# MATERIALS SCIENCE

# Surface molecular pump enables ultrahigh catalyst activity

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The performance of electrocatalysts is critical for renewable energy technologies. While the electrocatalytic activity can be modulated through structural and compositional engineering following the Sabatier principle, the insufficiently explored catalyst-electrolyte interface is promising to promote microkinetic processes such as physisorption and desorption. By combining experimental designs and molecular dynamics simulations with explicit solvent in high accuracy, we demonstrated that dimethylformamide can work as an effective surface molecular pump to facilitate the entrapment of oxygen and outflux of water. Dimethylformamide disrupts the interfacial network of hydrogen bonds, leading to enhanced activity of the oxygen reduction reaction by a factor of 2 to 3. This strategy works generally for platinum-alloy catalysts, and we introduce an optimal model PtCuNi catalyst with an unprecedented specific activity of  $21.8 \pm 2.1 \, \text{mA/cm}^2$  at  $0.9 \, \text{V}$  versus the reversible hydrogen electrode, nearly double the previous record, and an ultrahigh mass activity of  $10.7 \pm 1.1 \, \text{A/mg}_{Pt}$ .



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#### INTRODUCTION

Improvements in performance of costly catalysts (e.g., platinum) are critical for the storage and conversion of renewable clean energy (1). Intensive efforts have been devoted to searching for catalysts with higher catalytic activity (2-5) and longer-term stability (6, 7). A generally adopted strategy is through the strain effect (2, 8) or the ligand effect (3, 9) to achieve an optimal binding energy between catalyst surface and reaction species, which can lower the overpotential of the reaction and enhance the activity. Despite considerable progress made so far, further enhancement is required. Meanwhile, such guidance has been largely motivated by theoretical considerations in vacuum, while microkinetics at the catalystelectrolyte interface was rarely taken into quantitative consideration. A long-standing issue for many electrochemical reactions is the insufficient reactant supply at the catalyst surface. For example, the low oxygen solubility  $(2.56 \times 10^{-4} \text{ M})$  in 0.1 M HClO<sub>4</sub> and small equilibrium constants for initial physisorption (10, 11) impede the already sluggish oxygen reduction reaction (ORR) kinetics. Quantitative understanding of such solid-electrolyte interfaces with gases and other reactants can be gained through simulations with the INTERFACE force field (IFF) (12) which reaches a high accuracy of ~0.03 kcal/mol (11, 13). Advances to address this challenge in experiments have been made through the development of a high oxygen permeability ionomer (14) or hydrophobic ionic liquids with a higher bulk O<sub>2</sub> solubility (ca. 2.4 times compared to that in 0.1 M HClO<sub>4</sub>) (15–17) with success, although at a higher developmental or material cost. In parallel, timely removal of reactant water is of equal importance to accelerate the reaction (18, 19), following Le Chatelier's principle. Recently, the addition of cations (20), pyridine (21), melamine (22), melamine-formaldehyde polymer (23), and tetrahexylammonium cations (24) in the electrolyte has been shown to effectively improve the ORR kinetics of pure Pt. These studies attributed the activity enhancement to the destabilization or weakening of OH<sub>ad</sub> adsorption on the Pt surface, which promotes the ORR activity following the Sabatier principle, whereby final protonation could be the rate-determining step. However, such ligand enhancement mechanisms remain elusive to verify, given the weak attraction of oxygen to Pt in aqueous solution (11), and more in-depth analysis is necessary.

Herein, we demonstrated that a ligand can act as a surface molecular pump to substantially accelerate reactant supply and product removal in ORR (Fig. 1) (25). We propose a concept of "molecular pump" to illustrate the net effect of dimethylformamide (DMF) surface modification, showing that DMF pulls in more O<sub>2</sub> through increased physisorption of O<sub>2</sub> on the surface and releases more H<sub>2</sub>O per unit time (decreased H<sub>2</sub>O adsorption time). The local Pt-O<sub>2</sub> concentration,  $c_{Pt-O2}$ , could be approximately doubled and the surface molecular pump allows for reduced surface water concentration and retention time, leading to faster outflux of water. Together, the surface molecular pump can further enhance the specific activity (SA) of electrocatalytic reactions (e.g., ORR). Our experimental studies revealed that adsorbed DMF molecules, as a surface molecular pump, can effectively improve the SA of Pt/C by twofold in 0.1 M HClO<sub>4</sub> electrolyte. Our molecular dynamics (MD) simulations with the IFF suggest that physisorption of DMF on the Pt surface can accelerate O<sub>2</sub> transport from the electrolyte toward the Pt surface, lower the oxygen Gibbs physisorption energy and prolong the oxygen contact time. DMF thus greatly affects the physisorption behavior of O<sub>2</sub> toward Pt, which was recently related to the overall chemical reaction process (11). Reactions require milliseconds at the nanometer scale to occur and therefore strongly depend on preceding physisorption equilibria (Fig. 1). Because of a much shorter timescale of nanoseconds and low binding free energies of O<sub>2</sub> below -0.6 kcal/mol (-0.025 eV), physisorption has an immense effect on the interfacial microenvironment, which

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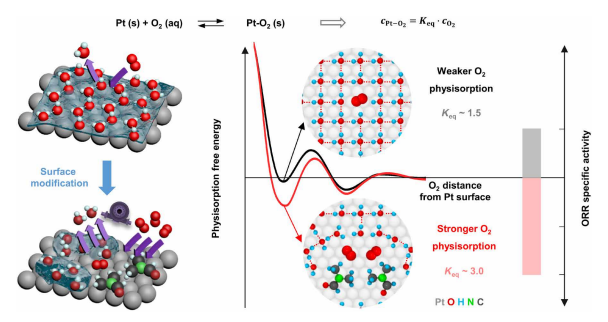


Fig. 1. Schematic illustration of the boosted ORR kinetics by surface modification of Pt. The application of DMF as an oxygen trap and a surface molecular pump can facilitate higher interfacial oxygen concentration ( $c_{Pt-O2}$ ) and lower interfacial water concentration by breaking the network of hydrogen bonds in water adlayers on the Pt surface (left). The surface molecular pump lowers the physisorption free energy of  $O_2$  on the surface and shifts the physisorption equilibrium constant toward higher ORR specific activity (right, length of light red bar). The specific activity is shown in arbitrary units for simplicity and a color legend for the atoms is included on the lower right.

directly correlates with the ORR kinetic current (11, 26). Our MD simulations show that DMF physisorption on the Pt surface leads to increased interfacial O<sub>2</sub> concentration and reduced water concentration on the Pt surface, which collectively boosts the ORR activity by way of Le Chatelier's principle (Fig. 1). We further applied this approach to a model Pt-alloy catalyst (PtCuNi), achieving an unprecedented ORR SA of  $21.8 \pm 2.1 \,\text{mA/cm}^2$  at 0.9 V versus the reversible hydrogen electrode (RHE), which nearly doubles the previously reported record in the literature (27), leading to an ultrahigh mass activity (MA) of  $10.7 \pm 1.1 \,\text{A/mg}_{Pt}$ .

