Controlled release of hydrogen isotope compounds and tunneling effect in the heterogeneously-catalyzed formic acid dehydrogenation

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The hydrogen isotope deuterium is widely used in the synthesis of isotopically-labeled compounds and in the fabrication of semiconductors and optical fibers. However, the facile production of deuterium gas (D₂) and hydrogen deuteride (HD) in a controlled manner is a challenging task, and rational heterogeneously-catalyzed protocols are still lacking. Herein, we demonstrate the selective production of hydrogen isotope compounds from a combination of formic acid and D₂O, through cooperative action by a PdAg nanocatalyst on a silica substrate whose surface is modified with amine groups. In this process, D₂ is predominantly evolved by the assist of weakly basic amine moieties, while nanocatalyst particles in the vicinity of strongly basic amine groups promote the preferential formation of HD. Kinetic data and calculations based on semi-classically corrected transition state theory coupled with density functional theory suggest that quantum tunneling dominates the hydrogen/deuterium exchange reaction over the metallic PdAg surfaces.
Hydrogen isotopes, such as deuterium gas (D₂) and hydrogen deuteride (HD), are minor components of naturally occurring molecular hydrogen, and have applications as fine chemicals owing to their unique physical, quantum, and nuclear properties. In particular, D₂ is often employed for the synthesis of a diverse array of deuterium-labeled compounds, which have been recognized as indispensable research tools in the life, environmental, and material sciences because of their non-radioactive and stable characteristics. Unfortunately, commercially available D₂ gas is quite expensive, since it is produced by the electrolysis of D₂O with the investment of large amounts of energy. The reactions of various metals, including Na, Fe, and Mg, with D₂O have also been traditionally employed to generate this gas on a laboratory scale, but these reagents have serious drawbacks in that they are toxic, produce large amounts of waste, and require high reaction temperatures.

A number of metal-catalyzed H/D exchange reactions between H₂ and D₂O, especially those based on Ir, Rh, and Ru, have been developed based on the investigation of hydrogenases. Sajiki and co-workers explored Pd/C-catalyzed D₂ generation via H/D exchange reactions using a H₂/D₂O combination, and applied the D₂ that was generated in situ to the synthesis of D-labeled compounds. More recently, Fujita and co-workers developed a catalytic method to produce D₂ from deuterated methanol (CD₃OD) and D₂O using an Ir complex bearing functional bipyridylidate ligands in the presence of sodium hydroxide.

Formic acid (HCOOH, FA) has been the subject of significant research effort because it is one of the most promising hydrogen storage materials, due to its low toxicity, nonflammable liquid state at room temperature, and high hydrogen content (43.8 g H₂/L). Various metals, including Na, Fe, and Mg, with D₂O have also been traditionally employed to generate this gas on a laboratory scale, but these reagents have serious drawbacks in that they are toxic, produce large amounts of waste, and require high reaction temperatures.

Interestingly, hydrogen isotope compounds have been evolved via the H/D exchange reaction during the metal-catalyzed dehydrogenation of FA in D₂O. In an early study, Puddephatt and co-workers demonstrated that a mixture of H₂, HD, and D₂ gases in a molar ratio of 1:2:1 can be generated by reactions between HCOOD and D₂O or between DCOOH and H₂O, catalyzed by a binuclear Ru complex. Subsequently, the distribution of D during the decomposition of deuterated FA analogs (i.e., HCOOD, DCOOH, and DCOOD) in D₂O in the presence of [Rh(μ-(Cp*)(H₂O))(H₂)]²⁻ and the heterodinuclear iridium–ruthenium complex [Ir(Cp*)(H₂O)(bpm)Ru(bpy)](SO₄)₂ (Cp* = pentamethylcyclopentadienyl, bpm = 2,2'-bipyridine, bpy = 2,2'-bipyridine) was explored by Fukuzumi et al. A convenient means of producing D₂ and various deuterated compounds using a half-sandwich Ir complex with 4,4'-dihydroxy-2,2'-bipyridine ligands was also demonstrated by Himeda and co-workers. Although some progress has been achieved in conjunction with the use of transition metal complexes, these processes often require high catalyst concentrations and expensive deuterated FA to obtain reasonable levels of activity and selectivity. In addition, the use of homogeneous catalysts on an industrial scale is extremely challenging due to the difficulties involved in recovering the precious metals and ligands from the reaction mixture. For these reasons, systems based on heterogeneous catalysts in association with undeuterated FA and D₂O would be highly beneficial, but such processes have not yet been researched, in spite of the obvious practical advantages.

Our group has previously reported that the presence of basic functional groups on support materials in the vicinity of active Pd-based alloy nanoparticles (NPs) promotes the production of high-purity H₂ during the dehydrogenation of FA. Herein, we describe a straightforward method for the production of hydrogen isotope compounds from a combination of FA and D₂O using a PdAg nanoparticle catalyst on a substrate that has been surface modified with amine groups. This work also demonstrates that the selectivity of this process can be tuned by varying the basicity of the amine moieties. The H/D exchange reactions between FA and D₂O is found to involve a quantum tunneling effect, as evidenced by the large kinetic isotope effect (KIE) value, and based on semi-classically corrected transition state theory (SC-TST) coupled with density functional theory (DFT) calculations.

