Decamethylruthenocene Hydride and Hydrogen Formation at Liquid|Liquid Interfaces

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Supporting Information

ABSTRACT: The formation and the dissociation of metal hydrides are key steps within the hydrogen evolution reaction (HER) pathway for photochemical water splitting, but also impacts a wide range of other catalytic, industrial, and biochemical reactions. Herein, we describe our recent work studying HER at the interface between two immiscible electrolyte solutions (ITIES), between water|1,2-dichloroethane. This is a unique platform for evaluating the kinetics/thermodynamics for metal-lucene hydride formation using decamethylruthenocene. In this approach, an aqueous acid serves as the proton source and is pumped across the ITIES via an externally applied potential or the use of a phase transfer catalyst. Simulated curves developed using COMSOL Multiphysics software and compared to experimental ones, indicate a modified EC′ (electrochemical—chemical) mechanism for the decamethylruthenocene hydride formation. In the proposed pathway, decamethylruthenocene hydride is metastable in 1,2-dichloroethane and persists on the time scale of the recorded cyclic voltammograms long enough to transfer to the aqueous phase where it quickly dissociates. This is evidenced through an asymmetric, ion transfer wave observed experimentally and concluded to be hydride transfer. Shake-flask experiments with head space gas sampling demonstrated that hydrogen production was observed only when the biphasic system was positively polarized, to favor proton transfer, and decamethylruthenocene was photoactivated. This approach, combining electrochemical, simulation, and chromatographic methods, brings new insight into the factors that underlie the mechanism and rates of hydride formation/dissociation at soft interfaces.

1. INTRODUCTION

Transition metals are possessed of remarkable chemical properties such as multiple valencies, various coordination sites and a number of possible complex structures. As a consequence, they can be employed as catalysts owing to their exceptional and versatile reactivity and are capable of performing a wide range of chemical and biochemical reactions including hydrogenation, catalysis reduction, and water dissociation. Catalysts incorporating the transition metal ruthenium at their core have been found to be very effective owing to ruthenium’s eight oxidation states, its exceptional functional group tolerance, and its reasonable price relative to architecturally similar noble metal catalysts. Many organo-ruthenium catalyzed pathways have been explored and are well characterized, and they often involve the formation of a transition metal hydride as an intermediate. Metal hydrides are crucial steps that allow for the transfer of a free proton to an organic molecule, such as for hydrogenation or hydroformylation. The understanding and the control of proton transfer, and possibly proton coupled electron transfer, to and from the metal hydride is particularly important in photochemical water splitting and fuel cell electrocatalysis. Indeed, with rising energy demands, the depletion of fossil fuel reserves, as well as the resultant anthropogenic climate change owing to the use of carbon based fuels, society is on the cusp of a significant shift toward renewable energy sources. Organometallic catalysis is staged to play a key role to facilitate this change. Among future energy sources under consideration, molecular hydrogen (H2) appears very attractive as the only waste from burning H2 in air is water. Nevertheless,
H₂ production induces environmental impacts through the tremendous energy required for water electrolysis, a chief source of protons, and thus, it remains very expensive to produce owing to current methods based on platinum catalysts. Thus, an avid area of research is the development of alternative methods of H₂ production using more common, cheaper alternatives. In this way, investigation of the H₂ evolution reaction (HER) necessitates analysis of the key step: metal hydride formation.

The interface between two immiscible electrolyte solutions (ITIES), for example, at a water|1,2-dichloroethane (w|DCE) interface, provides an alternative electrochemical perspective from which to investigate the HER. Different charge transfer reactions can be monitored by controlling the potential difference between the two phases. Employing this utility, our group has developed an alternative strategy to produce H₂ that relies on electron reductants (e.g., metalloccenes), dissolved in an organic electrolyte solution and an acidic aqueous solution. In this way, product separation is inherent within the system, which is ultimately biomimetic. Upon polarization of the interface, either via the application of an external voltage or by dissolving a common ion in both phases, protons are pumped into the organic phase. Thus, the potential-dependent proton transfer step, with corresponding metal hydride formation, can be controlled and monitored electrochemically, thus facilitating the reactions kinetic/thermodynamic investigation. Furthermore, this methodology takes advantage of the fact that proton reduction in organic solvents, such as DCE, is thermodynamically more favorable than in aqueous media. Consequently, milder organic donors, with higher (more positive) redox potentials, can be effectively coupled to proton pumping across the interface to produce H₂. In our recent work, we demonstrated the production of H₂ by decamethylruthenocene (Cp²Ru(II); Cp*=C₅Me₅). Explorations of the mechanistic pathway revealed that the reaction could produce H₂ without an additional photosensitizer avoiding several electron transfer steps. Thus, coupling of the Cp²Ru(II) photocatalytic scheme with soft interfaces represents an interesting model platform to study the photoproduction of H₂ and is the major focus of this article.

