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Recent advances in uranium-containing polyoxometalates

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Uranium-containing polyoxometalates (U-POMs), an important class of actinide cluster compounds, are the products of the assembly of uranium or uranium and POMs. These cluster compounds illustrate the structural beauty and diversity of obtained U-POMs that can be achieved by the reaction of uranium or uranium and POMs. The structural diversity is revealed by the classification of U-POMs in this review, which are mainly divided into two parts, according to the main metal elements of the clusters, named U-containing heteropolyoxometalates (U-hetero-POMs) and U-oxo-clusters. U-hetero-POMs are further classified by development date from the end of the 20th century based on the basic building units, including Keggin-type, Dawson-type, Silverton-type, hybrid structures, and cryptand structures. U-oxo-clusters mainly include uranyl peroxide clusters, their derivatives, and U^{IV}-oxo-clusters and mixed-valence U-oxo-clusters. This review provides a comprehensive summary of the synthesis and structures of U-POMs as well as their characterization, properties, and applications. Finally, the development prospects and directions for U-POMs in the future are also elucidated.

Introduction

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Polyoxometalates (P

1.

Polyoxometalates (POMs) are the most representative metal oxide clusters that self-assemble in solution with controllable structures and give us a unique way to design and synthesize metal oxide materials from the micro- to macro-scale. The incomparable chemical compatibility of POMs makes them ideal models to assess the influence of various elements on the structures and properties across the periodic table of

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Ke Li

lates, polyoxometalate-based metal-organic frameworks, and coordination polymers.

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elements. Research on POMs has developed rapidly not only based on the fascinating structures but also the applications in various fields. 1-8 One notes that the research trends of POMs have evolved from structure-driven research to performance-driven research in the past decades. In contrast, actiniderelated POMs are less explored even if the first actinide-containing POMs were found more than 50 years ago. 9 The studies of actinide-containing POMs may be limited by the difficulties in obtaining them and the radioactivity of actinides due to the strategic significance of actinides related to national defense, energy sources, and other fields, and the development of actinide-containing POMs lags far behind other transition metal (TM)-based POMs. But there are also several investigations on actinide-containing POMs, involving Th, U, Np, Pu, and Am. 10-17 However, only U-related POMs are studied relatively more due to the stability of isotopes and the difficulty in acquiring the actinides.

Uranium is one of the most important strategic resources in the nuclear industry and exhibits abundant oxidation states from +2 to +6.18,19 Uranium chemistry already developed into the representative field of actinide chemistry. 20-23 The most stable state of +6 uranium atoms is to form linear uranyl ions (UO22+) combining two oxygen atoms with triple bonds. However, the two opposite oxygen atoms are inert under the most common conditions. One uranyl ion could coordinate with at most six oxygen atoms in a hexagonal bipyramid geometry. 24 While, 5-coordinated uranyl in pentagonal bipyramid geometry is more commonly found. Uranium atoms with a +4 oxidation state exhibit different coordination behaviors, which are more similar to other tetravalent transition metal ions such as Zr4+ and Hf4+.25-28 Tetravalent uranium also shows more abundant coordination modes and the coordination numbers could be 12 or higher.9 Thus, the combination of uranium atoms and traditional POMs has a strong coordi-



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nation basis. Besides, the polymerizable nature of uranyl ions and tetravalent uranium salts in aqueous solutions makes them easy to assemble into U-oxo-clusters, clusters built from a certain number of ${\rm UO_2}^{2^+}$ or ${\rm U}^{4^+}$ ions and bridging oxygen linkers, which is also a significant category of POM chemistry. One can see the trend that the research focus of U-containing POMs (U-POMs) has gradually turned into structure-based performance research not only due to the development tendency of chemistry but also the increasing numbers of new U-POMs prepared in recent years.

To better summarize the structural features, the U-POMs in this review are mainly divided into two parts, U-containing heteropolyoxometalates (U-hetero-POMs) and U-oxo-clusters. U-hetero-POMs are those POMs in which the early transition metal elements (W etc.) are the main part of the clusters and uranium is the heterometal ion. While U-oxo-clusters are those POMs in which uranium atoms are the main building units of clusters. Thus, the U-POMs discussed in this review are divided into three parts: (i) U-hetero-POMs, which are classified based on the building units, including Keggin, Dawson, Silverton, and so on; (ii) U-oxo-clusters reported in recent years, including uranyl peroxide clusters, POM-related uranium phosphates derivatives, U-oxo-clusters containing other transition metals, and UIV-oxo-clusters and mixedvalence U-oxo-clusters; (iii) other types of U-POMs, which are hard to classify. This review restricts our attention to U-hetero-POMs and recent U-oxo-clusters containing four or more essential uranium atoms. We focus on the development of U-POMs with respect to the aspects of syntheses, structures, properties, and applications. Finally, several directions for further research are proposed.

2. Synthesis and structure of U-POMs

2.1 U-hetero-POMs

U-hetero-POMs are built from traditional POM building units with different kinds of lacunary precursors or from simple starting materials (such as Na_2WO_4 , heteroatom salts, etc.). $UO_2(NO_3)_2$ and $UO_2(OAc)_2$ (OAc = acetate) are the most commonly used uranium sources and the employment of UCl_4 or $U(SO_4)_2$ may introduce considerable uncertainty for the synthesis of novel U-hetero-POMs due to the oxidation and richer bonding ability of U^{IV} . Those U-hetero-POMs based on Keggintype lacunary precursors are the most reported due to the natural stability of Keggin-type lacunary precursors. Consequently, the U-hetero-POMs here are divided up based on the basic building blocks, including Keggin-type, Dawson-type, Silverton-type, hybrid structures, and cryptand structures (Table 1).

