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Voltammetric Perspectives on the Acidity Scale and H⁺/H₂ Process in Ionic Liquid Media

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Abstract
Nonhaloaluminate ionic liquids (ILs) have received considerable attention as alternatives to molecular solvents in diverse applications spanning the fields of physical, chemical, and biological science. One important and often overlooked aspect of the implementation of these designer solvents is how the properties of the IL formulation affect (electro)chemical reactivity. This aspect is emphasized herein, where recent (voltammetric) studies on the energetics of proton (H⁺) transfer and electrode reaction mechanisms of the H⁺/H₂ process in IL media are highlighted and discussed. The energetics of proton transfer, quantified using the pKₐ (minus logarithm of acidity equilibrium constant, Kₐ) formalism, is strongly governed by the constituent IL anion, and to a lesser extent, the IL cation. The H⁺/H₂ process, a model inner-sphere reaction, also displays electrochemical characteristics that are strongly IL-dependent. Overall, these studies highlight the need to carry out systematic investigations to resolve IL structure and function relationships in order to realize the potential of these diverse and versatile solvents.
1. INTRODUCTION

Ionic liquids (ILs) possess physicochemical properties that are identical to those of molten salts. However, practical aspects regarding their use, maintenance, and handling are sufficiently different to warrant a distinction (1, 2). ILs are arbitrarily defined as any ionic compound with a melting point or glass transition below the boiling point of water at 1 atm pressure (i.e., 100°C) (3). The ILs discussed herein are of the so-called second/third generation air/water stable nonhaloaluminate class (2, 4, 5). Interest in ILs in both academia and industry for use in a range of applications has increased dramatically over the last 25 years (2, 6, 7). They are versatile solvents, and have been employed as replacements for volatile organic compounds in green synthesis (8, 9), separations (10), clean catalysis (11), and as electrolytes in electrochemical devices (1, 12, 13). In addition, ILs have found application in the biosciences, being employed as media for enzymatic reactions (14) and solubilization of biomolecules (15). In this review, a brief introduction to the general structure and physicochemical properties of ILs is presented, followed by an in-depth discussion on the energetics (thermodynamics) of proton transfer (i.e., equilibrium acidity) and electrochemical mechanisms of the H⁺/H₂ process in this class of solvent.

1.1. Structure of Ionic Liquids

Although all ILs are often treated collectively as a class of solvent, the physicochemical properties (e.g., melting point, viscosity, polarity, solvating ability) of each formulation are uniquely determined by the nature of the constituent cation and anion. The chemical structures and general properties of the cations and anions that constitute commonly encountered aprotic and protic ILs are considered below.

1.1.1. Aprotic ionic liquids.

Aprotic ionic liquids (AILs) are those which contain no dissociable protons in their structures. The constituent cations of AILs tend to be organic, bulky, and asymmetric, with extensively shielded or delocalized charge (i.e., they are weak Lewis acids). Altering the chain length of the alkyl substituents on the cation influences the physicochemical properties of the resulting IL (e.g., melting point, hydrophobicity, and viscosity), which allows fine-tuning to a particular application (3, 16). The anions of AILs tend to be inorganic with highly diffuse or delocalized charge. They are typically weak Lewis bases and poor hydrogen bond acceptors. The
hydrophilicity/hydrophobicity, and thus, the level of water-miscibility of an IL, are largely determined by the choice of anion (3, 17, 18).

The diffuse, shielded, or delocalized charge on the constituent anion and/or cation of an IL results in intermolecular forces that are relatively weak compared to traditional ionic compounds. This translates to a relatively low lattice energy (further lowered by inefficient packing due to cation asymmetry), which is why these compounds are liquid at or below 100°C. It is important to note, however, that the cohesive forces operating within an IL are strong relative to those operating in most conventional molecular solvents, and for this reason, ILs often possess negligible vapor pressures and relatively high viscosities (1, 3). The names, abbreviations, and structures of a range of commonly encountered AIL cations and anions are summarized in Figure 1.

Figure 1

1.1.2. Protic ionic liquids.
Protic ionic liquids (PILs) are formed through the transfer of a proton from a Brønsted acid (HA) to a Brønsted base (B):

\[ HA + B \rightarrow A^- + [BH]^+ . \]  

By definition, they contain dissociable (acidic) protons and possess intrinsic (anhydrous) proton conductivity. The proton activity in PILs depends on the identity of the constituent Brønsted acid/base, analogous to pH in aqueous systems, which allows some control over proton-coupled chemical/electrochemical processes (19–26).

When complete transfer of the proton from HA to B occurs, the properties of PILs are generally not discernible from those of AILs. This is often the case when HA is a strong acid and B is a strong base in aqueous media, for example, when triflic acid is combined with a tertiary
amine. When proton transfer from HA to B does not go to completion, the melt may contain a significant amount of neutral species (i.e., HA/B). This is often the case when HA is a weak aqueous acid and/or B is a weak aqueous base, for example, when a carboxylic acid is combined with imidazole or pyridine. Proton-donor (i.e., HA or BH\(^+\)) and -acceptor (i.e., B or A\(^-\)) sites establish a hydrogen-bonded cation/anion network, often resulting in water-like properties (23, 25). In addition, when the proton transfer is reversible (does not go to completion), the boiling temperature of [BH][A] can be below its decomposition temperature, thereby allowing the PIL to be distilled (23, 27, 28). The names, abbreviations, and structures of commonly encountered PIL cations and anions are also provided in Figure 1.

1.2. Physicochemical Properties of Ionic Liquids

One attractive feature of ILs is their great versatility; it has been estimated that \(10^{18}\) different ILs could be prepared by varying the constituent anion and cation (3). Indeed, they are commonly referred to as designer solvents because their physicochemical properties can be tuned to an extent by changing their constituent cation and/or anion. Although the only truly ubiquitous property shared by all ILs is intrinsic ionic conductivity, in the chemical literature, they are often claimed to possess a number of generic properties. These include variable degrees of water miscibility (hydrophobicity/hydrophilicity); high viscosity (usually in the range of 20 to 1,000 cP); negligible vapor pressure and nonflammability; high solvating ability, but low or no coordinating ability; high thermal, chemical, and electrochemical stability; large liquidus range; and they are nontoxic/environmentally friendly (1, 3, 7, 16, 29–31). In reality, the actual physicochemical properties of an IL are decided by the identity of its constituent cation and anion, and although some ILs do possess some of the above attributes, there are many exceptions (5, 30). The focus herein is the specific use of ILs as nonconventional electrolytes in electrochemical investigations, whereby the IL simultaneously functions as the solvent and salt (1, 32), with an emphasis on the energetics (thermodynamics) of proton transfer (i.e., minus logarithm of acidity equilibrium constant, \(K_a\), known as p\(K_a\)) and electrode reaction mechanisms of the H\(^+\)/H\(_2\) process.

2. PROTON TRANSFER IN IONIC LIQUID MEDIA
Proton (H\(^+\)) transfer is involved with many of the most important reactions in biology and chemistry (33, 34). For example, proton transfer and conduction play an important role in green energy conversion technologies, such as electrochemical CO\(_2\) fixation (electro-reduction) (28, 35, 36), methanol oxidation (37), and water (photo)electrolysis (38). As alluded to above, air/water stable (nonhaloaluminate) ILs, particularly PILs, have been identified as potential anhydrous proton conductors, which could be employed at temperatures well above 100°C (23, 39). Indeed, there are many applications in areas as broad as energy conversion and storage (e.g., proton-exchange membrane fuel cells, PEMFCs) (12, 13, 39–42), synthetic chemistry (e.g., hydrogenation/dehydrogenation) (8, 9), separations chemistry (15), and electrochemistry (e.g., electrochemical hydrogenation, gas sensors) (43) that would benefit from high-temperature, anhydrous proton conduction. A general discussion on the energetics (thermodynamics) of proton transfer is included below, followed by a summary highlighting recent electrochemical studies on the pK\(_a\) scale in IL media.

