Enhancing the Catalysis of Oxygen Reduction Reaction via Tuning Interfacial Hydrogen Bonds

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Abstract:

Proton activity at the electrified interface is central to the kinetics of proton-coupled electron transfer (PCET) reactions for making chemicals and fuels. Here we employed a library of protic ionic liquids in an interfacial layer on Pt and Au to alter local proton activity, where the intrinsic ORR activity was enhanced up to 5 times, exhibiting a
volcano-shaped dependence on the $pK_a$ of the ionic liquid. The enhanced ORR activity was attributed to favorable proton transfer kinetics for strengthened hydrogen bonds between the ionic liquid to the ORR product with comparable $pK_a$. This proposed mechanism was supported by in situ surface-enhanced Fourier-Transform Infrared Spectroscopy and our simulation of PCET kinetics based on computed proton vibrational wavefunction at the H-bond interface. These findings highlight opportunities in using non-covalent interactions of hydrogen bond structures and solvation environments at the electrified interface to tune the kinetics of ORR and beyond.

Understanding proton coupled electron transfer (PCET) processes are critical to control the reaction kinetics in bio$^1$, organic$^{2, 4}$, inorganic$^{5, 6, 7}$, environmental chemistry$^8$ and electrochemistry$^{4, 9}$. In particular, PCET steps have been used extensively in recent research on catalysts for making energy carriers or solar fuels$^{4, 10, 11, 12}$ including water splitting$^{4, 12, 13}$, and for converting energy carriers in fuel cells to generate work$^7, 9$ including oxygen reduction reaction (ORR)$^9$. Current thinking of the ORR mechanism$^9$ involves PCET steps on surface metal sites, which can involve one key intermediate on metals such as Au$^{15}$ for two-electron ORR ($O_2 + 2H^+ + 2e^- \rightarrow H_2O_2$, 0.68 $V_{\text{RHE}}$), -OOH$^{15, 16, 17}$, and three key intermediates on metals such as Pt$^{14, 18, 19}$, -OOH, -O and -OH for four-electron ORR ($O_2 + 4H^+ + 4e^- \rightarrow 2H_2O$, 1.23 $V_{\text{RHE}}$)$^{20}$. Conventional designs of catalysts involve the tuning of the surface electronic structure and covalent interactions with reaction intermediates$^{14, 20, 21}$ (surface binding or adsorption strength), which has led to significant advances in the intrinsic ORR activity by tuning strain$^{22, 23}$, the ligand effect$^{24, 25}$, and surface orientation$^{26, 27, 28}$ For example, Pt$_3$Ni(111)$^{29}$ is shown to exhibit intrinsic ORR activity ten times greater than Pt (111) at 0.9 $V_{\text{RHE}}$, and
polycrystalline PtGd\textsuperscript{30} exhibit five times enhancement than polycrystalline Pt.

Recent studies\textsuperscript{31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41} have shown that changing electrolyte compositions can significantly alter the kinetics of electrochemical reactions such as oxygen reduction reaction (ORR)\textsuperscript{31, 32, 34, 35, 38, 39} and hydrogen oxidation reaction\textsuperscript{33, 36, 37, 40, 41}, highlighting new opportunities in noncovalent interactions via the chemical physics of electrolytes to control activity. While the pH of electrolyte, which represents proton activity, does not greatly alter the ORR activity of metals such as Pt\textsuperscript{42}, increasing pH can significantly increase the ORR activity on weakly interacting metals such as Au\textsuperscript{43, 44, 45, 46}, where the activity in base is much higher than that in acid\textsuperscript{44}. Moreover, modifying Pt-based catalysts with protic ionic liquids\textsuperscript{32, 47, 48} is shown to enhance the ORR activity in acid, where ionic-liquid-modified PtNi nanoparticles have \textasciitilde3 times greater intrinsic ORR activity than pristine PtNi\textsuperscript{32}, and \textasciitilde20 times greater than commercial Pt/C\textsuperscript{48}. Although this increase in the ORR activity has been attributed to the increase of hydrophobicity at the interface between Pt and ionic liquids\textsuperscript{47, 49} and oxygen solubility in ionic liquids\textsuperscript{32, 50}, it is not apparent on how to design new catalysts to control the catalytic activity of ORR and other reactions using this strategy. Considering these ionic liquids used in these previous studies\textsuperscript{32, 47, 48} have different $pK_a$ from the acidic electrolyte, it is postulated that interfacial proton activity of ionic liquids can alter proton transfer or proton-coupled electron transfer (PCET) kinetics on metals such as Pt and Au, and thus ORR activity, which is the focus of this work.