# **RESULTS**

# Surface modification of commercial Pt/C catalysts

We first used Pt/C catalysts [40 weight % (wt %) Pt, HiSPEC 4000, Johnson Matthey] as a model system to explore the effect of surface molecular pumps. DMF was chosen as the surface ligand due to its stability in a wide potential window (28) and the strong physisorption of amides in close contact following a soft epitaxial pattern, which has been previously described (29). We treated the Pt/C catalyst in a pure DMF solution at 120°C for a varied period of time to allow gradual molecular adsorption on the surface (see the Supplementary Materials). We chose the elevated temperature because it can equilibrate an even distribution of DMF on the catalyst surface faster than at room temperature and enhance the sampling of surface coverage with DMF over a wide range of values. Cyclic voltammetry (CV) revealed that the Pt/C catalysts exhibited an initial electrochemically active surface area (ECSA) (based on hydrogen underpotential deposition,  $H_{upd}$ ) of 51.6  $\pm$  1.5 m<sup>2</sup>/g<sub>Pt</sub>, which decreased to  $21.8 \pm 0.2 \,\mathrm{m^2/g_{Pt}}$  after 48 hours of modification evidenced by a decreased H<sub>upd</sub> area (Fig. 2A). The size of Pt catalysts and carbon support did not show noticeable size changes upon surface

modification (figs. S1 and S2), indicating that the reduction of ECSA upon treatment can be attributed to the blocking of available Pt surface sites with increasing DMF surface coverage (defined as the reduced ECSA: the decreased  $H_{\rm upd}$  region normalized by the original  $H_{\rm upd}$  region of the catalyst, Fig. 2B). Our x-ray photoelectron spectroscopy (XPS) studies (fig. S3) revealed that the presence of N-species increased with longer modification time, indicating an increased amount of adsorbed DMF, which is consistent with the decreased ECSA (Fig. 2B). We did not observe apparent shift in the XPS peak position for both  $4f_{7/2}$  and  $4f_{5/2}$  peaks, suggesting that surface-adsorbed ligands do not induce notable electronic effects on Pt (fig. S3B).

We further evaluated the ORR activities of Pt/C catalysts using the linear sweep voltammetry (LSV) method (Fig. 2C). The SA of Pt/C catalysts increased from 0.48  $\pm$  0.06 to 0.89  $\pm$  0.04 mA/cm<sup>2</sup> after 24 hours of modification (surface coverage of 37.1%) and then decreased to  $0.55 \pm 0.02 \text{ mA/cm}^2$  after 48 hours of modification (surface coverage of 58.0%) (Fig. 2D). Accordingly, the MA of Pt/C catalyst increased from  $0.23 \pm 0.03$  to  $0.28 \pm 0.03$  A/mg<sub>Pt</sub> after 24 hours of modification (Fig. 2D). The electrochemical impedance spectroscopy did not show the ohmic resistance change for the catalyst before and after surface modification (fig. S4). We also used DMF to treat Pt disk electrode and found similar activity enhancement. In particular, we dropped 10 µl of 80°C DMF on a 5-mm-diameter Pt disk electrode and dried it in an 80°C oven. Then, we used ethanol to wash away residual DMF. Similarly, the CV curves showed a 22% areal decrease at the hydrogen underpotential region upon DMF treatment (fig. S5A). The half-wave potential of the surface-treated Pt disk is 0.818 V (vs. RHE), which is 44 mV larger than that of the untreated Pt disk (0.774 V) (fig. S5B). The SA of the surface-treated Pt disk is 0.53 mA/cm<sup>2</sup> (0.9 V versus RHE), which is 3.6 times higher than that of the untreated Pt disk (0.15 mA/

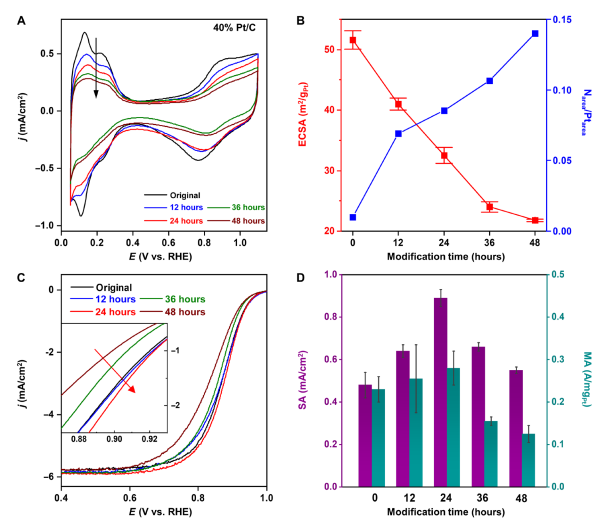


Fig. 2. Electrochemical measurements in 0.1 M HClO<sub>4</sub> electrolyte of the 40% Pt/C catalysts modified with DMF at different treatment times. (A) CV curves of Pt/C catalysts at the original stage (black) and after 12 hours (blue), 24 hours (red), 36 hours (green), and 48 hours (wine) of modification time. (B) ECSA and N/Pt ratio obtained from XPS peak integration as a function of modification time. (C) ORR LSV curves of Pt/C catalysts at the original stage (black) and after 12 hours (blue), 24 hours (red), 36 hours (green), and 48 hours (wine) of modification time. The inset of (C) is the enlarged potential window. (D) The evolution of SA (purple, left y axis) and MA (olive, right y axis) with modification time. The SAs of different stages:  $0.48 \pm 0.06$  mA/cm<sup>2</sup> (0 hours),  $0.64 \pm 0.03$  mA/cm<sup>2</sup> (12 hours),  $0.89 \pm 0.04$  mA/cm<sup>2</sup> (24 hours),  $0.66 \pm 0.02$  mA/cm<sup>2</sup> (36 hours), and  $0.55 \pm 0.02$  mA/cm<sup>2</sup> (48 hours). The MAs of different stages:  $0.23 \pm 0.03$  A/mg (0 hours),  $0.26 \pm 0.08$  A/mg (12 hours),  $0.28 \pm 0.04$  A/mg (24 hours),  $0.16 \pm 0.01$  A/mg (36 hours), and  $0.13 \pm 0.02$  A/mg (48 hours).

cm<sup>2</sup>) (fig. S5C). These intriguing enhancements of both SAs and MAs motivated us to conduct multi-scale computational studies to understand the role of DMF in modulating the ORR activity at catalyst-electrolyte interfaces.

### MD simulations of kinetics at catalyst-electrolyte interfaces

We carried out MD simulations with the IFF (12) (see Supplementary Materials for simulation details) on a representative Pt(111) slab in a 0.1 M HClO<sub>4</sub> electrolyte including explicit solvent molecules and ions to compare the dynamics and local concentration of O<sub>2</sub> and H<sub>2</sub>O at different DMF surface coverages from 0 to 100% over 50-ns timescales (fig. S6). IFF MD simulations for catalystelectrolyte interfaces were earlier shown to be of high reliability and efficient, suitable for analyzing realistically sized nanostructures and their dynamics (11). Without the addition of DMF (Fig. 3A), adsorbed O<sub>2</sub> molecules on the Pt(111) surface were surrounded by

approximately seven to eight  $H_2O$  molecules, forming a dense network of hydrogen bonds (blue circles). The circles with decreasing blue segments indicate that the number of  $H_2O$  molecules surrounding the  $O_2$  molecules adsorbed onto Pt considerably decreased for higher coverage with DMF molecules (Fig. 3, A to C). DMF molecules preferred to be flatly adsorbed on the Pt(111) surface with a physisorption energy of  $-17.3 \pm 2$  kcal/mol in an aqueous solution (Fig. 3, B and C, and figs. S6, B to E, and S7A), and  $-37.5 \pm 1$  kcal/mol in vacuum (fig. S7D). Both energy differences support the strong binding of DMF to Pt(111) and the permanent replacement of  $H_2O$  molecules on the surface, which have a lower adsorption energy of  $-6.6 \pm 1$  kcal/mol on the Pt(111) surface in vacuum and create the characteristic adlayer structure of  $H_2O$  in solution. It is unlikely that DMF is removed from the surface.