Results

Structural Characterization. Various amine-grafted mesoporous silica (SBA-15) substrates (termed SBA-15-Amine-x (SA-x), where x = 1–5) were synthesized using a surface modification technique. This silica possesses relatively large hexagonal channels ~9 nm in diameter. The surface modification was performed using a post-synthesis method in association with silane-coupling reagents containing different amine groups, as shown in Fig. 1. The SA-4 specimen, having Schiff base functional groups (-N = CH₃), was also synthesized by the further modification of SA-2 with formaldehyde. These substrates were impregnated with Pd and Ag (1 wt% Pd; Pd:Ag molar ratio = 1:1) simply by immersion in aqueous solutions of Pd(NH₃)₄Cl₂ and AgNO₃, followed by reduction with NaBH₄. The resulting catalysts are denoted PdAg/SA-x (x = 1–5) herein. The concentration of grafted amine functionalities in all samples was determined to be ≈0.3 mmol g⁻¹ substrate using thermogravimetric (TG) analysis. The presence of the amine groups was confirmed by the analysis of each specimen by Fourier transform-infrared (FT-IR) spectroscopy (Supplementary Fig. 1) and N 1s X-ray photoelectron spectroscopy (XPS) (Supplementary Fig. 2). The long-range ordering of hexagonally packed mesoporous structures in these materials was also demonstrated on the basis of low-angle X-ray diffraction (XRD) patterns (Supplementary Fig. 3).

The textual properties of the PdAg/SA-x as well as PdAg/S made with unmodified silica are summarized in Table S1. All samples generated N₂ adsorption/desorption isotherms having characteristic type IV curves with sharp capillary condensation steps and H1-type hysteresis loops. These features indicate highly ordered and well-preserved cylindrical mesopores (Supplementary Fig. 4). The Brunauer–Emmett–Teller (BET) surface areas (S_BET), pore diameters (D_pore) and pore volumes (V_p) for the members of the PdAg/SA-x series were all found to be slightly lower than those for the unmodified PdAg/S. During surface modification, the silane-coupling reagents were able to readily diffuse through the channels, which ensured homogeneous grafting, because the estimated molecular sizes of these reagents (up to 1 nm) were significantly smaller than the diameters of the channels.

Supplementary Figure 5 presents high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) images together with size distribution plots for the PdAg/S and PdAg/SA-x materials. The PdAg NPs on the PdAg/SA-x were evidently well dispersed and had a narrow size distribution, with average diameters in the range of 4.5–7.3 nm. These values are
smaller than the 7.5 nm NP size on the unmodified PdAg/S, and so amine functionalization evidently resulted in the formation of smaller NPs. This result indicates that variations in the basicity of the grafted amine groups may affect both the formation of Pd metal nuclei and the consecutive growth, ultimately producing different NP sizes.

The shapes of the Pd K-edge X-ray absorption near edge structure (XANES) spectra generated by the PdAg/SA-x and unmodified PdAg/S differ from that seen in the PdO spectrum, but resemble the spectrum of Pd foil (Supplementary Fig. 6A). The FT-extended X-ray absorption fine structure (FT-EXAFS) spectra of PdAg/SA-x and PdAg/S exhibit a single intense peak associated with Pd−Pd bonds that suggests the presence of metallic Pd (Supplementary Fig. 6B). However, the Pd−Pd bond lengths for these materials were slightly longer than that for Pd foil, indicating Pd−Ag bonds. The Ag K-edge XANES spectra obtained from the PdAg/SA-x and PdAg/S are similar to that of Ag foil, and the FT-EXAFS spectra of these specimens exhibit a single sharp peak ascribed to contiguous Ag–Ag bonds with lengths of 2.6–2.8 Å (Supplementary Fig. 7). This peak is shifted to a slightly shorter interatomic distance compared with that for pure Ag foil. These results present clear evidence for the formation of PdAg alloy NPs. In addition, the Pd 3d peaks in the XPS spectra produced by the PdAg/SA-x are slightly shifted to higher binding energies compared to those for the unmodified sample (Supplementary Fig. 8A). A similar shift of the Ag 3d peaks in the XPS spectra is also evident after modification (Supplementary Fig. 8B). Our experimental results suggest that surface amine functional groups were present at the peripheries of the loaded PdAg NPs to alter their electronic state. The alternation of the electronic state of the supported NPs by the organic modifier of the support materials have also been reported previously.

**Controllable release of hydrogen isotope compounds.** Utilizing the prepared catalysts having different amine functional groups, the dehydrogenation reaction was conducted in D₂O solutions containing 5 M HCOOH:HCOONa (9:1) at 343 K for 30 min. The selectivities for the resulting hydrogen isotope compounds (HD and D₂) are summarized in Fig. 2a, together with the turnover frequency (TOF) values based on the amount of Pd. No H₂ formation was observed, and the molar ratios of total hydrogen isotope compounds to CO₂ generated during the course of the reaction were close to 1 for all samples. The use of bare SA-5 without PdAg NPs does not show any activity. It should be noted that the present catalytic system limits the undesirable formation of CO to less than the detection limit of the gas chromatography (GC) system (<2 ppm). As expected, we succeeded in the controlled synthesis of these hydrogen isotope compounds by changing the surface grafted amine groups. As an example, the PdAg/SA-1 (having the strongly basic −NEt₂ groups) preferentially generated HD with 65% selectivity, while D₂ formation was predominant with 87% selectivity when using the PdAg/SA-5, which was modified with the weakly basic −PhNH₂ groups. The D₂ and HD selectivity appeared to be independent of the yield level and remained unchanged during the reaction, suggesting that the H−D exchange reaction with the produced hydrogen isotope compounds did not take place (Supplementary Fig. 9). In addition, the catalytic activity was also greatly affected by the properties of the supports. As an example, the TOF value obtained with PdAg/SA-5 was 65 h⁻¹, which was seven times that observed using the PdAg/SA-1.

