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Synthesis, Crystal Structure, and Adsorption Properties of Werner-type Cu(II) Complex $[\text{Cu}(\text{CF}_3\text{SO}_3)_2(4\text{-methylpyridine})_4]$

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A Werner-type Cu(II) complex, α - $[\text{Cu}(\text{CF}_3\text{SO}_3)_2(4\text{-mepy})_4]$ (α -PAC-1- CF_3SO_3 , PAC: porous assembly of coordination complex and 4-mepy: 4-methylpyridine), was synthesized, crystallographically characterized, and its adsorption properties were compared with those of the derivative α - $[\text{Cu}(\text{PF}_6)_2(4\text{-mepy})_4]$ (α -PAC-1- PF_6).

Porous assemblies of coordination complexes (PACs), which are formed by the assembly of discrete coordination complexes without internal spaces via intermolecular interactions (van der Waals, electrostatic, and hydrogen-bonding interactions) that are weaker than coordination bonds,^{1–3} have the potential to provide the following unique host–guest features: (i) diversity in assembled structures with guest molecules, resulting from weak and flexible intermolecular interactions;^{1h,1i} (ii) formation of host frameworks consisting of two or more different discrete coordination complexes;^{1j} (iii) facile fabrication of films because of high solubility;^{1k,1l} and (iv) polymorphism without guest molecules, which is controlled by a kind of preadsorbed guests.^{1c,1i} These characteristics of PACs could supply not only fundamental information about their pores but also unprecedented, useful porous functions.

The Werner-type complex $[\text{Ni}(\text{NCS})_2(4\text{-mepy})_4]$ (4-mepy: 4-methylpyridine) and its derivatives are well known as examples of PACs; their preparation and guest-inclusion abilities were first reported in 1957.^{1a} Among them, $[\text{Ni}(\text{NCS})_2(4\text{-mepy})_4]$ has been thoroughly studied because of its interesting guest-inclusion properties derived from pseudopolymorphism, such as a dense α -form, and guest-including β - and γ -forms. The α -form has no space for including guest molecules because of its dense packing structure, whereas the β - and γ -forms are channel-type and layer-type clathrates, respectively.¹ⁱ In particular, this complex easily affords an empty β -form that adsorbs several gases with type I isotherms.

We have recently found that a Cu(II)-based PAC, $[\text{Cu}(\text{PF}_6)_2(4\text{-mepy})_4]$ (PAC-1- PF_6), which is a quasi-Werner-type metal complex, has interesting guest-recognition abilities.⁴ The dense α -form, α -PAC-1- PF_6 , has inorganic PF_6^- anions covered with only fluorine atoms in the framework, and it adsorbs CO_2 gas with structural transformations; this is the first example of gas-adsorption properties in the dense α -form. On the other hand, when α -PAC-1- PF_6 is recrystallized from acetone/*n*-hexane and 2-butanone/*n*-hexane, some of the weakly coordinated PF_6^- anions are easily released from the axial sites and Lewis-base guests (acetone and 2-butanone) attach to these sites instead, forming the guest-including γ -form, γ - $\{[\text{Cu}(\text{PF}_6)_2(4\text{-mepy})_4][\text{Cu}(\text{PF}_6)(4\text{-mepy})_4(\text{acetone})]\cdot\text{PF}_6\cdot 4\text{acetone}\}$ (γ -PAC-1- $\text{PF}_6 \supset 2.5\text{acetone}$), γ - $\{[\text{Cu}(\text{PF}_6)_2(4\text{-mepy})_4][\text{Cu}(\text{PF}_6)(4\text{-mepy})_4(2\text{-butanone})]\cdot\text{PF}_6\cdot 3.5(2\text{-butanone})\}$ (γ -PAC-1- $\text{PF}_6 \supset 2.25(2\text{-butanone})$). The prefix “quasi-” derives from such labile axial bonds. These unprecedented phenomena have never been observed in any other Werner-type metal complexes. These results clearly indicate that Cu(II)-based PACs are useful for the construction of porous materials with multiguest-recognition properties.

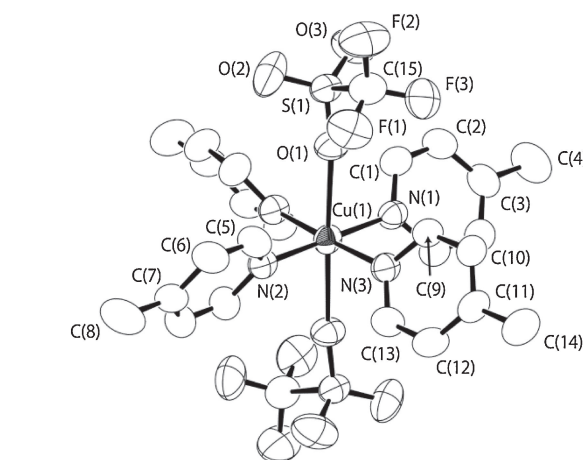


Figure 1. ORTEP drawing around the Cu(II) center in α -PAC-1- CF_3SO_3 . The hydrogen atoms are omitted for clarity.