### 2.1. Equilibrium Acidity

The acid dissociation constant (K\(_a\)) of a Brønsted acid, HA, formally defined as follows, is a quantitative measure of acid strength in solution:

\[
\text{HA} \rightleftharpoons \text{H}^+ + \text{A}^- \quad K_a = \frac{[\text{H}^+][\text{A}^-]}{[\text{HA}]},
\]

where H\(^+\) is taken to be a solvated proton species (vide infra). Although the acid is taken to be a neutral species (HA) in Equation 2, the expression for K\(_a\) is equally valid for a cationic acid species, BH\(^+\) (i.e., a protonated neutral base, B):

\[
\text{BH}^+ \rightleftharpoons \text{H}^+ + \text{B} \quad K_a = \frac{[\text{H}^+][\text{B}]}{[\text{BH}^+]}.
\]

It should be noted that, while the theory discussed herein refers to either HA or BH\(^+\) (or HA/H\(_2\) or BH\(^+\)/H\(_2\) couples, respectively; see below) as the weak acid, it is equally applicable for both species. Brønsted acidity/basicity is a well-established concept in aqueous media (44), where water, an amphoteric species, can act as a Brønsted base and accept a proton:
HA (or BH⁺) + H₂O(base) ⇌ A⁻ (or B) + H₃O⁺,  

or it can act as a Brønsted acid and donate a proton:

A⁻ (or B) + H₂O(acid) ⇌ HA(BH⁺) + OH⁻.  

From Equation 4, it is clear that the hydronium ion (also known as a hydrated/solvated proton), H₃O⁺, is the strongest acid that can exist in aqueous media. In other words, the process shown in Equation 4 effectively levels the acidity of all strong acids in aqueous media and is the origin of the aqueous pKₐ scale (45, 46), where pKₐ = -log₁₀Kₐ.

Many organic compounds contain acidic and/or basic functional groups that dictate their physical, chemical, and biological properties. Indeed, bond transformations in solution frequently involve the cleavage or formation of R–H bonds. As described above, pKₐ describes the free energy change of R–H bond heterolysis, making it a critically important thermodynamic parameter in synthetic chemistry and related fields (47–49). Equilibrium acidities (pKₐ) are media dependent, being dictated by the ability of the solvent to solvate each of the species outlined in Equation 2 (i.e., HA, A⁻, and H⁺) or Equation 3 (i.e., BH⁺, B, and H⁺). It follows that solvent acidity/basicity, dielectric properties, and the ability to donate/accept hydrogen bonds can all influence the pKₐ value of an acid in solution (49, 50). Although pKₐ data are most readily available in water (51, 52), pKₐ scales have been established in a range of nonaqueous solvents, including acetonitrile (53), dimethylsulfoxide (49, 54, 55), and 1,2-dichloroethane (53). A variety of methods (56) are typically employed to quantify pKₐ in conventional media, including potentiometry (57), spectrophotometry (49, 58), and voltammetry (50, 54, 55).

The protons released from the dissociation of a Brønsted acid in IL media must associate with (or be solvated by) the most basic component of the IL, most commonly the anion (A⁻Ⅱₐ):

HA + A⁻Ⅱₐ ⇌ A⁻ + HAⅡₐ,  

where HAⅡₐ is the strongest acid that may exist in a given IL. HAⅡₐ effectively levels the acidity of strong acids in IL media, comparable to H₃O⁺ in aqueous media. In other words, HAⅡₐ is the origin of the pKₐ scale in IL media, and for this reason, pKₐ data are not directly comparable for ILs with different constituent anions (45, 47, 48, 59–61). It should be noted that from this point
forward, H\(^+\) refers to the solvated proton species and is equivalent to HA\(_{\text{IL}}\) in the context of ILs or H\(_3\)O\(^+\) in the context of aqueous media.

### 2.2. The \(pK_a\) Scale in Ionic Liquids

Although electrochemical (voltammetric) studies are the main focus herein, the recent comprehensive work by Cheng and coworkers should be acknowledged (47–63). Using a spectrophotometric (overlapping indicator) method, these authors have established the equilibrium acidity of over 75 different acids (e.g., carbon acids, oxyacids, protonated amines, and sulfonamides) in a range of ILs. In all studies, \(pK_a\) (IL) data have been correlated to corresponding \(pK_a\) data available in molecular solvents (e.g., water, dimethylsulfoxide, and acetonitrile) and the gas phase (where available), to draw conclusions about the relative solvating properties (i.e., Lewis acidity/basicity) of ILs. In addition, in all studies it is noted that the identity of both the IL anion, and to a lesser extent, the cation was found to influence the \(pK_a\) of a given acid, as discussed further below.

The Angell group (24) first explored the concept of \(pK_a\) (proton thermodynamics) in the electrochemical context by studying hydrogen redox chemistry (i.e., the H\(^+\)/H\(_2\) process, described below) at a platinum (Pt) electrode in 10 PILs of widely varying proton transfer energies. A significant gap (ca. 0–800 mV) between the onset potential at which the hydrogen evolution reaction (HER) occurs and the potential at which the hydrogen oxidation reaction (HOR) occurs facilely was observed. This difference occurs because hydrogen oxidation involves the HA/H\(_2\) couple, whereas proton reduction involves the BH\(^+\)/H\(_2\) couple. It follows that the potential gap separating these two processes is proportional to the energetics for proton extraction from the protonated anion (i.e., HA in Equation 1) relative to the cation (i.e., BH\(^+\) in Equation 1), which is defined by the differences in proton free energy between the Brønsted couples HA/A\(^-\) and BH\(^+\)/B. Hence, the potential difference between the HA/H\(_2\) (HOR) and BH\(^+\)/H\(_2\) (HER) can be correlated to the difference between the aqueous \(pK_a\) values of HA and BH\(^+\) (i.e., \(\Delta pK_a\)), suggesting that aqueous \(\Delta pK_a\) data provide a good approximation to proton activity in PILs.

Kanzaki and coworkers (64, 65) recently established a \(pK_a\) scale (14 \(pK_a\) values from 12 compounds) in the neat PIL ethylammonium nitrate (EAN), which was determined using potentiometric titration. \(pK_a\) (EAN) values were consistently one unit greater than the corresponding value in water, regardless of the structure, charge (neutral, cationic, and anionic
acids were investigated), and/or hydrophobicity of the compounds. The authors attributed this to the stronger acidity of HNO₃ (i.e., HA_IL or H⁺; see Equation 6) in EAN than that of H₃O⁺ in water, rather than a difference in the solvation state of the ions. In addition, assuming that pKₐ(EAN) = pKₐ(H₂O) + 1 always applies, and from the autoprotolysis constant of EAN [pKₐp = 9.83, determined in a previous study (66)], it was concluded EAN is a solvent with stronger acidity and weaker basicity (approximately 10⁵ times weaker) than water. Furthermore, although not based on thermodynamically rigorous arguments, it was estimated that the pH scale in EAN lies in the range of −1 to 9 on the water-based pH scale, again suggesting that EAN is an acidic solvent relative to aqueous media.

The pKₐ values of weak acids dissolved in an IL can be conveniently calculated using dynamic electrochemical methods such as cyclic voltammetry from the following relationship (45, 50, 54, 55, 67):

\[
E^0'(BH^+ / H_2) - E^0'(H^+ / H_2) = \frac{RT}{F} \ln \frac{[B][H^+]}{[BH^+]} = -\frac{2.303RT}{F} pK_a.
\]

where \(E^0'(H^+/H_2)\) and \(E^0'(BH^+/H_2)\) are the formal reversible potentials for the \(H^+/H_2\) and \(BH^+/H_2\) processes (discussed below), respectively. \(R\) is the universal gas constant, \(T\) is the absolute temperature, and \(F\) is Faraday’s constant. Indeed, this method was used by Barhdadi et al. (45) to estimate the pKₐ value of the pyridinium cation in a range of ILs, where they reported a strong dependence on the identity of the IL anion. In their study, it was assumed that \(E_{1/2} \approx E^0\), where \(E_{1/2}\) is the reversible half-wave potential, introduced originally as a polarographic term, but in the present context taken to be the average of the reductive \(E_{p, red}\) and oxidative \(E_{p, ox}\) peak potentials of a reversible transient cyclic voltammogram (the midpoint potential, \(E_{mid}\)) (55, 68). Although this approximation is valid for simple (unimolecular) reactions in conventional media (equations describing the relationship between \(E^0\), \(E_{1/2}\) and \(E_{mid}\) can be found in 67, 68), the bimolecular nature of the \(H^+/H_2\) process and grossly different diffusion coefficients of \(H^+\) and \(H_2\) in IL media mean that this approximation introduces considerable systematic error into the determination of pKₐ (vide infra).

