Chemical Science

EDGE ARTICLE

Check for updates

Cite this: Chem. Sci., 2019, 10, 9125

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 22nd May 2019 Accepted 12th August 2019

DOI: 10.1039/c9sc02503k

rsc.li/chemical-science

Introduction

Tin oxide (SnO₂) has attracted increasing research attention due to its application in a variety of areas, including gas sensors,¹ catalysis,^{2,3} lithium batteries,^{4,5} solar cells^{6,7} and transparent electrodes.⁸ Some important factors, such as the size, composition, and structure, greatly influence the electronic and physicochemical properties of SnO₂.⁹⁻¹¹ Thus it is crucial to understand the binding mode and atomic connectivity of tin oxide materials at the molecular level, which will be beneficial for exploring the structure–property relationship and further achieving precise tuning of the physicochemical properties.

As molecular models of tin oxide materials, crystalline tinoxo clusters (TOCs) can provide precise atomic structural information by X-ray diffraction analysis. Accordingly, they are efficient molecular tools for building bridges between theoretical modeling, crystallography, and physical applications. Many research groups have made great efforts to explore the synthesis and structures of TOCs.^{12–22} One common way is to use organotin as precursors to react with carboxylate ligands in organic solvents, such as benzene, toluene, and alcohols.^{23–27} To date,

Assembly of high-nuclearity Sn₂₆, Sn₃₄-oxo clusters: solvent strategies and inorganic Sn incorporation[†]

Yu Zhu, Lei Zhang 🕩 * and Jian Zhang 🕩

A series of unprecedented high-nuclearity tin-oxo nanoclusters (up to Sn_{34}) with structural diversity have been obtained. The characteristics of the applied solvents had great influence on the assembly of these Sn-O clusters. Pure alcohol environments only gave rise to small clusters of Sn₆, whilst the introduction of water significantly increased the nuclearity to Sn₂₆, which greatly exceeds those of the known tin-oxo clusters (\leq 14); the use of aprotic CH₃CN finally produced the largest Sn₃₄ to date. Apart from the nuclearity breakthrough, the obtained tin-oxo clusters also present new structural types that are not found in previous reports, including a layered nanorod-like structure of Sn₂₆ and the cage-dimer structure of Sn₃₄. The layered Sn₂₆ clusters represent good molecular models for SnO₂ materials. Moreover, an electrode derived from TOC-17 with a {Sn₂₆} core shows better electrocatalytic CO₂ reduction activity than that from TOC-18 with Sn₃₄. This work not only provides an efficient methodology for the rational assembly of high-nuclearity Sn-O clusters, but also extends their potential applications in energy conversion.

the representative structural types of TOCs are characterized by cubic $[n-BuSn(O)O_2P(t-Bu)_2]_{4}$,²⁸ hexameric drum $\{R'SnO(O_2-CR)\}_{6}$,²⁹ ladder $[(R'Sn(O)O_2CR)_2R'Sn(O_2CR)_3]_2$,³⁰ and Keggin-type alkyltins.³¹⁻³³ However, despite these advances, the structural diversity of tin-oxo clusters is still less developed compared to transition-metal oxo clusters. And the TOC nuclearity also remains quite low, with the highest one being **Sn**₁₄.³⁴ Therefore, it is an attractive but challenging goal to synthesize TOCs with larger core nuclearity and more structural diversity. Moreover, although most of the driving forces of the research on TOCs originate from understanding the physical attributes of tin oxide, the studies on their applications (*e.g.* in catalysis) still remain very rare. Thus it is rather urgent to explore the physical application of TOCs to acquire the important Sn–O connectivity–activity relationship.