In this letter, we report the synthesis, crystal structure, and adsorption properties of a derivative of α -PAC-1- PF_6 , α - $[\text{Cu}(\text{CF}_3\text{SO}_3)_2(4\text{-mepy})_4]$ (α -PAC-1- CF_3SO_3), to investigate the effect of inorganic anions on the adsorption properties. The introduction of CF_3SO_3^- anions enhanced the intermolecular interactions through the oxygen atoms of the anions and inhibited gas adsorption with structural changes.

α -PAC-1- CF_3SO_3 was synthesized as follows: $\text{Cu}(\text{CF}_3\text{SO}_3)_2$ (362 mg, 1.0 mmol) and 4-mepy (466 mg, 5.0 mmol) were dissolved in acetone (20 mL) and excess *n*-hexane was added to the acetone solution, forming blue microcrystals of α -PAC-1- CF_3SO_3 . Single crystals suitable for X-ray diffraction analysis were obtained by recrystallization from CHCl_3 /*n*-hexane.

The crystal structure of α -PAC-1- CF_3SO_3 was determined by single-crystal X-ray diffraction analysis at 173 K.⁵ Figure 1 shows the ORTEP view around the Cu(II) ion. The Cu(II) ion has an elongated octahedral environment with four 4-mepy molecules in the equatorial plane and two CF_3SO_3^- anions at the axial sites. The Cu–O bond distance of 2.468(2) Å is considerably longer than the equatorial Cu–N values (2.014(4)–2.025(3) Å), indicating that the Jahn–Teller axis is formed along the O–Cu–O direction. The axial bonds in α -PAC-1- CF_3SO_3 are shorter than those in α -PAC-1- PF_6 (Cu–F bond dis-

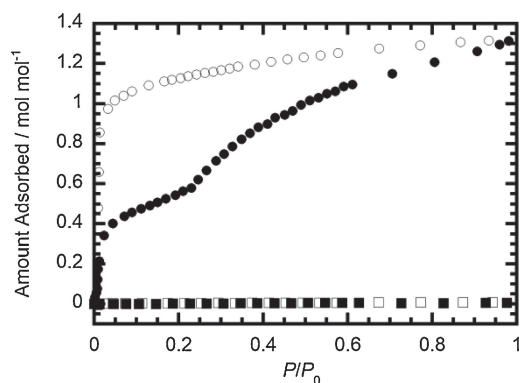


Figure 2. Adsorption (filled symbols) and desorption (open symbols) isotherms for CO₂ at 195 K on α -PAC-1-CF₃SO₃ (squares) and α -PAC-1-PF₆ (circles).

tances = 2.478(2), 2.528(2), 2.586(2), and 2.629(2) Å),⁶ which is a reasonable result because CF₃SO₃[−] anions have stronger Lewis basicity than PF₆[−] anions.⁷ Each mononuclear Cu(II) complex is densely packed, resulting in the assembled structure having no pores (accessible void space calculated using the PLATON program is 0%).⁸

The adsorption and desorption isotherms for CO₂ at 195 K on α -PAC-1-PF₆ and α -PAC-1-CF₃SO₃ were measured to compare the effects of inorganic anions. As shown in Figure 2, α -PAC-1-PF₆ shows CO₂ adsorption. Because α -PAC-1-PF₆ possesses no vacant space, the observed adsorption behavior with large hysteresis should be associated with gate-opening processes, in which adsorption occurs with structural transformations.⁴ On the other hand, α -PAC-1-CF₃SO₃ scarcely adsorbs CO₂ gas over the entire pressure range. In general, the gate-opening pressure depends on the strength of host–host and/or host–guest interactions. Weak host–host and strong host–guest interactions decrease the gate-opening pressure. First, host–host interactions were checked from the viewpoint of (1) the type of intermolecular interactions, and (2) their distances and angles. In α -PAC-1-CF₃SO₃, there are weak intermolecular hydrogen bonds between noncoordinated oxygen atoms of CF₃SO₃[−] anions and hydrogen atoms of 4-mepy ligands (O...C = 3.33, 3.40, and 3.42 Å, Figure 3). α -PAC-1-PF₆ bears slightly shorter intermolecular contacts between noncoordinated fluorine atoms of PF₆[−] anions and hydrogen atoms of 4-mepy ligands (F...C = 3.19–3.32 Å, see Supporting Information).¹⁴ However, considering the van der Waals radii of oxygen and fluorine atoms (1.52 and 1.47 Å, respectively) and the C–H(pyridyl)...O(F) angles (see Table S2),^{9,14} the slight difference observed in the intermolecular distances may not reflect a difference in the strength of the intermolecular interactions. On the other hand, as confirmed by DFT calculations for α -[Cu(A)₂(py)₄] (α -PAC-2-A, A = PF₆ and CF₃SO₃, py: pyridine), α -PAC-2-PF₆ bears an almost uniform negative potential over all the surface fluorine atoms of PF₆[−]. In contrast, the CF₃SO₃[−] anions in α -PAC-2-CF₃SO₃ have negative potentials on their oxygen atoms; their magnitude is higher than that on the fluorine atoms in α -PAC-2-PF₆.¹⁰ Moreover, a fluorine atom, with a small polarizability, weakens intermolecular interactions. These results imply that host–host interactions (including acid–base and van der Waals interactions) in α -PAC-1-PF₆ are weaker than those in α -PAC-1-