Building on this work, our group (59–61) reported a series of studies focusing on the energetics (thermodynamics) of proton transfer in IL media, predominantly AILs, which were probed using electrochemical (voltammetric) methods. Pt was selected as the electrode, as the
H⁺/H₂ and BH⁺/H₂ (or HA/H₂) processes are well defined and chemically reversible on this material. This allows for the accurate estimation of \( E_{1/2} \), which can be used to derive \( pK_a \) values, as described in detail below. In the first study, the nature of the solvated proton species (i.e., HA_IL; see Equation 6) was investigated in a range of AILs: \([C_2mim]⁺, [C_4mpyr]⁺, [S_{2,2,2}]⁺, [C_4dmim]⁺, [N_{1,1,2,3}]⁺, \) and \([N_{1,4,4,4}]⁺\), each with the common \([NTf_2]⁻ \) anion (H[NTf₂] was the proton source). Through correlation of the electrochemically derived proton diffusion coefficient [calculated using the convolution method (69–71) or through chronoamperometry (72)] with the self-diffusion coefficient of \([NTf_2]⁻ \) (derived using pulsed field gradient spin-echo nuclear magnetic resonance), it was unequivocally shown that H⁺ remains associated with the anion part of the IL, and the strongest acid that can possibly exist in an \([NTf_2]⁻ \) IL is therefore H[NTf₂] (45). In other words, H[NTf₂] is the origin of the \( pK_a \) scale in \([NTf_2]⁻ \) media, in accordance with the concepts outlined above.

This initial work shows that identity of the cation has very little influence on \( E^{0'}(H^+/H_2) \), which is a measure of the energetics of proton transfer in this medium, with the H⁺/H₂ process occurring at approximately –0.030 V versus ferrocene/ferrocinium (Fc/Fc⁺) in all investigated [NTf₂]⁻ AILs [it was assumed that the Fc/Fc⁺ couple possesses a solvent-independent formal potential (73–75)]. In a subsequent study (60), it was shown that \( E^{0'}(H^+/H_2) \) is strongly governed by the anion of the IL. For example, a value of approximately –0.34 V versus Fc/Fc⁺ was obtained in \([C_2mim][OTf] \) [H⁺ was from either H[OTf] or H[NTf₂], both strong acids in this media]. In contrast, two proton reduction processes were found in the PIL \([N_{H,2,2,2}][NTf_2] \); one occurred at approximately –0.030 V versus Fc/Fc⁺, corresponding to the H[NTf₂]/H₂ process (as above for the AILs), and one occurred at approximately –0.9 V versus Fc/Fc⁺, originating from the bulk reduction of the protonated PIL cation [i.e., the \((N_{H,2,2,2})⁺/H_2 \) process].

In later studies (60, 61), the \( pK_a \) of a range of NH [protonated amines and sulfonamides (60)] and OH [phenols, carboxylic acids and sulfonic acids (61)] acids were quantified in \([C_2mim][NTf_2] \). Building on the theory reported by Barhdadi et al. (45), the following relationships between \( E^{0'} \) and \( E_{1/2} \) were derived using the diffusion layer approximation method (55, 67, 76, 77):

\[
E^{0'}(H^+/H_2) = E_{1/2}(H^+/H_2) - \frac{RT}{2F} \ln \left( \frac{D_{H_2}}{\sqrt{D_{H^+}}} \right) - \frac{RT}{2F} \ln [H^+],
\]

8.
\[ E^{th} (\text{BH}^+ / H_2) = E_{1/2} (\text{BH}^+ / H_2) - \frac{RT}{2F} \ln \left( \frac{4 \sqrt{D_H}}{\sqrt{D_{\text{BH}^+}}} \right) + \frac{RT}{2F} \ln \left[ \text{BH}^+ \right], \quad 9. \]

where \( D \) is the diffusivity (diffusion coefficient). Combining Equations 7 to 9 gives

\[ - \frac{2.303RT}{F} pK_a = E_{1/2} (\text{BH}^+ / H_2) - E_{1/2} (\text{H}^+ / H_2) + \frac{RT}{2F} \ln \left( \frac{\sqrt{D_{\text{BH}^+}}}{D_{\text{H}^+}} \right) + \frac{RT}{2F} \ln \left[ \frac{(\text{BH}^+)(\text{H}^+)}{4} \right]. \quad 10. \]

Note that in the derivation of Equations 9 and 10, it was assumed that \( D_B = D_{\text{BH}^+} \). As alluded to above, assuming that \( E_{1/2} \approx E^{th} \) neglects the two logarithmic terms shown in Equation 10, which leads to considerable systematic error (approximately 3 \( pK_a \) units when working at millimolar concentrations) in the determination of \( pK_a \). As highlighted above, \( E_{1/2} \) values can be directly estimated from a cyclic or derivative cyclic voltammogram from the average of \( E_{p,\text{red}} \) and \( E_{p,\text{ox}} \) (i.e., \( E_{\text{mid}} \)), and \( D \) values can be derived electrochemically using the convolution method (69–71) or chronoamperometry (72). Indeed, it was demonstrated how all of the parameters required to calculate \( pK_a \) from Equation 10 could be derived from an IL solution containing a 2:1 mixture of \( \text{H}^+ \) and \( \text{B} \), as illustrated in Figure 2 for \( \text{H}[\text{NTf}_2] \) and pyridine in \([\text{C}_2\text{mim}][\text{NTf}_2]\). It was also shown how the homoassociation constant \( (K_{\text{homo}}) \) could be derived voltammetrically for the carboxylic and sulfonic acids (vide infra) (61).

Figure 2 (a) A cyclic voltammogram obtained from the reduction of 57.4 mM \( \text{H}[\text{NTf}_2] \) and 24.3 mM pyridine in \([\text{C}_2\text{mim}][\text{NTf}_2]\) at a 1.6-mm-diameter Pt macrodisk electrode with a scan rate of 100 mV s\(^{-1}\). The \( E_{1/2} \) values used in Equation 10 are calculated as follows: \( E_{1/2} \approx E_{\text{mid}} = (E_{p,\text{ox}} + E_{p,\text{red}})/2 \). (b) Experimental (lines) and Shoup-Szabo theoretical chronoamperograms (circles) obtained from the solution used in panel a at a 20-µm diameter Pt microdisk electrode by (i) stepping the potential to –0.323 V versus Fc/Fc\(^+\) (blue trace), (ii) stepping the potential to –1.023 V versus Fc/Fc\(^+\) (purple trace), and (iii) subtracting curve i from curve ii (red trace). The \( D \) values used in Equation 10 are obtained from the Shoup-Szabo fitting procedure. Abbreviation: Fc/Fc\(^+\), ferrocene/ferrocenium. Adapted with permission from Reference 60. Copyright 2015, American Chemical Society.