In the present work, the first HER by Cp²Ru(II) at the liquid/liquid interface is presented. The hydride formation was characterized voltammetrically and the biphasic mechanism elucidated through comparison of experimental and simulated cyclic voltammograms (CVs); the latter generated using a 1D geometry developed within COMSOL Multiphysics software. A modified electrochemical (E)—chemical (C)—or EC mechanism, was the primary pathway evaluated, where potential-dependent interfacial proton transfer (from w to DCE) and the formation of the metal hydride within the bulk DCE phase constituted the electrochemical and chemical steps, respectively.

2. THEORY

2.1. Mechanism of Hydrogen Evolution at a Liquid/Liquid Interface. At its core, the mechanism of metalloocene hydride formation, as recently presented, can be regarded as an interfacial complexation reaction; whereby, the proton takes on the role of the ligand and coordinates to the metal center of Cp²Ru(II). Interfacial complexation reactions have been widely studied and offer insight into a possible mechanism for the ruthenocene hydride formation, either by transfer through interfacial complexation/decomplexation (TIC/TID); transfer of the proton, followed by organic phase complexation (TOC); or aqueous phase complexation, then transfer (ACT). In order to elucidate a possible reaction pathway, along with the associated kinetics/thermodynamics, a model was constructed within COMSOL Multiphysics software utilizing a simplified 1D geometry as described in detail elsewhere. Similar simulations have been employed to examine complex systems occurring at microelectrodes, indicating possible challenges with the use of ultramicroelectrodes, as well as the mechanistic elucidation and detection of short-lived radical species coupled with scanning electrochemical microscopy. The diffusion of chemical species within the system was defined using Fick’s laws, while the electrochemical flux of ions across the ITIES was defined using the Butler–Volmer series of equations as has been described previously. Further simulation details, along with the terms and coefficients, have been provided in the Supporting Information (SI) and Table S1.

3. EXPERIMENTAL SECTION

3.1. Chemicals. All chemicals were used as received without further purification. All aqueous solutions were prepared with ultrapure water (Millipore Milli-Q, specific resistivity 182 MΩ cm). The solvents employed were 1,2-dichloroethane (DCE, ≥99.8%, Fluka), α,α,α-trifluorotoluene (TFT, ≥99%, Sigma-Aldrich), hydrochloric acid (HCl, 37%, Merck), and sulfuric acid (H₂SO₄, 98%, Merck). Decamethylruthenocene (Cp²Ru(II), 99%) was supplied by ABCR and stored under an argon atmosphere until use. Anhydrous lithium chloride (LiCl, ≥99%), anhydrous sodium sulfate (Na₂SO₄, ≥99%), and tetraethylammonium chloride (TEACl, ≥98%) were obtained from Fluka.

Lithium tetrakis(pentafluorophenyl)borate diethyl etherate ([Li(OET₂)₂]TB, Boulder Scientific) and bis(triphenylphosphoranylidene)ammonium chloride (BACl, ≥98%, Aldrich) were used to prepare bis(triphenylphosphoranylidene)-ammonium tetrakis(pentafluorophenyl)borate (BACl) by metathesis of equimolar solutions of BACl and Li(OET₂)₂ in methanol/water (2/1, v/v). The resulting precipitate was collected by filtration, washed, and recrystallized from acetone.

3.2. Water/DCE, Shake-Flask Methodology. All anaerobic experiments were prepared using aqueous and organic solutions previously saturated with nitrogen and transferred to a nitrogen filled glovebox; wherein, samples were either placed in a black box for dark conditions or illuminated with white-light using a 500 W xenon lamp (Oriel, Research Arc Lamp Source).

Two-phase, so-called “shake-flask”, experiments were performed in septum-sealed glass vials with a solution composed of an oil (o) phase of DCE and an aqueous (w) phase whose compositions are exemplified schematically in Vial 1, below:

<table>
<thead>
<tr>
<th>2.5 mM DMRc</th>
<th>0.1 M HCl</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 mM BATB</td>
<td>5 mM LiTBDDE</td>
</tr>
<tr>
<td>(DCE)</td>
<td>(w)</td>
</tr>
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</table>

(Vial 1)

After combining the two phases, a magnetic stirrer was added, the vial was sealed, and then stirred for 60 min at ambient temperature.

One milliliter portions of the headspace gas from shake-flask vials were sampled using a lock-in syringe with a push—pull valve (SGE Analytical Sciences) and subsequently analyzed via injection into a gas chromatograph (PerkinElmer GC) equipped with a 20 μL sample loop, a 5 Å molecular sieves
column, with argon used as carrier gas, and a 80/100 mesh thermal conductivity detector (TCD).