The H₂O/D₂O molar ratio in the reaction solution also significantly affected selectivity. Fig. 2b, c plots the selectivity for various hydrogen isotope compounds versus the D₂O mol%. In the case of the PdAg/SA-1, HD selectivity increased with increasing D₂O concentration, accompanied by a decrease in H₂, up to a D₂O concentration of 67 mol%, with no D₂ formation observed. The maximum HD selectivity of 80% was attained at 80 mol% D₂O without the formation of H₂, after which the HD selectivity decreased. In the reaction with PdAg/SA-5, the HD selectivity increased along with the D₂O mol%, while the D₂ selectivity was drastically enhanced at D₂O levels above 50 mol%. The highest D₂ selectivity was obtained at 100 mol% D₂O, with 87% selectivity. In contrast, the type of amine functional group, PdAg NP loading, reaction temperature, and ratio of FA to sodium formate did not show significant effect on selectivity (Supplementary Figs. 10–13), demonstrating that these parameters are minor factors in determining the selectivity.

The recyclability is a crucial point to consider when heterogeneous catalysts are employed for industrial applications. After the reaction, the PdAg/SA-x (x = 1 and 5) catalysts were easily separated from the reaction mixture and could be reused with retention of its activity and selectivity; preferential formation of hydrogen isotope compounds could be attained for at least five recycling experiments (Supplementary Fig. 14). More interestingly, the present catalyst provides a simple and efficient heterogeneous catalytic system for the in situ deuteration of organic compounds (Table 1 and Supplementary Fig. 15). As an example, the reaction of diphenylacetylene proceeded smoothly using PdAg/SA-5 and 5 M HCOOH:HCOONa (9:1) solution in D₂O, giving the deuterated product with 96% deuterated content in 99% chemical yield. The expected deuterated products were also obtained with excellent deuterated contents via addition reactions of deuterium to carbon–carbon double bonds and deuterium substitution reactions.

The observed variations in selectivity for hydrogen isotope compounds were because of determining the adsorption energy (E_ads) of FA based on DFT calculations for each SA-x. This was done because such values reflect the basicity of the amine functional groups, since the adsorption involves formation of an acid–base pair via N−H−O hydrogen bonding. As an example, the E_ads over SA-1 with −NH(C₂H₅)₂ groups was found to be ~215 kJ mol⁻¹, which was substantially stronger than the value of ~120 kJ mol⁻¹ over SA-5, having −PhNH₂ groups. It is worth noting that a good correlation between the HD selectivity and E_ads was observed. Specifically, the HD selectivity increased with increases in E_ads, as shown in Fig. 3a.
suggesting that the basicity of the amine functional groups plays a crucial role in determining selectivity.

We suspected that differences in the sizes of the PdAg NPs also affected selectivity. However, there was no significant correlation between selectivity and the average diameter of PdAg NPs on the SA-x series of samples, based on HAADF-STEM analysis (Fig. 3b). Unfortunately, this investigation was unable to accurately decouple the concomitant effect of the basicity of the various amine functional groups from that of the NP size. In order to further examine the effect of NP size on selectivity, a series of colloidal poly(vinylpyrrolidone)-capped Pd NPs were first synthesized under several different conditions and then deposited on the SA-5. The TEM images and size distribution plots shown in Supplementary Fig. 16 demonstrate the fabrication of size-controlled Pd/SA-5 specimens with particle sizes ranging from 2.8 to 16 nm. These materials were subsequently applied to the dehydrogenation of FA in D2O. As expected, the D2 selectivity was found to remain almost constant at ~90% regardless of the particle size, although a volcano-type variation in activity based on the particle size was observed, with a maximum TOF at 3.9 nm (Supplementary Fig. 17).

In addition, Pd-based alloy NPs supported on SA-5 were synthesized, using Au, Co, Cr, Ni, Zn, or Cu precursors as the second metal and employing the same impregnation method followed by reduction with NaBH4. Notably, these variations in the metal component did not greatly affect the D2 selectivity (Fig. 3c), although the activity varied significantly. This effect can presumably be ascribed to the synergic effect between the Pd and the second metal, as has been reported previously.

These results suggest that the selectivity for the production of hydrogen isotope compounds is governed by the grafted amine groups in the periphery of each active center, and not by the inherent ability of the metal NPs themselves.