Using the approach outlined above, the \( pK_a \) values of 20 weak acids, covering 18 orders of magnitude in acid strength (2.0 ≤ \( pK_a \) ≤ 19.5), were calculated and compared with data from
conventional solvents (acetonitrile, AN, and water, shown in Figure 3a and 3b, respectively), to gain insight into how the nature of the solvent [i.e., dielectric properties, Lewis acidity/basicity, hydrogen donating/accepting ability, etc. (18, 78)] influences equilibrium acidity. As shown in Figure 3a, with the exception of H[OTf], the pKₐ(IL) data correlate well with pKₐ(AN), with pKₐ values consistently 1–2 units lower in [C₂mim]NTf₂ compared to AN. On the other hand, as shown in Figure 3b, considering all of the acids in a single group, pKₐ(IL) data correlate poorly with pKₐ(H₂O), although pKₐ values in [C₂mim][NTf₂] are consistently several units higher than in H₂O, owing to the exceptionally high dielectric constant and Gutmann donor/acceptor numbers of the latter solvent. Dividing the acids into two groups, cationic and neutral, makes it clear that there is good correlation between pKₐ(aq) and pKₐ(IL) for acids of like charge (i.e., cationic versus cationic or neutral versus neutral) but not for acids of different charge (i.e., cationic versus neutral). This phenomenon was explained by the well-known preferential solvation of the water molecule toward oxyanions through hydrogen bonding (i.e., the acidity of neutral oxyacids are enhanced relative to protonated amines in aqueous media), which is absent in the neat, poorly hydrogen-bonding AIL. Finally, it was shown how the nature of the IL can have a significant influence on the acid/base properties of a given weak acid; taking tosylic acid as an example, it was shown that switching the cationic constituent from [C₂mim]+, a poor hydrogen bond donor, to [NH₂,2,2,2]+, a good hydrogen bond donor, suppresses anionic homoassociation (discussed below) and increases the acidity of this weak acid a thousandfold (i.e., pKₐ decreases from 6.2 to 3.3) (60, 61).

**Figure 3** (a) Plot of pKₐ(AN) versus pKₐ(IL) for a range of weak neutral acids. (b) Plot of pKₐ(aq) versus pKₐ(IL) for a range of nitrogen (triangles) and oxygen (squares) acids. Regression lines for the uncharged (red) and charged (purple) acids have been included solely to guide the reader’s eye. H[OTf] not considered when calculating the regression line in panel a. Abbreviations: AN, acetonitrile; An, aniline; aq, aqueous media; ClPyr, 2-chloropyridine; DBSA, di[bensensulfonyl]imide; DCAA, dichloroacetic acid; dClPyr, 2,4-dichloropyridine; GA, glycolic acid; H[OTf], triflic acid; H[OTs], tosylic acid; IL, ionic liquid; MA, malonic acid; Melm, N-methylimidazole; MePyrd, N-methylpyrrolidine; MSA, methanesulfonic acid; oHBA, o-hydroxybenzoic acid; oPD, o-phenylenediamine; PCP, pentachlorophenol; Pyr, pyridine; SACC, saccharin; TCAA, trichloroacetic acid. Adapted with permission from Reference 61. Copyright 2015, American Chemical Society.
3. THE H⁺/H₂ PROCESS IN IONIC LIQUIDS

The HER and HOR are technologically important processes (e.g., in electrolyzers and fuel cells, respectively) that have been characterized on many electrode materials in a range of media (79–81), predominantly aqueous (76, 82, 83):

\[ 2\text{H}^+ + 2e^- \rightarrow \text{H}_2. \]  \hspace{1cm} 11.

In addition, the fundamental reference potential scale in aqueous electrochemistry, the standard hydrogen electrode, is based on the H⁺/H₂ couple (67). Although Equation 11 is an apparently simple one electron per proton process, it is subject to significant kinetic barriers, necessitating the use of an electrocatalyst in practical (industrial) applications. It follows that different metal electrodes are well known to exhibit vastly different catalytic activities toward the HER/HOR in aqueous media (84, 85).

As mentioned above, ILs—particularly PILs—have attracted significant interest as anhydrous proton conductors for application in nonhumidified intermediate temperature fuel cells (12, 27, 31). In the past decade, numerous studies have been dedicated to identifying PIL formulations that display fast proton transport in addition to facile HOR and oxygen reduction reaction (ORR) kinetics (12, 22, 24, 25, 39–41). These studies have been addressed in a number of recent comprehensive reviews (12, 26, 42). Application-driven studies on the HER/HOR in IL media are not discussed here. Rather, recent fundamental studies on the mechanistic aspects of the H⁺/H₂ process in IL media are reviewed.

3.1. The Hydrogen Oxidation Reaction

The HOR is postulated to proceed via the Tafel-Heyrovsky-Volmer route in acidic aqueous media (80, 82, 86):

\[ \text{H}_2(\text{aq}) \rightarrow 2\text{H}_{\text{ads}}, \]  \hspace{1cm} 12.

\[ \text{H}_2(\text{aq}) \rightarrow \text{H}_{\text{ads}} + \text{H}^+(\text{aq}) + e^- , \]  \hspace{1cm} 13.

\[ \text{H}_{\text{ads}} \rightarrow \text{H}^+(\text{aq}) + e^- , \]  \hspace{1cm} 14.

where \( \text{H}_{\text{ads}} \) is a chemisorbed hydrogen atom. Thus, the formation of \( \text{H}_{\text{ads}} \) through dissociative chemisorption (Tafel reaction; Equation 12) or dissociative chemisorption and ionization to H⁺
The Heyrovsky reaction; \textit{Equation 13} is followed by the oxidation of $H_{\text{ads}}$ (Volmer reaction; \textit{Equation 14}) to form $H^*$. Compton and coworkers \cite{87, 88} performed the pioneering work on the mechanistic aspects of the HOR in IL media. In their first reports, the HOR was investigated in a range of AILs. On Pt, a broad (i.e., large cyclic voltammetric peak-to-peak separation, $\Delta E_p$) but chemically reversible process was observed in the $[\text{NTf}_2]^-$ and $[\text{OTf}]^-$ ILs, which the authors attributed to the following process:

$$H_2 - 2e^- + 2A_{\text{IL}}^- \rightleftharpoons 2HA_{\text{IL}}, \quad 15.$$  

The HOR is inhibited at strongly oxidative potentials [also observed in the study by Angell and coworkers \cite{24}, described above], which is attributed to blocking of the catalytically active sites on the Pt electrode surface by a platinum oxide ($\text{PtO}_x$) layer (i.e., the electrode is passivated), formed due to the presence of residual water in the IL. It was also shown that if the Pt electrode is preanodized at strongly oxidizing potentials, the kinetics of the HOR are greatly increased in the potential range where the surface oxide is not present (evidenced by a decreased $\Delta E_p$), presumably due to the formation of a catalytically active pristine Pt surface after $\text{PtO}_x$ formation/dissolution. In $[\text{BF}_4]^-$, $[\text{PF}_6]^-$, $\text{Cl}^-$, and $[\text{NO}_3]^-$ ILs, the HOR is only partially chemically reversible, which the authors attributed to instability of the protonated anion (formed during the HOR; see \textit{Equation 15}). In the case of $[\text{PF}_6]^-$ and $[\text{BF}_4]^-$, it is thought that the anion undergoes acid-catalyzed dissociation (decomposition) on the voltammetric timescale, for example:

$$H[\text{BF}_4] \rightarrow \text{HF} + \text{BF}_3. \quad 16.$$  

In the case of $\text{Cl}^-$, it is thought that $\text{HCl}$ (formed from hydrogen oxidation; see \textit{Equation 15}) reacts to form stable $H[\text{Cl}_2]^-$, as highlighted in a first principles simulation study by Del Popolo et al. \cite{89}. An analogous mechanism was also proposed for $[\text{NO}_3]^-$.

The identity of the IL anion, and to a lesser degree the IL cation, can have a significant influence on the kinetics of the HER/HOR, as proposed by Navarro-Suárez and coworkers \cite{90}, who investigated the HOR at Pt (100), Pt (110) and Pt (111) macroelectrodes (flame annealed, low-index Pt single crystals) in a range of imidazolium-based AILs. The activity toward the HOR (assessed from the onset potential of the HOR) was shown to increase in the order of Pt
(100) < Pt (110) < Pt (111) in [C₄mim][OTf], [C₅mim][EtSO₄], and [C₄mim][BF₄], whereas Pt (100) was found to be the most active surface in [C₂mim][NTf₂]. The variations in electrocatalytic activity of low-index Pt crystal surfaces were attributed to differences in anion adsorption at oxidative potentials (specifically, weaker adsorption at more densely packed surfaces leads to higher activity). In a recent study, Tang et al. (91) proposed that the HOR is catalyzed by the formation of Pt-bound NTf₂⁻ radicals in deaerated [C₄mpyr][NTf₂], a conclusion supported by experimental data (cyclic voltammetry) and density functional theory calculations.