Here we employ a library of protic ionic liquids in an interfacial layer to tune ORR kinetics of Pt and Au in acid. We hypothesize that the protic cations in the ionic liquids
can serve as intermolecular proton relay between proton in the bulk electrolyte and that near the metal surface (Scheme 1), where altering of protic cation with different proton activity (or $pK_a$) provides opportunities to examine how local proton activity near the active sites can influence the ORR kinetics. A library of protic cations with different $pK_a$ values were used (Scheme 1) \([\text{C}_4\text{C}_1\text{im}]^+\) $pK_a=23.3^{51}$, \([\text{TMPim}]^+\) $pK_a=21.3^{51}$, \([\text{MTBD}]^+\) $pK_a=15.0^{52}$, \([\text{DBU}]^+\) $pK_a=13.5^{52}$, \([\text{DEMA}]^+\) $pK_a=10.3^{53}$, \([\text{TEMEDA}]^+\) $pK_a=8.9^{54}$, \([\text{C}_4\text{Him}]^+\) $pK_a=7.1^{55}$ to influence the ORR activity of Au and Pt, which catalyzes ORR via the 2e$^{-55}$ and 4e$^{-56}$ pathway in acid, respectively. The ORR activity of Au and Pt was found to increase with the presence of protic cations, where the maximum enhancement was obtained when the $pK_a$ values of protic cations and the reaction intermediate in the ORR rate-limiting step are similar. The ORR activity enhancement can be attributed to the stronger hydrogen bond between protic cations and the rate-limiting ORR intermediate, which is supported by in situ attenuated total reflection surface-enhanced infrared absorption spectroscopy (ATR-SEIRAS). Further support came from predicted PCET kinetics based on computed proton vibrational wavefunction at the H-bond interface, where the stronger H-bond between protic cations and ORR intermediates led to faster proton tunneling kinetics, giving rise to greater ORR exchange current density. Our work reveals the role of local $pK_a$ and interfacial H-bond strength on the kinetics of PCET and electrochemical reactions, which highlight new opportunities to further enhance electrocatalytic activity of ORR, reduction of CO$_2$ and N$_2$, and beyond by tuning how local proton activity near the active sites.

$pK_a$-dependent ORR activity measurements of protic-cation-modified catalysts
The ORR activity of Au/C (6.0 nm Au) and Pt/C (TKK, 19wt% 1.7 nm Pt) with and without protic cations was measured using rotating disk electrode measurements, as shown in Fig. 1a and 1b, respectively. The ionic liquid layer thickness with protic cations on the surface of Au/C and Pt/C was estimated to ~1 nm (details in Fig. S1). The catalysts with protic cations showed enhanced activity compared to those without, as indicated by the positive shifts in the half-wave potential in the voltammetry data. ORR kinetic currents were extracted from the data in Fig. 1a, 1b and Fig. S2 using the Koutecky–Levich analysis, from which the specific and mass activity was obtained by normalizing the kinetic ORR current (Fig. S3) by the electrochemically surface area from CV measurements (Fig. S4-5) and metal mass (Fig. S6), respectively. The specific ORR activity for Au/C at 0.4 V_RHE can be enhanced up to ~5 times with protic cations relative to pristine Au/C shown in this work while that of Pt/C at 0.9 V_RHE can be increased by ~3 times compared to pristine Pt/C (TKK 19%, ~190 μA/cm²Pt) shown in this work and previous studies. Similar activity trends and enhancement were found for polycrystalline Au (~4 times at 0.3V_RHE, Fig. S7a) and Pt (~2 times at 0.9 V_RHE, Fig. S7b). The exchange current density of ORR was extracted for Au/C (O₂ + 2H⁺ + 2e⁻ => H₂O₂, 0.68 V_RHE) and for Pt/C (O₂ + 4H⁺ + 4e⁻ => 2H₂O, 1.23 V_RHE) using Butler-Volmer in Fig. S8 and Fig. S9 respectively. These observations suggest that the local proton activity at the metal/electrolyte interface can considerably influence the ORR kinetics.

The ORR activity of both Au/C and Pt/C was found to first increase and then decrease, exhibiting a volcano trend, as a function of the pKₐ value of protic cations, as shown in Fig. 1c and Fig. 1d, respectively. The maximum ORR activity enhancement for Au/C was found for [DEMA][NTf₂] with pKₐ of 10.3, which is similar to the ORR product on Au, H₂O₂ of pKₐ of 11.6. In acid, the first PCET step of ORR on Au (O₂ + H₃O⁺
+ e⁻ => OOHₐu + H₂O) has similar kinetics to the second step (OOHₐu + H₃O⁺ + e⁻ => H₂O₂ + H₂O), as indicated by the 0.09 eV difference in activation barrier from DFT²¹, ⁶³. As increasing pH is shown to promote the rate of the first step by several orders of magnitude⁴⁶, the second step becomes rate-limiting to the overall kinetics with increasing pH. Therefore, we propose that the ORR kinetics on Au with interfacial ionic liquids are limited by the PCET kinetics of OOHₐu to form H₂O₂ ( OOHₐu + H⁻ + e⁻ => H₂O₂ + N)¹⁵,²¹ (the detailed kinetic analysis depicted in Fig. S10). This hypothesis is in agreement with previous kinetic measurements of ORR in organic solvents, where the first electron transfer step (O₂ + e⁻ => O₂⁻)⁶⁴ is more than 10³ times faster than the second PCET step (O₂⁻ + e⁻ + H₂O => OOH⁻ + OH⁻)⁶⁵. Further support comes from the observation that OOHₐu has been detected as the stable ORR intermediate accumulating on Au by in situ ATR-SEIRAS¹⁶,¹⁷ and in situ surface enhanced Raman⁶⁶. Therefore, it is proposed that different pKₐ altered the kinetics of the second PCET step for ORR on Au in ionic liquid (OOHₐu + N-H⁻ + e⁻ => H₂O₂ + N) and tuned the overall ORR kinetics consequently. This hypothesis departs from the decoupled proton-electron transfer mechanism outlined by Koper,⁹,⁴¹ where increasing kinetics with increasing pH for one-electron reduction can be attributed to enhanced electron-transfer kinetics on the RHE scale.

