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Heterogeneous catalase-like activity of gold(i)–cobalt(III) metallosupramolecular ionic crystals†

Mihoko Yamada, a Nobuto Yoshinari, a Naoto Kuwamura, a Toru Saito, b Satoshi Okada, c Sai Prakash Maddala, c Koji Harano, c Eiichi Nakamura, c Kohei Yamagami, d Keisuke Yamanaka, e Akira Sekiyama, d Tomoyoshi Suenobu, f Yusuke Yamada g and Takumi Konno* a

Unique heterogeneous catalase-like activity was observed for metallosupramolecular ionic crystals \([\text{Au}_4\text{Co}^\text{III}_2(\text{dppe})_2(\text{dpen})_4\text{X}_n] \cdot ([\text{Xe}]_n;\text{dppe} = 1,2\text{-bis(diphenylphosphino)ethane}, \text{dpen} = \text{dipenicillaminate}; X_n = (\text{Cl}^-)_2, (\text{ClO}_4^-)_2, (\text{NO}_3^-)_2\text{ or SO}_4^{2-})\) consisting of \([\text{Au}_4\text{Co}^\text{III}_2] \text{complex cations, } [\text{X}]^+\text{, and inorganic anions, X}^-\text{ or X}^{2-}\). Treatment of the ionic crystals with an aqueous \(\text{H}_2\text{O}_2\) solution led to considerable \(\text{O}_2\) evolution with a high turnover frequency of \(1.4 \times 10^5 \text{ h}^{-1}\) for the heterogeneous cobalt complexes, which was dependent on their size and shape as well as the arrangement of cationic and anionic species. These dependencies were rationalized by the presence of cobalt(III) centers on the crystal surface and their efficient exposure on the \((111)\) plane rather than the \((100)\) plane based on morphological and theoretical studies.

Introduction

Heterogeneous catalysts have been developed for industrial applications because of their facile handling, high durability and reusability.1 Various approaches, such as increasing the surface area by decreasing the particle size,2 attachment to mesoporous carriers,3 introduction of heterogeneous elements,4 enlargement of dispersion4 and modification of the crystal surface, have been used to control and improve their catalytic activities.5 In particular, the catalytic activities of metals or metal oxides with high durability have been improved by controlling the size and shape of the crystals.6 Compared to metal and metal oxide catalysts, catalysts consisting of coordination compounds (metal complexes) have certain advantages with respect to the precise modulation of their structures and properties through modification of the ligands employed.7 However, the relationship between the size and shape of the crystals and the catalytic activities of coordination compounds has rarely been investigated.

One well-known catalytic reaction is the disproportionation of hydrogen peroxide into water and oxygen, which is similar to the catalase reaction in living organisms (eqn (1)).

\[
2\text{H}_2\text{O}_2 \rightarrow \text{O}_2 + 2\text{H}_2\text{O}
\] (1)

This catalase-like activity has been observed for many coordination compounds under homogeneous and/or heterogeneous conditions. However, only a few cobalt(III) complexes have been reported to show catalase-like activity under heterogeneous conditions.9 Furthermore, no reports exist on the catalase-like activity of cobalt(III) complexes except one report of low activity under homogeneous conditions.10 Herein, we report that metallosupramolecular ionic crystals containing cobalt(III) centers, \([\text{Au}_4\text{Co}^\text{III}_2(\text{dppe})_2(\text{dpen})_4\text{X}_n] \cdot ([\text{Xe}]_n; \text{dppe} = 1,2\text{-bis(diphenylphosphino)ethane}, \text{dpen} = \text{dipenicillaminate}, \text{X}_n = (\text{Cl}^-)\text{, (ClO}_4^-)_2\text{, (NO}_3^-)_2\text{ or SO}_4^{2-})\), which were recently synthesized and structurally characterized,11 exhibit high catalase-like activity under heterogeneous conditions. This class of ionic crystals, which we refer to as ‘non-coulombic ionic solids (NCIS)’, consists of \([\text{Au}_4\text{Co}^\text{III}_2] \text{hexanuclear complex cations, } [\text{X}]^+\text{, and inorganic anions, X}^-\text{ or X}^{2-}\), and adopts an unusual non-alternate arrangement of cationic and anionic species.
ClO₄ (5.0 mg) and a homogeneous saturated solution of [Au₄Cr₂(dppe)₂(D-pen)₄]Cl₂, which are isomorphous with [CoIII{AuI(PPh₃)(D-pen)}₂]⁺, show no signiﬁcant evolution of O₂ gas (Fig. S4†). Similar treatment of aqueous H₂O₂ under the same conditions for 60 min, had an activity comparable to that in the ﬁrst run, thus indicating reusability (Fig. S9†).