UV/vis spectra were recorded with an Agilent Cary 8453 photodiode array spectrophotometer employing a quartz cell with an optical path length of 1 cm.

3.3. Electrochemical Measurements. Charge transfer reactions between wDCE were characterized using a traditional four electrode, electrolytic cell, as described previously, with an interfacial area of ~1.53 cm², and controlled through a PGSTAT 30 potentiostat (Metrohm, CH). Two platinum counter electrodes, connected to the working and counter electrode leads of the potentiostat were positioned in the aqueous and organic phases, respectively, and supplied the current. An external potential was applied by means of Ag/AgCl reference electrodes placed in Luggin capillaries whose tapered openings were positioned proximal and on opposing sides of the ITIES. This configuration can be described through Cell 1 below:

<table>
<thead>
<tr>
<th>Ag</th>
<th>AgCl</th>
<th>10 mM LiCl (w ≈ ref)</th>
<th>5 mM Cp₂Ru⁺⁺</th>
<th>5 mM BATB (DCE)</th>
<th>zMHCl</th>
<th>AsCl</th>
<th>Ag</th>
</tr>
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<tbody>
<tr>
<td>(Cell 1)</td>
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</table>

The concentration of HCl (z) was varied from 0.001 to 0.010, and 0.100 M, or pH 3, 2, and 1, respectively. The measured Galvani potential difference across the wDCE interface, Δφₜ unsettled, was calibrated relative to the formal ion transfer potential of the tetraethylammonium cation (TEA), taken to be 0.019 V, except for the determination of the formal ion transfer potential of the decamethylruthenocenium ion ([Cp₂Ru⁺⁺]), Δφₜ unsettled, which was calibrated relative to the formal ion transfer potential (Δφₜ unsettled) of tetraethylammonium cation (TEA⁺), taken to be 0.160 V. The half-wave ion transfer potentials (Δφₜ unsettled) of TEA⁺ and TMA⁺ were determined using cyclic voltammetry (CV). Experiments were performed with aqueous and organic phases thoroughly degassed using nitrogen, under anaerobic conditions in a nitrogen filled glovebox, in the dark, and at an ambient temperature, unless otherwise stated.

4. RESULTS AND DISCUSSION

4.1. Cyclic Voltammetric Observations at wDCE Interfaces. Figure 1A depicts the CVs obtained at a scan rate of 0.050 V s⁻¹ using Cell 1 at varying pH from 3 to 2 to 1 for the red, blue, and green traces, respectively, as well as in the absence of Cp₂Ru⁺⁺ (black trace) at pH 3, which served as the blank. In the case of the blank, the potential window is limited by the transfer of protons from (w) to (o) at the positive end, and similarly chloride ion (Cl⁻) transfer at negative potentials.

Upon closer inspection, the blank curve, without Cp₂Ru⁺⁺, differs from the CVs obtained in the presence of Cp₂Ru⁺⁺ in two respects. First, at the positive end, the CV potential profile for the blank has an expected return peak at 0.413 V, indicative of protons transferring back from (o) to (w); however, this curve feature is absent in the CVs acquired in the presence of Cp₂Ru⁺⁺ indicating irreversible proton transfer. This is in good agreement with the results of Samec et al. for ferrocene derivatives, along with our recent results, using decamethylferrocene (Cp₂Fe⁺⁺) at a wDCE interface. For the latter, proton transfer is followed by [Cp₂Fe⁺⁺H]⁺ formation, which subsequently reacts with dissolved O₂ to generate [Cp₂Fe⁺⁺O₂]⁺ as well as H₂O₂ and H₂O; i.e., eliciting the oxygen reduction reaction (ORR). Therefore, for the present case, it is likely that protons are coordinating to Cp₂Ru⁺⁺ and generating [Cp₂Ru⁺⁺(H)]⁺.

Second, after addition of Cp₂Ru⁺⁺ and during the cathodic scan in Figure 1A, a peak-shaped wave emerges with a peak potential, Δφₜ unsettled, at approximately -0.280 V for pH 3. This ion transfer wave is likewise irreversible in that an analogous peak is not observed in the anodic scan. Fundamentally, this could be attributed to either transfer of a positively charged species, from (o) to (w), or that of a negatively charged species, from (w) to (o). An initial clue to the origin of this ion transfer peak was obtained when the CV potential range was manipulated (see Figure 1B) such that the initial CV was swept to moderate positive potentials while subsequent scans went to increasing positive potentials. The ion transfer wave at roughly -0.280 V was observed to increase in cathodic current intensity concomitantly with increases in the positive potential range. Therefore, it can be concluded that there is a correlation between the occurrence of this peak, along with its intensity, and the amount of protons transferred. Indeed, after consideration of the control experiments discussed vide infra, this ion transfer wave is attributed to the transfer of [Cp₂Ru⁺⁺(H)]⁺ from (o) to (w), the rate of formation of which is intimately linked to the quantity of protons transferred from (w) to (o).