The Walsh group (92) investigated the HOR at a polycrystalline Pt rotating disk electrode in the PIL [N₁H₁,2,2][OTf]. This reaction was found to become hindered at highly positive (oxidizing) potentials. This feature was shown by X-ray photoelectron spectroscopy to be attributable to the formation of a blocking PtO₃ layer, effectively passivating the electrode to the electrocatalytic process, which is in agreement with the earlier work by Angell’s group (24) and Compton’s group (87, 88). In a follow-up study (93), the role of adsorption in the electrocatalytic HER was investigated in a more extensive range of PILs, [N₁H₁,2,2][OTf], [N₁H₁,1,2][OTf], and [N₁H₁,2,2][NTf₂]. Voltammetric analysis revealed that a monolayer of hydrogen adsorbs (H_ads; see Equations 12–14) onto Pt during potential cycling in the [OTf]⁻ PILs but not the [NTf₂]⁻ one. The authors attributed the suppression of H_ads formation in the latter class of PIL to the adsorption of the [NTf₂]⁻ anion onto the electrode surface at relatively low oxidative potentials (i.e., blockage of catalytic sites by adsorbed [NTf₂]⁻).

The Walsh group (94) has also probed the HOR in [N₁H₁,2,2][OTf] under high mass transport conditions using hydrodynamic and ultramicroelectrode voltammetry. Whereas only a single, mass transport–limited HOR plateau is observed at a Pt rotating disk electrode, an additional plateau was detected when using a Pt ultramicroelectrode (i.e., the voltammetric wave is split into two processes), which resulted from the higher rates of mass transport achievable at the latter type of electrode. On this basis, three models were proposed to fit the experimental data. Two are based on the coexistence of dual reaction pathways (e.g., Volmer-Tafel versus Heyrovsky-Volmer are dominant in different potential regions; see Equations 12 to 14), and a third is based on the premise that underpotential deposited hydrogen atoms (H_UPD) can block the further adsorption and oxidation of H₂ at the Pt surface. Despite reasonably satisfactory theory-experimental fits for all three models, the H_UPD-blocking model best described the responses.
leading to the conclusion that H_{UPD} can act as a blocking species during electrocatalytic reactions in PIL media.

### 3.2. The Hydrogen Evolution Reaction

The HER is postulated to proceed via the Volmer reaction (Equation 17), followed by either the Tafel (Equation 18) or Heyrovsky (Equation 19) reaction in acidic aqueous media (80, 82, 86):

\[
H^+_{(aq)} + e^- \rightarrow H_{ads}, \quad 17.
\]

\[
2H_{ads} \rightarrow H_2_{(aq)}, \quad 18.
\]

\[
H_{ads} + H^+_{(aq)} + e^- \rightarrow H_2_{(aq)}, \quad 19.
\]

Thus, the formation of H_{ads} on the electrode surface by electron transfer (reduction) is followed by either surface migration and dimerization of H_{ads} to form H_2 or the direct formation of H_2 from H_{ads} and H^+ (solution) by electron transfer. The preferred reaction pathway depends on the identity of the metallic electrode and is thought to be governed by whether Equation 17 is sluggish or facile relative to H_{ads} migration across the electrode surface (80).

The Compton group (95) has investigated the HER from H[NTf_2] or HCl as the proton source in [C_4mim][NTf_2]. A chemically reversible proton reduction process is observed on Pt, which is thought to proceed via the combination of reactions outlined in Equations 17 to 19. On Au, the HER occurs (chemically) irreversibly at potentials negative of those on Pt, whereas on glassy carbon (GC), an irreversible, broad reduction process is observed at even more negative potentials and just prior to solvent breakdown. These results are in accordance with the relative order of hydrogen overpotentials of the electrode materials in aqueous media: GC >> Au > Pt.

On Pt, proton reduction from HCl is chemically reversible, occurring at potentials negative of the H[NTf_2]/H_2 process, in accordance with the relative acidities of HCl and H[NTf_2] (vide supra). These authors also postulate that H[Cl_2]^- forms as an intermediate during proton reduction at potentials where the concentrations of both HCl and Cl^- are significant.

In later studies, the Compton group (80, 96) investigated proton reduction (from H[NTf_2] as the proton source) on a range of metallic electrodes (Au, Mo, Ni, Ti, and Pt) in [C_2mim][NTf_2] [and with an expanded range of ILs in a follow-up study (81)]. Chemically and electrochemically irreversible proton reduction was observed on all of the metal surfaces except Pt, with no H_2
process being observed on the reverse cyclic voltammetric sweep within the investigated potential range. The charge-transfer coefficient ($\alpha$) was found to be $\leq 0.5$ on all metal surfaces (determined through mass transport corrected Tafel slope analysis), indicating that the Volmer reaction (Equation 17) is the rate-determining step in all cases. Finally, the heterogeneous kinetics of the HER (estimated by numerical simulation) was found to be strongly dependent on the identity of the metallic electrode, with the electrocatalytic trend being Pt > Mo > Au > Ni > Ti, which is different to that established in aqueous media: Pt > Ni > Au > Mo > Ti.

Compton and coworkers (97–99) have also investigated the proton reduction process from the weak aqueous acid, benzoic acid, on a Pt electrode in a range of ILs. Proton reduction from benzoic acid (i.e., the HA/H$_2$ process) occurred at potentials negative of the H$^+/H_2$ process (i.e., solvated proton reduction) and was postulated to proceed via a generic CE mechanism, where the electron transfer step (E; see Equations 11, 17–19) is preceded by a fast chemical (dissociation) step (C):

$$C_6H_5COOH \rightarrow C_6H_5COO^- + H^+. \quad 20.$$

A reductive prepeak was observed prior to the main reduction peak in all ILs examined, which was assigned to the formation of adsorbed H$^\bullet$ on the Pt electrode surface. The potential region in which proton reduction from benzoic acid occurs was also found to be dependent on the identity of the IL anion, consistent with earlier reports (81, 87, 88).

Our group (59) recently investigated the mechanism of the HER with H[NTf$_2$] as the proton source in a range of AILs ([C$_2$ mim]$^+$, [C$_4$ mpym]$^+$, [S$_{2,2,2}$]$_2^+$, [C$_{4}$dmim]$^+$, [N$_{1,1,2,3}$]$^+$, [N$_{1,4,4,4}$]$^+$, each with the common [NTf$_2$]$^-$ anion) by combining experimental cyclic voltammetric data at a Pt electrode with that from computational simulations. In this investigation, it was found that the voltammetric response was very sensitive to the history of the Pt electrode (i.e., cleanliness/surface state). As shown in Figure 4a, after the electrode has been freshly polished (cycle 1), a well-defined reductive response is obtained, which noticeably degrades on cycling the potential (or by leaving the electrode submerged in the IL at the open circuit potential), becoming flattened and less peak shaped (i.e., the electrode becomes deactivated). As demonstrated in Figure 4b, the initial voltammetric response can be recovered by mechanically polishing the electrode or by electrochemical conditioning at oxidative potentials (anodic polarization). In other words, when the oxidative limit is extended to 2.2 V (versus Fe/Fe$^+$), the
initially degraded response improves with cycling, indicating that the electrode is activated at oxidative potentials [previously reported in both conventional solvents (79) and other ILs (88)]. In a later study, it was shown that the degradation of the voltammetric HER response occurred more rapidly in [C2mim][OTf], suggesting that electrode deactivation is intrinsically linked to the IL anion (60).