To determine the effect of the arrangement of cationic and anionic species in the crystal on the appearance of the catalase-like activity, ionic crystals of [CoIII{AuI(PPh₃)(D-pen)}₂]ClO₄ (ref. 18) ([2]ClO₄, Fig. 1), which has the same coordination environment around the CoIII center as that in [1]Cl₂ but adopts a typical alternating arrangement of cations and anions in the crystal, were treated with aqueous H₂O₂ under the same conditions (Fig. 2). Remarkably, no evolution of O₂ gas was observed with this treatment, which was also the case during treatment of a saturated solution of [1]Cl₂ in H₂O with aqueous H₂O₂. Thus, the arrangement of cationic and anionic species plays a key role in the appearance of the heterogeneous catalase-like activity of the [1]Cl₂ and [1]NO₃₂ crystals.

Since the activity of heterogeneous catalysts often depends on the surface area, we examined the catalase-like activity of [1] Cl₂ and [1]NO₃₂ using crystals of different sizes (Fig. S10 and S11†). The use of small crystals (0.4 × 0.4 × 0.4 mm for [1]Cl₂ and 0.05 × 0.05 × 0.05 mm for [1]NO₃₂) led to greater evolution of O₂ gas compared to the use of large crystals (1.0 × 1.0 × 1.0 mm for [1]Cl₂ and [1]NO₃₂), which is expected from the larger surface area per unit mass. The TOFs at 15 min, which were calculated by considering the number of Co atoms on the crystal surface, were equivalent for the small and large crystals. Furthermore, a proportional relationship between the catalytic activities and the amounts of crystal used was observed, which is consistent with the surface-area dependency of the catalytic activities (Fig. S12†).

We observed that the crystal shapes of [1]ClO₄₂ and [1]SO₄ are cubic, unlike the octahedral shape of the [1]Cl₂ and [1]NO₃₂ crystals. To determine the difference in the catalase-like activities between the octahedral and cubic crystals, the
same size of crystals (ca. 1.0 × 1.0 × 1.0 mm) of these compounds were treated with aqueous H₂O₂ under the same conditions, eliminating the size-dependent effect (Fig. S13†). As expected, all the crystals showed O₂ evolution, which was confirmed by GC measurements (Fig. 3a). The TOFs at 15 min for the octahedral crystals [1]Cl₂ and [1]NO₃ were more than 5 times higher than those for the cubic crystals [1]ClO₄ and [1]SO₄ (Fig. 3b). No significant difference in the TOF values was observed between [1]Cl₂ and [1]NO₃ or between [1]ClO₄ and [1]SO₄. This result clearly indicates that the heterogeneous catalytic activity of [1]Xₙ is highly dependent on the crystal shape, rather than the type of anionic species in the crystal.

Crystal surface morphology and oxidation state of cobalt centers of [1]Xₙ

The crystal-shape dependency of the catalytic-like activity of [1]Xₙ can be discussed from the viewpoint of the surface morphology. Scanning electron microscopy (SEM) images of the crystal surfaces revealed that the crystal growth of the (111) plane is dominant for [1]Cl₂ and [1]NO₃, whereas growth of the (100) plane is dominant for [1]ClO₄ and [1]SO₄, as expected based on their macroscopic crystal shapes (Fig. 4). Thus, it is reasonable to assume that the catalytic activity on the (111) plane is higher than that on the (100) plane, considering the higher activity of the [1]Cl₂ and [1]NO₃ crystals. This tendency contrasts with the catalytic activities of platinum nanoparticles, for which the (100) plane has higher activity than the (111) plane; this phenomenon has been ascribed to a looser alignment of platinum atoms on the (100) plane. The surface structure of each crystal plane of [1]Xₙ, determined by single-crystal X-ray analysis, is shown in Fig. 5. The (111) plane is a d-pen-rich surface, and the cobalt centers are fully exposed by the easy dissociation of the carboxyl group of d-pen, as discussed below. On the other hand, the (100) plane is a dppe-rich surface, and the cobalt centers are much less exposed. Thus, the (111) plane is advantageous due to the exposure of the cobalt centers on the crystal surface, which leads to high catalytic activity.