2.1.1 Keggin-type structures. It is known that Keggin-POMs are the first and the most widespread kind of POMs to be studied. Stable Keggin-type lacunary precursors and Keggin-POM derivatives are also the most abundant. In most cases, polyvacant polyoxotungstates (POWs) are the most stable, and

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Table 1 Summary of U-hetero-POMs with crystal structures

Туре	Compounds	Building Units	Materials of POMs	Reaction Temp./°C	hd	Ref.
	-21-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1		()			
keggın	[Na ₂ (UO ₂) ₂ (FW ₉ O ₃₄) ₂] [(MIII) (IO) (MIX O) 112-	{PW9} (mir,)	{PW9}	r.t.		30
	$[(1N14)_2(0O_2)_2(VV9O_34)_2]$ [V(11O) (DXIO) (112-	(FW9)	(FW9)	~40		
	$[N_3(100_2)_2(100_3)_2]_{14-}$	\{\sqrt{\veepsilon}\}	(5 V 9) {SiW_c}	2 + 1		34
	[:\cdot\cdot\cdot\cdot\cdot\cdot\cdot\cdot	SiW.?	{SiW.2}	rt	0	•
	$\{K(OH_2)\}_A(UO_2)_A(OH)_2(SW_{10}O_{36})_A$	$\{SiW_{10}\}$	$\{\operatorname{SiW}_{10}\}$	r.t.	5.0	
	$[(00_2)_2(H_20)_2(\mathrm{SbW}_90_{33})_2]^{14}$	$\{\mathbf{s}_{\mathbf{M}}^{\mathbf{m}}\mathbf{q}\mathbf{s}\}$	$\{\mathrm{Sb^{III}W_9}\}$	Reflux	6.4	35
	$[(00_2)_2(H_2O)_2(TeW_9O_{33})_2]^{12}$	$\{ Te^{TV}W_9 \}$	$\{\text{Te}^{\text{IV}} \mathbf{W}_{9}\}$ or in situ	Reflux	1	
	Na ₁₂ [Na ₂ As ₂ W ₁₈ U ₂ O ₇₂]:35H ₂ O	$\{As^VW_9\}$	În situ	Reflux/5		36
	$(NH_4)_{13}[NH_4As_2W_{18}U_2O_{72}]\cdot17H_2O$	$\{\mathbf{As_{V}^{V}SA}\}$	In situ	Reflux/5		
	$ m K_{13}[KAs_2W_{18}U_2O_{72}]\cdot 10H_2O$	$\{As^VW_9\}$	In situ	Reflux/5	1	
	$Na_{14}[Na_2(UO_2)_2(A-\alpha-GeW_9O_{34})_2]\cdot 48H_2O$	{GeW ₉ }	$\{GeW_9\}$	80–90	5.9	37
	$Na_{14}[Na_{2}(UO_{2})_{2}(A-\beta-GeW_{9}O_{34})_{2}]\cdot 42H_{2}O$	{GeW ₉ }	$\{GeW_9\}$	r.t.		
	$[Na(UO_2)_2(H_2U)_4(BIW_9U_{33})_2]^{}$	$\{B1^{-1}W_9\}$	$\{BI^{-}W_9\}$	00	6	38
	(\bar{1}18\big \big \big \big \big \big \big \big	{AS W ₉ } {SiW ₂ }	$\{NH_4\}_{13}[NH_4AS_2W_{18}O_2O_{72}]$ ·1/H ₂ O $\{S_1W_2\}$	oo ama so	5.5	96 28
	$[(\mathbf{U}^{1})_{1}, (\mathbf{U}^{1})_{1}, (\mathbf{U}^{1})_{1}, (\mathbf{U}_{1}, \mathbf{O})_{1}](\mathbf{U}^{1}, \mathbf{O})_{2}]$	$\{As^{III}W_{q}\}$	$\{{ m As^{III}W_q}\}$	70	4.7	40
	$[(U^{\text{IV}})_3(U^{\text{VI}}O)_3(\mu_3-OH)_4(\mu_3-O)_4(AsW_9O_{33})_3]^{1.5}$	$\{As^{III}W_{9}\}$	$\{AS^{III}N_9\}$	70	4.7	
	$[({ m U}^{1})_{ m b}({ m \mu_3-OH})_4({ m \mu_3-O})_4({ m AsW}_9{ m O}_{33})_4]^{24}$	$\{AS^{III}W_{9}\}$	$\{As^{\Pi I}W_9\}$	70	4.7	
	$Na_{17}(Na@[(SbW_{9}O_{33})_{2}(UO_{2})_{6}(PO_{3}OH)_{6}]\} \cdot xH_{2}O$	{8Vds}	In situ	95	L	41
	$Na_{11}H(H_2O)_{31}[Na(UO_2)(A-PW_9O_{34})]_2.7H_2O$	$\{PW_9\}$	${\rm bm}_{6}$	85	5.1	31
	$Na_{12}(H_2O)_{29}[Na(UO_2)(A-PW_9O_{34})]_{27}7H_2O$	$\{PW_9\}$	{PW ₉ }	520	0.0 0.1	ć
	$Na_{10.5}H_{3.5}(H_2O)_{36}[Na(UO_2)(\alpha-5)W_9O_{34})]_2$ 2.5 H_2O	{SIW ₉ }	{SIW ₉ }		5.9	32
	$\ln (1/10^{-36} \ln (1/10^{-36}) \ln (1/10^{-34}) \ln (1/$	{SIW ₉ }	{SIW ₉ }	35	6.4	,
	$NN3_{11}\Pi_2(\Pi_2 \cup)_{30}[N3(\cup \cup_2)(S)W_9 \cup_{34})[2.8\Pi_2 \cup$	{6MIC}	{51W10}	65	5.9	55
Dawson	$Na_{32}[(UO_2)_{12}[\mu_2\cdot O]_4[\mu_2\cdot H_2O]_{12}(P_2\cdot W_1\cdot O_{56})_4]\cdot 77H_2O$	$\{P_2W_{15}\}$	$\{P_2W_{15}\}$	r.t.	I	42
	$[\mathrm{U}(lpha_1 - \mathrm{P}_2 \mathrm{W}_{17} \mathrm{O}_{61})_2]^{16-}$	$\{\mathbf{P_2W_{17}}\}$	$lpha_1 ext{-}\{ ext{P}_2 ext{W}_{17}\}$	09	4.7	43-45
	$\left[\tilde{\mathrm{U}} (\alpha_1 ext{-} \mathrm{P}_2 \mathrm{W}_{17} \mathrm{O}_{64}) (\alpha_2 ext{-} \mathrm{P}_2 \mathrm{W}_{17} \mathrm{O}_{64}) \right]^{16}$	$\{\mathbf{P}_2\mathbf{W}_{17}\}$	$lpha_1 - \left\{ \mathbf{P_2W_{17}} ight\}$	r.t.	I	
	$\left[\mathrm{U}(lpha_2 ext{-}\mathrm{P}_2 \mathrm{W}_{17} \mathrm{O}_{61})_2 ight]^{16}$	$\{\mathbf{P_2W_{17}}\}$	$lpha_2 - \{\mathbf{P_2W_{17}}\}$	r.t. or 80	4.75	
	$\{ K[UO_2(H_2O)]_2(SiW_9O_{33})_2 \}^{11}$	$\{(SiW_9)_2\}$	$\{^{6} ext{MiS}\}$	80	3.7	46
	$[\text{Li}(\text{H}_2\text{O})\text{K}_4(\text{H}_2\text{O})_3\{(\text{UO}_2)_4(\text{H}_2\text{O})_2\}_2[\text{PO}_3\text{OH})_2\text{P}_6\text{W}_3\text{cO}_{136}]^{23}$	$\{\mathbf{P_6W_{36}}\}$	$\{\mathrm{P_4W_{24}}\}$	50	4.0	47
	$[(\mathrm{UO}_2)_{7.2}(\mathrm{HCOO})_{7.8}(\mathrm{P_8W_{48}O_{184}})\mathrm{Cl_8}]^{41.4-}$	${ m P_8W_{48}} $	$\{P_8W_{48}\}$	160/r.t.	3.7	48
	$[(UO_2)_7(P_8W_{48}U_{184})]^{}$	$\{P_8W_{48}\}$	$\{P_8W_{48}\}$	Keriux	6.2 or 6.6/1 5	49
	$[({ m UO_2})_{ m g}({ m P_2W_{AS}O_{18A}})]^{24-}$	$\{P_8W_{48}\}$	$\{\mathrm{P_8W_{48}}\}$	Reflux	6.2 or	
					6.6/1.5	
	$[\{({ m Uo_2})_4({ m O_2})_4\}_2({ m P_8W_4}_8{ m O_{184}})]^{40}$	$\{\mathbf{P_8W_{48}}\}$	$\{\mathbf{P_8W_{48}}\}$	50	6.0/5.7	50
	$Na_3[H_{19}(UO_2)_2(\mu_2-O)(Se_2W_{14}O_{52})_2]\cdot 41H_2O$	$\{\operatorname{Se^{17}_2W_{14}}\}$	In situ	06	5.37	51
	$(\ln \Pi_4)_{10}[\Pi_4(36O)_2(OO_2)_2(\Pi_2O)_2(\Pi_236_2W_{14}O_{52})(36_2W_{14}O_{52})]$	(3c 2w ₁₄)	in stu	90	4.03	
Silverton	$[M_{a}UMo_{12}O_{42}]^{b-}, M = Th^{4+}, Er^{3+}, VO^{2+}, UO_{2}^{2+}, Ni^{2+}, Nd^{3+}, Cu^{2+}, Sc^{3+}, Fe^{3+}; a = 1 \text{ or } 2: b = 2 \text{ or } 4.$	$\{\mathbf{U^{IV}Mo_{12}}\}$	I	I	I	9, 15 and 52–59
Hybrid	$\left[({\rm UO}_2)_3 ({\rm H}_2 {\rm O})_4 {\rm As}_3 {\rm W}_{26} {\rm O}_{94} \right]^{17-}$	$\{As^{III}W_9\},$	$\left\{ \mathbf{As^{III}W_{9}}\right\}$	95	4.6	34
	[(IJO ₂),(H ₂ O) _E AS ₂ W ₂₀ O _{20,1} 19-	$\{(AS W_8)_2W\}$ $\{AS^{III}W_6\}$.	{AS ^{III} ,W _a ,}	06	6.6 + 0.2	09
	10, -2, -1, -2, -1, -2, -1, -1, -1, -1, -1, -1, -1, -1, -1, -1	$\{(As^{III}W_9)_2(W_2)\}$	(04 4)))
	$[(H_3 \text{SbW}_{17} O_{59}) \text{U}^{\text{N}} (\text{HW}_5 O_{18})]^{11}$	$\{\mathbf{W}_5\}, \{\mathbf{Sb^{III}}_{\mathbf{W}_17}\}$	In situ	09	5.5/4-4.5	61
	[(UU ₂)3(SeU ₃)3Na ₅ (H ₂ U)6(SeW ₆ U ₂₁)(SeW ₉ U ₃₃)3] ⁻	{se w ₆ }, {se w ₉ }	ın situ	90	2.5	79