**Figure 4** Cyclic voltammograms obtained from the hydrogen evolution reaction (proton source = H[NTf2]) in [C2mim][NTf2] at a 20-µm-diameter Pt microdisk electrode. (a) Ten cycles (ν = 100 mV s⁻¹) in the potential range 0.41 to −0.59 V (versus Fc/Fc⁺) demonstrate progressive deactivation of the electrode. (b) Four cycles (ν = 100 mV s⁻¹) in the potential range of 2.21 to −0.59 V (versus Fc/Fc⁺) demonstrate the activation of the electrode with anodic polarization. (c) Cyclic voltammograms obtained with [H⁺] = 99.2 mM and ν = 50, 100, 250, and 500 mV s⁻¹. (d) Cyclic voltammograms obtained with ν = 100 mV s⁻¹ and [H⁺] = 25.4, 44.5, 75.8, and 99.2 mM. Simulated cyclic voltammograms (circles) are also shown in panels c and d, obtained by combining Equations 22 and 23 with the following parameters:  

```
E₀(H⁺/H₂*) = −0.148 V, kₜ = 0.022 cm s⁻¹, α = 0.50 (Equation 22, apparent values); Kdim = 10⁸, kdim = 10¹⁶ M⁻¹ s⁻¹ (Equation 23); D_H⁺ = 3.1 × 10⁻⁷ cm² s⁻¹, D_H₂ = 2.2 × 10⁻⁵ cm² s⁻¹. E₀'(H⁺/H₂) = −0.030 V, derived from E₀(H⁺/H⁺) and Kdim. Abbreviations: Fc/Fc⁺, ferrocene/ferrocenium; IL, ionic liquid; PtOₓ, platinum oxide; UPD, underpotential deposited. Adapted with permission from Reference 59. Copyright 2014, American Chemical Society.
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**Figure 4c,d** shows representative cyclic voltammograms obtained at a 20-µm-diameter Pt microdisk electrode, which had previously been activated by stepping the potential to ≈2 V versus Fc/Fc⁺ for ≤ 1 ms. D_H⁺ (~10⁻⁷ cm² s⁻¹) and D_H₂ (~10⁻⁵ cm² s⁻¹) are hundredfold different in this medium, and as a result, the reduction (HER) and oxidation (HOR) processes are peak shaped (mass transport due to both planar and radial diffusion) and sigmoidal (mass-transport predominantly due to radial diffusion), respectively. The underpotential deposition of adsorbed hydrogen on the electrode is observed prior to the main reduction process (labeled in **Figure 4c,d**), consistent with the behavior of Pt surfaces in aqueous acidic media (82).

Also shown in Figure 4c,d are simulations of the voltammetric response. Although in reality, adsorption is thought to play an important role in the HER and HOR (80, 82, 86), for simplicity, the H⁺/H₂ reaction has been treated as a purely solution based process, simulated using the DigElch electrochemical simulation software (http://www.elchsoft.com/digielch/DigiElch7/).
A single step process, using the bimolecular capability of DigiElch (version 7) was initially considered:

\[ \text{H}^+ + \text{H}^+ + 2e^- \rightarrow \text{H}_2. \quad 21. \]

However, this was not found to emulate the experimental data over a wide range of conditions, requiring a different set of parameters for each investigated concentration or scan rate. Building on this, two mechanisms were considered, based on the Volmer-Tafel (Equations 17 and 18) and Volmer-Heyrovsky (Equations 17 and 19) routes, both of which were both found to give excellent agreement with the experimental data:

\[ \text{H}^+ + e^- \rightarrow \text{H}^+ \quad E^0(\text{H}^+/\text{H}^+)\), k_s, \alpha, \quad 22. \]

\[ \text{H}^+ + \text{H}^+ \rightarrow \text{H}_2 \quad K_{\text{dim}}, k_{\text{dim}}, \quad 23. \]

\[ \text{H}^+ + \text{H}^+ + e^- \rightarrow \text{H}_2 \quad E^0_{2}, k_{s,2}, \alpha_2, \quad 24. \]

where \( E^0 \), \( k_s \), and \( \alpha \) are the formal potential, standard heterogeneous electron-transfer rate constant at \( E^0 \) and charge transfer coefficient, respectively. As the Volmer-Tafel (Equations 22 and 23) route is the simpler mechanism from a simulation perspective (54, 55, 59, 60), it is solely considered herein. As discussed above, in reality, \( \text{H}^+ \) is likely to be a surface-confined species (82, 83), although for simplicity it has been treated as a solution-based (diffusing) species in the simulations. One consequence of treatment of the mechanism in this manner is that the process shown in Equation 23 can be regarded as empirically based, as are the parameters \( k_s \), \( \alpha \), \( K_{\text{dim}} \), \( k_{\text{dim}} \), and \( D_{\text{H}^+} \) (i.e., they are not physically meaningful) (54). Furthermore, it should be emphasized that the kinetic parameters, \( k_s \), and \( \alpha \), are apparent values and therefore should not be interpreted quantitatively.

Despite the use of a significantly simplified mechanism, the simulations are in excellent agreement with the experimental data in all investigated ILs over a wide range of scan rates (Figure 4c) and concentrations (Figure 4d). Regardless of the mechanism employed (i.e., Equations 22 and 23 in Figure 4 or Equations 22 and 24 in 59), the first electron transfer step (Volmer reaction, Equation 17) is assumed to be rate determining on Pt (i.e., \( k_{\text{dim}} \), or \( k_{s,2} \) were set to be fully reversible or nonlimiting on the voltammetric timescale), in accordance with the work by Compton and coworkers (80, 81, 96). The identity of the IL cation was found to have a minor
influence on $k_s$ (59), in contrast to a later study, where it was shown that the IL anion can significantly influence the kinetics of the HER at Pt (60), in agreement with the discussions above (90).

A single PIL, [N$_{H,2,2,2}$][NTf$_2$], was also investigated, and the HER mechanism (from H[NTf$_2$]) was found to be analogous to the AILs with respect to computational simulations. This is unsurprising, considering that the acidic constituent of the PIL, H[NTf$_2$], is a superacid and therefore, complete proton transfer to the basic constituent, [N$_{2,2,2}$], would be expected (23). As a result, the proton reduction is solely due to the H[NTf$_2$] introduced into the neat [N$_{H,2,2,2}$][NTf$_2$] (i.e., the H$^+$/H$_2$ process). The proton on the amine, [N$_{H,2,2,2}$]$^+$, can also be reduced. However, this species is a significantly weaker acid than H[NTf$_2$], and therefore, much more negative potentials (approximately $\approx$–0.9 V) must be reached before bulk proton reduction is observed (100).

The thermodynamics, kinetics, and mechanisms of the HER at a Pt electrode were also investigated in [C$_2$ mim][NTf$_2$] using range of nitrogen (R$_n$NH) acids (protonated amines or sulfonamides) as the proton source (60). Hydrogen evolution from weak nitrogen acids in the BH$^+$/H$_2$ process (the overall reaction shown in Equation 25) is a diffusion-controlled process that occurs in a potential region negative of the H$^+$/H$_2$ process, as shown in Figure 5a:

$$2\text{BH}^+ + 2\text{e}^- \rightarrow \text{H}_2 + 2B.$$  \[25\]

The $E^0'$ values of the investigated BH$^+$ acids span a potential window of approximately 0.9 V, representing a $pK_a$ range of approximately 5.3 to 19.5 (derived as in Equation 10). Interestingly, the $\Delta E_p$ value, which is a qualitative indicator of electron-transfer kinetics, was found to increase with decreasing $pK_a$ (compare 2-chloropyridinium and N-methylpyrrolidinium in Figure 5a) and increasing acid charge (compare diprotonated and monoprotonated o-phenylenediaminium in Figure 5a). In addition, $\Delta E_p$ was found to differ significantly when comparing structurally disparate acids (protonated amines versus sulfonamides) of comparable strengths (compare pyridinium and saccharin in Figure 5a). Although the exact reason for these observations is not known, it was speculated that because Equation 22 is the rate determining step, the activation energy required to break the N–H bond is related to the heterolytic bond dissociation energy (i.e., $pK_a$) and structural factors (i.e., charge and chemical nature of the covalent bond).