The maximum ORR activity enhancement for Pt/C occurred for [MTBD][NTf₂] with pKₐ of 15.0, which is similar to the ORR product in the rate-limiting step, water with pKₐ of 15.7 ⁶². Previous DFT studies have shown that PCET of adsorbed OH on Pt to form H₂O is rate-limiting (OHₚt + H⁺ + e⁻ => Pt + H₂O),¹⁴ which is supported by ambient pressure X-ray photoelectron spectroscopy¹⁸ and in situ electrochemical surface-enhanced Raman spectroscopy¹⁹. Therefore, these results show that the kinetics of ORR on Au and Pt could be regulated by the pKₐ of protic cations at the interface,
which can work as the proton donor near the active sites catalyzing the rate-limiting PCET, having the maximum ORR activity obtained with minimum $pK_a$ difference between the protic ionic liquid (proton donor) and rate-limiting ORR product (proton acceptor).

As the thermodynamic driving force of proton coupled electron transfer (PCET) reactions diminishes with minimized $pK_a$ difference between the proton donor and acceptor, we propose that the enhancement in the ORR activity can be attributed to difference in the interfacial hydrogen bond structure as predicted previously by PCET theory for homogeneous reactions, which will be examined by in situ ATR-SEIRAS experiments and computation below.

$pK_a$-dependent interfacial H-bond detected by in situ ATR-SEIRAS

In situ ATR-SEIRAS was performed on Au and Pt thin-film chemically deposited on Si prism, which detected ORR intermediates and revealed the interactions between protic cations in ionic liquids and ORR intermediates by monitoring the stretching frequency of H-bonded species of protic cations as influenced by the formation of ORR intermediates and product as a function of potential. Potential-dependent in situ ATR-SEIRA spectra were collected from [MTBD][NTf$_2$]-modified Au thin-film surface (estimated to have a thickness of ~ 50 nm for Au and ~20 μm for [MTBD][NTf$_2$]). Differential spectra subtracted by that collected at OCV are shown in Fig. 2, along with that of bulk [MTBD][NTf$_2$] and [MTBD][NTf$_2$] with water (0.5 M water). Two sharp peaks at 1632 cm$^{-1}$ and 1603 cm$^{-1}$ (Fig. 2a), resulting from the stretching mode of C=N-H$^+$ and C=N in the cation, respectively, grew with decreasing potential. The increased peak intensities can be attributed to having more protic cations on the Au surface with decreasing potential, which is in agreement with increasing peak intensities of C-N.
stretching\textsuperscript{69, 70} from protic cations at 1231 cm\textsuperscript{-1} (Fig. 2b). Similar peaks to those found on Au were found on [MTBD][NTf\textsubscript{2}]-modified Pt thin-film surface (Fig. S11), which also grew with decreasing potential.

Two new peaks, at 1263 cm\textsuperscript{-1} and 3238 cm\textsuperscript{-1} emerged (Fig. 2b and 2c) with decreasing potential, which can be assigned to the bending (H-O-O) of HOO-adsorbed on Au\textsuperscript{16, 17} and the stretching of N-H\textsuperscript{+}⋅⋅⋅OOH\textsubscript{Au} of [MTBD][NTf\textsubscript{2}], respectively. The assignment of the broad feature at 3238 cm\textsuperscript{-1} is supported by DFT calculations, where computed N-H\textsuperscript{+}⋅⋅⋅OOH\textsubscript{Au} stretching of [MTBD][NTf\textsubscript{2}] was determined to be 3267 cm\textsuperscript{-1} (Table S1), in good agreement with the experimental observation. This new broad peak at \sim3238 cm\textsuperscript{-1} is redshifted compared to the N-H\textsuperscript{+} stretching of bulk [MTBD][NTf\textsubscript{2}] with and without water added (\sim3400 cm\textsuperscript{-1} in Fig. 2c), indicating the presence of hydrogen bond interactions between protic cation and ORR intermediates such as [MTBD]N-H\textsuperscript{+}⋅⋅⋅OOH\textsubscript{Au}. These two peaks at 1263 cm\textsuperscript{-1} and 3238 cm\textsuperscript{-1} grew at the expense of the peak at 3583 cm\textsuperscript{-1}, attributable to the O-H stretching\textsuperscript{21} of water dissolved in the [MTBD][NTf\textsubscript{2}] in Fig. 2c and Fig. S11, which indicated reduction of water adsorbed on the surface with decreasing potential.