Туре	Compounds	Building Units	Materials of POMs	Reaction Temp./°C	Hd	Ref.
Cryptand	Cryptand $(NH_4)_{11}[U(H_2O)P_5W_{30}O_{110}]\cdot 12H_2O$	$\left\{ P_5W_{30}\right\}$	$\left\{ P_5W_{30}\right\}$	60-70/	1	63
	$\left[({\rm UO}_2)_3 ({\rm H_2O})_6 {\rm As_3W_{30}O_{105}} \right]^{15-}$	$\{({\rm A}{\rm S}^{\rm III}{\rm W}_9)_3({\rm W}_3)\}$	$\left\{ \mathrm{AS}^{\mathrm{III}}_{}4}\mathrm{W}_{40} ight\}$	140-180 90	$5.4/3.8 \pm$	09
	$(NH_4)_{17}[(UO_2)_3(H_2O)_6AS_3W_{29}V^{IV}O_{105}]$ -44 H_2O	$\{(AS^{III}W_9)_2(V^{IV}W_2)\}$	$[(UO_2)_3(H_2O)_5AS_3W_{29}O_{104}]^{19-}$ and	r.t.	7.	
	$[Na(UO_s)_3(OH)(H_sO)_6AS_4W_{a0}O_{140}(WO)]^{18-}$	$\left\{ \mathbf{A}\mathbf{S}^{\mathrm{III}}{}_{4}\mathbf{W}_{40} ight\}$	$egin{array}{l} VOSO_4 \ AS^{-1} W40 \end{array}$	r.t.	I	34
	$[\{U_2(\mu_2-OH)_3(\mu_2-O)\}AS_4W_{40}O_{140}]^{\frac{25}{25}}$	$\{ \mathbf{A} \mathbf{S}^{\mathrm{III}} \mathbf{W}_{4} \mathbf{W}_{40} \}$	$\{As^{III}_{}W_{40}^{}\}$	70	4.7	64
	$[\{U_2(\mu_2-0)(HCOO)_3\}AS_4W_{40}O_{140}]^{25-}$	$\left\{ \mathbf{A}\mathbf{s}^{\mathrm{III}}\mathbf{w}_{4}\mathbf{W}_{40} ight\}$	$\{As_{-4}^{III}W_{40}^{-1}\}$	70	4.7	
	$[U_4(\mu_2-0)_2(HCOO)_4]AS_4W_{40}O_{140}]^{20-1}$	$\left\{ \mathbf{A}\mathbf{s}^{\mathrm{III}}\mathbf{w}_{4}\mathbf{W}_{40} ight\}$	$\{As_{-4}^{III}W_{40}^{-1}\}$	70	4.7	
	$[(00_2)_{7.2}(HCOO)_{7.8}(P_8W_{48}O_{184})CI_8]^{41.4-}$	$\{\mathbf{P_8W_{48}}\}$	$\{P_8W_{48}\}$	160	3.7	48
	$[(00_2)_7(P_8W_{48}O_{184})]^{26}$	$\{\mathbf{P_8W_{48}}\}$	$\{P_8W_{48}\}$	Reflux	6.2 or	49
					6.6/1.5	
	$[({ m UO}_2)_8 ({ m P}_8 { m W}_{48} { m O}_{184})]^{24-}$	$\{\mathrm{P_8W_{48}}\}$	$\{\mathrm{P_8W_{48}}\}$	Reflux	6.2 or	
					6.6/1.5	
	$[\{({ m UO}_2)_4({ m O}_2)_4\}_2({ m P}_8{ m W}_{48}{ m O}_{184}]]^{40-}$	$\{\mathrm{P_8W_{48}}\}$	$\{\mathrm{P_8W_{48}}\}$	50	6.0/5.7	50