When approaching the catalyst surface to gain direct access to the metal, O<sub>2</sub> molecules have to overcome free energy barriers to

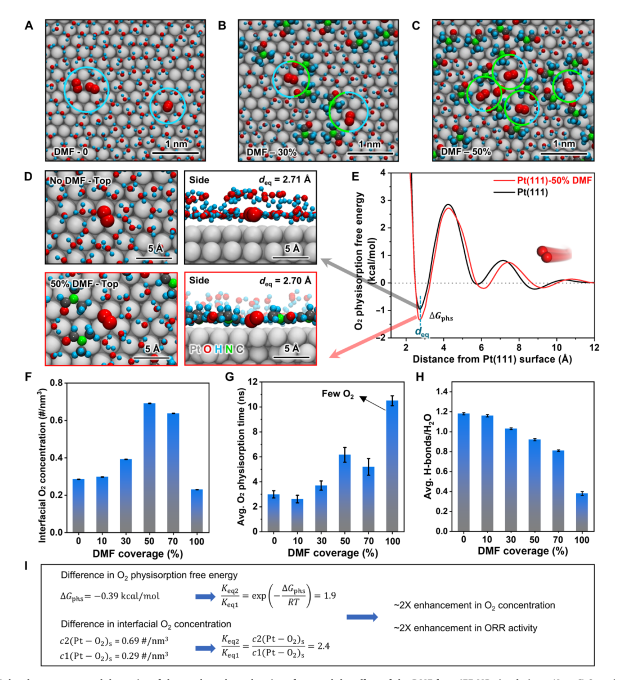


Fig. 3. Molecular structure and dynamics of the catalyst-electrolyte interface, and the effect of the DMF from IFF MD simulations. (A to C) Snapshots of the molecular conformations of  $O_2$ , water, and DMF on the Pt(111) surface at a DMF coverage of (A) 0%, (B) 30%, and (C) 50%. The circles represent the surrounding  $H_2O$  (blue) and DMF methyl groups (green) of adsorbed  $O_2$  molecules. (D) Top and side views of the Pt(111) and Pt(111)–50% DMF surface, showing how DMF disrupts the water adlayer integrity and  $O_2$  molecules are attracted by vdW interactions to both DMF and Pt. (E) Comparison of  $O_2$  physisorption free energy on Pt(111) and Pt(111)–50% DMF surface from SMD simulations.  $d_{eq}$  is the equilibrium distance for  $O_2$  contact with the Pt(111) surface. (F) The relationship between DMF surface coverage and interfacial  $O_2$  concentration (number of molecules/nm³). The cutoff of the interfacial layer was set as 0.5 nm from the Pt surface (30). (G) Average physisorption time of  $O_2$  molecules on the Pt(111) surface (cutoff, 3.5 Å). (H) Average number of H-bonds per  $H_2O$  molecule on the Pt(111) surface (cutoff, 3.5 Å). (I) Two methods of estimating the change of equilibrium constants  $K_{eq}$  and implications for the observed relative ORR activity. The upper equations show the change in the physisorption free energy  $\Delta G$  for 0 and 50% DMF coverage, and the lower equations show the change in the interfacial  $O_2$  concentration for 0 and 50% DMF coverage. Ions in the electrolyte are hidden for clarity. Error bars represent the standard error of mean (SEM).

penetrate the hydrogen-bonded network of water molecules on the Pt surface and replace between one and two surface-bound H<sub>2</sub>O molecules (Fig. 3, A, D, and E) (11). In the presence of the molecular pump DMF, O<sub>2</sub> molecules can more conveniently reach the Pt surface near the methyl groups of DMF molecules, enabling multiple van der Waals (vdW) interactions that increase O<sub>2</sub> interaction with and retention on the Pt surface (Fig. 3, B to E). The free energy barriers for physisorption are hereby small enough (<3 kcal/mol or <0.13 eV) and crossed within hundreds of picoseconds, without an effect on the statistical physisorption equilibrium beyond nanoseconds (Fig. 3E).

Thus, we analyzed the equilibrium properties such as the interfacial concentration and physisorption time of O<sub>2</sub> and H<sub>2</sub>O (figs. S8 and S9). Within an interfacial layer of 0.5 nm thickness (30), the O2 concentration increased from 0.29 to 0.69 nm<sup>-3</sup>, i.e., by a factor of 2.4, with a corresponding increase in DMF coverage from 0 to 50%; and decreased to 0.64 nm<sup>-3</sup> when DMF coverage reached 70%, then to 0.23 nm<sup>-3</sup> at 100% coverage (Fig. 3F). The average physisorption time of O<sub>2</sub> on the Pt (111) surface was prolonged to 6.2 ns at 50% DMF coverage compared to 3.0 ns for the pure Pt(111) surface (Fig. 3G), corresponding to an increase by a factor of 2.1. The average physisorption time at 10% DMF coverage (2.6  $\pm$  0.3 ns) is slightly lower than that at 0% DMF coverage (3.0  $\pm$  0.3 ns), which is within the statistical uncertainty of the simulation, and a clear trend toward higher coverage is seen. At 70% DMF coverage, a fraction of O<sub>2</sub> molecules has much-prolonged physisorption times (>20 ns) due to vdW interactions with DMF, and 47% of O2 molecules had barely any contact with the Pt(111) surface (fig. S8). As a result, the average O<sub>2</sub> physisorption time at 70% DMF coverage (5.2 ns) is shorter than that at 50% DMF coverage (6.2 ns) (Fig. 3G). Furthermore, 100% DMF coverage blocked the adsorption of O2, leading to a markedly decreased percentage of adsorbed O<sub>2</sub> (fig. S8). However, DMF molecules cannot fully cover the surface due to strong packing constraints and leave adequate spaces that can still trap a few O<sub>2</sub> molecules (figs. S6F, S8, and S10F). The exceptional attraction of few O<sub>2</sub> molecules with adsorption time of more than 40 ns resulted in an anomalous large average adsorption time of 10.85 ns (averaged among adsorbed O<sub>2</sub> molecules) at 100% DMF coverage (Fig. 3G and fig. S8).

In addition to local oxygen entrapment by increasing the DMF coverage from 0 to 100%, the interfacial H<sub>2</sub>O concentration decreased from 30.1 to 6.0 nm<sup>-3</sup> within the 0.5-nm surface layer (fig. S9A) and the average physisorption time of H<sub>2</sub>O declined markedly from 3.5 to 0.3 ns (fig. S9B). This change signifies that the surface molecular pump also facilitated product (water) removal, thereby speeding up the ORR rate. The average number of H-bonds per H<sub>2</sub>O molecule within the adlayer (see the Supplementary Materials for simulation details) also showed a continuous decrease from 1.18 to 0.38 with increasing DMF coverage, indicating a notable reduction of the H-bond network due to DMF functioning as a molecular pump (Fig. 3H). The analysis of hydrogen bonds assumes a maximum O(H) ··· O distance of 3.5 Å and a maximum deviation of 30° from the linear (H)O-H  $\cdots$  O(H<sub>2</sub>) angle (31). The average number of H-bonds per water molecule was calculated by dividing the total number of formed H-bonds by the total number of H<sub>2</sub>O molecules in the adlayer. In addition, the configuration of the adlayer network of water (blue shade in fig. S10) shows a sparser and increasingly incomplete pattern when the Pt(111) surface coverage by DMF increases from 0 to 100%, providing more space near the molecular

pump DMF for O2 adsorption (pale violet red shade). Our surface wettability test results showed that the contact angle increased with increasing treatment time for both Pt/C and PtCuNi catalysts (fig. S11), which indicated that the catalyst surface became more hydrophobic. This finding is consistent with our hypothesis that a modified catalyst surface facilitates water removal, thus favorably shifting reaction kinetics and improving ORR. We also investigated the interfacial  ${\rm H_3O}^+$  concentration in MD simulations (fig. S12). The interfacial  ${\rm H_3O}^+$  concentration gradually decreases from 0.55 to 0.17 nm<sup>-3</sup> with an increase of DMF coverage, which indicates a slower proton transport process due to the DMF attachment. It is worth noting that the interfacial H<sub>3</sub>O<sup>+</sup> concentration (0.49 nm<sup>-3</sup>) only declined by 10% at 30% DMF coverage. However, the interfacial H<sub>3</sub>O<sup>+</sup> concentration substantially decreased by 35% when DMF coverage reached 50%. In the experiments, we found that both Pt/C and PtCuNi catalysts first showed an activity increase and then an activity decrease. Our results indicate that the change of O<sub>2</sub> physisorption dynamics is the dominant factor in enhanced ORR activity. Another possible factor for the activity decrease could be reduced hydronium conduction; however, we do not expect that the proton-electron transfer process will be strongly influenced while the DMF surface coverage is less than 50%, or even somewhat higher.