Here, we reconsider the oxidation state of the cobalt centers by focusing on the crystal surface. Cobalt centers in [1]Xₙ have a +III oxidation state in the crystal as well as in solution. Consistent with this, in the partial fluorescence yield (PFY) mode of X-ray absorption spectroscopy (XAS), which is appropriate for bulk-layer observations, the crystal [1]Cl₂ exhibited a single L₃-edge peak at 780 eV due to cobalt(III) species (Fig. 6a). However, in the partial electron yield (PEY) mode of XAS, which is appropriate for surface-layer observations, a clear shoulder (mark A) was observed at a lower energy of the L₃-edge peak. The presence of this shoulder is indicative of the presence of cobalt(II) species; therefore, the cobalt species located on the crystal surface at least partially exist in a +II oxidation state, although cobalt(III) species are dominant in the bulk sample (Fig. 6b). This is supported by the EPR spectra of crystal [1]Cl₂ at 4 K, which showed a very weak, broad signal at approximately g = 6 that is assigned to cobalt(II). For [2]ClO₄, which does not exhibit catalase-like activity, the EPR-silent state suggests the absence of cobalt(II). It is quite reasonable that the aggregated cluster structure of inorganic anions in [1]Xₙ cannot be maintained on the crystal surface due to high repulsive energy.

Fig. 3 (a) Time-dependent profiles of the evolution of O₂ during treatment with a catalytic amount of 1.0 × 1.0 × 1.0 mm-sized crystals of [1]Cl₂ (5.2 mg), [1]NO₃ (5.1 mg), [1]ClO₄ (4.8 mg) or [1]SO₄ (5.3 mg) with 5% aqueous H₂O₂ (1.00 mL) at 298 K and (b) TOF calculated based on the number of surface Co atoms at 15 min.


Fig. 5 Side and top views of the arrangement of cationic supramolecular octahedrons ([1]²⁺)ₙ on the crystal surface of the (100) plane and the (111) plane.
among inorganic anions. This disruption of the aggregated anionic cluster structure should generate defects of inorganic anions, which in turn, leads to the reduction of cobalt(III) to cobalt(II) on the crystal surface to balance the total charge of the ionic crystal. Thus, the presence of cobalt species with a +II oxidation state, which are efficiently exposed at the (111) surface of the crystal, is key to the high catalase-like activity of the octahedral crystals [1]Cl₂ and [1](NO₃)₂.

The mechanism of the catalase-like reaction of [1]Xₙ

The mechanism of the catalase-like reaction in this system was estimated as follows, based on the mechanism proposed for the iron(III) center of native catalase but replacing the redox active site from iron to cobalt and the deprotonation site from histidine to carboxylate (Scheme 1). In the first step of this reaction, an H₂O₂ molecule approaches an octahedral cobalt(II) center, with the concomitant dissociation of a carboxyl group to form a five-coordinated cobalt(II) center. The carboxylate group acts as a base in the deprotonation of the H₂O₂ molecule, giving rise to a Co–OOH intermediate. The next step involves the formation of an oxo-cobalt(IV) center and the release of an H₂O molecule via a heterolytic O–O bond cleavage. Finally, another H₂O₂ molecule attacks the oxo-cobalt(IV) center to produce the original octahedral cobalt(II) center, with the concomitant release of H₂O and O₂ molecules during the formation of a hydroxo-cobalt(II) center and an HOO’ radical. Each transition state was calculated with the effect of solvent (H₂O) at the UB3LYP-D/6-31G* level, which revealed that the O–O bond cleavage is the rate-determining step (RDS) with an activation energy of 21.2 kcal mol⁻¹ relative to the substrate complex (Fig. S14†). We also calculated each transition state of the cobalt(II) model, in which the cobalt(III) center is replaced by a cobalt(II) center. In the cobalt(II) model, the activation energy of the RDS (O–O bond cleavage) was calculated as 26.7 kcal mol⁻¹, which is higher than that in the cobalt(II) model (Fig. S15†). This result indicates that the cobalt(II) model is preferable for this catalase-like reaction.

