Coordination nano-space as stage of hydrogen ortho—para conversion

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The ability to design and control properties of nano-sized space in porous coordination polymers (PCPs) would provide us with an ideal stage for fascinating physical and chemical phenomena. We found an interconversion of nuclear-spin isomers for hydrogen molecule \(\text{H}_2\) adsorbed in a Hofmann-type PCP, \(\{\text{Fe(pz)}[\text{Pd(CN)}_4]\}\) (pz = pyrazine), by the temperature dependence of Raman spectra. The ortho (o)–para (p) conversion process of \(\text{H}_2\) is forbidden for an isolated molecule. The charge density study using synchrotron radiation X-ray diffraction reveals the electric field generated in coordination nano-space. The present results corroborate similar findings observed on different systems and confirm that o–p conversion can occur on non-magnetic solids and that electric field can induce the catalytic hydrogen o–p conversion.
1. Introduction

Hydrogen molecule H₂ has two different nuclear-spin isomers, para (p) and ortho (o), owing to requirement of symmetry of wave function. Since interconversion between these isomers is a forbidden process in an isolated molecule, it takes a long time constant, 100 h. The o–p conversion accompanied by heat generation causes a boil-off problem in a cryogenic H₂ storage [1]. Many kinds of o–p conversion catalysis have been studied so far [2–13]. Several models such as the first-order [8] and second-order [9] perturbation models using paramagnetic and/or diamagnetic metals have been theoretically proposed for the o–p conversion catalysis. The conversion process is often facilitated on a solid surface. The distance between H₂ and atoms on the surface is crucial to reveal the mechanism. The atomic configurations of H₂ molecules on the surface have recently been visualized by a neutron powder diffraction [14] and scanning tunnelling microscope [15,16]. Here we adopt three-dimensional nano-pore surfaces of porous coordination polymers (PCPs) [17,18] as a platform for the o–p conversion, and report a structural dependence of nuclear-spin states for H₂ in the Hofmann-type PCP, {Fe(pz)[Pd(CN)₄]} (pz = pyrazine) [19,20], by a combination of adsorption isotherm, Raman spectroscopy and in situ synchrotron radiation X-ray powder diffraction (SR-PXRD). We have successfully confined and converted H₂ using the nano-space of PCP.

2. Material and methods

2.1. Synthesis of {Fe(pz)[Pd(CN)₄]}

The Hofmann-type PCP {Fe(pz)[Pd(CN)₄]·2H₂O} was synthesized by slow diffusion of the two solutions in a glass tube for more than 3 days at room temperature. First, one solution contained a mixture of (NH₄)₂Fe(SO₄)₂·6H₂O (0.10 mmol), pyrazine (0.10 mmol) and citric acid 0.2 mmol in 10 ml water was put in a glass tube. Then, K₂[Pd(CN)₄] (0.10 mmol) in water (5 ml)/ethanol (5 ml) mixture solvent was carefully poured on the first solution surface. The glass tube was kept undisturbed. Microcrystals suitable for X-ray diffraction were obtained from the interface between two solutions. The guest-free PCP {Fe(pz)[Pd(CN)₄]} was prepared by heating under vacuum at 423 K for 3 h.

2.2. Measurement of hydrogen gas adsorption isotherm

Hydrogen gas adsorption isotherm was measured with an automatic volumetric adsorption apparatus BELSORP-MAX (BEL Japan, Inc.) in an error of ±0.25%, which is caused by resolution of the pressure gauge. Normal hydrogen gas of 6-nine purity was supplied from a gas cylinder for isotherm, in situ SR-XRPD, in situ Raman measurements.

