.700943	-2.050050	0.118782
7	3.415366	-1.964286	-0.047346
6	1.179805	4.048599	-0.192331
1	1.307611	4.424084	-1.214076

1	1.230052	4.890260	0.500418
6	-0.094048	3.213237	-0.032364
1	-0.839665	3.420474	-0.804595
1	-0.563453	3.363229	0.948544

(6-H⁺)-HCO₂

E = -1323.44867446 a.u.

77	0.628032	0.028937	0.025070
7	-0.377814	1.871589	-0.141745
7	-1.517159	-0.467367	-0.205237
6	-1.668751	1.862461	0.019733
6	-2.364297	0.580923	-0.101603
6	-3.730077	0.467326	-0.134748
1	-3.182457	3.248790	0.103472
1	-4.407105	1.309857	-0.052361
6	-4.195169	-0.847748	-0.319985
6	-2.052956	-1.739135	-0.331324
6	2.093487	0.344658	-1.498078
6	1.607856	-1.004243	-1.588186
6	2.706382	0.511015	-0.194488
6	2.624413	-0.754911	0.482108
6	1.957948	-1.696540	-0.361562
6	2.008426	1.383554	-2.558349
1	1.967442	2.391536	-2.132444
1	1.122166	1.242993	-3.185823
1	2.891504	1.334411	-3.207854
6	0.927559	-1.614254	-2.762541
6	3.383270	1.729200	0.328388
1	0.464571	-0.849025	-3.394178
1	3.015957	2.636673	-0.162070
1	0.147226	-2.314727	-2.447301
1	3.219507	1.838435	1.406832
1	1.649869	-2.166031	-3.377696
1	4.466566	1.674276	0.160090
6	3.142682	-1.020407	1.849433
6	1.828028	-3.155631	-0.094986

1	2.669561	-1.897924	2.299260
1	1.481540	-3.355191	0.923933
1	2.980917	-0.161782	2.510597
1	1.132073	-3.631451	-0.789838
1	4.224248	-1.202866	1.805508
1	2.806084	-3.640211	-0.212633
8	-1.307741	-2.732372	-0.372411
7	-2.215287	3.055291	0.339466
8	-5.521773	-1.015306	-0.408575
1	-5.695824	-1.961133	-0.545610
7	-3.416905	-1.897602	-0.417435
6	-0.272261	-0.579906	2.878211
1	-0.599969	-0.249849	3.887899
8	-0.176515	-1.786614	2.637364
8	-0.038733	0.402315	2.088021
6	0.092040	3.221731	0.186991
1	0.492861	3.208745	1.212602
1	0.886972	3.550326	-0.488675
6	-1.176309	4.066270	0.082412
1	-1.301698	4.484871	-0.924409
1	-1.215821	4.877375	0.811251

(6-H⁺)-HCO₂-TS

E = -1323.41460155 a.u.

77	-0.541332	-0.068477	0.000950
7	0.426049	1.800575	-0.092586
7	1.577415	-0.495342	0.128417
6	1.722123	1.819531	-0.175699
6	2.424184	0.548686	-0.023582
6	3.790355	0.427805	-0.005084
1	3.207222	3.220836	-0.081683
1	4.468708	1.265101	-0.123781
6	4.251375	-0.889768	0.164445
6	2.110621	-1.740168	0.422646
6	-2.662452	-0.656918	-0.017665
6	-1.842699	-1.798960	-0.336880