Changing the pK\textsubscript{a} of protic cations was found to redshift the N-H\textsuperscript{+}⋅⋅⋅OOH\textsubscript{Au} stretching at ORR-relevant potentials (Fig. 3 and Fig. S12 and 13). The N-H\textsuperscript{+}⋅⋅⋅OOH\textsubscript{Au} stretching frequency decreased from 3429 cm\textsuperscript{-1} for [C\textsubscript{4}Him][NTf\textsubscript{2}] with pK\textsubscript{a} of 7.1 in Fig. 3c, 3238 cm\textsuperscript{-1} for [MTBD][NTf\textsubscript{2}] with pK\textsubscript{a} of 15.0 in Fig. 3c, to \sim3000 cm\textsuperscript{-1} for [DEMA][NTf\textsubscript{2}] with pK\textsubscript{a} of 10.3 in Fig. 3b, at 0.2 V\textsubscript{RHE}. These assignments are supported by DFT calculations (Table S1), which also show the two fine peak features at \sim3000 cm\textsuperscript{-1} for [DEMA][NTf\textsubscript{2}] with the maximum ORR activity, having the feature at 2985 cm\textsuperscript{-1} to the stretching of [DEMA]N-H\textsuperscript{+}⋅⋅⋅OOH\textsubscript{Au} and the feature at 3024 cm\textsuperscript{-1} to [DEMA]N-H\textsuperscript{+}⋅⋅⋅OHOH. Further support to these assignments came from the observed peak growth
as expected for increasing ORR intermediates with decreasing potential (Fig. S12, S13 and S14). Weakening the N-H\(^+\) (proton donor) stretching of N-H\(^+\)...OOH\(_{Au}\) from [C₄Him][NTf\(_₂\)] to [DEMA][NTf\(_₂\)] indicates stronger hydrogen bond interactions in N-H\(^+\)...OOH\(_{Au}\) as shown from previous IR spectroscopy studies of hydrogen-bonded (H-Bond) species\(^{72,73}\), suggesting the strongest hydrogen bond found for [DEMA][NTf\(_₂\)] with a similar \(pK_a\) (10.3) to H\(_2\)O\(_₂\) (11.6). This observation is in agreement with general trends that the H-bond would become stronger when the difference between the \(pK_a\) value of proton donor and acceptor decreases as shown from thousands of H-bond structures from the Cambridge Structural Database including the N-H\(^+\)...O, N-H\(^+\)...N etc.\(^{74}\) Therefore, the maximum ORR activity found on Au in presence of [DEMA][NTf\(_₂\)] is associated with the strongest H-bond interaction between N-H\(^+\)...OOH (Fig. 3g). Considering the rate of ORR on Au is limited by PCET from OOH\(_{Au}\) to form H\(_2\)O\(_₂\) on Au in ionic liquids (OOH\(_{Au}\) + N-H\(^+\) + e\(^-\) = H\(_2\)O\(_₂\) + N), strengthening H...OOH facilitated by protic cations with comparable \(pK_a\) such as [DEMA]N-H\(^+\) would enhance ORR kinetics.

Similar to Au, the maximum ORR activity found on Pt in presence of [MTBD][NTf\(_₂\)] is associated with the strongest H-bonding interaction between N-H\(^+\) and OH of species such as N-H\(^+\)...OH\(_{Pt}\) among the protic cations examined in this study, as shown in Fig. 3d-f, Fig. 3h and Fig. S15-17. This assignment is in agreement with the detection of Pt-OH as the dominant ORR intermediate by \textit{in situ} SERS and XPS\(^{18,19}\) and previous DFT findings that PCET of OH adsorbed on Pt is rate-limiting for ORR\(^{68,75}\). Further support came from the agreement between experimentally observed and DFT calculated (Table S1) wavenumber for N-H\(^+\) stretching, where the computed value for [MTBD]N-H\(^+\)...OH\(_{Pt}\), [DEMA]N-H\(^+\)...OH\(_{Pt}\) and [C₄Him]N-H\(^+\)...OH\(_{Pt}\) was found to be 3089 cm\(^{-1}\), 3188 cm\(^{-1}\) and 3335 cm\(^{-1}\), respectively. Like Au, the N\(^+\)-H stretching for the protic
cation with the maximum ORR activity has one additional feature to [MTBD]N-H’⋯OH\textsubscript{Pt} at 3089 cm\textsuperscript{-1}, which can be attributed to [MTBD]N-H’⋯OH\textsubscript{2} at 3215 cm\textsuperscript{-1} as supported by DFT (Table S1), revealing the presence of the ORR intermediate (OH\textsubscript{Pt}) and final product (H\textsubscript{2}O) in the rate-limiting step. Considering the PCET from OH\textsubscript{Pt} to form H\textsubscript{2}O (OH\textsubscript{Pt} + N-H\textsuperscript{+} + e\textsuperscript{-} = H\textsubscript{2}O + N) is rate-limiting, strengthening H⋯OH facilitated by protic cations with comparable pK\textsubscript{a} such as [MTBD]N-H\textsuperscript{+} would enhance ORR kinetics.