One alternative positively charged species that may be present in DCE is the cationic [Cp₂Ru⁺⁺(H)]⁺ species. Therefore, the latter was synthesized by electrolysis, as described previously, and its formal ion transfer potential (Δφₜ unsettled) determined through CV to be 0.005 V (see Cell S1 and Figure S1 in the SI). Owing to the disparity between this value recorded for [Cp₂Ru⁺⁺]⁺ and that found using Cell 1, one can conclude that the transfer wave at roughly -0.280 V is not owing to [Cp₂Ru⁺⁺(H)]⁺. Two further possibilities are the transfer of [Cp₂Ru⁺⁺(H)]⁺ from (o) to (w), or potential interactions between Cp₂Ru⁺⁺ and Cl⁻ that may facilitate Cl⁻ transfer from (w) to (o). In order to differentiate between these
two possibilities, HCl was replaced by 5 mM of Li$_2$SO$_4$ and subsequently, one drop of 0.5 M KCl solution was added (see Cell S2 and Figure S2 in the SI). In the case of Cell S2 with Li$_2$SO$_4$ the CV was not affected by the presence of Cp$_2$Ru(II), this indicates that Li$^+$ does not coordinate to Cp$_2$Ru(II) nor do protons within its hydration sphere, as was recently discovered for Cp$_2$Fe(II).$^{23,31}$ After addition of KCl, the potential window response narrowed owing to the reversible K$^+$ and Cl$^-$ transfer at the positive and negative limits, respectively. No ion transfer wave was observed in the region of $\sim$0.280 V and a return peak was observed in the current—potential profile at the positive potential limit; therefore, one can conclude that Cl$^-$ does not interact with Cp$_2$Ru(II), at least in a manner observable by CV. However, when HCl (Cell 1) was replaced by H$_2$SO$_4$ (see Cell S3 and Figure S3 in the SI) the obtained CV generated a similar profile to that illustrated in Figure 1A, i.e., irreversible proton transfer at the positive potential limit, as well as an asymmetric ion transfer wave at $\sim$0.280 V. Therefore, proton interaction with Cp$_2$Ru(II) generating [Cp$_2$Ru(H)]$^+$, plays a key role in the observed CVs.

To further explore the formation of [Cp$_2$Ru(IV)(H)]$^+$, CVs were performed using Cell S4 and have been plotted in Figure S4 of the SI such that the concentration of Cp$_2$Ru(II) in DCE was varied, while HTB has been added to the organic phase. When the concentration of HTB is roughly equivalent to that of Cp$_2$Ru(II) ($c = 165 \mu M$), the proton return peak is relatively unchanged relative to the blank trace (without HTB or Cp$_2$Ru(II)). However, the transfer wave at $\Delta \phi \approx 0.280$ V is still present. Therefore, one can conclude that [Cp$_2$Ru(IV)(H)]$^+$ is generated in the organic phase using dissolved HTB and that one equivalent of the acid is sufficient to fully convert Cp$_2$Ru(II) to [Cp$_2$Ru(IV)(H)]$^+$. Furthermore, as illustrated in Figure S4, the magnitude of the proton return peak decreases with increasing Cp$_2$Ru(II) concentration as the HTB concentration remains static. One can consider as the Cp$_2$Ru(II) concentration increases the rate of complexation increases concomitantly (more Cp$_2$Ru(II) are free to coordinate to protons) and the rate of dissociation decreases. This accounts for the decrease in magnitude of the proton return peak with a simultaneous increase in magnitude of the [Cp$_2$Ru(IV)(H)]$^+$ transfer wave as the concentration of Cp$_2$Ru(II) increases (Figure S4). However, it was not possible to study the intensity of the current for the wave relative to [Cp$_2$Ru(IV)(H)]$^+$ transfer quantitatively because the latter is highly sensitive to the proton concentration of the organic phase, which is not stable owing to ITIES polarization and the presence of HTB.

4.2. Shake-Flask Experiments. In order to further investigate the phenomena observed voltammetrically, shake-flask experiments were performed using the wDCE solution composition described for Vial 1 in section 3.2. Two main conditions were varied: with/without (i) light exposure and (ii) electrochemical polarization of the wDCE interface. In order to polarize the interface, a phase transfer catalyst was employed, as previously described,$^{12-14}$ using TB$^-$ as the common ion initially dissolved in the aqueous phase. TB$^-$ spontaneously transfers from (w) to (o) and fixes the Galvani potential difference across the ITIES at positive values ($\Delta \phi \approx 0.50$ V). Simultaneously, protons are driven to the organic phase; by which means they are available to react with Cp$_2$Ru(II).