Upon consideration of the above results as well as the theoretical studies, we determined a possible reaction pathway...
leading to the selective formation of hydrogen isotope compounds from the dehydrogenation of FA in D_2O, as illustrated in Fig. 4. In this process, regardless of the basicity of the amine groups, the reaction of HCOOH with 1 is promoted by the amine groups in the periphery of the PdAg NPs as a result of the acid–base interaction, affording a Pd formate species along with a proton-scavenged amine group (2). The Pd formate species, for which the trans-AgO-PdH(O)-bridged configuration is the lowest energy structure, subsequently undergoes isomerization followed by β-hydride elimination to release CO_2 and a Pd hydride species (3). Intermediate 3 undergoes H–D exchange reactions in the presence of D_2O at both the Pd and basic amine sites, but to differing extents according to the basicity of the amine groups. In the case of PdAg/SA-1, which has strongly basic –NH_2 groups, four preferentially undergoes an H–D exchange reaction at the Pd site with a barrier of \( E_{\text{ad}} = 140.2 \text{ kJ mol}^{-1} \) to afford 5. Subsequently, an H–D exchange reaction proceeds at the basic amine site with a barrier of 212.5 kJ mol\(^{-1}\) to produce 6. The intermediates 5 and 6 release HD and D_2, respectively. Thus, the substantially higher activation energy for the formation of 6 compared to that for 5 contradict the occurrence of the reversed reaction, which also explains the preferential formation of HD. In contrast, the activation energies for the formations of 8 and 9 from 7 via the H–D exchange reaction at Pd and weakly basic –PhNH_2 groups in the case of PdAg/SA-5 were calculated to be 140.2 and 149.8 kJ mol\(^{-1}\), respectively. From an energetic point of view, the H–D exchange reaction would therefore be expected to occur simultaneously at both sites, leading to the preferential formation of intermediate 9 and ultimately favoring D_2 generation. The \( E_a \) for the H–D exchange reaction at the strongly basic –N(C_2H_5)_2 site is substantially larger than that at the weakly basic –PhNH_2 site, which are reflected by the basicity determined by the \( E_{\text{ad}} \) of FA in Fig. 3a; the \( E_a \) increased with increasing the \( E_{\text{ad}} \) of FA. It can therefore be concluded that the selectivity for hydrogen isotope compounds is determined by the basicity of the surface amine groups, which ultimately influence on the H–D exchange reaction at the basic sites in the peripheries of the active centers. In addition, the preliminary DFT calculations determined the \( E_a \) for the release of hydrogen from 5 or 6 and 8 or 9 were 52.7 and 13.0 kJ mol\(^{-1}\), respectively, indicating rate-determining step is the H–D exchange reaction at the basic amine site with D_2O. These results also suggest that hydrogen release is accelerated in the presence of weakly basic amines, which accounts for the higher TOF obtained from PdAg/SA-5 as compared to that for PdAg/SA-1, as shown in Fig. 2a.

To improve our understanding of this mechanism, H/D KIE values were examined in trials using H_2O and D_2O. Intriguingly, large KIE values of 10 at 323 K and 23 at 313 K were obtained when using PdAg/SA-2 and PdAg/SA-5, respectively. Abnormally large KIE values at room temperature or higher generally indicate the participation of a quantum tunneling effect in a proton transfer reaction, and this can be unambiguously confirmed from the temperature dependence of KIE. Specifically, values for the Arrhenius parameters \( E_a(D) - E_a(H) \), \( A_A/A_D \), and KIE of >5 kJ/mol.
mol$^{-1}$, $<1$, and $>9$ at 298 K provide evidence for the involvement of a quantum tunneling effect$^{27,29}$. The Arrhenius plots for H$_2$O and D$_2$O obtained in the present study are provided in Fig. 5. The resulting $E_a$(D)$-E_a$(H) and $A_H/A_D$ values were 19.7 kJ mol$^{-1}$ and 0.005 for PdAg/SA-2 and 9.4 kJ mol$^{-1}$ and 0.06 for PdAg/SA-5, respectively. Our experimental data therefore meet the above requirements, and indicate the contribution of a quantum tunneling effect to the present catalytic cycle.

**Tunneling effect calculations.** Quantum mechanical tunneling of atoms, which originates from wave-particle duality, has been observed not only in chemical reactions, but also in many other fields of chemistry including astrochemistry and biochemistry$^{40-42}$. In particular, the tunneling of hydrogen species such as protons, hydrogen atoms and hydride ions is an important aspect of acid–base, transfer and proton-coupled electron transfer reactions$^{43-48}$. As an example, hydrogen atoms have been frequently observed to tunnel through various metal surfaces, such as Cu(001)$^{49}$, Ru(0001)$^{50}$ and Ni(100)$^{51}$, and the motion of hydrogen atoms is known to be enhanced by tunneling at low temperatures. Proton hopping between oxygen atoms on the Brønsted acidic sites of zeolites and heteropolyacids has also been found to be associated with tunneling pathways$^{52,53}$. In addition, it has been demonstrated that reactions based on heterogeneous catalysis, such as CO oxidation and NH$_3$ formation on metal surfaces, are also affected by tunneling$^{54-58}$.