Conclusions

We showed that [1]Xₙ ionic crystals, which consist of Au₄CoIII₂ complex cations ([1]⁺) and inorganic anions (X = Cl⁻, NO₃⁻, ClO₄⁻, and SO₄²⁻), act as heterogeneous catalysts for a catalase-like reaction. To our knowledge, this is the first example of cobalt(II) coordination compounds that exhibit catalase-like activity under heterogeneous conditions. Of note is the remarkably high catalytic activity, which is dependent on the crystal shape (octahedral vs. cubic) rather than the charge (−1 vs. −2) and the geometry of the inorganic anions. This is due to the unusual arrangement of cations and anions in the crystal [1]Xₙ, leading to the generation of cobalt(II) centers on the crystal surface, which are efficiently exposed on the (111) plane with readily dissociated carboxyl groups compared to the (100) plane. This study demonstrated that the oxidation state and exposure of metal centers on the crystal surface that are required for heterogeneous catalytic activity can be controlled via the arrangement of cationic and anionic components as well as the crystal shape and morphology. These results provide significant insight into the future design and creation of various heterogeneous catalysts based on ionic crystals.

Experimental

Materials

Chemicals were purchased from commercial sources and used without further purification. Aqueous solutions of hydrogen peroxide (30 wt%) were purchased from Wako Pure Chemical Industries. Ultra-pure water was obtained from a Millipore Direct-Q3 UV water purification system, wherein the electronic conductance was 18.2 MΩ cm⁻¹. [Au₄CoIII₂(dppe)₂(D-pen)₄]Cl₂ ([1]X₄n), [XI₄, X = (Cl⁻)₂, (ClO₄⁻)₂, (NO₃⁻)₂ or SO₄²⁻), [Au₄Cr₂(dppe)₂(D-pen)₄]Cl₂, [CoIII(AuI[PPh₃]₄)(D-pen)₄]ClO₄ ([2]ClO₄) and [Au₄(dppe)₄(HPen)₄] were prepared according to the methods reported in the literature.¹¹⁻¹⁸

Catalase-like catalytic disproportionation of hydrogen peroxide

A typical procedure for the catalase-like catalytic disproportionation of hydrogen peroxide is as follows. A vial containing 1.0 mL of 5 wt% aqueous H₂O₂ and another vial containing [1]Cl₂ (5.0 mg) were sealed with rubber septa. The two vials were carefully deaerated by bubbling Ar for 15 min. The aqueous H₂O₂ solution was transferred to the vial containing crystals via pipette, and the solution was mixed with the [1]Cl₂ solution with a magnet stirrer. The reaction mixture was then stirred at room temperature. After a known reaction time, 1.0 mL of the reaction solution was taken out with a pipette, added to 1.0 mL of 10 vol% (v/v) aqueous HCl (0.1 M), and the absorbance at 230 nm was measured by a UV spectrophotometer.

a Teflon tube to start the reaction without stirring at 298 K. At each reaction time, 100 μL of Ar gas was injected into the vial, and the same volume of gas in the headspace was then sampled by a gastight syringe and quantified on a Shimadzu GC-17A gas chromatograph (GC) [Ar carrier gas, capillary column with molecular sieves (Agilent Technologies 19094PMS0, 30 m × 0.53 mm, 313 K) equipped with a thermal conductivity detector (TCD). Microscopic observation of the crystal sizes and filtration treatment were performed as needed.

ICP-AES measurements
ICP-AES was measured on a Shimadzu ICPS-8100. The samples were obtained by filtration of the reaction mixture at 20–30 min in conditions similar to the reaction for GC measurements.

XPS measurements
XPS spectra were measured on a Kratos Axis 165x with a 165 mm hemispherical electron energy analyzer. The incident radiation was a Mg-Kz X-ray (1253.6 eV) at 120 W (12 kV, 10 mA), and a charge neutralizer was turned on for acquisition. Each sample was mounted on a stainless-steel stage with double-sided carbon scotch tape. The binding energy of each element was corrected using the C 1s peak (284.5 eV) from the residual carbon.

NMR measurements
$^1$H and $^{31}$P NMR spectra were measured on an ECA 500 spectrometer at 298 K. The NMR spectra were calibrated with TMS (methyl-d$_4$) = 0 ppm as an internal standard.

PXRD measurements
High-quality powder X-ray diffractions were recorded at room temperature, in transmission mode [synchrotron radiation $\lambda = 0.999139(2)$ Å; $0^\circ \leq 2\theta \leq 30^\circ$; step width = 0.006°; data collection time = 30 s] on a diffractometer equipped with a MYTHEN microstrip X-ray detector (Dectris Ltd.) at the SPring-8 BL02B2 beamline. The crystals were loaded into glass capillary tubes (diameter = 0.3 mm), and the samples were rotated during the measurements.