To confirm the more favorable O<sub>2</sub>-DMF configurations compared to O<sub>2</sub>-H<sub>2</sub>O configurations near the Pt surface, we further analyzed the stabilization energy (IFF-based simulations) of small clusters consisting of Pt, DMF, H<sub>2</sub>O, and O<sub>2</sub>, which represent the local environment of molecular pumps at the catalyst-electrolyte interface (fig. S13, A and B). The simulations with IFF have a unique accuracy of better than 0.03 kcal/mol (13) and the results corroborate that an increasing number of DMF molecules in the cluster increase the oxygen stabilization energy (fig. S13C). Hereby, coordination of O2 with one DMF "pump" molecule replaces approximately two H<sub>2</sub>O molecules and increases the stabilization energy by  $-0.24 \pm 0.02$  kcal/mol (fig. S13D). Therefore, DMF stabilizes O<sub>2</sub> physisorption on Pt and, at the same time, destabilizes H<sub>2</sub>O physisorption, which leads to a higher interfacial O<sub>2</sub> concentration and lower interfacial H<sub>2</sub>O concentration, creating a surface molecular pump that accelerates the ORR kinetics (32).

In particular, the MD simulations of Pt(111) with 50% DMF (Fig. 3, D and E) showed the highest interfacial O<sub>2</sub> concentration  $(0.69 \text{ nm}^{-3})$  within a 0.5 nm distance from the surface (30) and a long O<sub>2</sub> physisorption time (6.2 ns), equal to a higher equilibrium concentration of O2 on the Pt surface and leading to accelerated ORR kinetics. The expected increase in the negative physisorption free energy of O<sub>2</sub> in the presence of DMF is quantitatively shown by the free energy profiles of O<sub>2</sub> molecules using steered molecular dynamics (SMD) simulations for two control cases of 0 and 50% DMF coverage (Fig. 3, D, E, and I, and figs. S14 and S15), respectively. The simulation models consider full hydration of oxygen molecules while traveling from solution to the Pt surface, as well as changes in hydration upon proximity to ions in solution or coordination with DMF on the surface, and realistic concentrations of all present electrolytes. The distance of the O<sub>2</sub> molecule from the Pt surface is measured between the center of mass of an  $O_2$  molecule and the z coordinate of surface layer Pt atoms. Hereby, 10 replicas with a slow pulling speed of 0.01 Å/ns and a total simulation time of 4 μs were used to fully sample the oxygen interactions at the interface in high accuracy (see Materials and Methods). The mobility of DMF molecules, such as surface hopping and in-plane rotation, and si-

multaneous rearrangements of the networks of hydrogen-bonded water molecules notably affect the  $O_2$  dynamics at distances < 0.6 nm from the surface. The O<sub>2</sub> molecules ultimately adsorbed to Pt (111) in coordination with DMF molecules in all replicas of the SMD simulations, showing dynamic O<sub>2</sub>-DMF coordination numbers, and that coordination with DMF molecules is the preferred position for O<sub>2</sub> adsorption (Fig. 3, B to D). These results are consistent with independent IFF-based calculations of O2 physisorption energies to small Pt(111)/DMF/water clusters, whereby an increase in O2 coordination number with DMF from 0 to 4 increased the net stabilization energy from 0 to -0.96 kcal/mol (fig. S13D). The global free energy minimum represents direct contact of O<sub>2</sub> with the Pt surface and was found at 2.72 Å distance from the Pt(111) surface atoms ( $d_{eq}$ ), and the total physisorption free energy ( $\Delta G$ ) of O<sub>2</sub> was, as expected, lower at -1.34 kcal/mol for Pt(111)-50% DMF compared to -0.95 kcal/mol for pure Pt(111) in water (Fig. 3E). The gain in O<sub>2</sub> physisorption free energy of -0.39 kcal/mol in the presence of DMF (denoted as  $\Delta G_{\text{phs}}$  in Fig. 3, E and I) is consistent with the independent IFF-based calculations of the gain in O<sub>2</sub> physisorption energies to small clusters (fig. S13D). Although the small cluster estimates are somewhat simplistic without entropy contributions and full water dynamics, both results affirm that DMF promotes the adsorption of O<sub>2</sub> near the Pt(111) surface. The peaks at ~4.2 Å represent the energy barriers associated with O2 penetration through the network of Hbonds in the closest adlayer of H<sub>2</sub>O molecules and the simultaneous replacement of H<sub>2</sub>O by O<sub>2</sub>. The maximum energy barrier for the transport of O<sub>2</sub> from bulk electrolyte (at infinity) to the surface is lower for the 50% DMF-modified surface. The physisorption free energy ( $\Delta G_{\rm phs}$ ) is directly related to the thermodynamic equilibrium constant  $\left[K_{\rm eq}=\exp\left(-\frac{\Delta G_{\rm phs}}{RT}\right)=\frac{c({\rm Pt-O_2})_s}{c({\rm O_2})_{\rm aq}}\right]$  (Figs. 1 and 3I). Accordingly, an increase in  $K_{eq}$  by a factor of 2 using DMF can effectively increase the availability of adlayer  $O_2$  for ORR, c(Pt -O<sub>2</sub>)<sub>s</sub>. Although the enhanced O<sub>2</sub> coverage is still low on an absolute scale, a small difference of 1 RT (~0.59 kcal/mol at 298.15 K) can change the coverage by approximately 2.7 times. The difference in free energy minima of  $\Delta G_{\rm phs}$  of -0.39 kcal/mol (Fig. 3E) increases the equilibrium constant  $K_{\text{eq}}$  by a factor of ~2.0  $\left[\frac{K_{\text{eq2}}}{K_{\text{eq1}}} = \exp\left(-\frac{\Delta G_{\text{phs}}}{RT}\right)\right]$ semiquantitatively matching the enhancement factor in ORR SA observed in the experiments (from 0.48 to 0.89 mA/cm<sup>2</sup>, a factor of 1.9) (Figs. 2D and 3, F, G, and I). This enhancement is also independently validated by the interfacial O2 concentration results (Fig. 3, F and I), where the interfacial O<sub>2</sub> concentration at 50% DMF coverage is 2.4 times of that on pure Pt(111), equal to an increase of the equilibrium constant  $K_{\rm eq}$  by a factor of ~2.4 according to the definition  $\left[\frac{K_{\text{eq}2}}{K_{\text{eq}1}} = \frac{c2(\text{Pt-O}_2)_s}{c1(\text{Pt-O}_2)_s}\right]$ . This order of magnitude, while comparable to thermal motion, is statistically significant and critical over time like in other equilibrium and kinetic processes, such as conformation equilibria in chain molecules (33). The interfacial O<sub>2</sub> concentration begins to decrease when the DMF coverage reaches 70%, which is attributed to a substantially reduced number of available Pt surface sites for O<sub>2</sub> adsorption. These observations agree with our experimental results and the observation that excessive surface modification will lead to a lower activity (Fig. 2D)

## The universality of molecular pumps on catalyst surfaces

Our experimental results and MD analyses suggested that the proper surface molecular pump on Pt facilitates the influx of O<sub>2</sub> and outflux of water, which leads to increased O2 surface concentration and reduced H<sub>2</sub>O concentration, and sustainably higher ORR activity. We note that this surface molecular pump follows Le Chatelier's principle, which is fundamentally different and may work independently from strategies used in modifying the electronic structures of the catalyst surface. Therefore, we expect that such effects can work synergistically to enhance the SA of Pt-alloy catalysts beyond the limits using the Sabatier principle. To this end, we prepared the PtCuNi catalyst using a modified method reported previously (36) and applied the same surface modification protocol. The overall average size, composition, and elemental distribution did not show noticeable change before and after modification (figs. S16, A to E, and S17, A to E). Similar to Pt/C, the ECSA of modified PtCuNi catalysts decreased from 70 to 49 m<sup>2</sup>/g<sub>Pt</sub> after 24 hours of modification and further decreased to 36 m<sup>2</sup>/g<sub>Pt</sub> after 48 hours of modification (Fig. 4A and fig. S18A). We noted that the morphology, size, composition, and structures of the modified catalysts remained overall unchanged compared to the original PtCuNi catalysts (figs. S19, A to E, and S20, A to E), suggesting that the decreased ECSA is dominantly attributed to increased DMF coverage, which is consistent with the Pt/C model system discussed above (Fig. 2A).