\textit{pK}_a-dependent PCET kinetics in ORR facilitated by protic cations

Here we compute the exchange current densities of PCET kinetics (Fig. 4a) from select protic cations in the rate-limiting ORR step on Au (N-H’⋯OOH\textsubscript{Au} + e\textsuperscript{-} => H\textsubscript{2}O\textsubscript{2} + N) and Pt (N-H’⋯OH\textsubscript{Pt} + e\textsuperscript{-} => H\textsubscript{2}O + N) using reported methods\textsuperscript{5,7,76} While the minimum point of proton potential is close to proton donor side (e.g. N atom of N-H’⋯OOH\textsubscript{Au}) before electron transfer, the reaction intermediate (such as adsorbed OOH or OH) accepts one electron from electrode and lowers the proton potential, enabling proton transfer to form product, where proton is assumed to transferred simultaneously with electron\textsuperscript{5,76}. The transferring proton was treated by quantum mechanics\textsuperscript{76} and the proton vibronic states were computed from the proton potentials of [MTBD]N-H’⋯OOH\textsubscript{Au} (Fig. S18a), [DEMA]N-H’⋯OOH\textsubscript{Au} (Fig. S18b), [C\textsubscript{4}Him]N-H’⋯OOH\textsubscript{Au} (Fig. S18c), [MTBD]N-H’⋯OH\textsubscript{Pt} (Fig. S18d), [DEMA]N-H’⋯OH\textsubscript{Pt} (Fig. S18e) and [C\textsubscript{4}Him]N-H’⋯OH\textsubscript{Pt} (Fig. S18f), where the wavefunctions and energy levels of different quantum states are shown in Fig. S19-S24. The PCET reaction was described regarding the transition between vibrational states\textsuperscript{76} of reactant (e.g. N-H’⋯OOH\textsubscript{Au}) and product (e.g.
H$_2$O$_2$), where the exchange current density was proportional to the product of the vibronic coupling squared. Here we assume that the electronic coupling is unchanged for ORR with different protic cations as the distance between reaction intermediate (adsorbed OOH and OH) and electron donor (electrode surface, Au and Pt) remains constant. Therefore, we will focus on the H-bonding structure effect on overall PCET kinetics in this part, the exchange current density is proportional to the product of the vibronic coupling of proton squared (denoted as $S_{\nu\mu}^2$), the Boltzmann probability of different proton vibrational states ($P_{\nu}$) and the Boltzmann probability overcoming the activation free energy ($\Delta G^\ddagger$) in Fig. 4b. As summarized in Table S10 and S11, where one or two states dominate the overall kinetics, showing much higher rate constant than others, the differences in $\Delta G^\ddagger$ of contributing states of [MTBD]N-H$^+$···OOH$_{\text{Au}}$, [DEMA]N-H$^+$···OOH$_{\text{Au}}$, [C$_4$Him]N-H$^+$···OOH$_{\text{Au}}$ are within 0.05 eV. Similarly, the differences in $\Delta G^\ddagger$ (Table S11) of contributing states of [MTBD]N-H$^+$···OH$_{\text{Pt}}$, [DEMA]N-H$^+$···OH$_{\text{Pt}}$, [C$_4$Him]N-H$^+$···OH$_{\text{Pt}}$ are within 0.04 eV. These results indicate the effect of $\Delta G^\ddagger$ on the PCET kinetics is insignificant in this work, while the enhancement in ORR exchange current density mainly resulted from the H-bonding structure, regulating the tunneling kinetics of PCET and tuning the pre-exponential factor in the rate constant expression.

Our computational results show that the enhancement in the ORR activity of Au and Pt by the presence of protic cations with select $pK_a$ can be attributed to greater $P_{\nu}S_{\nu\mu}^2$ of protons across the contributing states (Fig. 4e and 4f). $S_{\nu\mu}^2$ was quantified by the integral overlap of proton vibrational wavefunctions between reactant (oxidized state, before
ET) and product (reduced state, after ET) (Table S8) indicating the vibronic coupling of proton through PCET process and \( P_v \) as the Boltzmann probability of different quantum states is shown in Table S9. As shown in Fig. 4c, the energy of reactant higher states for \( \text{N-H}^+\cdots\text{OOH}_{\text{Au}} \) than the ground state (state 0) is much higher (0.35-0.4 eV in Table S8), so the (0, 0) transition is the contributing state for overall kinetics, the PCET kinetics are dominated by the vibronic coupling of protons across the ground states of reactant to product (0,0), having \( P_v \) of ground state close to 1 in Table S8. \( P_vS_{\nu_2}^2 \) and exchange current density \( (J_0) \) for \([\text{DEMA}]\text{N-H}^+\cdots\text{OOH}_{\text{Au}} \) was found higher than \([\text{C}_4\text{Him}]\text{N-H}^+\cdots\text{OOH}_{\text{Au}} \) and \([\text{MTBD}]\text{N-H}^+\cdots\text{OOH}_{\text{Au}} \) in Fig. 4d and Table S10, which is in agreement with experimental results (Fig. 1). In contrast to Au, as the energy of state 1 for reactant of \( \text{N-H}^+\cdots\text{OH}_{\text{Pt}} \) is only 0.1-0.2 eV higher than the ground state (Table S9), and proton wavefunction of reactant in state 1 distributes more on oxygen side (Fig. 4e) which can generate much higher vibronic coupling in (1, 0) transition state than (0, 0) to compensate the decrease in \( P_v \) of state 1, leading (1, 0) transition become contributing state for PCET kinetics in \( \text{N-H}^+\cdots\text{OH}_{\text{Pt}} \). Since the \( S_{\nu_1} \) (0,1) being similar for different ionic liquids in Table S9, the PCET kinetics are governed by the \( P_v \) of protons across the state 1. \( P_vS_{\nu_2}^2 \) and exchange current density \( (J_0) \) for \([\text{MTBD}]\text{N}^+\cdots\text{OH}_{\text{Pt}} \) was found the highest, in agreement with experimental exchange current density trends, as shown in Fig. 4f and Table S11. Therefore, increasing ORR activity of Au and Pt by the presence of protic cations can be attributed to greater \( P_vS_{\nu_2}^2 \) of protons across the contributing states, peaked at \([\text{DEMA}]\text{N-H}^+\cdots\text{OOH}_{\text{Au}} \) with comparable \( pK_a \) between \([\text{DEMA}]\text{N-H}^+ \) and \( \text{H}_2\text{O}_2 \), and \([\text{MTBD}]\text{N-H}^+\cdots\text{OH}_{\text{Pt}} \) with comparable \( pK_a \) between
[MTBD]N-H⁺ and H₂O, respectively. The predicted J₀ is in good agreement with those measured experimentally (Table S10 and S11). As the Pν and Sνμ reflect the properties of H-bonds vibronic states between selected protic cations and ORR intermediates, we reason that the kinetics of PCET relevant step on Au and Pt can be tuned by altering H-bond structures at the interface. This concept reflects the “Polanyi rules” discovered in fundamental reaction dynamic studies, which describes that vibrational energies play the most important role in the late-barrier atom-diatom reactions (H + X₂ => HX + X, X=Cl, Br)⁷⁷, ⁷⁸. This mechanism stands apart from previous studies on homogeneous reaction rates such as nitrogen reduction⁷⁹, hydrogen evolution⁸⁰,⁸¹ and CO₂ reduction⁸², where pKₐ-dependent kinetics has been attributed largely to the driving force of proton (chemical potential) and electron (electro-potential tuned by redox potential of metal center ) and pKₐ-dependent activation energy. Therefore, we propose that the stretching frequency and PνSνμ² of a H-bond, describing the vibrational features of intermediates, can be considered as descriptors for PCET kinetics. This study highlights the important role of vibrational features on reaction kinetics and new opportunities into electrolyte and interface tuning to control reaction kinetics.