SEM measurements
SEM observations were performed on an FEI Magellan 400L instrument at $5 \times 10^{-5}$ Pa. The working distance was set to 1.9–2.7 mm. Secondary electrons were collected with an Everhart-Thornley detector. The observations were performed at a beam landing voltage of 200 or 1000 V, and 200 V imaging was performed by applying 800 V of the beam deceleration bias onto the sample against the primary electron beam accelerated at 1 kV.

EPR measurements
The sample for EPR was prepared by flowing N$_2$ for 15 min through a Teflon tube to the crystals in a quartz EPR tube (2.0 mm i.d.). EPR spectra of the solutions were obtained on a JEOL X-band spectrometer (JES-FA200) at 4 K. The g value was calibrated using a Mn$^{2+}$ marker. The EPR spectra were obtained under non-saturating microwave power conditions. The magnitude of modulation was chosen to optimize the resolution and signal-to-noise (S/N) ratio of the observed spectra.

XAS measurements
The XAS measurements were carried out at BL-11 of the Synchrotron Radiation Center at Ritsumeikan University, Japan. In this beamline, so-called varied-line-spacing plane gratings were employed to supply monochromatic photons with $h\nu = 40$–1000 eV. The Co L$_{2,3}$-edge XAS spectra ($h\nu = 760$–810 eV), were taken simultaneously in the partial fluorescence yield (PFY) and partial electron yield (PEY) modes with a photon energy resolution of $\sim$300 meV. For the Co L$_{2,3}$-edge XAS measurements, the luminescence with $h\nu = 700$–950 eV, including the Co L lines, was detected as a signal. On the other hand, in PEY mode, the microchannel plate (MCP) for detecting the Auger and secondary electrons was set in the 45°-depression to the photon propagation. On the front of the MCP, a gold mesh was installed to enable the application of a voltage. We applied the voltage of $\sim$550 V to the mesh for the Co L$_{2,3}$-edge measurements in order to suppress a strong background caused by the C, N and O K-edge absorptions in the spectra. The powder-like single-crystalline samples of [1]Cl$_2$ and [2]ClO$_4$ were thinly expanded on the conductive carbon tape attached on the sample holder in air before transferring them to the vacuum chamber. We repeatedly measured the spectra on the same and different sample positions to confirm the data reproducibility with neither serious radiation damage nor a sample-position dependence of the Co L$_{2,3}$-edge XAS spectra. The measurements were performed at room temperature. Energy values were calibrated with the top of the L$_3$-edge peak of LiCo$^{3+}$O$_2$ (780.32 eV).

Theoretical calculations
Geometry optimizations were carried out at the UB3LYP-D/6-31G* level of theory for each transition state, as implemented in Gaussian 09, revision C.01. The effect of solvent (H$_2$O) was included implicitly using the polarizable continuum model (PCM) with the cavity constructed by the universal force-field (UFF) bond radii. The charges and spin multiplicities of the Co center models in the initial state were $-2$ and $4$ for the Co$^{II}$ center model and $-1$ and $1$ for the Co$^{III}$ center model, respectively.

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Notes and references


12 The saturated solution of [1]Cl2 (0.18 mM) was produced by standing crystals [1]Cl2 in H2O overnight (Table S1 in the ES†).

13 The O2 evolution was completely stopped when the crystals of [1]Cl2 were removed from the reaction mixture (Fig. S3†). The ICP-AES showed that only a quite small portion of the crystals (<0.2% for [1]Cl2, <0.03% for [1]([NO3]2) was dissolved during the reaction (Table S2†).


16 The number of Co atoms at the crystal surface was evaluated based on the X-ray analysis, assuming that each crystal has a perfect octahedral shape with uniform size.

17 Representatives of the reported best catalasa-like activities are summarized in Table S3† in comparison with those of [1]Cl2 and [1][NO3]2.


19 Gas leak with overpressure can cause saturation of O2 evolution considering saturation at a similar level regardless of the different reaction rates and reusability of the catalyst. Partial cracking of crystals may also affect the reaction rate.


22 The ratio of CoII : CoIII was calculated to be 11% : 89% by simulated fitting of PEY from CoII and CoIII standards to reproduce the XAS spectra.