We further evaluated the ORR activity of surface-modified Pt-CuNi catalysts (fig. S18B) and found that the optimal surfacemodified PtCuNi catalysts exhibited an unprecedented high SA of  $21.8 \pm 2.1 \text{ mA/cm}^2$  and an ultrahigh MA of  $10.7 \pm 1.1 \text{ A/mg}_{Pt}$  at 0.9 V versus RHE (Fig. 4A), which are ca. 2.6 times and 1.9 times those of original PtCuNi catalysts  $(8.2 \pm 0.4 \,\mathrm{mA/cm}^2)$  and  $5.7 \pm 0.3 \,\mathrm{A/cm}^2$ mg<sub>Pt</sub>) and ca. 45.4 times and 46.5 times those of Pt/C catalysts  $(0.48 \pm 0.06 \,\mathrm{mA/cm^2})$  and  $0.23 \pm 0.03 \,\mathrm{A/mg_{Pt}}$ . The half-wave potential of surface-modified PtCuNi is 0.952 V versus RHE, which is 19 mV higher than that of unmodified PtCuNi (0.933 V versus RHE). The SA of our surface-modified PtCuNi catalysts outperforms all current state-of-the-art catalysts and nearly doubles the SA performance of previously reported J-PtNWs (11.5 mA/cm<sup>2</sup>) (27) (Fig. 4B and Table 1). We also evaluated the SA at 0.95 V versus RHE and found that the activity trend remained unchanged. The optimal surface-modified PtCuNi catalysts still showed the highest SA of  $1.67 \pm 0.04 \text{ mA/cm}^2$ , which is about 2.78 times that of unmodified PtCuNi  $(0.6 \pm 0.02 \text{ mA/cm}^2)$  (fig. S18C). The enhancement factor is close to the 2.6 times evaluated at 0.9 V versus RHE.

We further evaluated the durability of the surface-modified Pt-CuNi catalysts and original PtCuNi catalysts by conducting 20,000 cycles of accelerated degradation test (ADT) and performing postmortem characterizations (figs. S21 to S23). After ADT, the surfacemodified PtCuNi catalysts still showed a highly respectable SA and MA of 12.3 mA/cm<sup>2</sup> and 6.6 A/mg<sub>Pt</sub>, respectively (Fig. 4C and Table 1), exceeding the initial performance of unmodified PtCuNi catalysts, which is also much higher than the as-prepared PtCuNi catalysts after ADT (SA of 5.3 mA/cm<sup>2</sup> and MA of 3.5 A/mg<sub>Pt</sub>). Despite the similar size after ADT (figs. S22 and S23), the surfacemodified PtCuNi catalyst showed ECSA retention of 111%, which is higher than that of the original PtCuNi catalysts (93%). This difference indicates that about 30% of adsorbed DMF was stripped after 20,000 cycles and the remaining DMF still provides considerable activity enhancement. This high number may be associated with partial stripping of the surface-adsorbed molecules (DMF) during

(34, 35).

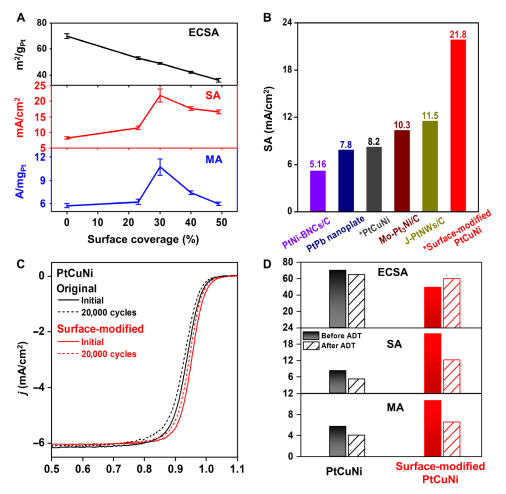


Fig. 4. Performance of PtCuNi catalysts with and without DMF. (A) The evolution of ECSA (black), SA (red), and MA (blue) with DMF surface coverage at 0.9 V versus RHE. (B) Comparison of SA for optimal surface-modified PtCuNi catalysts and state-of-the-art ORR catalysts. References: PtNi-BNCs/C (50), PtPt nanoplate (51), Mo-Pt<sub>3</sub>Ni/C (52), and J-PtNWs (27). \*PtCuNi and \*Surface-modified PtCuNi catalysts are from this work. (C) ORR stability LSV curves for original PtCuNi catalysts (black) and optimal surface-modified PtCuNi catalysts (red) before (black solid line and red solid line) and after (black dashed line and red dashed line) 20,000 cycles. (D) ECSA, SA, and MA comparisons of PtCuNi catalysts and surface-modified PtCuNi catalysts before and after ADT.

**Table 1. RDE performance of surface-modified PtCuNi catalysts in comparison to those in several representative reports.** The row in boldface shows the to-date highest reported specific activity and mass activities, along with the electrochemical active surface area and ADT cycles. BOL, the beginning of life; ADT, accelerated degradation test; NA, not available.

Catalyst	SA (mA/cm²)		MA (A/mg <sub>Pt</sub> )		ECSA (m <sup>2</sup> /g <sub>Pt</sub> )	ADT cycles
	BOL	ADT	BOL	ADT	_	
Surface-modified PtCuNi (this work)	21.8	12.3	10.7	6.6	49	20,000
As-prepared PtCuNi (this work)	8.2	5.3	5.7	3.5	70	20,000
Pt <sub>3</sub> Ni nanoframe (53)	8.4*	NA	5.7	NA	67.2	10,000
PtNi-BNCs/C (50)	5.16	5.15*	3.52	3.47	68.2	50,000
fct-Pt-Co@Pt octahedron ( <i>54</i> )	9.16	7.21	2.82	2.23	30.8	30,000
PtPb nanoplate (51)	7.8	7.0	4.3	3.97	55.0	50,000
PtNi/Ni-B/C (55)	9.05	NA	5.3	NA	59	5000
J-PtNWs (27)	11.5	10.9*	13.6	12.0	118	6000

<sup>\*</sup>Calculated from the literature.

long-term ADT cycling, exposing more available Pt sites. At the beginning of the reaction, the introduction of the surface molecular pump sacrifices ECSA while boosting SA (or turnover frequency per site) to achieve a high MA. With the progress of the reaction, while the SA of the catalyst decreases with stripped ligands, its ECSA recovers to maintain a decent MA after ADT. This suggests that the surface-modified catalysts can not only show improved catalytic activity but also retain a higher activity after ADT compared to the unmodified ones. Using nuclear magnetic resonance (NMR), we did not find the existence of DMF molecule in electrolyte before and after stability test (fig. S24), which indicated that DMF surface modification is unlikely to poison the electrolyte. Therefore, applying this surface modification approach to the same ORR catalyst enables high activity operation for a longer period. This molecular pump concept is broadly applicable. We found that the surface-modified Pt/C also showed 2.5 times enhanced hydrogen oxidation reaction (HOR) SA and noticeable enhanced MA (fig. S25), which suggested that modified surface can facilitate the mass transport for HOR, where hydrogen gas is pumped to Pt surface and water as the product is quickly removed.