**Conclusions**

This study shows that ORR activity forms a volcano relationship with pKₐ of ion liquids (serving as proton donor) on the surface of Au and Pt in acid. The optimized pKₐ for proton donor is around 15 and close to the pKₐ value of water for Pt while the optimized pKₐ for proton donor on Au is around 11 and close to the pKₐ value of H₂O₂.

*In situ* ATR-SEIRAS provides direct evidence for red-shifted stretching frequency of
X-H with decreasing $\Delta pK_a$, which is associated with enhanced ORR activity. The effect of the H-bond structure on the kinetics of PCET is examined by the Boltzmann probability ($P_\nu$) and vibronic coupling of proton ($S_{\nu\mu}$) in different quantum states. Our results reveal that stronger H-bond can increase proton tunneling kinetics ($P_\nu S_{\nu\mu}^2$) by $\sim 10-10^3$ times whereas the activation free energy ($\Delta G^\ddagger$) of reaction remains largely unchanged. This work shows compelling evidence for $pK_a$-dependent hydrogen bond structures and their impact on the kinetics of proton tunneling and the rate-limiting PCET of ORR on Pt and Au, where altering the $pK_a$ of proton donor at the catalyst surface can change the H-bonding interaction with ORR intermediates. Our findings highlight new opportunities beyond conventional catalyst design strategies of surface electronic structure tuning to control catalytic activity by tuning H-bond structures and/or solvation environments at the electrified interface.

**Material and Methods**

Pt/C and Au/C and Ag/C catalysts:

Pt/C catalysts was supplied by Tanaka Kikinzoku (TKK TEC10E20A), with weight fraction 19%. The Au nanoparticles were synthesized following a reported approach. Briefly, Tetralin (10 mL), Oleylamine (OAm) (10 mL), and HAuCl$_4$·3H$_2$O (0.1 g) were mixed at room temperature and magnetically stirred for 10 min under N$_2$ atmosphere to make precursor solution. Then, the reducing solution of 0.5 mmol of TBAB, tetralin (1 mL), and OAm (1 mL) was injected into the precursor solution, the reaction was carried out with water-ice bath to maintain the temperature at 2 °C for 1 h. After reaction, the nanoparticles were wash with acetone and collected by centrifugation. As prepared Au nanoparticles and carbon black were dispersed in solvent 1:1 isopropanol and hexane by 20-minute ultrasonicate treatment to synthesize the Au/C catalysts, resulting in 30wt.%
Au loading. The products were collected by centrifugation. To remove the residual surfactant, the loaded nanoparticles were heated at 200 °C for 0.5 h under the Ar atmosphere.

Synthesis of ionic liquids

The \([\text{C}_4\text{C}_1\text{im}][\text{NTf}_2]\)(1-butyl-3-methylimidazolium bis(trifluoromethanesulfonylimide) CAS No: 174899-83-3) was purchased from Sigma Aldrich. \([\text{TMPim}][\text{NTf}_2]\) 1,3-Bis(2,4,6-trimethylphenyl)imidazolium bis(trifluoromethanesulfonylimide) was synthesized by the reported anion exchange reaction.\(^{84}\) \(\text{Li}[\text{NTf}_2]\) aqueous solution was added to \([\text{TMPim}]\text{Br}\) slowly, which also dissolved in water. After the reaction, the IL would form a separated phase beneath the water phase. The product was washed by ultrapure water, and add 0.1M \(\text{AgNO}_3\) to test the trace residual \(\text{Br}^-\) in the aqueous layer. The ionic liquid was washed until the \(\text{AgNO}_3\) test was negative. The resulting product was dried by high vacuum at 80°C for 24 h.

\([\text{MTBD}][\text{NTf}_2]\)(1,3,4,6,7,8-Hexahydro-1-methyl-2H-pyrimido[1,2-a]pyrimidinebis(trifluoromethanesulfonylimide), \([\text{DBU}][\text{NTf}_2](2,3,4,6,7,8,9,10-Octahydropyrimidol[1,2-a]azepine \quad \text{bis(trifluoromethanesulfonylimide),}\]
\([\text{DEMA}][\text{NTf}_2](\text{N,N-diethylmethylamine} \quad \text{bis(trifluoromethanesulfonylimide,}\]
\([\text{TEMEDA}][\text{NTf}_2] \quad (\text{N,N,N',N'-tetramethylethlenediamine}\]