2.3. Synchrotron radiation X-ray powder diffraction experiment

The in situ SR-XRPD experiment of H₂ adsorption was performed at SPring-8 BL44B2. The large Debye-Scherrer camera with an imaging plate as a detector was used to measure powder profiles. The brown sample powders were carefully crushed with a micro spatula to obtain the powders with homogeneous grain size. The powder sample was loaded into a boron-silicated glass capillary with 0.4 mm internal diameter. The capillary was mounted on the goniometer head connected to the gas pressure control system. All the powder diffraction profiles were collected with a 0.01° step. The wavelengths of an incident X-ray were 0.511 and 0.5496 Å. These are slightly lower energy of the K-edge of Pd to minimize fluorescence scattering from the sample. We determined the wavelength by using a standard powder sample of CeO₂ by NIST. The X-ray diffraction intensities were collected from 2.0° to 68.95° in 2θ (0.511 Å) and 2.0° to 59.40° in 2θ (0.5496 Å), which correspond to d > 0.451 Å and d > 0.5496 Å d-spacing range. The beam size was 0.5 × 1.0 mm². The exposure time of all the data was for 12 min at each measurement. The temperature of the sample was controlled with a He open-flow cooling device, Helijet, by Agilent Technologies, Inc.

2.4. Raman spectroscopy

In situ microscopic Raman spectroscopy of H₂ gas adsorption was carried out by an iHR320 spectrometer (HORIBA Jobin Yvon Ltd) in the temperature range of 20–77 K under the temperature control condition.
3. Results and discussion

Figure 1a shows the H₂ adsorption isotherms at 77, 65, 35 and 20 K. A unit pore volume which corresponds to one unit cell of \( \text{Fe(pz)}[\text{Pd(CN)}_4] \) is shown as an inset. The adsorption amount of H₂ increased with decreasing temperature. The amounts at 77, 65, 35 and 20 K under 80 kPa are 2.5, 2.7, 3.3 and 3.5 H₂ per pore, respectively. Approximately one extra H₂ is adsorbed per pore by lowering the temperature.

Figure 1b,c shows powder diffraction profiles of (b) H₂ adsorbed and (c) desorbed states at 35 K (open squares) and 80 K (filled squares), respectively. All powder profiles show tetragonal symmetry with orientational disorder of the pz pillar ligands. There is a notable different change between the profiles of the H₂ adsorbed and desorbed states. In the H₂ desorbed state, peak positions and relative intensities at 80 K are almost identical to those at 35 K indicating no structural change, as shown in figure 1c. The lattice constants of the H₂ adsorbed state showed an anisotropic negative thermal expansion from 80 to 35 K, as shown in figure 1d. The expansion is clearly recognized as peak shifts in figure 1b.

Accurate structure in charge density level is crucial to understand H₂ adsorption property of the present system. We optimized an experimental condition to reveal the detailed structure in the H₂ adsorption process. We measured the data with high counting statistics, approximately 1 000 000 counts, and wide \( d \)-spacing range, \( d > 0.45 \) Å, at 35 and 65 K in the H₂ adsorbed and desorbed states, respectively. We determined accurate charge densities, \( \rho(\mathbf{r}) \) [21], and electrostatic potentials, \( \phi(\mathbf{r}) \) [22], by combining the maximum entropy method (MEM) and Rietveld refinement [23].
Figure 2 shows the $\rho(r)$ and $\phi(r)$ in desorbed state as contour maps on the (001) plane. There is no charge density peaks in the pore space at both 35 and 65 K, indicating no residual gas. Local minima of the $\phi(r)$ was found at the pore centre at 35 and 65 K. Figure 3a,b shows the $\rho(r)$ in the H$_2$ adsorbed states at 35 and 65 K as an isosurface with structure model. There are $\rho(r)$ peaks showing H$_2$ in the pore. As no overlap of the $\rho(r)$ was observed between the adsorbed H$_2$ molecules and framework, H$_2$ molecules are physically trapped in the pore. We consider that adsorbed H$_2$ behaves as not atom and ion but a molecule. The peak at 65 K was found at the centre of the pore on the (001) plane (a green sphere, figure 3b). We also found eight charge density peaks between the pz rings along $a$- and $b$-axes at 65 K (blue spheres, figure 3b), whereas the central peak at pore extended to diagonal (a red area, figure 3a) and the peaks between the pz rings disappeared at 35 K.