#### **DISCUSSION**

To further demonstrate the practical applicability of the proposed surface modification strategy, we performed ORR activity and stability measurements at 65°C. At 65°C, the surface-modified Pt/C exhibited an SA of 2.00 mA/cm² and an MA of 0.37 A/mg, which are much higher than those of unmodified Pt/C (an SA of 0.57 mA/cm² and an MA of 0.26 A/mg) (fig. S26). The MA of modified Pt/C only decreased by 8.1% after ADT, while the MA of unmodified Pt/C decreased by 42.3% (fig. S26C). The after-ADT MA of modified Pt/C (0.34 A/mg) is still higher than that of unmodified Pt/C (0.26 A/mg). We also found that this surface modification strategy works in MEA (fig. S27). In the MEA test, we found that a commercial surface-modified PtNi showed an MA of 0.64 A/mg at 0.9 V versus RHE, which is higher than that of unmodified PtNi (0.5 A/mg).

In addition, in the practical catalyst system with ionomer, we believe that DMF as a polar hydrophilic aprotic additive at the Pt surface can modulate the surface environments. Two hydrophobic methyl substituents (-CH<sub>3</sub>) inhibit interaction with water molecules to disrupt the water network near the Nafion-Pt interface, which facilitates water diffusion and oxygen diffusion. On the other hand, the hydrophilic aldehyde group (-HC=O...) may favor the hydrogen bonding connection with sulfonate groups (...H-O=S) of the Nafion ionomer, stabilizing the adsorbed DMF on the Pt surface. We assume that the paired aldehyde groups and sulfonate groups may reduce the hydrophilicity of the Nafion-Pt interface, further promoting water diffusion and oxygen diffusion.

In summary, we report that surface modification can greatly enhance the ORR activity of Pt-based catalysts by interrupting the surface water adlayer network and hence facilitating the influx of  $\rm O_2$  and outflux of  $\rm H_2O$  at the catalyst-electrolyte interface, functioning as a surface molecular pump to promote the rate of the reaction. In particular, combining experimental design and IFF MD simulations provides a quantitative explanation. IFF MD simulations alone reach 10 times higher accuracy than other MD methods to date due to incorporation of deep chemical knowledge into the IFF parameters, full interpretability, and validation. The simulations showed

that DMF-modified Pt surfaces lower the O<sub>2</sub> physisorption free energy to larger negative values, enabling better penetration of the interfacial water layer. The result equates to a two- to threefold increase in local O2 concentration and hence a two- to threefold increase in SA toward ORR, following Le Chatelier's principle. We further showed that this effect can be used independently and in conjunction with the Sabatier principle (strain-ligand effect). Accordingly, we demonstrated a record-high SA of 21.8  $\pm$  2.1 mA/cm<sup>2</sup> (at 0.9 V versus RHE) on DMF-modified PtCuNi catalyst. Unlike current studies that focus on modifying surface hydrophobicity to alter intermediate binding energy or reaction pathways (37), our strategy aims to enhance oxygen availability and expedite water removal, thus driving the reaction toward a more efficient ORR process. This study opens an exciting avenue to modulate catalytic activities through designing and constructing local environments at the molecular level that enhance reaction kinetics at the catalyst-electrolyte interfaces. Future studies may also illuminate the role of ionomers for the dynamics of oxygen and hydronium ions at the catalyst

#### **MATERIALS AND METHODS**

#### Chemicals

Platinum(II) acetylacetonate [Pt(acac)2, 97%], nickel(II) acetylacetonate [Ni(acac)<sub>2</sub>, 95%], copper (II) acetate [Cu(AC)<sub>2</sub>, 97%], perchloric acid (HClO<sub>4</sub>, 70%, PPT grade), glucose, oleylamine [OAm; CH<sub>3</sub>(CH<sub>2</sub>)<sub>7</sub>CH=CH(CH<sub>2</sub>)<sub>7</sub>CH<sub>2</sub>NH<sub>2</sub>, >70%], 1-octadecene[ODE;  $CH_2$ = $CH(CH_2)_{15}CH_3$ ; technical grade, >90%], and 5 wt % Nafion were all purchased from Sigma-Aldrich. Commercial Pt/C catalyst (40 wt % Pt) was purchased from Alfa Aesar. DMF (≥99.8%), ethanol (200 proof), and cyclohexane (C<sub>6</sub>H<sub>12</sub>; analytical reagent, >99.5%) were obtained from EMD Millipore and Decon. Isopropanol (≥99.5%) was purchased from Thermo Fisher Scientific. All reagents were used as received without further purification. Carbon black (Vulcan XC-72) was received from Carbot Corporation and annealed in 200°C air before used. The deionized water (18 megohm cm) was obtained from an ultrapure purification system (Milli-Q advantage A10). Ultrahigh-purity CO was purchased from Airgas Inc.

# **Preparation of PtCuNi catalysts**

The synthesis of PtCuNi catalysts is similar to a previous report (36). In a typical preparation of PtCuNi catalysts, Pt(acac)<sub>2</sub> (20 mg), Cu(AC)<sub>2</sub> (6 mg), Ni(acac)<sub>2</sub> (6 mg), glucose (135 mg), OAm (3 ml), and ODE (2 ml) were added into a vial (volume, 35 ml). The mixture was ultrasonicated for 1 hour and then purged with ultrahighpurity carbon monoxide (CO) for 5 min and heated from room temperature to 170°C and kept for 12 hours in an oil bath. The colloidal products were collected by centrifugation and washed with cyclohexane/ethanol (v/v = 1.5) mixture two times. The PtCuNi catalysts were then loaded onto 20 mg of carbon black (Vulcan XC-72) by sonication for 2 hours. The as-prepared PtCuNi catalysts were suspended in 20 ml of cyclohexane/ethanol (v/v = 1:1) mixture. To load the sample onto carbon black, 20 mg of carbon black was added and sonicated for 2 hours. The resulting catalysts were collected by centrifugation, washed with cyclohexane/ ethanol mixture, and annealed at 130°C for 6 hours in a homebuilt tube furnace with an Ar flow of 100 standard cubic centimeters per minute.

# Application of surface-modified PtCuNi catalyst and Pt/C catalyst

The resulting dried PtCuNi catalysts or as-received Pt/C catalysts were first dispersed in DMF with a concentration of 1 mg/ml and ultrasonicated for 20 min. The resulting solution was heated with magnetic stirring in an oil bath at 120°C for the desired time to absorb DMF. After the modification, the resulting catalysts were washed with IPA three times and further dried in a vacuum dryer.

## Structure and composition characterization

High-resolution transmission electron microscope (HRTEM) images, energy-dispersive x-ray spectroscopy (EDX) line-scan file, and the high-angle annular dark-field scanning transmission electron microscope (HAADF-STEM) images were taken on a JEM-ARM300F Grand ARM Transmission Electron Microscope operated at 300 kV. TEM samples were prepared by dropping ethanol dispersion of catalysts onto carbon-coated aluminum TEM grids (Ted Pella, Redding, CA). The Pt loading of catalysts was determined by the inductively coupled plasma atomic emission spectroscopy. The Pt loading is determined before electrochemical measurement. XPS tests were carried out on a Kratos AXIS Ultra DLD spectrometer. All samples were electrochemically activated before XPS tests. The NMR measurement was performed on a Bruker AV400 NMR Spectrometer. 3-(Trimethylsilyl)propionic-2,2,3,3-d4 acid sodium salt was used as the internal standard.

#### **Electrochemical measurements**

A three-electrode cell system was used to conduct all electrochemical tests. The working electrode was a glassy carbon rotating disk electrode (RDE) with a 0.196 cm<sup>2</sup> glassy carbon geometry area from Pine Instruments. The counter and reference electrodes are Pt coil and Ag/AgCl (3 M Cl<sup>-</sup>). For activity measurement, Pt loadings were 10.2 and 7.85 μg/cm<sup>2</sup> for Pt/C catalysts and all PtCuNi catalysts, respectively. CV activations were performed in N<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub> electrolyte with a potential scan rate of 100 mV/s. The ECSA determined by H<sub>upd</sub> was calculated by integrating the hydrogen adsorption charge on the CV curve by taking a value of 210 μC/cm<sup>2</sup> for the adsorption of a hydrogen monolayer. The ORR was measured in an O<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub> electrolyte with a potential scan rate of 20 mV/s. The measurement temperature was set at 25°C using a water bath five-neck flask. The current density of the ORR polarization curve was iR-corrected during the measurements. The kinetic current was determined by the Koutecký-Levich equation. This kinetic current was then normalized by the ECSA to obtain the SA. The accelerated durability test (ADT) was performed at room temperature in O<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub> solutions by applying CV sweep between 0.6 and 1.0 V versus RHE at a sweep rate of 100 mV/s for 20,000 cycles. The ORR activity was measured after 20,000 cycles of ADT. The impedance spectra were recorded with an AC amplitude of 10 mV rms by sweeping the frequency from 100 kHz to 1 Hz at 40 points per decade (data collection on logarithmic scale).