Atom tunneling can be experimentally demonstrated by constructing a non-linear Arrhenius plot based on data acquired at low temperatures, where the rate constant is independent of temperature, or (as described above) by unusual KIE values. However, the effect of tunneling on reaction rates can also be directly confirmed by theoretical calculations$^{59-62}$. In an effort to theoretically demonstrate the participation of tunneling pathways in the present heterogeneously-catalyzed dehydrogenation of FA using PdAg NPs in the vicinity of amine groups, we adopted the semi-classically corrected TST (SC-TST) developed by Fermann and Auerbach$^{63}$, together with a modification by Bhatia and Sholl$^{64,65}$. A tunneling crossover temperature, $T_c$, was calculated using the equations

$$T_c = \frac{h\nu_{1m}\Delta E_{ZP}/k_B}{2\pi\Delta E_{ZP} - h\nu_{1m}\ln 2},$$  (1)

$$\Delta E_{ZP} = \left( E^0 + \sum_{j=1}^{N-1} \frac{h\nu_j^2}{2} \right) - \left( E_A + \sum_{i=1}^{N} \frac{h\nu_i^2}{2} \right),$$  (2)

where $\Delta E_{ZP}$ is the activation energy barrier (i.e., the difference in energies between the TS and the local minimum, including the SCF energy ($E^0$ or $E_A$), and the zero-point energy, as defined in Eq. (2)). $N$ is the number of degrees of freedom for the system, $h$ and $k_B$ are the Planck and Boltzmann constants, $\nu_j$ is the harmonic frequency for the local minimum, and $\nu_{1m}$ and $\nu_j$ are those for the TS (the former being an imaginary frequency).

Within the SC-TST framework, the reaction rate constant at temperature $T$ is given by Eq. (3), using the TST rate constant, $k_{TST}^0$ ($T$), and the tunneling correction factor, $\Gamma(T)$, which are expressed by Eqs. (4) and (5), respectively. In Eq. (4), $\Delta E$ is the SCF energy difference between the TS and the local minimum.

$$k_{SC-TST}^0 (T) = k_{TST}^0 (T) \cdot \Gamma(T),$$  (3)

$$k_{TST}^0 (T) = \prod_{j=1}^{N} \nu_j f(\nu_j/2k_B T) \cdot \exp \left( \frac{-\Delta E}{k_B T} \right),$$  (4)

$$\Gamma(T) = \frac{\exp(\Delta E_{ZP}/k_B T)}{1 + \exp(2\pi\Delta E_{ZP}/h\nu_{1m})} \cdot \frac{1}{2} \int_{-\infty}^{\infty} \exp \left( \frac{h\nu_{1m}}{k_B T} \right) \text{sech}^2 \left( \frac{\theta}{\lambda} \right) \text{d} \theta,$$  (5)

The production of hydrogen isotope compounds from an FA/ D$_2$O combination using PdAg NPs together with amine groups evidently comprises a series of H–D exchange reactions between D$_2$O and the PdAg NPs and/or amine groups. Here, five
Similar trends are observed in the case of the Pd7Ag6 cluster lower than those for path I, and are below room temperature. Eq. (1) using a Pd2Ag2 cluster model. Below these values, temperature independent below transitions occurs, but rather a point where the relative change in mechanism between quantum tunneling and classical is almost temperature independent below \( T_c \) because of the increase in the tunneling correction factor with decreasing temperature. Owing to the quantum tunneling effect, an Arrhenius fit of the rate data affords different activation energies depending on the temperature range investigated. An activation energy of 46.2 kJ mol\(^{-1}\) was obtained from the \( k_{\text{TST}} (T) \) data between 300 to 400 K, whereas the values of \( k_{\text{SC-TST}} (T) \) from 100 to 200 K gave an activation energy of 0.8 kJ mol\(^{-1}\). It should be noted that \( k_{\text{SC-TST}} \) is also larger than \( k_{\text{TST}} \) at \( T_c \) by a factor of more than 35 (\( \Gamma (320) = 35.1 \)). Therefore, \( T_c \) is not a distinct point at which a change in mechanism between quantum tunneling and classical transitions occurs, but rather a point where the relative dominance of the two mechanisms rapidly changes. For reaction pathways II–V, the temperature independence of \( k_{\text{SC-TST}} (T) \) suggests that the tunneling mechanism is predominant (Supplementary Fig. 18). Although (as noted) the \( T_c \) values are lower than those for path I, and are below room temperature. Similar trends are observed in the case of the Pd2Ag2 cluster model, as summarized in Supplementary Table 3 and Supplementary Fig. 19. Here, the \( T_c \) for path I was determined to be 257 K, which is higher than those for paths IV and V. These theoretical investigations strongly suggest that a tunneling mechanism occurs primarily in path I (i.e., direct H/D exchange over the PdAg surface) and plays a crucial role in the overall reaction at temperatures as high as 320 K. The tunneling probability depends to different degrees on the mass of the moving particle, the barrier height, and the barrier width. Especially, the barrier width mainly determines the probability of tunneling in chemical reactions: the narrower a barrier is, the more likely tunneling is. As shown in Supplementary Fig. 20, our calculation results reveal that the \( T_c \) correlates well with \( \nu_{\text{Im}} \) not \( \Delta E_{\text{ZP}} \). The absolute value \( \nu_{\text{Im}} \) is the curvature at the vicinity of the transition state in the reaction coordinate direction, which significantly influence on the barrier width: the larger a value \( \nu_{\text{Im}} \) is, the narrower a barrier is. The representative reaction paths used to assess the tunneling effect proceed at the same time; thus, the \( T_c \) of the reaction with highest \( T_c \) can be experimentally observed. The lower \( T_c \) in paths II–V suggests that the H–D exchange at the basic amine cites are largely dominated by classical transitions; thus, the effect of basicity of the amine in the HD/D2 ratio can be experimentally appeared.