The assembly of tin-oxo clusters highly depends on the reactivity of organotin precursors, which is further influenced by organic ligands, reaction temperature, solvent and so on. Therefore, to prepare a diverse range of high-nuclearity TOCs, it is necessary to develop new synthetic approaches. During our recent research in titanium-oxo clusters,^{35–37} high-temperature assembly reactions were widely used for the preparation of high nuclearity Ti–O clusters. Herein, we introduce this method into the field of tin-oxo clusters, and make the following indispensable modifications: (1) mixed alcohol–water solvents were applied to influence the aggregation rate of Sn atoms and the nucleation rate of tin-oxo clusters; (2) the aprotic solvent CH₃CN was then used in some cases instead of the general protic solvent to affect the configuration of basic building blocks, as



View Journal | View Issue

State Key Laboratory of Structural Chemistry, Fujian Institute of Research on the Structure of Matter, Chinese Academy of Sciences, Fuzhou, Fujian 350002, P. R. China. E-mail: LZhang@fjirsm.ac.cn

[†] Electronic supplementary information (ESI) available: Experimental details, images of the structures, IR and UV-vis spectra, PXRD patterns, TG curves and the calibration curve for formate and faradaic efficiencies of H_2 . CCDC 1563679, 1563680, 1563685 and 1895007–1895010. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c9sc02503k

well as their way of connecting; (3) finally, inorganic Sn atoms without any alkyl group or phenyl group were introduced into the reaction system to supply more bridging Sn atoms to promote the formation of high nuclearity TOCs. With these synthetic strategies, a series of unprecedented high-nuclearity TOCs have been successfully obtained whose atomic structures were characterized by single crystal X-ray diffraction analysis (Table 1). They possess a much higher number of Sn atoms (26, 34) than the known TOCs (\leq 14). Meanwhile, they also present new structural types different from previous TOCs, including the rod-shaped Sn₂₆ and cage-dimer Sn₃₄. Furthermore, the application of Sn₂₆ or Sn₃₄ derived electrode in electrocatalytic CO₂ reduction was investigated for the first time. Nuclear magnetic resonance (NMR) spectroscopy analysis indicated that formate was obtained as the only liquid reduction product. And the corresponding formate faradaic efficiency (FE) was found to be cluster dependent, with the highest value of 41.90% on the Sn₂₆ derived electrode.

Results and discussion

In the initial stage of our research, mixed methanol-isopropanol was used as the solvent for reactions between butyltin hydroxide oxide and 2,6-pyridinedicarboxylic acid/NaOH or phenylphosphonic acid, resulting in the formation of two Sn_6 clusters (TOC-12 and TOC-13). Structural analysis indicated that they were both composed of two O-capped {Sn₃O₄} units (Fig. 1a and b; Table 1). Different from the back-to-back linking mode of two O-capped $\{Sn_3O_4\}$ units via PP ligands in TOC-13, the Ocapped $\{Sn_3O_4\}$ units in TOC-12 are linked by a Na atom in a face-to-face fashion to form a sandwich-like $\{Sn_3NaSn_3\}$ architecture. It is notable that methanol plays a bridging role in the O-capped $\{Sn_3O_4\}$ units of TOC-12 and TOC-13, implying that the strong coordination ability and small steric hindrance of methanol may be in favour of the formation of the O-capped {Sn₃O₄} unit. In order to obtain different building units and change the aggregation/nucleation rate of tin atoms, water which is crucial for the growth of oxo clusters was directly introduced into the reaction system. Consequently, a new Sn_{12} TOC was prepared namely [(n-BuSn)₁₂(OH)₁₈O₄(BPA)₂(H₂- BPA_4]·6H₂O (TOC-14) (H₃BPA = 2,2-bis(hydroxymethyl)

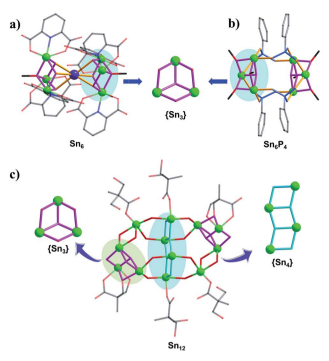


Fig. 1 The ball-and-stick representations of TOC-12 (a), TOC-13 (b) and TOC-14 (c). Atom color code: green Sn; purple Na; red O; black C; dark blue N; light blue P.

propionic acid). As shown in Fig. 1c, **TOC-14** is constructed from two different building blocks, the O-capped $\{Sn_3O_4\}$ and ladder $\{Sn_4O_4\}$ units, indicating that the introduction of water is an effective strategy to prepare different structural types of TOCs with higher nuclearity.