Photographs taken of the reaction vial for the case where both light illumination and ITIES polarization were used are displayed in Figures 2A–C. Once solubilized in DCE, Cp$_2$Ru(II) produced a clear organic phase (Figure 2A). After 1 h of irradiation, the DCE phase exhibited a pink color (Figure 2B) with a strong UV/vis absorption band at 500 nm (see Figure S5 of the SI), which agrees well with the absorption band for [Cp$_2$Ru(III)]$^+$ as shown previously.$^{18}$ If the reaction was allowed to stir for 180 min, then the solution turned purple, which agrees with the appearance of [Cp$_2$Ru(IV)(C$_{6}$M$_{6}$C$_{6}$H$_{12}$)]$^+$ (Figure 2C) after prolonged photoirradiation.$^{18}$ In summary, [Cp$_2$Ru(III)]$^+$ was seen to transition from Ru(II) (colorless), to Ru(III) (pink), and finally to Ru(IV) (purple).

Additionally, after 60 min of stirring, the headspace gas was sampled, as described in section 3.2, and analyzed using gas chromatography; chromatograms have been plotted in Figure 2D for each case as indicated inset. For the case of both ITIES electrochemical polarization and light irradiation, a peak corresponding to H$_2$ gas was observed after $\sim$2 min; however, no H$_2$ evolution was observed for the cases where either the phase transfer catalyst was omitted or the vial was kept in the dark. This indicates that Cp$_2$Ru(II) is a possible photoactive electron donor for H$_2$ evolution under biphasic conditions and that this HER is predominately performed in the bulk organic phase; therefore, Cp$_2$Ru(II) does not undergo appreciable interfacial electron transfer to aqueous protons.

The amounts of H$_2$ evolved were recorded as a function of time using a calibrated GC with wDCE (red squares) as well as wtrifluorotoluene (wTFT, navy circles) liquidliquid systems for comparison (Figure 3). The solution compositions were as outlined in Vial 1 but with an initial excess of [Li(OEt$_2$)$_2$]TB present in (w) relative to Cp$_2$Ru(II) in (o), and therefore, an excess of protons transferred from (w) to (o) as tetraakis-(pentfluorophenyl)borate diethyl etherate acid [H(OEt$_2$)$_2$]TB. The quantity of H$_2$ evolved was observed to reach a plateau.
after 180 min at \(\sim 5.0 \mu\text{mol}\). This value is twice the predicted stoichiometric amount of \(\text{H}_2\) estimated to be 2.5 \(\mu\text{mol}\) (Figure 3, dashed line) if each \(\text{Cp}_2\text{Ru}^{(II)}\) molecule acts as a single electron donor. Ruthenium is a well-known catalyst for hydrogenation reactions such as the Noyori asymmetric hydrogenation, which have become of interest in CO and \(\text{CO}_2\) reduction reactions to methanol. The first step in the activation of the ruthenium catalyst often involves the formation of the metal hydride, where a proton coordinates to the metal center. Organo-ruthenium complexes have also been used for C–H bond activation, as demonstrated by the Ackermann group for alkyne annulations. In this way, it is possible for the ruthenium core to extract a proton from one of its ligands (e.g., a proton from one of the methyl groups on the cyclopentadienyl rings), thus providing the second equivalent of protons needed to facilitate \(\text{H}_2\) evolution. In other words, the permethylated cyclopentadienyl (\(\text{Cp}^*\)) group is converted from \((\eta^5\text{-C}_5\text{Me}_5)\) into the form of a methylene(cyclopentadienyl) ligand \((\eta^5\text{-C}_5\text{Me}_5\text{CH}_2)\). Indeed, these results are in good agreement with our recent report studying the reaction mechanism of the photoproduction of \(\text{H}_2\) with \(\text{Cp}_2\text{Ru}^{(II)}\) in a single acidified organic phase.