Temperatures not mentioned in the original text are room temperature (r.t.) by default except for the unverifiable Silverton structures.

the basic mono-, bi-, and trivacant Keggin-type precursors $\{XW_{11}\}, \{XW_{10}\}, \{XW_{9}\}$ tolerate various heteroatoms including B, Si, Ge, P, As, Sb, Bi, S, Se, Te, etc. Thus, Keggin-type U-hetero-POWs are also the most studied. The first Keggin-type U-hetero-POW for which a single-crystal structure was recorded might be monovacant {GeW₁₁}-based Cs₁₂[U (GeW₁₁O₃₉)₂]·13-14H₂O reported by Tourné and co-workers in 1980 (Fig. 1a).²⁹ Most of the Keggin-type U-hetero-POMs based on trivacant precursors show similar sandwich structures in which two trivacant {XW₉} sandwich a tri- or tetranuclear cluster containing 2 to 4 UO₂²⁺ or U⁴⁺. The empty position in the sandwich was occupied by H₂O or counter cations such as Na⁺, K⁺, and NH₄⁺ (Fig. 1b and c). Pope synthesized the first series of typical sandwich-type U-hetero-POWs with the general formula of $[M_2(UO_2)_2(PW_9O_{34})_2]^{12-}$ (M = Na, K, NH₄) in 1999 (Fig. 1b and c).³⁰ $[Na_2(UO_2)_2(PW_9O_{34})_2]^{12-}$ represented the first typical sandwich structure, in which two {PW₉} sandwiched a {Na₂(UO₂)₂} cluster to form the symmetrical U-hetero-POWs (Fig. 1b). $[K_2(UO_2)_2(PW_9O_{34})_2]^{12-}$ and [(NH₄)₂(UO₂)₂(PW₉O₃₄)₂]¹²⁻ showed another typical asymmetrical sandwich structure in which two uranyl ions and K⁺ or NH₄⁺ were sandwiched with a triangular configuration (Fig. 1c). Recently, our group also synthesized five $\{Na_2(UO_2)_2(XW_9O_{34})_2\}$ -based (X = P/Si) U-POMs with different 2D/3D structures, which revealed the significant influence of counter cations on the spatial structures of U-POMs even if the structures of $\{Na_2(UO_2)_2(XW_9O_{34})_2\}$ were similar. ^{31–33} Our work demonstrates that different reaction conditions provide the assembly conditions for Na-H2O clusters with different coordination environments and different dimensions, which result in different spatial structures.