By further optimizing the reaction conditions, especially the amount of used water, **TOC-15** and **TOC-16** with nuclearities of **Sn**₂₆ were successfully obtained in a mixed solvent system of methanol–isopropanol–water (Table 1, Fig. 2). To the best of our knowledge, the numbers of Sn atoms (26) in these TOCs far exceed the value (≤ 14) in reported tin-oxo clusters to date. The core size of these nanoclusters is $\sim 1.6 \times 1.0$ nm. As shown in Fig. 2, different from the usual cage structures of reported TOCs, **TOC-15** presents a rod-shaped cluster core that can be

Table 1 Summary of the composition and reaction conditions of the obtained TOCs ^a				
Compound Formula		Sn source	Solvent	Nuclearity
TOC-12	NaH[(<i>n</i> -BuSn) ₃ (PDC) ₃ (OCH ₃)(OH) ₃] ₂ ·6CH ₃ OH	Butyltin hydroxide oxide	Methanol–isopropanol	Sn ₆
TOC-13	$(n-\mathrm{BuSn})_6\mathrm{O}_2(\mathrm{PP})_4(\mathrm{OCH}_3)_6$	Butyltin hydroxide oxide	Methanol-isopropanol	Sn ₆
TOC-14	$[(n-BuSn)_{12}(OH)_{18}O_4(BPA)_2(H_2BPA)_4] \cdot 6H_2O$	Butyltin hydroxide oxide	Methanol-water	Sn ₁₂
TOC-15	$[(n-BuSn)_{22}Sn_4(OH)_{26}O_{22}(TZAC)_6] \cdot 4Cl \cdot 2(TZAC) \cdot 2(CH_3OH) \cdot 8H_2O$	Butyltin hydroxide oxide	Methanol–isopropanol– water	Sn ₂₆
TOC-16	$[(n-BuSn)_{22}Sn_4(OH)_{26}O_{22}(IANO)_6] \cdot 6(IANO) \cdot 6(CH_3OH) \cdot 4H_2O$	Butyltin hydroxide oxide	Methanol-isopropanol- water	Sn ₂₆
TOC-17	$[(n-BuSn)_{22}Sn_4(OH)_{26}O_{22}(IANO)_6]][(n-BuSn)_2(OH)_2(IANO) Cl_4]_2 \cdot 4Cl \cdot 6CH_3OH \cdot 10H_2O$	Butyltin hydroxide oxide and SnCl4	Methanol–isopropanol– water	Sn ₂₆
TOC-18	$[(n-BuSn)_{34}Na_2(OH)_{14}O_{40}(PA)_8] \cdot 2(PA) \cdot 8H_2O$	Butyltin hydroxide oxide	Acetonitrile	Sn ₃₄

^{*a*} *n*-Bu = butyl group; PDC = 2,6-pyridinedicarboxylate; PP = phenylphosphonate; $H_3BPA = 2,2$ -bis(hydroxymethyl)propionic acid; HTZAC = 1*H*-tetrazole-1-acetic acid; HIANO = isonicotinic acid-*N*-oxide; HPA = 2-picolinic acid.