\text{bis(trifluoromethanesulfonylimide),} \quad \text{[C}_4\text{Him}][\text{NTf}_2] \quad \text{(1-butyl-imidazolium bis(trifluoromethanesulfonylimide)}\) were prepared through neutralization reactions in accordance with a previous report.\(^{85}\) The MTBD, DBU, DEMA, TEMEDA and C$_4$Him aqueous solution was added to HNTf$_2$ solution slowly with cooling by ice bath. The products would form a separated phase beneath the water phase which was washed by ultrapure water several times and dried by high vacuum at 80 °C for 24 h.
Synthesis of ionic-liquid-modified catalysts

The ionic-liquid-modified catalysts were synthesized by a reported protocol\textsuperscript{47}. Briefly, 90 mg M/C (M=Pt, Au, 19wt.% TKK Pt/C, as prepared 30wt.% Au/C were adopted) was mixed with 10mL iso-propyl alcohol solution containing a calibrated amount of IL (corresponding to a final IL loading of 16 wt%) with stirring at room temperature. After a 20-minute ultrasonic treatment, the solvent was evaporated under a low vacuum condition firstly and then dried under high vacuum at 80 °C for 24 h.

Electrochemical characterization

All the electrochemical measurements were performed using Biologic SP-300 potentiostat with a three-electrode electrochemical system. The working electrode was a glassy carbon rotating disk electrode (0.196 cm\textsuperscript{2}). The slurry was deposited on the working electrode, which consisted of 5 mg M/C (M=Pt, Au) and 5 mL mixture of deionized water, isopropyl alcohol, and 5 wt % Nafion solution in a volume ratio of 4:1:0.025. The loading of Pt was controlled at 20 μg·cm\textsuperscript{-2} and the loading of Au and Ag were controlled at 40 μg·cm\textsuperscript{-2}. The electrolyte was prepared from Milli-Q water (18 MΩ·cm) and HClO\textsubscript{4} (70%, 99.999% trace metals basis). The SCE potential scale was calibrated with the RHE scale using H\textsubscript{2} electro-oxidation. All measurements were collected under Ar or O\textsubscript{2} saturation (ultra-high-grade purity, Airgas). The specific kinetic activity ($J_k$ (mA/cm\textsuperscript{2}Au or Pt)) of Au/C and Pt/C samples was calculated from Koutecky-Levich equation and electrochemical surface area determined by cyclic voltammetry, the detailed methods were depicted in SI.

\textit{In situ} Surface Enhanced Infrared Absorption Spectroscopy (SEIRAS) Experiments.

For the \textit{in situ} SEIRAS measurements, we used a working electrode composed of a
thin (~50 nm) Au or Pt film deposited on the Si prism (radius 22 mm, Pier optics) by
electroless deposition \(^{86, 87, 88}\). The prism was then assembled to a three-electrode cell
along with an Ag/AgCl reference electrode and a platinum wire counter electrode. The
SEIRAS experiments were conducted on a FT-IR Vertex 70 (Bruker) FT-IR equipped
with an MCT detector. The optical path was fully filled with N\(_2\) gas. The FTIR spectra
were acquired in the attenuated total reflection (ATR) mode using a single-reflection
ATR accessory (Pike Vee-Max II, Pike Technologies) at an incident angle of 68 degrees.
The spectral resolution was 4 cm\(^{-1}\) and the scan velocity was 7.5 kHz. Each spectrum
was measured by superimposing 256 interferograms. During each experiment, O\(_2\) was
bubbled through the electrolyte, and the prism surface was then cleaned by cyclic
voltammetry between 0 and 1.0 V\(_{\text{RHE}}\). After cleaning, the spectra were collected while
applying potentiostatic potentials. All spectra were presented in the form of absorbance
according to log(I\(_0\)/I), where I\(_0\) and I are the spectra of background (at open circuit
potential) and at the potentiostatic potential, respectively. Details of \textit{in situ} SEIRAS
were described elsewhere \(^{71, 86, 87}\).

\textbf{FTIR frequency calculations}

The computed the vibrational frequency shifts for hydrogen-bonded species were
simulated by DFT from the vibrational frequencies of individual ionic liquid cations,
and with interaction of individual cations with a H-bond from a nearby OH or OOH, in
an explicit solvation model (PCM), and 2-Pentanone (dielectric constant=15.5) was
used as solvent \(^{89}\). We used the B3LYP functional and 6-311++G** basis set, as
implemented in the Gaussian (g16) suite \(^{90}\).

\textbf{Proton Vibration Wavefunction Calculation}
The vibrational wavefunctions and energies were calculated from a series of proton potentials determined at different H-bonded species calculated for FTIR frequency, which represent the observed interfacial H-bond structure at equilibrium states. The 1D proton potentials were scanned using a reported method \(^7\). Then, the 1D vibrational wavefunctions and energies were calculated from the proton potential by solving the one-dimensional Schrödinger equation numerically using the Fourier Grid Hamiltonian Multiconfigurational Self-Consistent-Field (FGH-MCSCF) method \(^5, 91, 92\). The Boltzmann probability \(P_\nu\) was calculated from energies of different states and the \(S_{\nu\mu}\) was obtained by integrating the overlap of reactant and product’s wavefunction\(^{69,93}\).

The calculation details was presented in Supplementary methods.

**Author contributions:** Y.S.H. and T.W conceived the idea and designed the experiments. T.W. carried out experiments on catalyst synthesis, electrocatalytic tests and data analysis. Y.Z. and T.W. performed the \textit{in situ} surface-enhanced FT–IR spectroscopy measurements. Y.Z. and T.W performed the DFT calculations and analysis. T.W. and B.H. analyzed electrochemical data. B.C. synthesized nanoparticles and performed TEM test. R.R.R., L.G. and S.G.S. participated in the discussion and interpretation of experimental and theoretical data. Y.S.H. and T.W. wrote the manuscript. All of the authors discussed the results and commented on the manuscript.