#### **Computational protocols**

The atomic/molecular models were prepared using the Materials Studio program (38) under three-dimensional (3D) periodic boundary conditions, including the Pt(111) surfaces,  $H_2O$  molecules (flexible simple point charge, SPC water model), DMF molecules,  $O_2$  molecules, perchlorate anions, and hydronium cations.

The IFF was used for Pt atoms,  $O_2$  molecules, perchlorate anions, the flexible SPC water model, and hydronium cations using the 12-6 Lennard-Jones (LJ) format [same as Chemistry at Harvard Macromolecular Mechanic (CHARMM) and Consistent Valence Force Field (CVFF)], along with CHARMM36 parameters for DMF molecules. IFF uses a classical Hamiltonian and includes exceptionally accurate parameters for inorganic and organic compounds with systematic validation of chemical bonding (polarity), structures, energies, and physical-chemical interpretability (12). IFF parameters for FCC metals are thermodynamically consistent LJ parameters that enable the reproduction of lattice parameters (0.1% accuracy), surface energies (3% accuracy), and hydration energies (~5 to 10% accuracy) relative to experiments, surpassing DFT methods by multiples (11, 39-41). The IFF parameters for molecular O<sub>2</sub> reproduce the liquid density, vaporization energy (at boiling point and 1 atm), and solvation free energy (298.15 K) within 3% deviation from experimental measurements down to an accuracy of 0.03 kcal/mol (13), also one to two orders of magnitude more accurate than other force fields and quantum methods (42). Multiple studies have demonstrated that the dynamics at realistic temperatures and surface properties of metals and their interfacial interactions with solvents can be reproduced in about 95% agreement with experimental data, which is a significant improvement over earlier methods and opens up exciting perspectives on molecular engineering of catalysts (11, 29, 40, 41, 43).

To construct models with different levels of DMF surface coverage, initially a Pt(111) slab with six Pt layers (600 Pt atoms) and a box size of  $27.744 \times 24.027 \times 45.000 \text{ Å}^3$  was built. The zone above the Pt layers was filled with up to 175 DMF molecules, corresponding to a slab of liquid DMF of approximately 30 Å thickness, excluding water, to correlate with the conditions of surface modification of Pt catalyst in DMF. The simulation temperature was set to 393.15 K (equivalent to 120°C, treatment temperature). MD simulations were carried out in the isothermal-isobaric ensemble (NPT) at a pressure of 1 atm for 8 ns to reach equilibrium, as previously demonstrated for similar systems (44, 45). The equilibrium configurations were used to determine the number of DMF molecules per unit area corresponding to monolayer coverage. For this purpose, using multiple equilibrium configurations, different numbers of DMF molecules near monolayer coverage were placed flat on the Pt surface and the zone above was kept as a vacuum. MD simulations were carried out in the canonical (NVT) ensemble for 8 ns at 298.15 K, the electrochemical measurement temperature. The structure with the highest number of DMF molecules lying flat on the Pt(111) surface was defined as one monolayer coverage and determined to be 20 DMF molecules for the chosen slab, equal to 33.3 Å surface area per DMF molecule. This average value was used to determine the number of DMF molecules needed to represent different surface coverages in follow-on simulations.

Physisorption of  $O_2$  molecules on the pure and DMF-modified Pt(111) surfaces was investigated using models of supersaturated  $O_2$  solutions in contact with the surface that facilitate extensive adsorption-desorption statistics. The models featured a box size of  $69.360 \times 62.470 \times 53.000 ~\text{Å}^3$  with 3900 Pt atoms (six atomic layers of Pt), 5557 H<sub>2</sub>O molecules (flexible SPC model), 100  $O_2$  molecules (equivalent to 1.0 M concentration), and 10 hydronium ions and perchlorate anions (equivalent to pH 1.0). This concentration of  $O_2$  was still low enough to avoid correlations between neighbor  $O_2$  molecules. Five model systems with different surface coverages by

DMF molecules were built to represent conditions in experiments. The surface coverage in these models was 0.0, 0.1, 0.3, 0.5, and 0.7 with a corresponding number of 0, 13, 39, 65, and 91 DMF molecules for the chosen box size, respectively. Subsequently, the start structures were subjected to a brief energy minimization and to MD simulations to analyze adsorption using the Nanoscale Molecular Dynamics (NAMD) program (46). First, simulations were first equilibrated for 1 ns in the NPT ensemble at a pressure of 1 atm, and then continued for 50 ns in the NVT ensemble to sample O2 contacts, using equilibrium lattice parameters from the NPT period. All simulations involved 3D periodic boundary conditions, a time step of 1 fs, a summation of LJ interactions with a spherical cutoff at 1.2 nm, a summation of electrostatic interaction using the particle mesh ewals (PME) method with a tolerance of 10<sup>-4</sup> (high accuracy) and temperature of 298.15 K consistent with electrochemical measurement conditions. 3D snapshots were collected every 1 ps to collect abundant data for the physisorption-desorption dynamics of the 100  $O_2$  molecules on the Pt(111) surfaces with various amounts of DMF (5 million independent data points for each surface coverage).

In the following, we describe the analysis of average physisorption time, interfacial concentration of O<sub>2</sub> and H<sub>2</sub>O, hydrogen bonds per water molecule, and free energy profile of O<sub>2</sub> binding to the surface. The equilibrium trajectories of 50 ns duration were analyzed across the time window from 10 to 50 ns for all models, equal to 40,000 snapshots for each system (equal to 4 million data points for 100 O<sub>2</sub> molecules), using self-developed Python code with the MD Analysis package (47, 48). O<sub>2</sub> molecules (or H<sub>2</sub>O molecules) at 3.5 Å or less vertical distance from Pt atoms on the surface were categorized to be in an adsorbed state (26). For the calculation of the average physisorption time, we only considered the O2 molecules that were encountered within 3.5 Å distance from the surface Pt atoms (defined as adsorbed O2 molecules) and excluded those O2 molecules that had no contact with the surface. The total physisorption time equals the sum of adsorption time of all O<sub>2</sub> molecules within 3.5 Å distance. The average physisorption time was obtained by dividing the total physisorption time by the number of adsorbed O<sub>2</sub> molecules (hereby, trends would not change if minimum physisorption times up to 100 ps were required).

The interfacial concentrations of  $O_2$  and  $H_2O$  are crucial for electrochemical reactions and were defined with a slightly larger cutoff of 5 Å above the top atomic layer of the Pt(111) surface, which includes all molecules in the first layer and in the intermediate space to the second molecular layer (but excluding the second molecular layer) (11). The interfacial concentration of  $O_2$  and  $H_2O$  was defined as the average number of  $O_2$  and  $H_2O$  molecules per unit volume within the interfacial layer with an uncertainty equal to the SEM. All simulations were repeated three times and averaged to gain high accuracy and assure reproducibility.

The number of hydrogen bonds per water molecule were counted within the nearest molecular layer of water atop the Pt(111) surface. The maximum are two hydrogen bonds per  $\rm H_2O$  molecule, equal to four shared hydrogen bonds, as found in ice or bulk water, with a strength of about 4.7 kcal/mol per hydrogen bond (equal to approximately half the vaporization energy of water). The criterion of identification of a hydrogen bond (H) O–H···O( $\rm H_2$ ) in water was a distance of less than 3.5 Å between the two oxygen atoms and an O–H···O angle within  $\rm 180^{\circ} \pm 30^{\circ}$  (31). The average number of H-bonds was calculated by dividing the total number of H-bonds by the total number of  $\rm H_2O$ 

molecules in the adlayer, including periodic images across box boundaries.