We found three H$_2$ sites, what we call sites-I, -II and -III. The site-I is the centre of the pore, 1/2, 1/2, 0; the site-II between the pz rings, 0.41, 0, 0 and site-III is 1/4, 1/4, 0. The schematic structures of H$_2$ sites were shown in figure 3c,d as red (site-III), blue (site-II) and green (site-I) spheres. The charge densities of H$_2$ at 65 K correspond to the mixture of the sites-I and -II. The H$_2$ densities at 35 K are interpreted as the mixture of the sites-I and -III. The nearest inter-site distances of the sites-I · · · II and sites-I · · · III are 3.681 Å, and 2.060 Å, respectively. There should be repulsion between H$_2$ molecules owing to the electrostatic interaction and quantum zero-point motion [24,25]. The H$_2$ · · · H$_2$ distance in the pore has been estimated by a quantum mechanical calculation using a two harmonic potential model [14]. The study suggested that 3.0 Å is about the shortest distance between two H$_2$ molecules. In the case of hexagonal H$_2$ crystal at 4.2 K, the intermolecular distance is 3.776 Å [26]. These facts provide us a clue to construct possible H$_2$ arrangements.

The sites-I and -II can coexist in a pore and the arrangement was achieved at 65 K (figure 3d). The maximum amount of H$_2$ per pore in this arrangement is three, which is consistent with the adsorbed H$_2$ amount of 2.7. Charge densities at site-I were larger than those at site-II. A number of 2.7 H$_2$ per pore was achieved by combining fully occupied site-I and partially occupied site-II. Therefore, an increase of H$_2$ uptake from 77 to 65 K should occur at site-II.

The main site of H$_2$ at 35 K is site-III. The charge densities at sites-I and -II decreased. The charge density at site-II was almost empty. The hydrogen molecules at sites-I and -II moved to site-III from 65 to 35 K. Site exchanges occurred only with cooling from 65 to 35 K. It is impossible to coexist sites-I, -II and -III in one pore, since the inter-site distances are too short for H$_2$ · · · H$_2$. In addition, site-II cannot coexist with site-III of an adjoining occupied pore because of the too short H$_2$ · · · H$_2$ distance (2.43 Å). The
H₂ molecules should give and receive intermolecular repulsions. A direction of resultant repulsion force $E$ can be observed. The maximum amount of H₂ per pore for site-III is four. In the rest of the pores, H₂ mainly occupied site-I. Approximately 75% of H₂ was located at site-III at 35 K from charge densities and adsorption isotherms. It can be easily calculated by an equation: $4x + (1 - x) = 3.3$, $x \sim 0.76$, where $x$ is the occupancy at site-III.

Lattice expansion with the H₂ uptake was mainly caused by the intermolecular repulsion between H₂ at site-III. The H₂ · · · H₂ distance of 3.599 Å in site-III is smaller than that in the hexagonal H₂ crystal. The H₂ molecules should give and receive intermolecular repulsions. A direction of resultant repulsion force is [110] which should expand a- and b-axes. There is no expansion of the c-axis indicating that four H₂ molecules were located on the (001) plane without a shift along the c-axis. The Fe–N₉CN bond length on the (002) plane was elongated in the lattice expansion, while the change of Pd-C was within errors.

