アカデミックプログラム [B講演] | 07. 無機化学:口頭B講演

**=** 2025年3月27日(木) 15:55~17:15 **=** [C]C304(第2学舎 2号館 [3階] C304)

[[C]C304-2vn] 07. 無機化学

座長:河底秀幸、森合達也

#### ● 日本語

15:55 ~ 16:15

[[C]C304-2vn-01]

フラックス法による層状カルコゲナイド化合物の単結晶育成と剥離応用

〇林 文隆 $^1$ 、横塚 雄斗 $^1$ 、Chokradjaroen Chayanaphat $^1$ 、工藤 諒大 $^1$ 、山田 哲也 $^1$ 、手嶋 勝弥 $^1$  (1. 信州大学)

### ● 英語

16:15 ~ 16:35

[[C]C304-2vn-02]

ラムダ型五酸化三チタンの蓄熱特性評価と金属置換体の合成

〇清木 陸 $^1$ 、藤澤 聖斗 $^1$ 、Akhmad Fadilla $^1$ 、久保田 智子 $^1$ 、大越 慎一 $^2$ 、所 裕子 $^1$  (1. 筑波大学、2. 東京大学)

### ● 英語

16:35 ~ 16:55

[[C]C304-2vn-03]

X線分光と第一原理計算による多元素合金触媒の電子構造-物性相関の定量評価

〇中村 雅史 $^1$ 、Dongshuang Wu $^1$ 、向吉 恵 $^1$ 、草田 康平 $^{1,2}$ 、林 博之 $^3$ 、鳥山 誉亮 $^4$ 、山本 知一 $^4$ 、村上 恭和 $^{4,5}$ 、河口 彰吾 $^6$ 、伊奈 稔哲 $^6$ 、久保田 佳基 $^7$ 、家路 豊成 $^8$ 、小島 一男 $^8$ 、田中 功 $^3$ 、北川 宏 $^1$  (1. 京大院理、2. 京大白眉セ、3. 京大院工、4. 九大URC、5. 九大院工、6. JASRI/SPring-8、7. 阪公大院理、8. 立命大SRC)

### ● 英語

16:55 ~ 17:15

[[C]C304-2vn-04]

Mild, Facile Synthesis of Perovskite Oxynitride Nanoparticles Towards High-entropy Thermoelectric Materials

OSimon David Moore<sup>1</sup>, Mari Takahashi<sup>1</sup>, Shinya Maenosono<sup>1</sup> (1. School of Materials Science, Japan Advanced Institute of Science and Technology)

### フラックス法による層状カルコゲナイド化合物の単結晶育成と剥離応用

(信州大)○林 文隆・横塚 雄斗・Chokradjaroen Chayanaphat・工藤 諒大・山田 哲也・手嶋 勝弥

Flux Growth and Exfoliation Applications of Chalcogenide Compounds Single Crystals (*Shinshu University*) oFumitaka Hayashi, Yuto Yokotsuka, Chayanaphat Chokradjaroen, Ryota Kudo, Tetsuya Yamada, Katsuya Teshima

Transition metal dichalcogenides (TMD) have been attractive for their semiconducting character. TMD materials are represented by the chemical formula  $MX_2$ , M refers to a transition metal atom (e.g., Mo, W) and X is a chalcogen atom (e.g., S, Se). TMD can be produced in an atomic-scale thickness with direct band gap and suitable spin-orbit coupling through chemical vapor deposition and exfoliation, which make it favorable for especially electronics applications. In general, it is challenging to grow large, high-quality parent  $MX_2$  crystals, and achieving scalable crystal growth is highly desirable. Herein, we report the flux growth of  $MoX_2$  (X = S, Se, Te) single crystals from chloride and molybdate-based fluxes for the first time. When examining the holding temperature using NaCl flux, large  $MoS_2$  crystals exceeding 50  $\mu$ m were obtained, although size variation was observed at 1100 °C. In contrast, using  $K_2MoO_4$  flux resulted in relatively uniform  $MoS_2$  crystal particles with an approximate size of 20  $\mu$ m. The grown  $MoX_2$  crystals were exfoliated using a hexane-butyllithium solution.

Keywords: Transition metal dichalcogenides; Nanosheet; Exfoliation; Two-dimensional materials

謝辞:本研究の一部は,内閣府 SIP 事業(JPJ012307)および科研費基盤研究 B(24K01234)の援助のもとに遂行された。記して謝意を表する。

## Evaluation of heat-storage properties of lambda-type trititanium pentoxide and synthesis of metal-substituted lambda-type trititanium pentoxide

(¹Dep. of Materials Sciences, Faculty of Pure and Applied Sciences, Univ. of Tsukuba, ²Dep. of Chemistry, School of Science, Univ. of Tokyo) ○Riku Seiki,¹ Akito Fujisawa,¹ Akhmad Fadel Fadilla,¹ Tomoko Kubota,¹ Shin-ichi Ohkoshi,² Hiroko Tokoro¹ **Keywords**: titanium oxides, heat-storage properties, pressure-induced phase transition, metal substitution