Rusu studied the properties of four Keggin-type compounds, including isotope exchange, spectroscopy, thermal analyses, magnetic susceptibility, and paper electrophoresis in 1999 and 2000. ^{65–67} However, only the formulas, *i.e.*, $Na_{12}[(UO)_3(H_2O)_6(PW_9O_{34})_2]\cdot 21H_2O$, $K_{12}[U^{IV}(BW_{11}O_{39}H)_2]\cdot 23H_2O$, $K_{4}[U^{IV}GeW_{11}O_{39}]$, and $K_{12}[U^{IV}(GeW_{11}O_{39})_2]$, were provided without crystal data. Then, David prepared two sandwich-type structures with formulas of $Na_{14}[(U^{IV}O)_3(GeW_9O_{34})]\cdot 23H_2O$ and $Na_{6}[U_3^{IV}(SbW_9O_{33})_2]\cdot 22H_2O$, but crystal data were also not provided. ^{68,69} The utilization of $U(SO_4)_2$, prepared from $UO_2(OAc)_2$ hydrate, should be a key factor for the formation of these U^{IV} -hetero-POWs.

Compared with trivacant precursors, there are much fewer Keggin-type bivacant precursors. As a result, there were only two cases based on $\{\gamma\text{-SiW}_{10}\}$, which were also reported by Pope in 2002.³⁴ In this work, Pope reported five Keggin-derivative U-hetero-POWs, among which $[\text{Na}_2(\text{UO}_2)_2(\text{SiW}_9\text{O}_{34})_2]^{14-}$ showed a typical sandwich structure based on trivacant $\{\text{SiW}_9\}$ (Fig. 1b). Another two tetramers $[\{M(\text{OH}_2)\}_4(\text{UO}_2)_4(\text{OH})_2$ (SiW₁₀O₃₆)₄]²²⁻, M = Na/K, are the only two bivacant precursor-based structures (Fig. 1k and 1). The arrangement of four uranyl ions in the two tetrameric structures was similar to the $\{\text{As}_4\text{W}_{40}\}$ -based structures reported by Duval in 2018 (section 2.1.5).⁶⁴ However, it could be found that Na⁺ and K⁺ in the two structures displayed different arrangements. Na⁺ ions are adja-

Fable 1 (Contd.)

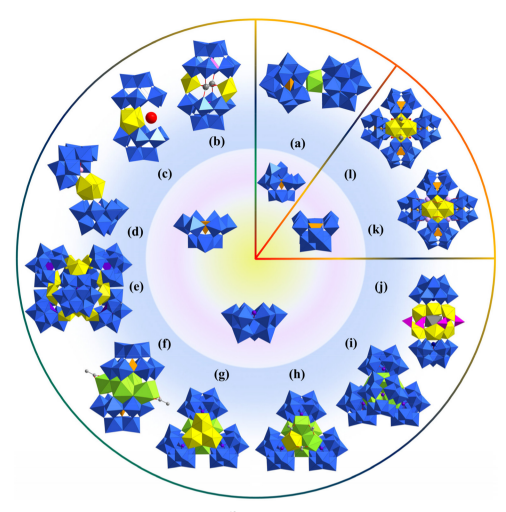


Fig. 1 View of Keggin-type U-hetero-POWs. (a) $[U(GeW_{11}O_{39})_2]^{12-}$; typical structures of (b) $\{Na_2(UO_2)_2(XW_9O_{34})_2\}$ (X = Si, P, Ge, As^V), (c) $\{M_2(UO_2)_2(XW_9O_{34})_2\}$ $\{M = H_2O, NH_4^+, K^+; X = Si, P, As^V\}$, $\{M = W_2O_1, W_3O_2\}$ $\{M = W_3O_2, W_3O_3\}$ $\{M = W_3O_3, W_$ $(NH_4)_{18}[(NH_4)_{12}(UO_2(H_2O))_6(UO_2(\mu_2-H_2O))_6(\alpha-AsW_9O_{34})_6]\cdot 74H_2O; \quad (f) \quad [U_4(\mu_3-O)_2(SiW_9O_{34})_2(CH_3COO)_2]^{10-}; \quad (g) \quad [(U^{IV})_{5.5}(U^{VI}O)_{0.5}(\mu_3-OH)_4(\mu_3-O)_4O(H_2O)_{0.5}(\mu_3-OH)_4(\mu_3 [(U^{IV})_3(U^{VI}O)_3(\mu_3-OH)_4(\mu_3-O)_4(AsW_9O_{33})_3]^{15-}; \qquad (i) \qquad [(U^{IV})_6(\mu_3-OH)_4(\mu_3-O)_4(AsW_9O_{33})_4]^{24-};$ $(OAc)_2(AsW_9O_{33})_3]^{19-};$ (h) $[(SbW_9O_{33})_2(UO_2)_6(PO_3OH)_6])^{17-}; (k) [\{K(OH_2)\}_4(UO_2)_4(OH)_2(SiW_{10}O_{36})_4]^{22-}; (l) [\{Na(OH_2)\}_4(UO_2)_4(OH)_2(SiW_{10}O_{36})_4]^{22-}. Color codes: W, blue; U^{VI}, blue;$ yellow; U^{IV}, light green; Na, dark grey; K, lavender; P, magenta; Si and Ge, orange; As, purple; Te^{IV}, pink; Sb^{III}, dark blue; O, red; C, light gray.

cent to each {SiW₁₀} but K⁺ ions are located at the positions in middle of two {SiW₁₀}.