6	-2.430522	0.357085	-0.996133
6	-1.505008	-0.173474	-1.983170
6	-1.145967	-1.493320	-1.566848
6	-3.640978	-0.580939	1.099846
1	-3.753340	0.442141	1.474275
1	-3.354054	-1.227895	1.934076
1	-4.626445	-0.913212	0.746632
6	-1.892678	-3.110302	0.369476
6	-3.120784	1.674977	-1.050949
1	-1.888149	-2.977167	1.457852
1	-3.288208	2.078263	-0.045515
1	-1.040486	-3.743375	0.106339
1	-2.547061	2.411518	-1.622613
1	-2.809119	-3.655034	0.106240
1	-4.100833	1.570480	-1.534510
6	-1.045268	0.524800	-3.214765
6	-0.208605	-2.400099	-2.282972
1	-0.035119	0.210445	-3.500049
1	0.543019	-1.833905	-2.844121
1	-1.031033	1.612006	-3.076971
1	0.314743	-3.063808	-1.588093
1	-1.711587	0.309418	-4.060362
1	-0.757452	-3.024844	-2.999002
8	1.361330	-2.682095	0.737540
7	2.269950	3.035989	-0.423166
8	5.577308	-1.079994	0.120879
1	5.750054	-2.024945	0.265019
7	3.470347	-1.924866	0.363723
1	-0.292626	0.108518	1.681638
6	-1.173869	0.564969	2.938123
8	-1.281933	1.748951	2.842586
8	-1.346144	-0.465744	3.511747
6	1.203450	4.015166	-0.147553
1	1.255651	4.866684	-0.828087
1	1.285292	4.379827	0.884444
6	-0.050687	3.164895	-0.345027
1	-0.422683	3.223052	-1.378954
1	-0.870327	3.431419	0.328036

(6-H⁺)-Hydride

E = -1134.92536388 a.u.

77	0.616490	-0.001763	-0.335448
7	-0.416283	1.824570	-0.229525
7	-1.507405	-0.510318	-0.221249
6	-1.703590	1.814274	-0.071200
6	-2.370035	0.518291	-0.059835
6	-3.727722	0.362661	0.084084
1	-3.231035	3.154809	-0.240873
1	-4.415257	1.190410	0.216596
6	-4.159126	-0.973262	0.069297
6	-2.023605	-1.794298	-0.309509
6	1.975904	-1.692690	-0.021767
6	1.528975	-1.276559	1.297963
6	2.688942	-0.586775	-0.602644
6	2.617269	0.515535	0.310818
6	1.932055	0.063236	1.513436
6	1.966254	-3.086092	-0.553791
1	1.791272	-3.106016	-1.635372
1	1.190754	-3.693154	-0.077882
1	2.932688	-3.573635	-0.366251
6	0.755677	-2.138193	2.234856
6	3.421748	-0.617277	-1.898937
1	-0.022250	-2.699043	1.703689
1	2.933962	-1.284639	-2.617192
1	0.272006	-1.546938	3.019269
1	3.487226	0.378185	-2.350248
1	1.414300	-2.868432	2.722620
1	4.445657	-0.984112	-1.747268
6	3.284921	1.837321	0.149080
6	1.718132	0.892461	2.732173
1	2.738236	2.631425	0.669883
1	1.492764	1.934789	2.475987
1	3.368880	2.119111	-0.906224
1	0.891462	0.511705	3.341048

1	4.299976	1.808403	0.567384
1	2.617620	0.904114	3.362301
8	-1.271773	-2.753279	-0.565420
7	-2.281692	3.034864	0.095648
8	-5.469332	-1.194816	0.255154
1	-5.617405	-2.153839	0.217511
7	-3.366913	-2.003396	-0.117255
1	0.314765	0.092081	-1.936858
6	0.029199	3.219740	-0.152806
1	0.491395	3.390550	0.830659
1	0.777109	3.434848	-0.921607
6	-1.265523	4.015690	-0.325537
1	-1.425838	4.299203	-1.373439
1	-1.299942	4.916235	0.290466

(6-2H⁺)

E = -1133.73655042 a.u.

77	-0.533000	0.003977	-0.011652
7	0.450732	1.808253	-0.085901
7	1.514096	-0.538620	-0.055875
6	1.761092	1.794737	0.006557
6	2.417993	0.493035	0.002326
6	3.756479	0.312827	0.040804
1	3.304863	3.153521	-0.077220
1	4.454614	1.144351	0.087252
6	4.255496	-1.056992	0.043938
6	2.030149	-1.839962	-0.113256
6	-2.242987	-0.324330	1.230681
6	-1.886041	-1.610789	0.659446
6	-2.622776	0.564079	0.168671
6	-2.398825	-0.132689	-1.060198
6	-1.972950	-1.490448	-0.751997
6	-2.276359	-0.018949	2.681493
1	-2.125074	1.049606	2.867344
1	-1.504763	-0.578425	3.220695
1	-3.250440	-0.301305	3.102964