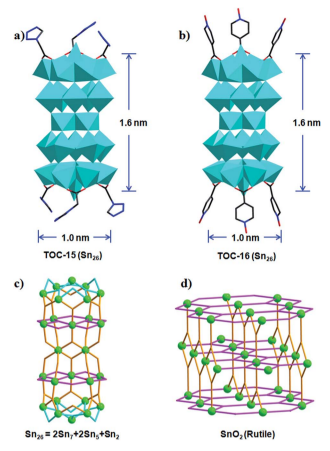


Fig. 2 Polyhedral representations of the Sn₂₆ cluster in TOC-15 (a) and TOC-16 (b). The ball-and-stick representations of Sn₂₆ (c) and SnO₂ (rutile) (d). Polyhedral color code: blue SnO₅C/SnO₃C/SnO₆. Atom color code: green Sn.

considered to be made up of five tin-oxo layers, including two planar $\{Sn_5\}$ units, two capped $\{Sn_7\}$ moieties and one $\{Sn_2\}$ dimer. The $\{Sn_7\}$ and $\{Sn_5\}$ units are linked together by five μ_3 -O atoms to form a $\{Sn_{12}\}$ moiety. And two symmetry-related $\{Sn_{12}\}$ moieties are further held together through the central $\{Sn_2\}$ dimer to give the Sn_{26} core of TOC-15, which is further protected by six TZAC ligands. Such a layered structure type is found for the first time in the area of TOCs, and is reminiscent of the tinoxo layers in SnO₂ (Fig. 2c and d). By changing the functional ligands, other Sn_{26} clusters of TOC-16 were obtained with the six labile coordination sites decorated with IANO ligands (Fig. S6†).

Although **TOC-15** and **TOC-16** present interesting highnuclearity layered structures, their yields are unfortunately quite low (2–4%), which greatly limits further applications. To obtain some mechanistic information of such synthetic shortages, we analyzed their structures in more detail. It is interesting to find that **TOC-15** and **TOC-16** contain pure inorganic SnO₆ nodes without butyl groups. Such completely O-coordinated Sn atoms should be derived from the Sn–C bond cleavage of the applied butyltin hydroxide oxide. Considering the difficulty of Sn–C bond cleavage under the applied low-temperature conditions, it will benefit the assembly of such Sn–O cores if isolated Sn ions could be incorporated into the reactions. For this aim, the inorganic SnCl_4 precursor was further introduced into the synthetic reaction of **TOC-16**. As expected, **TOC-17** was successfully isolated in a much higher yield (~60%). As presented in Fig. S8,† **TOC-17** is composed of the same Sn_{26} moiety in **TOC-16** and two additional {Sn}₂} dimers which are made up of two Sn atoms bridged by two oxygen atoms and one IANO ligand. The successful preparation of **TOC-17** demonstrates that the strategy of introducing additional inorganic Sn atoms is indeed helpful for the formation of high-nuclearity TOCs.

Based on the above results, we can clearly see that the assembly of Sn-O clusters is greatly influenced by the applied solvent conditions. Pure alcohol environments gave rise to small clusters of Sn_6 , and the introduction of water could significantly increase the nuclearities to Sn₂₆ (Table 1).³⁸ However, as a whole, the above used solvents are all protic ones. If an aprotic solvent, e.g. CH₃CN, could be applied, the different solvent environment may change the configuration of basic Sn-O building units, as well as their way of connecting. Following this consideration, the reaction of butyltin hydroxide oxide with 2-picolinic acid and NaOH was carried out in pure CH₃CN. As a consequence, TOC-18 with a nuclearity of Sn34 and a core size of \sim 2.6 \times 1.1 nm was successfully synthesized, which is the largest Sn-O cluster reported to date. As exhibited in Fig. 3a, **TOC-18** is made up of two unprecedented $\{Sn_{12}\}$ and $\{Sn_{22}\}$ cages linked together by a Na atom and PA ligands. Compared with the typical football cage $\{(RSn)_{12}O_{14}(OH)_6\}^{2-}$, the $\{Sn_{12}\}$ in TOC-18 displays an asymmetric cage-like structure (Fig. 3b). The $\{Sn_{22}\}\$ cage captures a central Na heteroatom *via* six oxygen atoms (Fig. 3c). From an architectural point of view, the $\{Sn_{22}\}$ cage can be considered to consist of four subunits with different