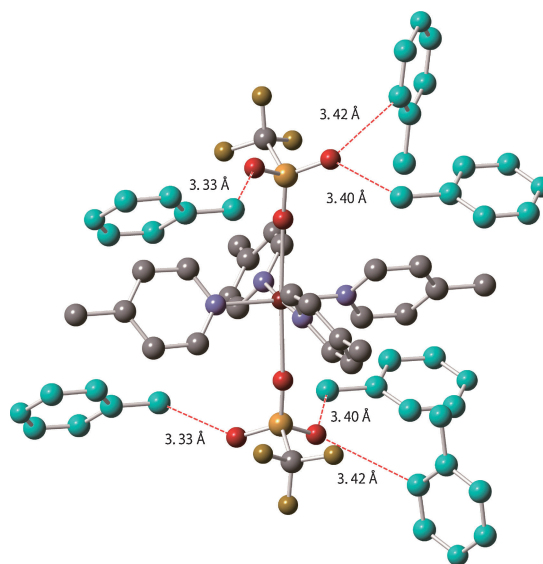


Figure 3. View of the intermolecular interactions in α -PAC-1-CF₃SO₃, in which the red dashed lines indicate hydrogen bonds. The 4-mepy molecules of other mononuclear complexes are represented in sky blue. The hydrogen atoms are omitted for clarity.

CF₃SO₃. Next, host–guest interactions were compared. It is known that a CO₂ molecule interacts with these fluorinated anions via weak acid–base interactions in ionic liquids.¹¹ As mentioned above, the Lewis basicity of CF₃SO₃[−] anions is stronger than that of PF₆[−] ones. Furthermore, the results of DFT and MP2 calculations show that the binding energy in the CF₃SO₃[−]–CO₂ complex is slightly higher than that in PF₆[−]–CO₂.¹² Hence, the host–guest interaction between the [Cu(PF₆)₂(4-mepy)₄] unit and CO₂ is probably weaker than that between the [Cu(CF₃SO₃)₂(4-mepy)₄] unit and CO₂. This comparison of host–host and host–guest interactions indicates that host–host interactions predominantly influence gate-opening adsorptions in these PACs.

A similar tendency has been observed in the two-dimensional Cu(II) porous coordination polymers (PCPs) [Cu(A)₂(4,4′-bpy)₂]_n (A = PF₆ and CF₃SO₃, 4,4′-bpy: 4,4′-bipyridine).^{10,13} Both PCPs have similar two-dimensional frameworks with weakly coordinated inorganic anions, PF₆[−] and CF₃SO₃[−], at the axial sites and show two kinds of adsorption events: micropore filling in open voids and gate-opening adsorption with expansion of the layers. The gate-opening pressures for [Cu(PF₆)₂(4,4′-bpy)₂]_n are considerably lower than those for [Cu(CF₃SO₃)₂(4,4′-bpy)₂]_n. These results can also be explained by the difference in the strengths of host–host interactions.

In conclusion, we achieved the synthesis and crystallographic characterization of the Werner-type Cu(II)-based PAC α -[Cu(CF₃SO₃)₂(4-mepy)₄] (α -PAC-1-CF₃SO₃). The weak hydrogen-bonding interactions between the oxygen atoms of the CF₃SO₃[−] anions and the hydrogen atoms of the 4-mepy ligands make the assembled structure rigid, resulting in the suppression of CO₂ adsorption with structural transformations. This finding could help in designing flexible PCPs that show gate-opening adsorption. In addition, although α -PAC-1-CF₃SO₃ did not

show porosity for CO₂ gas, other organic vapors, such as alcohols, ketones, and benzene, may adsorb on α -PAC-1-CF₃SO₃ with structural transformations.⁴ Further work is therefore in progress to isolate the guest-including β - and γ -PAC-1-CF₃SO₃ and check the vapor-adsorption properties of α -PAC-1-CF₃SO₃.

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