Afterward, Pope reported two similar sandwich structures based on $\{Sb^{III}W_9\}$ and $\{Te^{IV}W_9\}$ (Fig. 1d).³⁵ In both structures of $[(UO_2)_2(H_2O)_2(SbW_9O_{33})_2]^{14-} \quad \text{and} \quad [(UO_2)_2(H_2O)_2(TeW_9O_{33})_2]^{12-},$ the trivacant {XW₉} only offered four of the six unsaturated terminal oxygen atoms to coordinate with two uranyl ions. Thus, the obtained sandwich structure showed a different "open" configuration (Fig. 1d). Besides, [(UO₂)₂(H₂O)₂(TeW₉O₃₃)₂]¹²⁻ could also be in situ synthesized from simple raw materials including TeO₂, UO2(NO3)2·6H2O, Na2WO4·2H2O and NH4Cl. In 2005, David systematically studied and compared the spectra of several sandwich U-hetero-POWs with the general formulas of $[(UO)_3(H_2O)_6(XW_9O_{34})_2]^{12/14-}$ (X = P/Si/Ge) and $[U_3(XW_9O_{33})_2]^{6-}$ (X = As/Sb/Bi).⁷⁰ This work revealed the shifting law of bonds in these sandwich U-hetero-POWs and trivacant precursors $\{XW_9\}$.

Subsequently, Khoshnavazi also in situ synthesized three sandwich uranium-containing polytungstoarsenates using HAc, $Na_2WO_4 \cdot 2H_2O$, $Na_2HAsO_4 \cdot 7H_2O$, and $UO_2(OAc)_2 \cdot 2H_2O$. The addition of NaCl, NH₄Cl, and KCl yielded $Na_{12}[Na_2As_2W_{18}U_2O_{72}]\cdot 35H_2O$, $(NH_4)_{13}[NH_4As_2W_{18}U_2O_{72}]\cdot 17H_2O$, and K₁₃[KAs₂W₁₈U₂O₇₂]·10H₂O, respectively. These U-hetero-POWs exhibited similar structures with the general formula $[M_2(UO_2)_2(PW_9O_{34})_2]^{12-}$ (M = Na, K, NH4) reported by Pope as mentioned above (Fig. 1b and c). A Na salt based on {BiW₉}, which was similar to K⁺ and NH₄⁺ of [M₂(UO₂)₂(PW₉O₃₄)₂]¹²⁻, was also successfully obtained by Alizadeh and the formula was $[Na(UO_2)_2(H_2O)_4(BiW_9O_{33})_2]^{13}$. In 2011, Mohadeszadeh Keggin-{As^VW₉}-based synthesized a rare $(NH_4)_{18}[(NH_4)_{12}(UO_2(H_2O))_6(UO_2(\mu_2-H_2O))_6(\alpha-AsW_9O_{34})_6]\cdot 74H_2O,$ by the reaction of $(NH_4)_{13}[NH_4As_2W_{18}U_2O_{72}]\cdot 17H_2O$ and (NH_4) $Ce(SO_4)_4 \cdot 2H_2O$ in a pH = 3.5 solution (Fig. 1 and 2).³⁹ This hexameric polyanion is composed of six {AsVW₉}, six fourcoordinated uranyl ions, and two trinuclear clusters $\{(UO_2)_3\}$. Each {As W₉} is coordinated with three single uranyl ions and one $\{(UO_2)_3\}$ to form a cage configuration.

Four years later, Duval reported four {SiW₉}-based POMs, among which a tetranuclear $\{U_4^{IV}(\mu_3-O)_2\}$ -based sandwich structure was constructed from two {SiW₉} moieties (Fig. 1f).²⁸ It should be noted that $\left[U_4(\mu_3-O)_2(SiW_9O_{34})_2(CH_3COO)_2\right]^{10-}$ was synthesized using UCl₄ in a pH = 4.7 acetate buffer, instead of uranyl salts. However, one must admit that the combination of uranyl and acetate buffer is very difficult to control due to the fast formation and precipitation of Na[UVIO2(OAc)3]. The success of these U-hetero-POWs should be attributed to the usage of UCl₄ for assembly and quick separation before the formation of Na[UVIO2(OAc)3]. After this work, Duval also synthesized three multi-sandwich structures using a similar method (Fig. 1g-i).40 Similar hexanuclear uranium clusters with different ratios of substitutional-disordered UIV/UVI were stabilized by the coordination of three or four {As^{III}W₉}, $\big[\big(U^{IV} \big)_{5.5} \big(U^{VI} O \big)_{0.5} \big(\mu_3 \text{-OH} \big)_4 \big(\mu_3 \text{-O} \big)_4 O \big(OAc \big)_2 \big(AsW_9 O_{33} \big)_3 \big]^{19-},$ $[(U^{IV})_3(U^{VI}O)_3(\mu_3\text{-OH})_4(\mu_3\text{-O})_4(AsW_9O_{33})_3]^{15-}$, and $(\mu_3\text{-OH})_4(\mu_3\text{-O})_4(\text{AsW}_9\text{O}_{33})_4$]²⁴⁻. The former two compounds were obtained from the same solution at different time. The degree of oxidation with time from UIV to UVI could be revealed from the formulas. By diluting the concentration of acetate buffer and raw materials, crystals of Na₂₄[(U^{IV})₆(µ₃-OH)₄(µ₃-O₄(AsW₉O₃₃)₄]·42H₂O were obtained without Na[U^{VI}O₂(OAc)₃]. In 2021, Kong and co-workers synthesized a novel sandwich structure $Na_{17}\{Na@[(SbW_9O_{33})_2(UO_2)_6(PO_3OH)_6]\}\cdot xH_2O \ (x \approx 1)$ 46), which was built from Keggin-{SbW₉} and the central {Na@ (UO₂)₆(PO₃OH)₆} cluster (Fig. 1j). ⁴¹ Two {SbW₉} offered all six unsaturated terminal oxygen atoms for coordination with six uranyl ions. $Na_{17}\{Na@[(SbW_9O_{33})_2(UO_2)_6(PO_3OH)_6]\}\cdot xH_2O$ also has the biggest uranium clusters among those Keggin-type sandwich U-hetero-POWs. Besides, this compound is also synthesized by an in situ method from SbCl₃, Na₂WO₄, NaOAc, H_3PO_4 , and $UO_2(NO_3)_2 \cdot 6H_2O$.