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Scheme 1. Tuning local proton activity for ORR catalyzed on the surface of metallic catalysts by adding a thin layer of protic ionic liquids, where the $pK_a$ value can range from 7.1 to 23.3\textsuperscript{51, 52, 53, 54, 55}. The anion of all ionic liquids used was NTf$_2$ in this work.
**Fig.1** $pK_a$-dependent ORR activity on Au/C and Pt/C measured in O$_2$-saturated 0.1 M HClO$_4$, with a scan rate of 10 mV/s and the rotation speed is 1600 rpm. (a and b) Background and IR corrected ORR polarization curves of ionic-liquids-modified Au/C (a), Pt/C (b). (c and d) The relationship between the enhancement in ORR specific kinetic currents on ionic-liquids-modified Au/C (c), Pt/C (d) as a function of the $pK_a$ value of protic cations in ionic liquids. Error bars represent standard deviations (SDs) of at least three independent measurements. The loading of Pt was controlled at 20 μg·cm$^{-2}$ and the loading of Au were controlled at 40 μg·cm$^{-2}$. 0.05wt% Nafion was added to the catalytic layer.
Fig. 2 *In situ* ATR-SEIRAS measurement on a [MTBD][NTf$_2$]-modified Au electrode in oxygen-saturated 0.1 M HClO$_4$. (a) C=N stretching region, (b) O-O-H bending region and (c) X-H (X=N, O) stretching region were obtained during potential steps swept from 0.2 V$_{RHE}$ to 0.6 V$_{RHE}$ in 0.1 M HClO$_4$. The blue curves show the IR spectrum of pristine [MTBD][NTf$_2$] and the purple curve shows the IR spectrum of pristine [MTBD][NTf$_2$] with 0.5M water. The cumulative number of 256 was used at a 4 cm$^{-1}$ resolution. Spectra were subtracted with respect to a reference spectrum obtained at OCV in 0.1 M HClO$_4$. 
Fig. 3 *In situ* ATR-SEIRAS measurement on an ionic-liquid-modified Au and Pt electrode in 0.1 M HClO₄. X-H (X=N, O) stretching region of (a) [MTBD][NTf₂]-modified Au, (b) [DEMA][NTf₂]-modified Au, (c) [C₄Him][NTf₂]-modified Au, (d) [MTBD][NTf₂]-modified Pt, (e) [DEMA][NTf₂]-modified Pt and (f) [C₄Him][NTf₂]-modified Pt. The spectra were acquired at 0.2 V_RHE for Au and 0.5V_RHE for Pt. The peak
position of the H-bonded species indicates the strength of H-bond forming during the ORR process; (g, h) the relationship between the stretching frequency of the H-bonding species and the ionic liquid enhancement for ORR catalyzed by Au/C (g) and Pt/C (h), which is defined by \( \frac{J_{\text{IL-}M/C}}{J_{M/C}} \) (\( J_{\text{IL-}M/C} \) is the ORR current density of ionic-liquid-modified sample, \( J_{M/C} \) is the current density of pristine Pt/C or Au/C). Error bars represent SDs of at least three independent measurements. The relationship indicates stronger H-bond would lead to higher ORR activity. The curve only serves as a guide to eyes.
Fig. 4 Evaluating the structural effect of interfacial H-bond by calculating the vibrational wavefunction and energy levels for proton vibrational states of H-bonded species. (a) A scheme of the PCET reaction at H-bonded interface. The red and blue curves depict proton potential and proton wavefunction (shadowed curve) in ground state of reactant and product as functions of the proton coordinate. Proton is close to the proton donor in the reactant. After the H-bonded species get one electron from metal electrode, it will transfer to the proton acceptor in the product. The reaction kinetic is related with the integral overlap of wavefunctions between the reactant and product (denoted as $S_{\nu\mu}$) and Boltzmann probability of different vibronic states (denoted as $P_{\nu}$). (b) the expression of current density exhibits that the $J_0$ (exchange current density) is proportional to $P_{\nu}S_{\nu\mu}^2$, which is dominated by H-bonding structure. (c, e) Schematic depiction of free energy surface for PCET reaction in $\text{NH}^+\cdots\text{OHH}_{\text{Au}}$ and $\text{NH}^+\cdots\text{OH}_{\text{Pt}}$. The Marcus parabolas of reactant (red) and product (blue) in the center frame illustrate the nonadiabatic transitions occur at the collective solvent coordinates across the
intersection (black points) of reactant and product parabolas. Proton potentials and proton wavefunctions in contributing state of reactant and product as functions of the proton coordinate are depicted in left (reactant) and right (product) frame. For NH$^+\cdots$OOH$_{\text{Au}}$ (c), the reactant ground state is the contributing state. For NH$^+\cdots$OH$_{\text{Pt}}$ (e), the 1$^\text{st}$ state of reactant is the contributing state. (d, f) The $P_{\nu'S'\mu'}$ (in light color) of the contributing state for different ionic liquid cations interacting with OOH (d) and OH (f), comparing with the $J_{0,s}$ (specific exchange current density) (in dark color) extracted from Butler-Volmer equation for ORR on Pt (E, $\alpha$ is fixed to 1)$^{56,91}$ and Au (E, $\alpha$ is fixed to 0.4).