<COMP: PLEASE INSERT FIGURE 5 HERE>
**Figure 5** (a) Cyclic voltammograms (normalized to unity with the reduction peak current, \(I_{p,\text{red}}\)) showing the hydrogen evolution reaction (HER), with the following proton sources (top to bottom): H[NTf₂], α-phenylenediaminum (diprotonated), 2,4-dichloropyridinium, 2-chloropyridinium, di(benzenesulfonyl)amide, anilinium, α-phenylenediaminum (monoprotonated), saccharin, pyridinium, \(N\)-methylimidazolium, and \(N\)-methylpyrroolidinium. The pKₐ values calculated using Equation 10 are indicated in the figure. (b) Comparison of simulated (red and blue lines) and experimental (black line) cyclic voltammograms showing the HER, with 38.7 mM 2,4-dichloropyridinium as the proton source. The simulations were carried out using the CE mechanism only (Equations 22, 23, and 26; red line) or the CE + DR mechanism (Equations 22, 23, 26, and 27). Simulation parameters are as follows: \(E^0(\text{H}^+/\text{H}^+) = -0.148\) V, \(k_s = 0.022\) cm s⁻¹, \(\alpha = 0.50\) (Equation 22, apparent values); \(k_{\text{dim}} = 10^4\), \(k_{\text{dim}} = 10^{16}\) M⁻¹ s⁻¹ (Equation 23); \(E^0(\text{BH}^+/\text{H}^+) = -0.513\) V versus Fc/Fc⁺ (pKₐ = 6.2), \(k_{\text{assoc}} = 5 \times 10^8\) M⁻¹ s⁻¹ (Equation 26); \(k_{\text{DR}} = 0.058\) cm s⁻¹, \(\alpha_{\text{DR}} = 0.50\) (Equation 27, apparent values); \(D_{\text{H}^+} = 3.1 \times 10^{-7}\) cm² s⁻¹, \(D_{\text{H}_2} = 2.2 \times 10^{-5}\) cm² s⁻¹, \(D_{\text{BH}^+} = 2.4 \times 10^{-7}\) cm² s⁻¹. \(E^0(\text{BH}^+/\text{H}_2) = -0.030\) V and \(E^0(\text{BH}^+/\text{H}_2) = -0.395\) V, derived from \(E^0(\text{H}^+/\text{H}_2)\), \(E^0(\text{BH}^+/\text{H}^+)\) and \(k_{\text{dim}}\). All experiments were carried out at a 1.6-mm-diameter Pt macrodisk electrode in [C₂mim][NTf₂] with a scan rate of 100 mV s⁻¹. Adapted with permission from Reference 60. Copyright 2015, American Chemical Society.

In order to simulate the BH⁺/H₂ process, two pathways were considered. The first is the CE pathway, whereby proton reduction via the mechanism described by Equations 22 and 23 is preceded by dissociation of BH⁺:

\[
\text{BH}^+ + \overset{\text{K}_s \cdot k_{\text{dissoc}}}{{\text{H}^+ + \text{B}}} \quad \text{K}_s \cdot k_{\text{dissoc}} \cdot k_{\text{assoc}}, \quad \text{(26)}
\]

where \(k_{\text{dissoc}}\) and \(k_{\text{assoc}}\) are the dissociation and association rate constants, respectively. The second is a direct reduction (DR) pathway, where BH⁺ is discharged at the electrode surface without prior dissociation:

\[
\text{BH}^+ + e^- \overset{\text{K}_s \cdot k_{\text{DR}} \cdot \alpha_{\text{DR}}}{{\text{H}^+ + \text{B}}} \quad E^0(\text{BH}^+/\text{H}^+) \cdot k_{\text{DR}} \cdot \alpha_{\text{DR}}, \quad \text{(27)}
\]

Again, it should be emphasized that \(k_s\) and \(\alpha\) are apparent values and therefore should not be assumed to have direct physical significance. In all simulations, \(k_{\text{assoc}}\) was set to \(5 \times 10^8\) M⁻¹ s⁻¹, which should approximate the diffusion-controlled limit for a bimolecular reaction in [C₂mim][NTf₂] (78, 101). Simulations considering only the CE (Equations 22, 23, and 26) pathway or the CE + DR (Equations 22, 23, 26, and 27) pathways are shown in Figure 5b for the monoprotonated amine, 2,4-dichloropyridinium. Contrary to the mechanism proposed for benzoic acid (97–99), the CE pathway alone cannot support the mass transport–limited currents
observed experimentally due to the physical limitation of $k_{\text{dissoc}}$ ($k_{\text{dissoc}} = K_{a}k_{\text{assoc}} \approx 300 \text{ s}^{-1}$), necessitating the inclusion of the parallel $DR$ pathway in the simulations. Further simulations revealed that, under the experimental conditions, the $CE$ pathway could only attain a diffusion-controlled current when $pK_{a} < 4$ [confirmed in a related study (61)]. The $DR$ pathway becomes insignificant when $pK_{a} < 2$, which is in qualitative agreement with the studies by Evans and coworkers (54, 55) on the reduction of weak acids in dimethylsulfoxide.

In a related study (61), the thermodynamics, kinetics, and mechanisms of the HER at a Pt electrode were again investigated in the same IL, [C$_{2}$mim][NTf$_{2}$], but this time using a range of oxyacids (i.e., phenols, carboxylic acids, or sulfonic acids) as the proton source. Triflic acid, H[OTf], a well-known superacid in aqueous media, was shown to be a weak acid ($pK_{a} = 2.0$, derived using Equation 10) in [C$_{2}$mim][NTf$_{2}$] (conversely, H[NTf$_{2}$] was shown to behave as a strong acid in [C$_{2}$mim][OTf]), giving rise to a diffusion-controlled process that occurs in the potential region negative of the H$^{+}$/H$_{2}$ process, as demonstrated in Figure 6:

\[
2\text{HA} + 2e^{-} \rightarrow \text{H}_2 + 2\text{A}^{-}. \tag{28}
\]

The HA/H$_{2}$ process, using H[OTf] as the proton source, was successfully simulated using the $CE$ mechanism highlighted above (Equations 22, 23, and 26), in line with the theoretical predictions established in 60.

<COMP: PLEASE INSERT FIGURE 6 HERE>

**Figure 6** Cyclic voltammograms ($\nu = 100 \text{ mV s}^{-1}$, normalized to unity with the reduction peak current, $I_{p,\text{red}}$) showing the hydrogen evolution reaction obtained at a 1.6-mm-diameter Pt macrodisk electrode in [C$_{2}$mim][NTf$_{2}$], with the following proton sources (top to bottom): H[NTf$_{2}$], H[OTf], naphthalene-2-sulfonic acid, H[OTs], methanesulfonic acid, trichloroacetic acid, dichloroacetic acid, malonic acid, $\alpha$-hydroxybenzoic (salicylic) acid, pentachlorophenol, and glycolic acid. The $pK_{a}$ values calculated using Equation 10 are indicated in the figure.

Abbreviation: Fc/Fc$^{+}$, ferrocene/ferrocenium. Adapted with permission from Reference 61. Copyright 2015, American Chemical Society.

Proton reduction from a range of sulfonic or carboxylic acids (5.9 $\leq pK_{a} \leq 17.8$) occurs in two steps, as illustrated in **Figures 6** and 7a. The electron stoichiometry of the two steps was confirmed to be 1:1 through semi-integration (69, 102). Based on this finding and what is known about oxyacids in aprotic media (103), including ILs (48), it was proposed that splitting of the proton reduction wave is caused by intermolecular hydrogen bonding between HA and its
conjugate base, $A^-$, forming the stable intermediate species, $[HA_2]^-$. This is a well-documented phenomenon (104, 105) known as homoassociation or homoconjugation, and in the present case, it occurs in the diffusion layer adjacent to the electrode surface in the potential region where the concentration of both species is comparable. Therefore, proton reduction from any (oxy)acid that undergoes homoassociation is expected to proceed via an $ECE$ mechanism, where $C$ is the homoassociation step, and the two processes observed voltammetrically are attributable to:

\[
2HA + e^- \rightarrow \frac{1}{2} H_2 + [HA_2]^-, 