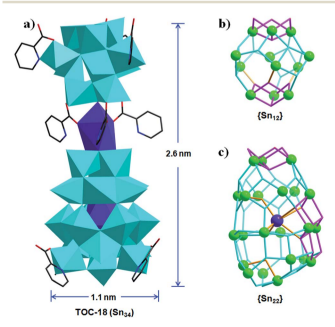


Fig. 3 (a) The polyhedral representation of TOC-18. (b) and (c) The ball-and-stick representations of $\{Sn_{12}\}$ and $\{Sn_{22}\}$ moieties in TOC-18. Polyhedral color code: blue $SnO_5C/SnO_4C/SnO_4NC$; purple NaO_7/NaO_6 . Atom color code: green Sn; purple Na; red O; black C; dark blue N.

numbers of Sn atoms, including a top Sn_3 moiety, two middle Sn_6 circles and a bottom Sn_7 base.

Recently, SnO₂ has shown interesting energy conversion applications through electrocatalytically reducing CO2 to formate.² Considering the similar layered characteristics of the Sn₂₆ cluster core and rutile SnO₂, the obtained Sn₂₆ may also have potential application in the electrocatalytic CO₂ reduction reaction (CO₂RR). Therefore, carbon paper with TOC-17 modification was used as the working electrode to study its catalytic activity towards the CO₂RR. A linear sweep voltammetry (LSV) test was conducted in Ar or CO₂ saturated 0.5 M KHCO₃ solution, respectively. As shown in Fig. 4a, the LSV curve of the TOC-17 derived electrode in the CO₂ saturated electrolyte exhibits the onset potential at approximately -0.69 V. Beyond this onset potential, the current density continuously increases and reaches 6.73 mA cm⁻² at -1.159 V, which is obviously higher than that in the Ar saturated electrolyte. This indicates that the TOC-17 derived electrode may possess high catalytic activity towards the CO₂RR. In order to identify and quantify the reduction products, gas chromatography (GC) and nuclear magnetic resonance (NMR) spectroscopy were applied to analyze the gas and liquid phase products during the CO₂RR process. The obtained results indicate that formate was the only liquid CO₂RR product, with the highest faradaic efficiency (FE) of 41.90% at -1.196 V (Fig. 4). Meanwhile, only a small amount of CO was detected in the gas phase products. Powder X-ray diffraction analysis further confirmed that the structures of TOC-17 remained rather intact after electrolysis (Fig. S46[†]). For comparison, the CO2RR application of the cage-dimer structure TOC-18 was also studied. Although presenting higher nuclearity and also producing formate as the main product, the activity of the TOC-18 derived electrode was significantly lower than that modified with TOC-17. Therefore, these results indicate that the

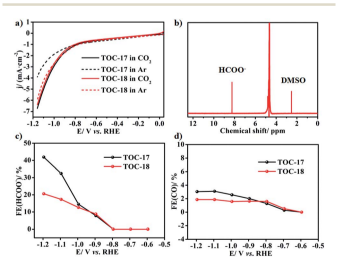


Fig. 4 (a) Linear sweep voltammetry (LSV) of TOC-17 and TOC-18 in Ar or CO₂ saturated 0.5 M KHCO₃ solution. (b) ¹H NMR spectrum of the KHCO₃ catholyte after 3600 s of CO₂ reduction on the TOC-17 derived electrode, E(RHE) = -1.096 V. (c) and (d) The faradaic efficiencies (FE) of formate and CO for TOC-17 and TOC-18 derived electrodes at various potentials.

layered Sn–O structures might be beneficial for electrocatalytic CO₂RR applications.