6	-1.543435	-2.811790	1.466428
6	-3.146042	1.946670	0.328659
1	-0.762724	-2.586723	2.202189
1	-2.687557	2.454195	1.184267
1	-1.195327	-3.638722	0.842665
1	-2.966185	2.552546	-0.565280
1	-2.429032	-3.149165	2.019888
1	-4.229254	1.919769	0.500900
6	-2.609595	0.400599	-2.428660
6	-1.714624	-2.529884	-1.781787
1	-1.925587	-0.068477	-3.143574
1	-1.013519	-2.166916	-2.542523
1	-2.453409	1.484250	-2.461989
1	-1.302727	-3.443476	-1.346513
1	-3.635228	0.196324	-2.763075
1	-2.651002	-2.784774	-2.294527
8	1.217334	-2.784103	-0.235505
7	2.321600	3.002677	0.114815
8	5.489106	-1.269611	0.124242
7	3.361322	-2.071520	-0.037646
6	1.293540	4.008892	-0.188300
1	1.408590	4.358005	-1.221362
1	1.368503	4.866932	0.482008
6	0.005566	3.204581	0.009872
1	-0.754808	3.425496	-0.744202
1	-0.434774	3.373408	1.001629

(6-2H⁺)-HCO₂

E = -1322.97103036 a.u.

77	0.587861	0.039079	0.058123
7	-0.487247	1.844019	0.213566
7	-1.499946	-0.518529	-0.129874
6	-1.781260	1.800462	0.115991
6	-2.409754	0.495274	-0.063251
6	-3.754415	0.313532	-0.152388
1	-3.345990	3.059514	0.502565

1	-4.455864	1.142047	-0.104504
6	-4.235643	-1.036927	-0.346590
6	-1.998060	-1.812049	-0.239762
6	1.790314	0.517814	-1.639987
6	1.453309	-0.878647	-1.692112
6	2.577657	0.740023	-0.439685
6	2.706899	-0.516005	0.227885
6	2.017452	-1.527216	-0.533176
6	1.465922	1.533241	-2.679000
1	1.524364	2.550829	-2.278902
1	0.458423	1.384848	-3.083357
1	2.174594	1.462948	-3.514256
6	0.696066	-1.539340	-2.789126
6	3.197679	2.025638	-0.016960
1	0.026625	-0.831400	-3.289612
1	2.638901	2.888118	-0.394874
1	0.092902	-2.372098	-2.415565
1	3.245761	2.106780	1.074625
1	1.390472	-1.933100	-3.542327
1	4.223026	2.101894	-0.401389
6	3.427200	-0.754677	1.506251
6	2.082025	-2.992985	-0.274575
1	2.954318	-1.554367	2.084828
1	1.960100	-3.222423	0.789032
1	3.458418	0.147057	2.126798
1	1.307569	-3.533715	-0.825682
1	4.462810	-1.055050	1.300175
1	3.056678	-3.387631	-0.591072
8	-1.182396	-2.770253	-0.218082
7	-2.395867	3.009860	0.152580
8	-5.467533	-1.263460	-0.487805
7	-3.326933	-2.045443	-0.370908
6	0.104234	-0.986073	2.951605
1	-0.174659	-0.796686	4.012086
8	0.407101	-2.132861	2.606214
8	0.062430	0.072638	2.232478
6	-0.085322	3.248355	0.328812
1	0.683211	3.375537	1.097823

1	0.332505	3.586775	-0.630939
6	-1.397141	3.962997	0.663520
1	-1.489228	4.936701	0.178489
1	-1.520092	4.095960	1.745936

(6-2H⁺)-HCO₂-TS

E = -1322.94007791 a.u.