**Discussion**

We succeeded in the controlled synthesis of hydrogen isotope compounds during the dehydrogenation of FA in D2O by tuning the H–D exchange reaction, using specific amine-functionalized PdAg nanocatalysts. The grafted amine groups were found to be indispensable to this process, and the reaction selectivity showed a close correlation with the basicity of the amine groups, with the D2 selectivity increasing with decreasing basicity. This study provides not only advanced insights into the preferred catalyst architecture but also demonstrates an ideal heterogeneous catalyst system with potential applications to this type of transformation. Large KIE values indicative of the quantum tunneling effect was observed for the reaction of FA in H2O vs. D2O. Theoretical investigations also emphasized the involvement of a quantum tunneling mechanism, predominantly during the direct H–D exchange reaction over the PdAg surface.
Fig. 6 Tunneling effect calculations. a Reaction paths used to assess the tunneling effect. (I) direct H/D exchange over the PdAg surface, (II) direct H/D exchange on an NEt₃ group, (III) direct H/D exchange over a PhNH₂ group, (IV) H/D exchange over PdAg in the vicinity of an NEt₃ group and (V) H/D exchange over PdAg in the vicinity of a PhNH₂ group. b Arrhenius plots as calculated for path I

Table 2 Parameters for the calculation of tunneling effect

| Path I | Path II | Path III | Path IV | Path V |
|--------|---------|----------|---------|--------|
| ΔE_T (kJ mol⁻¹) | 132.9 | 215.6 | 143.3 | 72.7 | 86.3 |
| ν_m (cm⁻¹) | 1376.4 | 731.7 | 535.3 | 490 | 886.7 |
| T_c (K) | 320 | 168 | 123 | 113 | 206 |

Methods

Materials. Pd(NH₃)₄Cl₂ was obtained from the Sigma-Aldrich Chemical Co. AgNO₃, FA, and 3-aminopropyltriethoxysilane were purchased from Nacalai Tesque, Inc., and aminophenyltrimethoxysilane was obtained from Manchester Organic. All commercially available chemicals were used as received.

Synthesis of the SA-x series. The mesoporous silica support SBA-15 was prepared using a surfactant self-assembly approach as follows: 4.0 g of Pluronic P123 was dissolved in 30 g of deionized water and 120 mL of 2 M HCl solution with stirring at 313 K; 8.5 g of TEOS was added into the above solution with stirring for 24 h. The mixture was aged at 333 K overnight without stirring, and then the product was recovered by vacuum filtration and washed with deionized water. The resulting white powder was dried at 383 K overnight and calcined at 823 K for 10 h.

Synthesis of PdAg/SA-x. A quantity of SA-x (0.3 g) was mixed with 50 mL of an aqueous solution containing Pd(NH₃)₄Cl₂ (10 mM, 2.85 mL) and AgNO₃ (10 mM, 2.85 mL), followed by stirring at room temperature for 1 h. The suspension was evaporated under vacuum and the resulting powder was dried overnight. Subsequently, each sample was reduced with NaBH₄, giving PdAg/SA-x (1 wt% Pd; Pd:Ag molar ratio = 1:1). Analysis using inductively coupled plasma (see Characterization section for details) clearly indicated that the desired amount of each metal species was loaded onto each support.

Characterization. Powder XRD patterns were recorded using a Rigaku Ultima IV diffractometer with Cu Kα radiation (λ = 1.54056 Å). Nitrogen adsorption–desorption isotherms were acquired at 77 K using a BELSORP-max system (MicrotracBEL Corp.). Prior to analysis, samples were degassed at 423 K for 3 h under vacuum to remove physisorbed water. Specific surface areas were calculated using the BET method based on the nitrogen adsorption data. Inductively coupled plasma optical emission spectrometry data were obtained using a Nippon Jarrell Ash ICAP-575 Mark II instrument. TG analysis was performed by a Rigaku Thermo plus EVOZ TG8121 system from room temperature to 973 K at a heating rate of 10 K min⁻¹ in air flow (10 mL min⁻¹). FT-IR spectra were obtained with a Hitachi FTIR-2300T system (MicrotracBEL Corp.). Prior to analysis, samples were mixed with KBr, followed by compression into thin disk-shaped pellets. XPS was performed with a Shimadzu ESCA-3000 system, using Mg Kα X-ray radiation (hv = 1253.6 eV) as the excitation source. The binding energy values for each spectrum were calibrated relative to the contaminant C 1s core level binding energy of 284.5 eV. Pd and Ag K-edge XAFS spectra were acquired using a fluorescence-yield collection technique at the 01B1 beamline at SPring-8, JASRI, Harima, Japan, in conjunction with a Si (111) monochromator (proposal numbers 2018A1144 and 2018B1082). Data reduction was performed using the REX2000 program (Rigaku).
Dehydogenation of FA. A 0.1 g quantity of catalyst was transferred into a 30 mL reaction vessel containing 1.8 mL of D₂O after which the vessel was sealed with a rubber septum. The reaction mixture was bubbled with Ar for 15 min, followed by the addition of 0.2 mL of a 5 M solution of HCOOH/HCOONa (9:1) in D₂O, with magnetic stirring at 343 K. The amount of H₂, HD, and D₂ generated were determined by GC using a Shimadzu GC–2014 instrument with He as the carrier gas and a Shinya OGO–SP column held at 77 K, in conjunction with a thermal conductivity detector. TOFs in units of h⁻¹ were calculated as the total moles of all hydrogen isotopes divided by total mole Pd in the catalyst after 30 min.