\]

\[
[HA_2]^- + e^- \rightarrow \frac{1}{2} H_2 + 2A^-.
\]

These equations account for the 1:1 electron stoichiometry observed experimentally.

Figure 7 (a) Comparison of simulated (circles) and experimental (lines) cyclic voltammograms showing the HER obtained in [C$_2$mim][NTf$_2$] from (bottom to top): 41.6 mM H[OTs], 41.6 mM H[OTs] + 44.0 mM [C$_2$mim][OTs], and 41.2 mM H[OTs] + 86.1 mM [C$_2$mim][OTs]. Simulations were performed using the $DR$ mechanism described by Equations 23, 27, and 31 using the following parameters: $K_{\text{dim}} = 10^4$, $k_{\text{dim}} = 10^{16}$ M$^{-1}$ s$^{-1}$ (Equation 23); $E_0^0(\text{HA}/\text{H}^*) = -0.512$ V versus Fc/Fc$^+$ ($pK_a = 6.2$), $k_{S,DR} = 0.09$ cm/s, $\alpha_{DR} = 0.4$ (Equation 27, apparent values); $K_{\text{homo}} = 950$, $k_{\text{homo}} = 5 \times 10^8$ M$^{-1}$ s$^{-1}$ (Equation 31); $D_{H_2} = 2.2 \times 10^{-5}$ cm$^2$ s$^{-1}$; $D_{HA} = D_A^- = 2.6 \times 10^{-7}$ cm$^2$ s$^{-1}$, $D_{[HA_2]^ -} = 1.85 \times 10^{-7}$ cm$^2$ s$^{-1}$. $E_0^0(\text{HA}/\text{H}_2) = -0.394$ V, derived from $E_0^0(\text{HA}/\text{H}^*)$ and $K_{\text{dim}}$. (b) Cyclic voltammograms obtained from the HER, with H[OTs] as the proton source in (bottom to top): neat [C$_2$mim][NTf$_2$], [C$_2$mim][NTf$_2$] + 5.2 vol% PG and neat [N$_{H,2,2,2}$][NTf$_2$]. All experiments were carried out at a 1.6-mm-diameter Pt macrodisk electrode with a scan rate of 100 mV s$^{-1}$. Abbreviations: Fc/Fc$^+$, ferrocene/ferrocenium; HER, hydrogen evolution reaction; PG, propylene glycol. Adapted with permission from Reference 61. Copyright 2015, American Chemical Society.

In order to simulate the HA/[HA$_2$]$^-$/H$_2$ processes, the previously established $DR$ mechanism (Equations 23 and 27) was combined with the homoassociation equilibrium reaction:

\[
HA + A^- \rightarrow [HA_2]^-, K_{\text{homo}}, k_{\text{homo}}.
\]

Taking tosylic acid (H[OTs]) as the model acid species, and by assuming that $k_{\text{homo}}$ occurs at the diffusion limited rate ($5 \times 10^8$ M$^{-1}$ s$^{-1}$), the proton reduction response was simulated by assuming...
a $K_{\text{homo}}$ value of approximately $10^3$, as shown in Figure 7a. In addition, adding A$^-$ (e.g., [OTs]$^-$) to a HA (e.g., H[OTs]) solution is expected to drive the equilibrium shown in Equation 31 to the right, forming [HA$_2$]$^-$ at the expense of HA. This was observed experimentally and was predicted by the simulations, also shown in Figure 7a. Since the driving force for anionic homoassociation (i.e., $K_{\text{homo}}$) is intermolecular hydrogen bonding, this phenomenon is predicted to be suppressed through the addition of hydrogen-bonding solvents, promoting the formation of solvent–solute hydrogen bonds in preference to solute–solute. This was confirmed experimentally through the addition of ~5% (v/v) propylene glycol, as shown in Figure 7b. Alternatively, the hydrogen bond donating/accepting ability can be enhanced by varying the constituent cation/anion of the IL, demonstrated through the use of [NH$_{2,2,2}$][NTf$_2$] ([NH$_{2,2,2}$]$^+$ is a relatively good hydrogen bond donor; see Figure 7b) or [C$_2$ mim][N(CN)$_2$] ([N(CN)$_2$]$^-$ is a relatively good hydrogen bond acceptor; not shown).

Despite the limitations in the simulations (simplified mechanisms using empirical rather than physically significant parameters), these studies (59–61) unveiled significant physical insights into the HER mechanism in IL media. Notably, although the characteristics of the H$^+$/H$_2$ process are relatively IL independent for a given anion (59), the thermodynamics, kinetics, and mechanisms of the BH$^+$/H$_2$ or HA/H$_2$ processes are strongly dependent upon the nature of the conjugate base (B or A$^-$) and the Lewis acid/base properties of the IL medium (60, 61).

4. CONCLUSIONS AND PROSPECTS

This review highlighted and discussed recent (electrochemical) studies on the energetics (thermodynamics) of proton transfer and electrode reaction mechanisms of the HER and HOR processes. The p$K_a$ scale has been the focus of many recent studies, with methods such as spectrophotometry, potentiometry, and voltammetry shown as robust methods for the quantification of (equilibrium) acidity in IL media. Although still in its infancy, especially when considering that each IL formulation will possess its own unique p$K_a$ scale (largely governed by the constituent anion), this work offers valuable insights into the influence of solvent (IL) properties (i.e., Lewis acidity/basicity, hydrogen bonding, polarity, etc.) on equilibrium acidity, expanded beyond conventional molecular solvent media. Furthermore, building on the so-called designer qualities of ILs, understanding the relationship between structure (i.e., anion and cation...
properties) and function (i.e., proton availability or solvating ability) is an important step for tailoring these solvents for particular applications.

The H⁺/H₂ process, although characterized extensively in aqueous media, has only been investigated in a limited range of ILs and almost exclusively on Pt electrodes over the past decade. The mechanism of this model inner-sphere process in ILs is proposed to be analogous to that in aqueous media, occurring via a combination of the well-established Volmer, Tafel and Heyrovsky reactions. The thermodynamics (i.e., \( E^\circ \)) of the H⁺/H₂ process is IL dependent, as highlighted in the studies of pKₐ in this media, being predominantly governed by the identity of the constituent IL anion. However, the electrode kinetics of this process show a complex and generally poorly understood IL dependence and display considerably different voltammetric characteristics (e.g., voltammetric \( \Delta E_p \) values) in even closely related IL formulations. ILs were identified early on for their potential as electrolytes in a range of electrochemical devices (e.g., PILs in fuel cells). Therefore, further work is necessary in this field to develop an understanding of how the characteristics of an IL formulation will influence the thermodynamics, kinetics, and mechanisms of inner-sphere reactions, such as the H⁺/H₂ process. The ultimate goal, from an applications (i.e., fuel cell) perspective, would be to develop an IL formulation that facilitates facile HOR and ORR kinetics and fast anhydrous proton transport over a wide temperature range.

In conclusion, ILs are a diverse and versatile class of solvent that have garnered considerable attention over the past two decades as replacements for volatile molecular solvents in a range of applications. Nevertheless, as research in the field has matured over this time frame, the limits to our knowledge on fundamental processes (e.g., thermodynamics, kinetics, and mechanisms) in this type of medium have become apparent. Moving forward, developing an understanding of the factors that influence (electro)chemical reactivity in ILs, as exemplified in the studies summarized in this review, will undoubtedly facilitate the widespread adoption of these designer solvents in fields as diverse as energy technology to the biological sciences.

**DISCLOSURE STATEMENT**

The authors are not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.
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