[Introduction] The lambda-type trititanium pentoxide ( $\lambda$ -Ti<sub>3</sub>O<sub>5</sub>, Fig. 1) was reported as a pressure-responsive heat-storage material to preserve thermal energy in the long-term <sup>[1,2]</sup>. In this study, we developed a synthesis method for  $\lambda$ -Ti<sub>3</sub>O<sub>5</sub> simply by preparing a precursor using titanium chloride as a starting material and evaluated the heat-storage properties <sup>[3]</sup>. Additionally, using the developed method, we synthesized manganese-substituted  $\lambda$ -Mn<sub>x</sub>Ti<sub>3-x</sub>O<sub>5</sub> and investigated the effect of Mn substitution for the heat-storage property.

[Experiment] Sample 1: A mixed solution of  $H_2O$ ,  $NH_3$ , and  $TiCl_4$  was prepared and stirred in a round bottle flask. The precipitation was extracted from the solution by centrifugation. Then the obtained precursor was sintered under a hydrogen flow of  $0.5 \, \mathrm{dm^3 min^{-1}}$  at  $1100 \, ^{\circ}\mathrm{C}$  for 5 h. Samples 2, 3, 4, 5: A mixed solution of  $H_2O$ ,  $NH_3$ , and  $TiCl_4$ ,  $MnCl_2 \cdot 4H_2O$  was prepared and stirred. The molar ratios of Mn to Ti were 3 % (2), 5 % (3), 7 % (4), and 10 % (5), respectively. The precipitation was extracted, washed with ethanol, and heated. The obtained precursors were sintered under a hydrogen flow.

[Result] X-ray fluorescence (XRF) measurement indicated that the composition formula of 1 was  $Ti_{3.00(5)}O_{5.00(5)}$  (Calc.: Ti 64.22, O 35.78 wt%; Found: Ti 64.53, O 35.47 wt%). XRD pattern with Rietveld analysis indicated that the obtained 1 was a single phase of  $\lambda$ - $Ti_3O_5$  (monoclinic, C2/m; a = 9.8332(2) Å, b = 3.78568(7) Å, c = 9.9688(2) Å,  $\beta = 91.259(2)$ °) and the crystallite size was estimated  $57 \pm 3$  nm. The heat-storage properties of 1, the pressure threshold ( $P_{th}$ ) was ca. 300 MPa and the transition enthalpy ( $\Delta H_{trans}$ ) was 7.78 kJ mol<sup>-1</sup> at 462 K. The relationship between the crystallite size and the heat-storage properties demonstrated that a reduction in crystallite size and an increase in ratio of surface atoms intensify the influence of surface energy on the Gibbs free energy and consequently it decreased the  $\Delta H_{trans}$ . The relationship between Mn-substituted ratio to Ti atoms and lattice constants (a, b, c) of 2, 3, 4, 5 were shown in Fig. 2. These lattice constants were changed depending on the Mn substitution ratio. The heat-storage property of 2, 3, 4, 5 will be presented in the oral presentation.

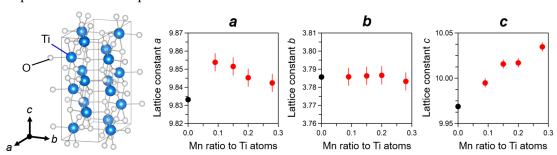


Fig. 1: Crystal structure Fig. 2: Three lattice constants versus Mn substitution ratio to Ti atoms plots of  $\lambda$ -Ti<sub>3</sub>O<sub>5</sub>. (left: a, center: b, right: c). Black plots are 1 and red plots are 2, 3, 4, 5.

### References

[1] S. Ohkoshi, et. al., *Nature Chem.* **2010**, *2*, 539. [2] H. Tokoro, et. al., *Nature Commun.* **2015**, *6*, 7037. [3] T. Kubota, R. Seiki, et. al., *Materials Advances*, **2024**, *5*, 3832.

# Quantitative Analysis of Electronic Structure–Property Relationship in Multi-Element Alloy Catalysts with X-ray Spectroscopy and *Ab Initio* Calculation

(¹Grad. Sch. Sci., Kyoto Univ., ²Hakubi Center, Kyoto Univ., ³Grad. Sch. Eng., Kyoto Univ., ⁴URC, Kyushu Univ., ⁵Grad. Sch. Eng., Kyushu Univ., ⁶JASRI/SPring-8, ¬Grad. Sch. Sci., Osaka Metro. Univ., ⁵SRC, Ritsumeikan Univ.) ⊙Masashi Nakamura¹, Dongshuang Wu¹, Megumi Mukoyoshi¹, Kohei Kusada¹.², Hiroyuki Hayashi³, Takaaki Toriyama⁴, Tomokazu Yamamoto⁴, Yasukazu Murakami⁴.⁵, Shogo Kawaguchi⁶, Toshiaki Ina⁶, Yoshiki Kubota⁻, Toyonari Yaji⁶, Kazuo Kojima⁶, Isao Tanaka³ and Hiroshi Kitagawa¹