In situ deuteriation in the FA–D₂O system. A 0.1 g quantity of the PdAg-SA-5 was placed into a 30 mL reaction vessel containing 1.8 mL of D₂O and 0.1 mmol of the substrate, after which the vessel was sealed with a rubber septum. A 0.2 mL quantity of a 5 M HCOOH/HCOONa (9:1) solution in D₂O was added and the reaction mixture was magnetically stirred at 353 K. After 24 h, the reaction mixture was passed through a membrane filter to remove the catalyst. The filtrate was mixed with Et₂O (10 mL) and the aqueous layer was extracted with Et₂O (2 × 10 mL), dried over MgSO₄, filtered, and concentrated in vacuo to give the analytically pure deuterated products. The yield was determined by GC/MS spectrum (MS) equipped with capillary column and the D contents of the products were determined by ¹H nuclear magnetic resonance (NMR) spectroscopy on the basis of the integration of the aromatic protons.

Computational method. FA adsorption energy values, E_ads, were calculated using DFT, employing the DMol3 program in the Materials Studio 17.2 software package. A generalized gradient approximation exchange-correlation functional proposed by Perdew, Burke, and Ernzerhof was combined with the double numerical basis set plus polarization functions. A supercell slab consisting of a 4 × 4 surface unit cell with three atomic (111) surface layers was adopted, with the bottom two layers fixed at the corresponding bulk positions and the top layer allowed to relax during geometry optimization. The slab was separated by a vacuum space with a height of 30 Å. Allowed to relax during geometry optimizations. The slab was separated by a

...and the deuteriation of aryl amines in water. Chem. Eur. J. 17, 3963–3967 (2011).

...constrained water-soluble Rh- and Ru-phosphine complexes. Green Chem. 5, 213–217 (2003).

...a facile Ru–H heterolytic activation and intramolecular proton transfer assisted by basic n–centers in the ligands. J. Am. Chem. Soc. 127, 15364–15365 (2005).

...kinetic studies on the first dihydrogen aquacomplex. [Ru (H₂)(H₂O)]²⁺: formylation under H₂ pressure and catalysis. H/D isotope exchange in water. Inorg. Chem. Acta 359, 1795–1806 (2006).

...Vignais, P. H/D exchange reactions and mechanistic aspects of the hydrogenses. Coord. Chem. Rev. 249, 1677–1690 (2005).

...Kirta, T. et al. Facile and convenient method of deuterium gas generation using a Pd/C-catalyzed H₂–D₂ exchange reaction and its application to synthesis of deuterium-labeled compounds. Chem. Eur. J. 14, 3371–3379 (2008).

...Enomoto, A., Kajita, S. & Fujita, K. Convenient method for the deuteriation of deuterium gas catalyzed by an iridium complex and its application to the deuteriation of organic compounds. Chem. Lett. 48, 106–109 (2019).

...Kempinger, J. & Huang, K.-W. Formic acid as a hydrogen energy carrier. ACS Catal. 2, 188–195 (2017).

...Grasemann, M. & Laurenczy, G. Formic acid as a hydrogen source—recent developments and future trends. Energy Environ. Sci. 5, 8171–8181 (2012).

...Joó, F. Breakthroughs in hydrogen storage—formic acid as a sustainable storage material for hydrogen. ChemSusChem 1, 805–808 (2012).

...Bulushev, D. A. & Ross, J. R. Heterogenous catalysts for hydrogenation of CO₂ and bicarbonates to formic acid and formates. Catal. Rev. 60, 566–593 (2018).

...Gunasekar, G. H., Park, K., Jung, K.-D. & Yoon, S. Recent developments in the catalytic hydorgenation of CO₂ to formic acid/formate using heterogeneous catalysts. Inorg. Chem. Front. 3, 882–895 (2016).

...Mori, K., Sano, T., Kobayashi, H. & Yamashita, H. Surface engineering of a supported PdAg catalyst for hydrogenation of CO₂ to formic acid: elucidating the active Pd atoms in alloy nanoparticles. J. Am. Chem. Soc. 140, 8902–8908 (2018).

...Mori, K., Taga, T. & Yamashita, H. Isolated single-atomic Ru catalyst bound on a layered double hydroxide for hydrogenation of CO₂ to formic acid. ACS Catal. 7, 3147–3151 (2017).

...Singh, A. K., Singh, S. & Kumar, A. Hydrogen energy future with formic acid: a renewable chemical hydrogen storage system. Catal. Sci. Technol. 6, 12–40 (2016).