Additionally, the \(\text{wDCE}\) biphasic system was compared to one using \(\text{wTFT}\). TFT has been previously proposed as an alternative solvent to study charge transfer across the interface, as well as for \(\text{H}_2\) and \(\text{H}_2\text{O}_2\) formation in biphasic systems, owing to its lower toxicity and price relative to DCE. The amount of \(\text{H}_2\) evolved over time has been plotted in Figure 3 (nary circles); however, the maximum yield approaches, but never exceeds, the quantitative yield calculated: 2.5 \(\mu\text{mol}\) (Figure 3, dashed line) if each \(\text{Cp}_2\text{Ru}^{(II)}\) molecule acts as a single electron donor. Ruthenium is a well-known catalyst for hydrogenation reactions such as the Noyori asymmetric hydrogenation, which have become of interest in CO and \(\text{CO}_2\) reduction reactions to methanol. The first step in the activation of the ruthenium catalyst often involves the formation of the metal hydride, where a proton coordinates to the metal center. Organo-ruthenium complexes have also been used for C–H bond activation, as demonstrated by the Ackermann group for alkyne annulations. In this way, it is possible for the ruthenium core to extract a proton from one of its ligands (e.g., a proton from one of the methyl groups on the cyclopentadienyl rings), thus providing the second equivalent of protons needed to facilitate \(\text{H}_2\) evolution. In other words, the permethylated cyclopentadienyl (\(\text{Cp}^*\)) group is converted from \((\eta^5\text{-C}_5\text{Me}_5)\) into the form of a methylene(cyclopentadienyl) ligand \((\eta^5\text{-C}_5\text{Me}_5\text{CH}_2)\). Indeed, these results are in good agreement with our recent report studying the reaction mechanism of the photoproduction of \(\text{H}_2\) with \(\text{Cp}_2\text{Ru}^{(II)}\) in a single acidified organic phase.

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4.3. Mechanistic Analysis Using Simulated Voltammetric Curves. In order to consolidate and further evaluate the voltammetric curves and shake-flask data, a theoretical model was constructed using COMSOL Multiphysics software (details in the SI and Section 2 vide supra). The experimental data leads one to conclude that proton transfer is coupled to a bulk, organic phase reaction; whereby, protons are consumed in the organic phase and unavailable to undergo ion transport back across the ITIES, effectively generating an irreversible ion transfer process. With this in mind, the proposed mechanism (Figure 4) includes a potential-dependent proton transfer step followed by formation of the \([\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+\) species. The ion transfer wave observed experimentally, at roughly \(-0.280\text{ V}\) (Figure 1B), is dependent on the concentration of protons transferred (Figure 1B); therefore, one can conclude that it is associated with \([\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+\) species. However, since this transfer wave is asymmetric (lacks a comparable return wave) and, as demonstrated by the shake-flask experiments for HER, is slow, relative to the time scale of the CV experiment, one can also conclude that the \([\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+\) formed is not yet fully dissociated. Therefore, a potential-dependent \([\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+\) transfer step was explored in the model (rather than a \([\text{Cp}_2\text{Ru}^{(III)}(\text{H})]^+\) transfer step), followed by dissociation of the metal hydride in water, as the HER is not thermodynamically favored in water, and finally partitioning of the neutral \(\text{Cp}_2\text{Ru}^{(II)}\) species back to the organic phase.

The effect of the rate of hydride formation was explored by systematically altering \(k_{\text{dl}}\) (the rate of \([\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+\) formation in the organic phase, see Figure 4), and the resultant CVs have been plotted in Figure 5; all other parameters were maintained as given in Table S1. As \(k_{\text{dl}}\) increases from \(10^2\text{ M}^{-1}\text{s}^{-1}\) up to \(10^6\text{ M}^{-1}\text{s}^{-1}\) (Figure 5A,B) the ion transfer \([\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+\) peak increases in magnitude from 0 to \(-230\mu\text{A}\). Simultaneously, the on-set potential for proton transfer decreases, and at \(k_{\text{dl}}\) equal to \(10^6\text{ M}^{-1}\text{s}^{-1}\) an anodic peak was generated at \(-0.410\text{ V}\) (Figure 5B). This shift in the potential of proton transfer is owing to the facilitated proton transfer, where protons coordinate to the \(\text{Cp}_2\text{Ru}^{(II)}\) metal center. Three mechanisms at the liquidliquid interface have been recognized and are described in Section 2.1. Owing to the hydrophobicity of \(\text{Cp}_2\text{Ru}^{(II)}\) and the dependence of the transfer wave at \(-0.280\text{ V}\) on the amount of protons transferred, as demonstrated in Figure 1B, as well as the need for a phase transfer catalyst in the shake flask experiments (Figure 2B), a spontaneous ACT mechanism can be excluded. The effective ACT pathway included in Figure 4 is dependent on the hydride formation in the organic phase. Only then is a potential-dependent \([\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+\) transfer step possible; whereby, the externally
applied Galvani potential difference is the driving force. For simplicity, the TIC/TID mechanism was not initially considered; however, it is a possible mechanism, but unlikely since the shake-flask experiments required electrochemical polarization in order to elicit H₂ production.