77	-0.637667	0.033076	0.091730
7	0.369993	1.862200	0.362960
7	1.423549	-0.381828	-0.343509
6	1.655272	1.889298	0.167776
6	2.308521	0.646177	-0.219171
6	3.653777	0.512882	-0.381660
1	3.091809	3.317116	-0.093779
1	4.335507	1.350792	-0.263154
6	4.155724	-0.796683	-0.732979
6	1.960391	-1.654084	-0.499881
6	-2.727160	-0.534064	0.384720
6	-2.069140	-1.541916	-0.415389
6	-2.674154	0.704895	-0.320888
6	-2.004643	0.476425	-1.589436
6	-1.645865	-0.907774	-1.640337
6	-3.373141	-0.772391	1.704769
1	-3.442372	0.150449	2.290137
1	-2.816170	-1.508453	2.295195
1	-4.391295	-1.160590	1.568510
6	-2.057389	-3.000395	-0.108148
6	-3.302634	1.980791	0.121806
1	-1.824349	-3.186715	0.946242
1	-3.170041	2.145169	1.197309
1	-1.321460	-3.536659	-0.713719
1	-2.887393	2.843910	-0.408756
1	-3.043420	-3.440632	-0.309087
1	-4.381739	1.961829	-0.079377
6	-1.800529	1.492504	-2.658109
6	-0.954014	-1.585341	-2.770258

1	-0.940339	1.243536	-3.288831
1	-0.348907	-0.879453	-3.349533
1	-1.628877	2.489692	-2.236361
1	-0.296149	-2.383861	-2.413636
1	-2.682009	1.560529	-3.309448
1	-1.689919	-2.030935	-3.451938
8	1.187363	-2.643485	-0.392020
7	2.249064	3.084181	0.419858
8	5.377394	-0.975406	-0.987524
7	3.276257	-1.832367	-0.758556
1	-0.144192	-0.298441	1.689071
6	0.725950	-1.353769	2.556336
8	1.794937	-0.834161	2.628443
8	-0.057681	-2.210628	2.822788
6	1.162480	4.073938	0.492817
1	1.037566	4.572273	-0.477326
1	1.361971	4.832206	1.252524
6	-0.041582	3.191129	0.827060
1	-0.959475	3.519719	0.331932
1	-0.237778	3.146106	1.907884

(6-2H⁺)-Hydride

E = -1134.44688057 a.u.

77	0.584816	0.000413	-0.328116
7	-0.488149	1.806364	-0.185438
7	-1.512718	-0.563734	-0.177932
6	-1.785639	1.760912	-0.144544
6	-2.418693	0.449593	-0.101710
6	-3.764573	0.260156	0.000133
1	-3.307108	3.066273	0.241040
1	-4.462782	1.091241	0.055220
6	-4.244491	-1.099713	0.062435
6	-2.017658	-1.857493	-0.224484
6	2.667220	-0.530435	-0.639897
6	2.000112	-1.649906	-0.030898
6	2.593955	0.576513	0.265191

6	1.948174	0.119278	1.487851
6	1.576731	-1.233022	1.296317
6	3.370130	-0.548066	-1.953322
1	3.402838	0.449107	-2.405200
1	2.879645	-1.225408	-2.660636
1	4.405661	-0.892131	-1.828230
6	2.028709	-3.051292	-0.542144
6	3.233573	1.908122	0.065715
1	1.851249	-3.092198	-1.622919
1	3.141805	2.248197	-0.972256
1	1.271207	-3.673248	-0.056177
1	2.786477	2.671052	0.712295
1	3.009042	-3.509396	-0.351077
1	4.304250	1.863678	0.306732
6	1.735879	0.960454	2.698699
6	0.862195	-2.109987	2.266081
1	0.968774	0.537228	3.355590
1	0.398615	-1.527997	3.069478
1	1.423400	1.977899	2.432616
1	0.075107	-2.687642	1.767369
1	2.661319	1.051433	3.282935
1	1.555811	-2.825046	2.727579
8	-1.211243	-2.805273	-0.431467
7	-2.404933	2.972461	-0.212668
8	-5.472771	-1.343050	0.218963
7	-3.338413	-2.106704	-0.057075
1	0.233974	0.104132	-1.919781
6	-0.095571	3.212321	-0.315999
1	0.763208	3.449502	0.318440
1	0.190929	3.404879	-1.360833
6	-1.362081	3.968414	0.083152
1	-1.508928	4.888766	-0.485141
1	-1.361556	4.210641	1.154213