We observed structural change of the H₂ arrangement by SR-PXRD and investigated nuclear-spin states of the adsorbed H₂ by Raman spectra. Temperature dependence of Raman spectra in the H₂ adsorbed states are shown in figure 3e. In the desorbed state at 77 K (spectrum 5; black), Raman bands at 420, 470, 485, 545 cm⁻¹ were observed and assigned to the vibrational motions of host framework. Three Raman-active bands were newly appeared at 307 (peak A), 328 (peak B) and 609 cm⁻¹ (peak C) and were assigned to the adsorbed H₂ due to the gas dosing in the same temperature of 77 K (spectrums 4; purple). We found some changes in the intensity and peak position for the Raman bands with cooling from 77 to 20 K. Since selection rules derived from intermolecular interactions, the rotational quantum number $J$, the coupling of nuclear-spin and angular momentum, and the arrangement of H₂, Raman excitation bands contain a great deal of information. In this study, the Raman excitations corresponding to rotational quanta can be observed. The rotational quanta of H₂ is given by $S_0(J) = E_{0,J+1} - E_{0,J}$ where $E_{0,J}$ is the rotational energy in the vibrational ground state ($v = 0$; $v$ is the vibrational quantum number). The quanta of H₂ have been investigated both theoretically and experimentally [3]. It is reported that at the low temperature, only the $J = 0$, $S_0(0) \approx 354$ cm⁻¹, and $J = 1$, $S_0(1) \approx 587$ cm⁻¹, levels are populated and assigned to p- and o-H₂, respectively. We observed two peaks A and B around 300 cm⁻¹ corresponding to the $S_0(0)$ spectra of p-H₂ and a broad peak C around 600 cm⁻¹, corresponding to the $S_0(1)$ spectra of o-H₂. The peaks A and C decreased with cooling and peak B shifted and increased at 35 K. Relative intensities of the peaks A, B and C were estimated by the profile fittings as shown in figure 3f. Figure 3g shows temperature dependence of relative intensities of peaks A, B and C. The intensities were
normalized by the intensity of a peak from host framework at 680 cm$^{-1}$. The position of peak B changed from 328(1) cm$^{-1}$ at 77 and 65 K to 333(1) cm$^{-1}$ at 35 and 20 K.

Temperature dependences of the Raman intensities for A and C were very similar indicating the same site occupation of nuclear isomers in figure 3g. Peak A was the most red-shifted from the gas phase values. The electrostatic potential around site-I was almost spherical as shown in figure 2d,e. Site-I is only one minima of the electrostatic potential in the pore. There can be the highest interaction between the adsorbed H$_2$ and the host framework at site-I. Therefore, peak A was assigned to p-H$_2$ at site-I, and peak C was also assigned to o-H$_2$ at site-I. Site-I was occupied by a mixture of o- and p-H$_2$ at 77 and 65 K. The ratio of o- to p-H$_2$ from Raman intensities at site-I is approximately 3:2 which is the midpoint between equilibrium ratios at 77 K (1:1) and at 300 K (3:1) of normal H$_2$. Relative peak intensities of A and C did not change from 77 to 65 K in figure 3g. This is also consistent with a full occupation of site-I at 77 and 65 K.

Peak B at 35 K was easily assigned to site-III from Raman spectra and the structure. Therefore, peak B at 77 and 65 K can be assigned to p-H$_2$ at site-II. Relative intensities of peak B increased from 77 to 65 K consistent with a partial occupation of site-II. In addition, peak B did not shift from 77 to 65 K which also support our assignment of Raman peaks.

The peak broadening and shift similar to peak B was recently observed in the Raman study of H$_2$ in ZnO [27]. In this case, the Raman bands of framework changed with cooling and the shift was interpreted by the interaction between rotational mode of confined H$_2$ (at around 320 cm$^{-1}$) and the framework phonons (at 379 cm$^{-1}$). In our case, Raman bands of framework were not observed in the vicinity of peak B and also showed no change. Therefore, the shift of peak B was assigned to the migration of H$_2$ between site-II and site-III from the experimentally determined structures.

The Raman peaks around 600 cm$^{-1}$ had almost disappeared within experimental noises at 35 and 20 K indicating that o-H$_2$ was converted to p-H$_2$. The intensity ratio between peak C and a sum of the peaks A and B from profile fittings was less than 7(3)% at 35 K and 1.2(9)% at 20 K. The result supports that most of the H$_2$ at sites-II and -III are p-H$_2$. We propose two principal processes to induce the conversion from o-H$_2$ to p-H$_2$ in the present PCP: (i) trapping of H$_2$ at sites-II and -III, and (ii) site exchange from site-I and -II to site-III by cooling.