In order to elucidate this point further, a two-step EC’ (electrochemical, E, chemical, C’) mechanism, with a potential-dependent ion transfer step (E) of a generic species (E*c) followed by an irreversible complexation step (C’, see equations S1 and S2 of the SI), was employed within a similar simulation geometry. In this comparatively simple system, the neutral ligand (L) is considered hydrophobic and does not undergo transfer across the ITIES with the same being true of the ion–ligand complex (IL*c). The resulting CVs are given in Figure S6 with the formal ion transfer potential of E*c, Δϕ*c, set arbitrarily equal to 0.250 V. As the rate of complexation increases, the half-wave potential decreases for the transfer of E*c and the ion transfer wave becomes asymmetric (the “cathodic” component disappears). This is in good agreement with the theory of facilitated ion transfer as has been previously established.21,22,43,44 This also serves to further illustrate that the lack of an observable proton transfer return peak within the vicinity of the interface decreases, the return peak for proton transfer re-emerges, and the cathodic wave for [Cp²Ru(IV)(H)]⁺ transfer, (a) to (w), at ~0.200 V decreases concomitantly. This demonstrates that the rate of complexation is high and effectively irreversible, i.e., negligible decomplexation.

Subsequently, the analogous reaction was investigated in the bulk aqueous phase by first examining the rate of decomplexation, k_{d2} (Figure 4). Figure 7 illustrates that as k_{d2} increases, from 50 to 10⁵ s⁻¹, the half-wave potential of the cathodic wave shifts to more positive potentials, from −0.447 to −0.197 V, for the red and blue curves, respectively, such that k_{d2} = 10 s⁻¹ provides a half-wave potential of −0.280 V.

Figure 5. Simulated CVs generated using the mechanism illustrated in Figure 4 and using the parameters listed in Table S1, except that k_{d2} has been varied as indicated and that c_{Ru,aq}⁰ was maintained at 1 mM.

Figure 6. Simulated CVs employing the mechanism drawn in Figure 4, while k_{d2} was varied as indicated, inset; all other parameters are the same as those listed in Table S1.

Figure 7. (A) CV traces obtained using the simulation described in Section 2. All parameters are the same as those described in Table S1 (c_{Ru,aq}⁰ = 1 mM); however, here k_{d2} has been altered as indicated, inset. (B) Plot of the [Cp²Ru(IV)(H)]⁺ transfer peak current intensity versus k_{d2} from [A].

Half-wave potentials, Δϕ*ν/ν, have been calculated using the [Cp²Ru(IV)(H)]⁺ transfer peak potential, Δϕ*ν⁺ and the following relationship: Δϕ*ν/ν = Δϕ*ν⁺ + (0.028 V)/z;45 however, the ion transfer peaks in Figure 7 demonstrate a peak broadening with increasing k_{d2}. For k_{d2} equal to 50, 100, 1000, 10³, and 10⁵ s⁻¹, the peak width at −4 μA increases from 0.075 V to 0.080, 0.090, 0.105, and 0.120 V, respectively. Additionally, the peak current intensity increases negatively to 10⁵ s⁻¹, after which it decreases (Figure 7B). The formal ion transfer potential for [Cp²Ru(IV)(H)]⁺ was approximated to be −0.500 V, which is just beyond the polarizable potential window (PPW). The simulated curves in Figure 7A illustrate that the transfer potential of [Cp²Ru(IV)(H)]⁺ is effected in a similar way by the change of the aqueous decomplexation rate as the proton transfer potential was by k_{d2}. That is, with increasing k_{d2} the half-wave potential of [Cp²Ru(IV)(H)]⁺ shifts...
to more positive potentials. However, while altering $k_{b_{2}}$ influences $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ transfer, it does not alter the proton transfer potential; likewise, changing $k_{d_{2}}$ does not impact the potential of $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ transfer (Figure 6). Indeed, results obtained experimentally with Cell S4 (see S1) show that when $\text{Cp}_2\text{Ru}^{(II)}$ concentration, and consequently, $k_{d_{2}}$ and $k_{b_{2}}$, vary in the organic phase, the potential of $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ transfer remains constant. Moreover, the potential of the proton transfer appeared dependent on $k_{d_{2}}$ and $k_{b_{2}}$. The potentials switch to more negative values when more $\text{Cp}_2\text{Ru}^{(II)}$ are present because the transfer is facilitated by the proton complexation with the metallocene. However, like as was shown by simulation, the transfer potential is not expected to be dependent on $k_{d_{2}}$ and $k_{b_{2}}$ because the quantity of protons provided by $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ is negligible compared to the amount of free protons in the aqueous phase. Therefore, the resulting simulated CVs are in good agreement with the phenomena observed experimentally (see Figure S4).