...Meinnell, D., Sponholz, P., Junge, H. & Reller, M. Formic acid as a hydrogen storage material - development of homogeneous catalysts for selective hydrogen release. Chem. Soc. Rev. 45, 3954–3988 (2016).

...Gao, Y., Kuncheria, J. K., Jenkins, H. A., Puddrefft, R. J. & Yap, G. P. A. The interconversion of formic acid and hydrogen/carbon dioxide using a bimetallic ruthenium complex catalyst. J. Chem. Soc. Dalton Trans. https://doi.org/10.1039/B004234J, 3212–3217 (2000).

...Fukuzumi, S., Kobayashi, T. & Suenobu, T. Efficient catalytic decomposition of formic acid for the selective generation of H₂ and H/D exchange with a water-soluble rhodium complex in aqueous solution. ChemSusChem 1, 827–834 (2008).

...Fukuzumi, S., Kobayashi, T. & Suenobu, T. Unusually large tunneling effect on highly efficient generation of hydrogen and hydrogen isotopes in pH-selective decomposition of formic acid catalyzed by a heterodinuclear iridium–ruthenium complex in water. J. Am. Chem. Soc. 132, 1436–1497 (2010).

...Wang, W.-H. et al. Highly efficient D₂ generation by dehydrogenation of formic acid in D₂O through H⁺/D⁺ exchange on an iridium catalyst: application to the synthesis of deuterated compounds by transfer deuterogeneration. Chem. Eur. J. 18, 9397–9404 (2012).

...Mori, K., Dojo, M. & Yamashita, H. Pd and Pd–Ag nanoparticles within a macrocyclic basic resin: an efficient catalyst for hydrogen production from formic acid decomposition. ACS Catal. 3, 1114–1119 (2013).

...Mori, K., Tanaka, H., Dojo, M., Yoshizawa, K. & Yamashita, H. Synergic catalysis of PdCu alloy nanoparticles within a macrocyclic basic resin for hydrogen production from formic acid. Chem. Eur. J. 21, 12085–12092 (2015).

...Mori, K., Naka, K., Masuda, S., Miyawaki, K. & Yamashita, H. Palladium copper chromium ternary nanoparticles constructed in situ within a basic resin: enhanced activity in the dehydrogenation of formic acid. ChemCatChem 9, 3456–3462 (2017).

...Mori, K. et al. Phenylamine-functionalized mesoporous silica supported PdAg nanoparticles: a dual heterogeneous catalyst for formic acid/CO₂-mediated enhanced hydrogen delivery. Chem. Commun. 53, 4677–4680 (2017).

...Masuda, S., Mori, K., Futamura, Y. & Yamashita, H. PdAg nanoparticles supported on functionalized mesoporous carbon: promotional effect of surface
amine groups in reversible hydrogen delivery/storage mediated by formic acid/CO$_2$. ACS Catal. 8, 2277–2285 (2018).

34. Martin, M., Mori, K., Fujikawa, K., Akiyama, W.-S. & Yamashita, H. Amine-functionalized MIL-125 with imbedded palladium nanoparticles as an efficient catalyst for dehydrogenation of formic acid at ambient temperature. J. Phys. Chem. C 117, 22805–22810 (2013).

35. Navlani-García, M., Mori, K., Nozaki, A., Kawaihara, Y. & Yamashita, H. Investigation of size sensitivity in the hydrogen production from formic acid over carbon-supported Pd nanoparticles. ChemistrySelect 1, 1879–1886 (2016).

36. Tedesee, K. et al. Hydrogen production from formic acid decomposition at room temperature using a Ag-Pd core-shell nanocatalyst. Nat. Nanotechnol. 6, 302–307 (2011).

37. Qin, Y.-l, Wang, J., Meng, F.-z, Wang, L.-m & Zhang, X.-b. Ef...

38. Martis, M., Mori, K., Ahn, W.-S. & Yamashita, H. Amine-functionalized MIL-125 with imbedded palladium nanoparticles as an efficient catalyst for dehydrogenation of formic acid at ambient temperature. J. Phys. Chem. C 117, 22805–22810 (2013).

39. Bercaw, J. E., Chen, G. S., Labinger, J. A. & Lin, B.-L. Hydrogen tunneling in protonation of platinum(ii) and palladium(ii) methyl complexes: mechanistic implications. J. Am. Chem. Soc. 130, 17654–17655 (2008).

40. McMahon, R. J. Chemical reactions involving quantum tunneling. Science 299, 833–834 (2003).

41. Meissner, J. & Kästner, J. Atom tunneling in chemistry. Angew. Chem. Int. Ed. 55, 5400–5413 (2016).

42. Scheiner, P. R. Tuning electronic properties of chemical reactions: the third reactivity paradigm. J. Am. Chem. Soc. 139, 15276–15283 (2017).

43. Pan, Z., Horner, J. H. & Newcomb, M. Tuning in C–H oxidation reactions by an o xo(IV) oxorhodium radical cation: direct measurements of very large H/D kinetic isotope effects. J. Am. Chem. Soc. 130, 7776–7777 (2008).

44. Sierka, M. & Sauer, J. Proton mobility in chabazite, faujasite, and ZSM-5...