We investigated structural difference between sites-I, -II and -III using the charge densities and electrostatic potentials experimentally determined. The receiving electric fields of H$_2$ from the [Fe(pz)][Pd(CN)$_4$] framework can be estimated from electrostatic potential in the desorbed state as shown in figures 2d,e. We calculated the electric fields, $E(r)$, of sites-I, -II and -III by the gradient of the electrostatic potential, $\nabla \phi(r) = -E(r)$. Figure 4a,b shows absolute electric fields, $|E(r)|$, along with [110] direction. The $E(r)$ for the site-I is 0.0 V m$^{-1}$ because of the inversion symmetry. The $E(r)$ for sites-II and -III are $7.38 \times 10^{10}$ V m$^{-1}$ and $7.62 \times 10^{10}$ V m$^{-1}$, respectively. There is a huge difference of $E(r)$ between site-I and sites-II, -III.
Recently, fast o–p conversions of the order of one or a few minutes have been observed on the H₂ physisorbed PCP [28] and amorphous solid water [29]. The several mechanisms; spin-orbit-electric dipole-hyperfine contact (SOPY) [29], ion-molecule electron exchanges (XY) and charge transfer [30,31], have also been proposed theoretically for these fast o–p conversions of non-magnetic insulating catalysts. Since measurement time of the present SR-PXRD and Raman spectrum were about 10 min, an upper limit of the o–p conversion time in the present PCP is proposed at 10 min. Most of the recently proposed mechanisms, such as SOPY, XY and charge transfer, can be applied to the present system. For example, the E(r) gradient from sites-I to -III can be applied to SOPY. In order to determine the conversion process experimentally, more conversion time resolution is required.

In this study, we observed a correlation between the structure and nuclear-spin state of H₂ in the three-dimensional Hofmann-type PCP, [Fe(pz)][Pd(CN)₄]₂ providing regular coordination nano-space by a combination of the X-ray charge density study and Raman spectrum. Nano-space constructing by PCPs would offer the design of the desirable electric field. The electric field induced processes for such catalytic hydrogen o–p conversion were proposed using the electric field obtained from X-ray diffraction data. X-ray diffraction and Raman spectra of a bulk powder sample provide us with quantitative time and space averaged information and demonstrate availability of the coordination nano-space as a stage for H₂ o–p conversion. There are still other possible mechanisms for the o–p conversion such as magnetic impurity effects. In this case, mixing of residual paramagnetic high-spin state was estimated to be less than 1.4%. Quantitative information in one molecule size scale with high time resolution is highly desired in the future. Recent advanced X-ray sources such as X-ray-free electron laser would enable us to access such a study.

Data accessibility. The datasets supporting this article have been uploaded as part of the electronic supplementary material. The material contains all powder diffraction data, Raman spectra and adsorption isotherm data used in this study.

Authors’ contributions. T.K. carried out the synthesis. T.K., Y.K., J.K. and E.N. carried out SR-PXRD measurements. T.K. and E.N. interpreted and discussed the results and structural analysis. A.H., S.K. and M.O. carried out adsorption isotherm and Raman spectrum measurements. A.M., A.H. and M.O. carried out magnetic measurements. H.T. and J.K. calculated and analysed the electrostatic potential. T.K., A.H., Y.K., J.R., S.K., M.T. and E.N. were involved in discussions. All authors commented on the paper, which was written by T.K., A.H., E.N. and M.T. All authors gave final approval for publication.

Competing interests. We have no competing interests.

Funding. This work was partly supported by ENEOS Hydrogen Trust Fund. M.O. thanks the MEXT, Japan for a Grant-in-Aid for Scientific Research Program (B) (grant no. 15H03787).

Acknowledgements. The SR powder experiments using Helijet were supported by Mr M. Takemoto (Japan Synchrotron Radiation Research Institute) and Mr S. Yamamoto (RIKEN). The synchrotron radiation experiments were performed at BL44B2 in SPring-8 with the approval of RIKEN (proposal no. 20120066 and 20110054).

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