MS.80.5

Acta Cryst. (2011) A67, C178

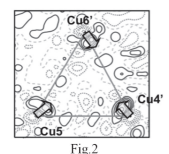
Photoexcited state crystallography of luminescent hexanuclear d¹⁰ metal complexes

Yoshiki Ozawa, Toru Ishida, Koshiro Toriumi Departament of Material Science, University of Hyogo (Japan). E-mail: ozawa@sci. u-hyogo.ac.jp

Hexanuclear metal complexes of Cu(I) or Ag(I) $[M_6(R\text{-pyt})_6]$ (M = Cu, Ag; pyt = pyridinethiolato) (Fig. 1) give intense luminescence under UV illumination in the solid-state. The nature of the emission has been assigned to a triplet multi-metal cluster-centered (3 CC) transition state like as in the halogen bridged Cu(I) tetranuclear cubane-like cluster complexes. [1,2] On the photo-excited states, the metal-cluster core is expected to be shrunk because of an electron migration to the CC orbital, that has a bonding-character. To figure out the photo-emission process by direct observation of the molecular distortion, we have performed single crystal X-ray structure analyses at the photo-excitation states.

X-ray diffraction experiments under photo-irradiation were performed at the SPring-8 BL02B1 station. A single crystal of the copper(I) complex with ethyl-pyridinethiole ligands [Cu₆(Et-pyt)₆] (Et-pytH=6-ethylpyridine-2-thione) was mounted on the vacuum camera at 25 K was used for X-ray diffraction data collection under UV laser (325/442 nm, 0.03/0.1 W) irradiation. Photo-difference Fourier syntheses at the section of the triangle Cu₃ plane perpendicular to the virtual 3-fold axis of the molecule show that two of three Cu atoms move toward to the remaining Cu atom (Fig. 2). This indicates that the contraction of Cu-Cu atomic distance will occur at the photo-excitation state resulting in shrinkage of the metal cluster core. We have also performed the same experiments for the silver complex [Ag₆(Et-pyt)₆]. The observed peaks and holes of charge densities in the photo-difference Fourier maps are similar to that in the Cu crystal, although their amplitude is below the significant levels.





[1] H. Xie, I. Kinoshita, T. Karasawa, K. Kimura, T. Nishioka, I. Akai, K. Kanemoto, *J. Phys. Chem. B*, **2005**, *109*, 9339-9345. [2] P.C. Ford, A. Vogler, *Acc. Chem. Res.* **1993**, *26*, 220-226.

Keywords: photochemistry, accurate measurement, synchrotron radiation

MS.81.1

Acta Cryst. (2011) A67, C178

A-site ordered perovskite-structure oxides with functional properties

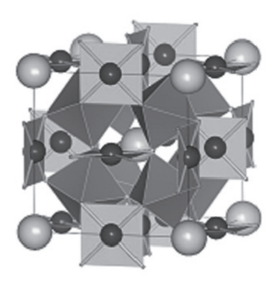
Yuichi Shimakawa, Institute for Chemical Research, Kyoto University, JST, CREST, Kyoto, (Japan). E-mail: shimak@scl.kyoto-u.ac.jp

A very unusual ordered arrangement of the A-site ions in the simple ABO_3 perovskite produces $AA'_3B_4O_{12}A$ -site ordered perovskites. The A site accommodates alkaline metals, alkaline-earth metals, and

lanthanides like those in the simple perovskites. At the originally 12-fold coordinated A' site, transition-metal ions such as Cu^{2+} and Mn^{3+} form square coordinated units that align perpendicular to each other. The presence of the transition-metal ions at both A' and B sites produces A'-A' and and/or A'-B interactions in addition to B-B interaction usually seen in the simple perovskite materials. Competitive and/or cooperative interplay of these interactions gives rise to diverse and intriguing physical properties [1].

Two new *A*-site ordered perovskite-structure oxides, CaCu₃Fe₄O₁₂ [2] and LaCu₃Fe₄O₁₂ [3], are highlighted. They were synthesized under high-pressure and high-temperature conditions. The compounds contain unusually high valence states of iron: Fe⁴⁺ in CaCu₃Fe₄O₁₂ and Fe^{3.75+} in LaCu₃Fe₄O₁₂. Instabilities of the high oxidation states at low temperatures are resolved in CaCu₃Fe₄O₁₂ by charge disproportionation from Fe⁴⁺ to Fe³⁺ and Fe⁵⁺, and in LaCu₃Fe₄O₁₂ by charge transfer between *A*-site Cu and *B*-site Fe ions. The charge disproportionation and the charge transfer are accompanied by significant changes in structural, transport, and magnetic properties.

We also show a few examples of new A-site ordered perovskites with intriguing physical properties. BiCu₃Mn₄O₁₂ [4] is a ferrimagnetic metal. The half metallic nature of the material produces spin-polarized conduction electrons leading to large magnetoresistance under a low applied field. In CaCu₃Ge₄O₁₂ - CaCu₃Ti₄O₁₂ - CaCu₃Sn₄O₁₂ sysystem [5], we can see very unusual A '-site magnetis, in which either ferromagnetic or antiferromagnetic behavior of A '-site Cu²⁺ (S = 1/2) spins can appear within the same structural framework.



Crystal structure of A-site ordered perovskite

[1] Y. Shimakawa, Inorg. Chem. 2008, 47, 8562.
[2] I. Yamada, et al., Angew. Chem. Int. Ed. 2008, 47, 7032.
[3] Y.W. Long, et al., Nature 2009, 458, 60.
[4] K. Takata, et al., Phys. Rev. B 2007, 76, 024429.
[5] Y. Shimakawa, et al., J. Phys. Soc. Jpn. 2008, 77, 113702.

Keywords: A-site ordered perovskite-structure oxides, crystal structure, physical property

MS.81.2

Acta Cryst. (2011) A67, C178-C179

Electron crystallography for li-based battery materials

Joke Hadermann, Artem M. Abakumov, Nellie R. Khasanova, Evgeny V. Antipov, Gustaaf Van Tendeloo, Artem M. Abakumov, Mellie R. Khasanova, Wastery V. Antipov, Moscow, Nellie R. Khasanova, Ewgeny V. Antipov, Gustaaf Van Tendeloo, Artematic Mellie R. Khasanova, Wastery Van Tendeloo, Artematic Mellie R. Khasanova, Wastery Van Tendeloo, Artematic Mellie R. Khasanova, Wastery Van Tendeloo, Artematic Mellie R. Khasanova, Nellie R. Khasanova, Mellie R. Kh