Keywords: Multi-Element Alloy, Catalyst, XANES, DFT, Structure-Property Relationship

Although alloying is a powerful strategy to develop catalysts with desired properties, it is not self-evident how the inter-element interactions in alloys affect their catalytic properties. The recent prosperity of multi-element alloy (MEA) catalysts makes the problem more challenging<sup>1</sup>. Limited investigations into individual alloys are not enough to break through the situation. To discuss the interactions among different combinations of elements in a single framework, we propose that the quantitative analyses of element-selective electronic structures by X-ray spectroscopy<sup>2</sup> and *ab initio* calculation are prospective.

We synthesized a 5-element MEA of platinum-group metals (PGMs; Ru, Rh, Pd, Ir and Pt) and 6-element MEAs with an additional base metal (BM; Fe, Co, Ni, Ga, In or Sn), and investigated their performance in CO<sub>2</sub> hydrogenation. As a result, their CO selectivities ranged from 30 to 100%. The electronic structures of PGMs were quantified in two different approaches; XANES spectra were measured on the MEA catalysts to precisely evaluate the complicated real samples while *ab initio* calculations were performed on binary alloys to resolve the individual contributions of elemental interactions. Then, the correlations among the electronegativities of constituents, the quantities characterizing electronic structures (charges estimated by calculation or characteristic values of XANES spectra) and CO selectivity were examined. The results indicated that alloying PGMs with BM induced electron transfer following electronegativities, and the consequent modulation of the PGM valence led to the change in adsorption energies of intermediates and the selectivities (**Fig.**).

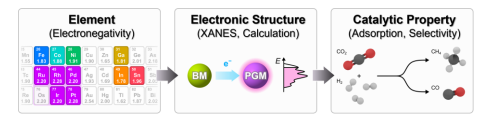


Fig. Schematics of element–electronic structure–property relationship in the MEA catalysts.

1) Y. Yao et al., Science, 2022, 376, eabn3103., 2) J. Chen, et al., J. Phys. Chem. C, 2022, 125, 2327.

### Mild, Facile Synthesis of Perovskite Oxynitride Nanoparticles Towards High-entropy Thermoelectric Materials

(School of Materials Science, Japan Advanced Institute of Science and Technology)

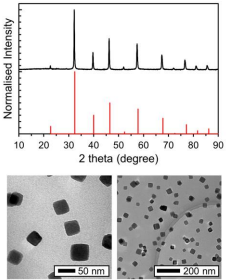
OSimon David Moore, Mari Takahashi, Shinya Maenosono

**Keywords**: Chemical synthesis; Strontium Titanate; Nitridation; Sustainable; High entropy nanoparticles

Currently commercially available thermoelectric (TE) materials – which can convert directly from a temperature difference to an electric current – have high conversion efficiency (represented by the ZT value), but make use of toxic or rare elements. In contrast, oxides, including perovskites, are known to be non-toxic and stable to high temperatures. However, their ZT values are low due to poor electrical conductivity, large band gap and high thermal conductivity. Increasing ZT generally involves the introduction of dopant elements such as La, Bi or Pb, which harms the sustainability of such materials. In this research, we focus on the synthesis of highly sustainable perovskite nanoparticles for future application as TE materials.

SrTiO<sub>3</sub> nanoparticles (NPs) (Fig. 1) were synthesized using a mild, facile gel-sol method,

followed by characterization by XRD, TEM, EDS, XPS, and UV-vis. Synthesis also included the introduction of different alloying elements, such as Ca and Mn, to the Sr and Ti sites. The introduction of multiple alloying elements is known to increase the configurational entropy, leading to benefits such as significantly reduced thermal conductivity. Following the synthesis of alloyed SrTiO<sub>3</sub> NPs a nitridation step was performed by mixing with urea at mild temperatures to substitute a portion of the O atoms with N and thus reduce the optical band gap. [4] The produced oxynitride NPs had a measured indirect optical band gap of 2.2 eV, significantly reduced from the 3.1 eV of the SrTiO<sub>3</sub> NPs.



**Figure 1:** XRD pattern and TEM images of assynthesised SrTiO<sub>3</sub> NPs. Red is the reference pattern for cubic SrTiO<sub>3</sub> (PDF: 00-005-0634)

#### References

[1] Shi, X. L. et al. *Nano Energy* **2020**, *78*, 105195. [2] Cardona, M. *Phys. Rev.* **1965**, *140*, 2A, A651-655. [3] Ma, J. et al. *J. Energy Storage* **2024**, *90*, 111890. [4] Atkinson, I. et al. *J. Photochem. Photobiol. A: Chem.* **2019**, *368*, 41-51.