2.1.2 Dawson-type structures. Dawson-type POMs are the second main category of POMs among thousands of derivative structures. However, the number of reported Dawson-type U-POMs to date is much lower than that of Keggin-type. Fewer Dawson-type precursors may be responsible for the lower number of Dawson-type U-hetero-POMs given the lacunary precursor synthesis strategy for the great majority of U-hetero-POMs. The first Dawson-type U-hetero-POWs were reported by Rusu and co-workers without crystal structures in 1999.⁷¹ But they provided the formulas, i.e., Na₁₈[(UO)₃(P₂W₁₅O₅₆)₂] and $Na_{18}[(UO)_3(H_2O)_6(P_2W_{15}O_{56})_2]\cdot 27H_2O$. The study of spectroscopy and isotope exchange experiments was carried out. The first Dawson-type U-hetero-POW with a definite structure was reported by Pope in 2003 (Fig. 3).42 The obtained $Na_{32}[(UO_2)_{12}(\mu_3-O)_4(\mu_2-H_2O)_{12}(P_2W_{15}O_{56})_4]$ -77H₂O product was synthesized using the vapor diffusion of CH3CN into an aqueous solution of trivacant Dawson-{P2W15} and UO₂(NO₃)₂·6H₂O. In this polyanion, four groups of trinuclear uranyl clusters $\{(UO_2)_3(\mu_3-O)(\mu_2-H_2O)_3\}$ adopted a near-ideal centrosymmetric arrangement around the center of symmetry. Each trivacant Dawson- $\{P_2W_{15}\}$ offered six oxygen atoms to coordinate with three $\{(UO_2)_3(\mu_3-O)(\mu_2-H_2O)_3\}$. Thus, the tetrahedral structure was formed by the sandwiching of four

Dawson- $\{P_2W_{15}\}\$ and four $\{(UO_2)_3(\mu_3-O)(\mu_2-H_2O)_3\}$. Solvent water molecules occupied the central hole in the polyanion.

In 2003, Antonio and Pope both reported a series of 1:2 sandwich structures based on UIV and monovacant Dawson- $\{P_2W_{17}\}\$ units with a similar formula of $[U(P_2W_{17}O_{61})_2]^{16-43,44}$ Pope's work represented the case of diastereomers in U-POMs caused the two isomers by of $\{P_2W_{17}\},$ $[U(\alpha_1-P_2W_{17}O_{61})_2]^{16-}$, $[U(\alpha_1-P_2W_{17}O_{61})(\alpha_2-P_2W_{17}O_{61})]^{16-}$, and $[U(\alpha_2-P_2W_{17}O_{61})_2]^{16}$ (Fig. 4). P-NMR revealed the interconversion relationships of these compounds. Moisy also reported similar Dawson-type $An^{IV}(\alpha_2-P_2W_{17}O_{61})_2$ in 2009, including $\left[U(\alpha_2-P_2W_{17}O_{61})_2\right]^{16-}$ in the *syn*-conformation.⁴⁵

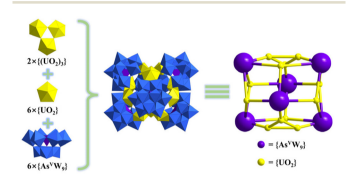


Fig. 2 Structural representation of $[(NH_4)_{12}(UO_2(H_2O))_6(UO_2(\mu_2-\mu_2))]$ $H_2O))_6(\alpha$ -AsW $_9O_{34})_6]^{18-}$. Color codes are the same as in Fig. 1.

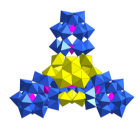
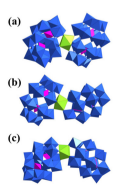


Fig. 3 View of $[(UO_2)_{12}(\mu_3-O)_4(\mu_2-H_2O)_{12}(P_2W_{15}O_{56})_4]^{32-}$. Color codes are the same as in Fig. 1.



of (a) $[U(\alpha_1-P_2W_{17}O_{61})_2]^{16-}$, (b) $[U(\alpha_1-P_2W_{17}O_{61})]$ $(\alpha_2 - P_2 W_{17} O_{61})^{16-}$, and (c) $[U(\alpha_2 - P_2 W_{17} O_{61})_2]^{16-}$. Color codes are the same as in Fig. 1.