Conclusions

In summary, we have successfully synthesized a series of unprecedented high-nuclearity tin-oxo clusters by solvent dependent synthetic strategies. These obtained high-nuclearity TOCs, with core sizes ranging from 1.6 to 2.6 nm, possess a much higher number of Sn atoms (26, 34) than previously known ones (≤ 14). Moreover, new structural types of layered nanorods and cage-dimers were also prepared. The applied solvent environments and Sn sources have proven to play crucial roles in the assembly of these high-nuclearity tin-oxo clusters. The introduction of water into alcohol greatly increased the cluster nuclearity; the incorporation of inorganic Sn ions significantly increased the yields of layered structures, while the application of aprotic CH₃CN produced the largest Sn_{34} to date. Moreover, electrocatalytic CO_2 reduction studies confirmed that the electrodes derived from the layered Sn₂₆ cluster presented better performance than those derived from the Sn₃₄ cage-dimer. Therefore, these results afford effective synthetic strategies for the assembly of high-nuclearity TOCs and also extended their potential applications.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work is supported by the National Key Research and Development Program of China (2018YFA0208600), NSFC (21673238), the Strategic Priority Research Program of the Chinese Academy of Sciences (XDB20000000) and the Natural Science Foundation of Fujian Province (2017J06009).

Notes and references

- 1 J. S. Jang, S. Yu, S. J. Choi, S. J. Kim, W. T. Koo and I. D. Kim, *Small*, 2016, **12**, 5989.
- 2 F. W. Li, L. Chen, G. P. Knowles, D. R. MacFarlane and J. Zhang, *Angew. Chem., Int. Ed.*, 2017, **56**, 505.
- 3 L. Fan, Z. Xia, M. J. Xu, Y. Y. Lu and Z. J. Li, *Adv. Funct. Mater.*, 2018, **28**, 1706289.
- 4 W. J. Dong, J. J. Xu, C. Wang, Y. Lu, X. Y. Liu, X. Wang, X. T. Yuan, Z. Wang, T. Q. Lin, M. L. Sui, I. W. Chen and F. Q. Huang, *Adv. Mater.*, 2017, **29**, 1700136.
- 5 G. D. Park, J. K. Lee and Y. C. Kang, *Adv. Funct. Mater.*, 2017, 27, 1603399.
- 6 J. Wei, F. W. Guo, X. Wang, K. Xu, M. Lei, Y. Q. Liang,
 Y. C. Zhao and D. S. Xu, *Adv. Mater.*, 2018, 30, 1805153.
- 7 X. K. Zhang, Y. C. Rui, Y. Q. Wang, J. L. Xu, H. Z. Wang,
 Q. H. Zhang and P. M. Buschbaum, *J. Power Sources*, 2018,
 402, 460.
- 8 B. R. Koo, D. H. Oh, D. H. Riu and H. J. Ahn, *ACS Appl. Mater. Interfaces*, 2017, **9**, 44584.