The profiles of the free energy of O<sub>2</sub> physisorption were obtained by SMD simulation with the large-scale atomic/molecular massively parallel simulator (LAMMPS) program (49). We used LAMMPS for SMD calculations because it has better options for SMD and analyzing the results than NAMD. Otherwise, all results from LAMMPS and NAMD are expected to be equal (within the statistical errors) due to using the same IFF energy expression, parameters, and simulation settings (23). We carried out SMD simulations using Pt(111) slabs with and without DMF, respectively. The box size was  $24.969 \times 24.027 \times 65.550 \text{ Å}^3$ , including six atomic layers of Pt atoms at the bottom, with periodic boundary conditions. DMF molecules were placed flatly on the Pt(111) surface at 50% surface coverage in the same manner as previously described. The model system contained 1000 flexible SPC H<sub>2</sub>O molecules and only one O2 molecule. Hydronium ions were not included as they can interact with the O<sub>2</sub> molecule during the pulling process and cause poor sampling. The O2 molecule was initially placed at least 12 Å away from the Pt(111) surface in the z direction. It was coupled via a harmonic spring to a virtual point closer to the surface, which was pulled with a constant velocity toward the Pt(111) surface. The free energy was sampled using a pulling velocity of 1 Å/ns at distances larger than 6 Å. When the O<sub>2</sub> molecule moved closer than ~6 Å to the surface atomic layer of Pt, the pulling velocity of the virtual point was decreased to 0.03 Å/ns for the model with water and 0.01 Å/ns for the model with DMF to ensure better convergence (the accuracy is high and results are identical for any value below a specific critical speed, and above a critical speed the results lose accuracy). Because the DMF molecules are mobile, including hopping and free rotation on the Pt surface, simultaneous rearrangements of the interfacial water networks occur. The given pulling velocities were slow enough to achieve equilibrium sampling of O2 interactions within the metal-water interfacial region, especially while passing through the immediate water adlayers that have strong interactions with both the metal surface and their neighbor molecules. Using a pulling velocity of 0.01 Å/ns or less then mimics a reversible process and serves to obtain an accurate free energy profile. To avoid movement of the platinum slab, the bottom atomic layer of the slab was constrained in the pulling direction (z direction) while allowing mobility in the other directions. In summary, the SMD simulations captured the approach of O<sub>2</sub> molecules from solution to physisorption on the surface for the entire electrodeelectrolyte interface including the Pt(111) surface, surface-adsorbed DMF molecules, explicit H<sub>2</sub>O molecules, O<sub>2</sub> molecules, and extensive dynamics over a period of 400 ns for each replica (4 µs for all replicas). The simulation models fully consider the hydration of oxygen molecules while traveling from solution to the Pt surface, as well as changes in hydration upon proximity to ions in solution and coordination with DMF on the surface. The distance of the  $O_2$  molecule from the Pt surface was measured between the center of mass of an O2 molecule and the z coordinate of the surface layer Pt atoms.

Specifically, the two model systems for SMD were first preequilibrated for 1 ns in the NPT ensemble, and then the SMD protocol was carried out in the NVT ensemble. The trajectory frames and corresponding free energies were collected every 1 ps to generate a high-quality free energy profile. The Visual Molecular Dynamics (VMD) program was used to determine the distance between the center of mass of the  $\rm O_2$  molecule and the Pt(111) surface atomic layer in each frame. The calculated free energies were then plotted versus the distances between  $\rm O_2$  and the Pt (111) surface layer. We noticed that a slower pulling speed changes the simulated free energies. Below a certain threshold, the results converge to constant free energies, become trustworthy, and allow full sampling of the metal-DMF-water-oxygen interfacial structure. In earlier work (11), a faster speed (0.1 Å/ns) was sufficient because Pt surface modification and ions were not present. Convergence could then be reached to compare the adsorption free energies of O<sub>2</sub> on various Pt(hkl) facets. In the current work, the extensive local dynamics of the DMF molecules on the Pt(111) surface and simultaneous rearrangements of the water network required longer simulation times and many repeats to reach convergence of the free energy profiles of O<sub>2</sub>. Therefore, we used a pulling speed of 0.01 Å/ns (or slower) at distances <6 Å to favor accuracy over speed. The SMD simulations were repeated with the same settings for over 10 times for better results, about 400 ns simulation time for each replica, and equal to a total simulation time of 4 µs at a DMF coverage of 50%.

For the calculation of the O<sub>2</sub> stabilization energy, we used the Materials Studio graphical user interface (38) to cut out representative small molecular clusters from the snapshots of the regular MD trajectories, typically consisting of one O<sub>2</sub> molecule, the surrounding H<sub>2</sub>O and DMF molecules, and two atomic layers of Pt atoms below (to provide enough interactions with the adsorbed O<sub>2</sub> molecule). For each composition, two systems, namely, the surface system and the away system, were created with the same number and types of atoms (fig. S13A). For the surface system, the O<sub>2</sub> molecule was typically in contact with the Pt(111) surface, while the O<sub>2</sub> molecule in the away system was at least 20 Å detached from the Pt(111) surface. The calculations used IFF (not DFT, which would be associated with several times higher errors and uncertainties) and the Discover (or Forcite) program in Materials Studio to carry out 200 steps of energy minimization/geometry optimization for each cluster to obtain a physically justified start structure, followed by single-point energy calculations. The spherical cutoff distance of pairwise atom-atom LJ interactions was set to be 12 Å, and the accuracy of the summation of Coulomb interactions was 10<sup>-5</sup> kcal/mol using the Ewald method. Hereby, Materials Studio and the Discover program are more convenient and faster to create the clusters and conduct energy minimization and single-point energy evaluations than NAMD and LAMMPS. The results are identical among all three programs within statistical simulation errors when the same IFF energy expression and simulations parameters (including cutoff settings) are used (23). At least four independent clusters for each DMF content were analyzed in this manner to obtain an average O2 stabilization energy as well as a standard deviation. The O2 stabilization energy was calculated using the following equation

$$E_{\rm st} = E_{\rm sur} - E_{\rm away} \tag{1}$$

where  $E_{\rm st}$  is the  ${\rm O_2}$  stabilization energy,  $E_{\rm sur}$  represents the total energy of the surface system, and  $E_{\rm away}$  is the total energy of the away system (fig. S13A). Typical clusters are displayed in fig. S13B.

To calculate the physisorption energy of DMF on Pt(111), Cu(111), and Ni(111) surfaces, a two-box method was used. The box size was 27.744  $\times$  24.027  $\times$  100.000 ų for Pt(111), 28.115  $\times$  22.135  $\times$  100.000 ų for Cu(111), and 27.410  $\times$  21.580  $\times$  100.000 ų for Ni(111) systems. The first model system was called the desorbed system, where DMF molecules were detached from the metal surface. The second model system was the adsorbed system, where DMF molecules were in contact with the metal surface. All

systems included six metal atomic layers,  $6480 \text{ H}_2\text{O}$  molecules, and one DMF molecule. Upon completion of the simulations, the average total energies of the two individual boxes were corrected to the exact target temperature of 298.15 K utilizing the heat capacity of each system, which eliminates potential errors up to 5.0 kcal/mol in adsorption energies. The simulation settings were the same as described above for standard MD over 50 ns. The physisorption energy  $E_{\text{phys}}$  was calculated using the following equation

$$E_{\rm phys} = E_{\rm ads} - E_{\rm des} \tag{2}$$

where  $E_{\text{ads}}$  is the average energy of the adsorbed system and  $E_{\text{des}}$  represents the average energy of the desorbed system.

# **Supplementary Materials**

The PDF file includes:

Figs. S1 to S27 Legend for data S1

Other Supplementary Material for this manuscript includes the following:
Data S1

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