Moving forward, $k_{d_{2}}$ was iteratively altered while the other rate constants were maintained (Table S1 in the SI); the resultant, simulated CVs have been plotted in Figure 8. As $k_{d_{2}}$ increases, the stability of $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ in the aqueous phase increases, and as $k_{d_{2}}$ approaches $k_{b_{2}}$ the CVs become more reversible with an anodic wave becoming visible at 1000 M$^{-1}\text{s}^{-1}$ and continues to increase in anodic current intensity until $1 \times 10^{5}$ M$^{-1}\text{s}^{-1}$. Simultaneously, the half-wave potential for $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ decreased until $1 \times 10^{2}$ M$^{-1}\text{s}^{-1}$, where it equals approximately the arbitrary value set in the simulation parameters.

The simulated data in Figures 7 and 8 indicate that the aqueous reaction heavily favors the dissociation of $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$, therefore, based on the shift in the $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ half-wave potential and the asymmetry of the anodic peak (or lack thereof) values of $k_{d_{2}}$ and $k_{b_{2}}$ have been estimated to be 100 M$^{-1}\text{s}^{-1}$ and $1 \times 10^{5}$ s$^{-1}$, respectively.

5. CONCLUSIONS

We have demonstrated $\text{Cp}_2\text{Ru}^{(II)}$ can act as a sacrificial electron donor to produce H$_2$ at polarized interfaces under light illumination employing DCE or TFF solvents.

The formation of $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ at the wDCE interface was investigated through cyclic voltammetry and subsequent comparison to simulated CVs generated using COMSOL Multiphysics software in order to elucidate the kinetics/thermodynamics. The proposed pathway follows a modified EC mechanism that involves a potential-dependent proton transfer step (E), from (w) to (o), followed by favorable $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ formation in the DCE phase (C'). The HER is slow, however, and on the time scale of CV the $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ transfers as the Galvani potential difference is swept to negative potentials (an asymmetric cathodic wave was observed) and rapidly dissociates in water. This is evidenced by two curve features in the recorded CVs: an irreversible transfer wave at the limit of the PPW for protons, along with the irreversible transfer wave at roughly −0.280 V that has been herein identified as $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ transfer. This is the first time an irreversible ion transfer wave has been observed in relation to metallocene hydride formation at a liquid/liquid interface and is attributed to the slow rate of reaction relative to other metallocene analogues.

The shake-flask/GC experimental data corroborates that recorded using CV, as well as the postulated mechanism, in that $\text{Cp}_2\text{Ru}^{(II)}$ oxidation and H$_2$ production were only observed when both ITIES polarization and the samples were illuminated. This indicates that the $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ formation is a bulk organic phase reaction and also requires the formation of a $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ or $([\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+)_{n}$ excited species. The evaluation of this excited state species is underway utilizing electrogenerated chemiluminescence (ECL); however, it is beyond the scope of the present work.

Furthermore, the investigations on the effect of the rate of hydride formation explain the irreversible nature of the proton transfer when the hydride is stable and that the cathodic irreversible peak, observed in the CV and never seen before with other metallocenes, can be attributed to $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ transfer. The effect of the rate of hydride formation and dissociation was explored confirming that the rate of complexation is high ($\sim 10^{5}$ M$^{-1}\text{s}^{-1}$), as confirmed through the simulations, and irreversible in the DCE phase. In contrast, the absence of an associated anodic peak related to $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ transfer from (w) to (o), is owing to the spontaneous decomplementation of the hydride in water, which was determined to occur at a rate of $\sim 10^{6}$ s$^{-1}$, while a rate of aqueous complexation was also estimated to be 100 M$^{-1}\text{s}^{-1}$. Furthermore, the transfer potential of the hydride is affected by the rate of the decomplementation of the hydride in the aqueous phase; however, it does not alter the proton transfer potential. Likewise, changing the rate of formation of the hydride in DCE phase does not impact the potential transfer of $[\text{Cp}_2\text{Ru}^{(IV)}(\text{H})]^+$ transfer but does influence the proton transfer potential.

ASSOCIATED CONTENT

3 Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.5b08148.

Additional electrolytic cells; cyclic voltammetric determination of $[\text{Cp}_2\text{Ru}^{(III)}]$ by Cell S1; interaction of Li$_2$SO$_4$ and KCl with $\text{Cp}_2\text{Ru}^{(II)}$ Cell S2; voltammetric investigation of H$_2$SO$_4$ interaction with $\text{Cp}_2\text{Ru}^{(II)}$ through Cell S3; effect of HTB and $\text{Cp}_2\text{Ru}^{(II)}$ on the CV response using Cell S4; preparation of tetrakis(pentafluorophenyl)bora diethyl acid; UV/vis absorbance spectra for Vial 1 at t = 0 and 60 min; simulation details; simulation terms and coefficients; simulated
cyclic voltammograms generated using a simplified EC mechanism (PDF).

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