- 9 D. D. Zhu, J. L. Liu and S. Z. Qiao, Adv. Mater., 2016, 28, 3423.
- 10 B. H. Zhang, Z. H. Guo, Z. Zuo, W. Pan and J. T. Zhang, *Appl. Catal.*, *B*, 2018, **239**, 441.
- 11 P. Han, Z. J. Wang, M. Kuang, Y. F. Wang, J. N. Liu, L. F. Hu, L. P. Qian and G. F. Zheng, *Adv. Energy Mater.*, 2018, 8, 1801230.
- 12 G. L. Zheng, J. F. Ma, Z. M. Su, L. K. Yan, J. Yang, Y. Y. Li and J. F. Liu, *Angew. Chem., Int. Ed.*, 2004, **43**, 2409.
- 13 X. Xiao, K. Z. Shao, L. S. Yan, Z. M. Mei, D. S. Zhu and L. Xu, *Dalton Trans.*, 2013, **42**, 15387.
- 14 G. Prabusankar, B. Jousseaume, T. Toupance and H. Allouchi, *Angew. Chem., Int. Ed.*, 2006, **45**, 1255.
- 15 G. Prabusankar, B. Jousseaume, T. Toupance and H. Allouchi, *Dalton Trans.*, 2007, 3121.
- 16 V. Chandrasekhar, V. Baskar and J. J. Vittal, *J. Am. Chem. Soc.*, 2003, **125**, 2392.
- 17 B. Zobel, J. Costin, B. R. Vincent, E. R. T. Tiekink and D. Dakternieks, *J. Chem. Soc., Dalton Trans.*, 2000, 4021.
- 18 Y. P. Xie, J. F. Ma, J. Yang and M. Z. Su, *Dalton Trans.*, 2010, 39, 1568.
- 19 M. Mehring, M. Schürmann, H. Reuter, D. Dakternieks and K. Jurkschat, *Angew. Chem., Int. Ed.*, 1997, **36**, 1112.
- 20 J. Beckmann, D. Dakternieks, A. Duthie, F. S. Kuan, K. Jurkschat, M. Schürmann and E. R. T. Tiekink, *New J. Chem.*, 2004, 28, 1268.
- 21 T. Zöler and K. Jurkschat, Inorg. Chem., 2013, 52, 1872.
- 22 B. Glowacki, M. Lutter, D. Schollmeyer, W. Hiller and K. Jurkschat, *Inorg. Chem.*, 2016, 55, 10218.
- 23 S. Kundu and V. Chandrasekhar, *Cryst. Growth Des.*, 2015, 15, 5437.
- 24 K. C. K. Swamy, R. O. Day and R. R. Holmes, *J. Am. Chem. Soc.*, 1987, **109**, 5546.

- 25 V. Chandrasekhar, V. Baskar, K. Gopal and J. J. Vittal, *Organometallics*, 2005, 24, 4926.
- 26 S. Y. Song, J. F. Ma, J. Yang, L. L. Gao and Z. M. Su, Organometallics, 2007, 26, 2125.
- 27 M. C. Sharps, D. A. Marsh, L. N. Zakharov, J. E. Hutchison and D. W. Johnson, *Cryst. Res. Technol.*, 2017, 52, 1700081.
- 28 R. R. Holmes, K. C. K. Swamy, C. G. Schmid and R. O. Day, *J. Am. Chem. Soc.*, 1988, **110**, 7060.
- 29 K. C. K. Swamy, R. O. Day and R. R. Holmes, *Inorg. Chem.*, 1988, 27, 958.
- 30 R. R. Holmes, C. G. Schmid, V. Chandrasekhar, R. O. Day and J. M. Holmes, *J. Am. Chem. Soc.*, 1987, **109**, 1408.
- 31 S. Saha, D. H. Park, D. C. Hutchison, M. R. Olsen,
 L. N. Zakharov, D. Marsh, S. Goberna-Ferrón,
 R. T. Frederick, J. T. Diulus, N. Kenane, G. S. Herman,
 D. W. Johnson, D. A. Keszler and M. Nyman, *Angew. Chem.*, *Int. Ed.*, 2017, 56, 10140.
- 32 D. C. Hutchison, R. D. Stern, M. R. Olsen, L. N. Zakharov, K. A. Persson and M. Nyman, *Dalton Trans.*, 2018, 47, 9804.
- 33 H. Reuter, Angew. Chem., Int. Ed. Engl., 1991, 30, 1482.
- 34 Q. F. Wang, C. L. Ma, G. F. He and Z. Li, *Polyhedron*, 2013, **49**, 177.
- 35 J. X. Liu, M. Y. Gao, W. H. Fang, L. Zhang and J. Zhang, Angew. Chem., Int. Ed., 2016, 55, 5160.
- 36 S. Chen, W. H. Fang, L. Zhang and J. Zhang, Angew. Chem., Int. Ed., 2018, 57, 11252.
- 37 X. Fan, J. H. Wang, K. F. Wu, L. Zhang and J. Zhang, *Angew. Chem., Int. Ed.*, 2019, **58**, 1320.
- 38 X. F. Xu, W. Wang, D. L. Liu, D. D. Hu, T. Wu, X. H. Bu and P. Y. Feng, J. Am. Chem. Soc., 2018, 140, 888.