Chen and co-workers also reported a Dawson-like structure based on a half-open {Si₂W₁₈} unit in 2007, i.e., Na₅{K $[UO_2(H_2O)]_2(A-\alpha-H_3SiW_9O_{33})_2$. ⁴⁶ A $\{K[UO_2(H_2O)]_2\}$ cluster was captured by the open sites of half-open {Si₂W₁₈}, which was similar to Keggin-type dimeric structures. One year later, Kortz synthesized a novel horseshoe-shaped Dawson-type structure using $K_{16}Li_2[H_6P_4W_{24}O_{94}]\cdot 33H_2O$ ($\{P_4W_{24}\}$), $UO_2(NO_3)_2$, and pH = 4 LiOAc/HOAc buffer (Fig. 5a).⁴⁷ In the polyanion $[\text{Li}(H_2O)K_4(H_2O)_3\{(UO_2)_4(O_2)_4(H_2O)_2\}_2(PO_3OH)_2P_6W_{36}O_{136}]^{25-},$ two groups of tetranuclear uranyl peroxide clusters {(UO₂)₄(O₂)₄(H₂O)₂} were stabilized by the coordination of oxygen atoms offered by $HPO_4^{\ 2-}$ and "U"-shaped trimeric Dawson-{P₆W₃₆}, which was in situ assembled from the Dawson-type precursor {P₄W₂₄}. It is worth noting that the addition of 4-6 drops of H₂O₂ and low reaction temperature (50 °C), which are usually used for the synthesis of uranyl peroxide clusters, may be the key factors for the synthesis of these POMs. However, it is a pity that there were no new Dawson-type U-POMs until ten years later. In 2019, Duval reported the first Dawson- $\{P_8W_{48}\}$ -based structure, $[(UO_2)_{7.2}(HCOO)_{7.8}(P_8W_{48}O_{184})]$ Cl₈]^{41.4-}, among which chloride was assembled into the structures of U-hetero-POMs for the first time. 48 This compound is also the first U-POM synthesized using a hydrothermal method. The reaction of {P₈W₄₈}, LiCl, and UO₂(NO₃)₂⋅6H₂O in HCOONa buffer at 160 °C gave a mixture of insoluble residue and a clear Crystals solution. of $(K_{11.3}Li_{8.1}Na_{22})[(UO_2)_{7.2}(HCOO)_{7.8}]$ (P₈W₄₈O₁₈₄)Cl₈]·89H₂O could be obtained by the evaporation of this solution. It is known that there is a big typical central cavity within {P₈W₄₈} and various relatively light metal ions or clusters can be captured in this cavity. But all the uranyl ions in $[(UO_2)_{7.2}(HCOO)_{7.8}(P_8W_{48}O_{184})Cl_8]^{41.4-}$ are located on both sides of the {P₈W₄₈} ring with disorder instead of in the central cavity. The occupancies of U1 and U2 are 0.25 and 0.65, respectively. Korenev and co-workers also reported two similar {P₈W₄₈}-based structures in 2019, but the uranyl ions were even disordered.49 The formulas are $K_{24}[(UO_2)_8]$ $(P_8W_{48}O_{184})$] $\cdot 50H_2O \cdot 3LiCl$ and $(NH_4)_{16}H_8[(UO_2)_8(P_8W_{48}O_{184})]$. 50H₂O. About one year later, Kortz synthesized a new {P₈W₄₈}uranyl-peroxide-cluster-containing POW, $[\{(UO_2)_4(O_2)_4\}_2(P_8W_{48}O_{184})]\cdot 133H_2O$, among which two groups of tetranuclear uranyl peroxide clusters {(UO₂)₄(O₂)₄} were both

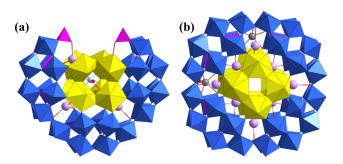
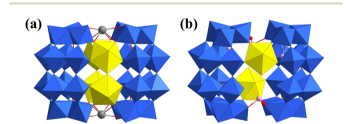


Fig. 5 Views of (a) $[Li(H_2O)K_4(H_2O)_3\{(UO_2)_4(O_2)_4(H_2O)_2\}_2(PO_3OH)_2$ $P_6W_{36}O_{136}]^{25-} \text{ and (b) } K_{18}Li_{22}[\{(UO_2)_4(O_2)_4\}_2(P_8W_{48}O_{184})]\cdot 133H_2O. \text{ Color}$ codes are the same as in Fig. 1; Li, pale blue.

located in the central cavity of {P₈W₄₈} (Fig. 5b).⁵⁰ This is also the first case in which such heavy atom clusters were assembled in the central cavity of {P₈W₄₈}. Several potassium ions were also captured in the central cavity.

Recently, our group and Zhang's group reported the first two uranium-containing selenotung states based on tetravacant Dawson-{Se₂W₁₄} (Fig. 6).⁵¹ The reaction of Na₂WO₄·2H₂O, Na₂SeO₃, and UO₂(NO₃)₂·6H₂O in an aqueous solution yielded two sandwich-type Dawson POWs, i.e., $Na_3[H_{19}(UO_2)_2(\mu_2-O)]$ $(NH_4)_{10}[H_4(SeO)_2(UO_2)_2(H_2O)_2]$ $(Se_2W_{14}O_{52})_2$]·41H₂O and $(H_2Se_2W_{14}O_{52})(Se_2W_{14}O_{52})]\cdot 66H_2O$. In the former one, two {α-Se₂W₁₄} sandwiched four halves of positionally disordered uranyl. In the latter polyanion, two $\{\gamma\text{-Se}_2W_{14}\}$ sandwiched two uranyl and two {Se^{IV}O}. It is also worth noting that this work represents two rare in situ synthesis cases among U-hetero-POMs. Different pH and counter cations ((CH₃)₂NH·HCl and NH₄Cl) were the significant factors of the one-pot synthesis procedures.

2.1.3 Silverton-type structures. The Silverton-type structures are one of the earliest U-hetero-POMs to be studied.9 All these Silverton-type U-hetero-POMs are based on the typical Silverton-type (UMo₁₂O₄₂)⁸⁻ anion. To the best of our knowledge, these Silverton-{UMo12}-based POMs were all reported by Torchenkova and co-workers from the former Soviet Union during 1981-1990 (Fig. 7). 15,52-59 Different metal ions, including Sc³⁺, VO²⁺, Fe³⁺, Ni²⁺, Cu²⁺, Nd³⁺, Er³⁺, Th⁴⁺, and UO₂²⁺,



of (a) $[(UO_2)_2(\mu_2-O)(Se_2W_{14}O_{52})_2]^{22-}$ $[(SeO)_2(UO_2)_2(H_2O)_2(H_2Se_2W_{14}O_{52})(Se_2W_{14}O_{52})]^{14-}. \quad Se^{IV},$ pink; other color codes are the same as in Fig. 1.

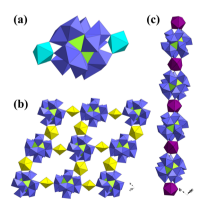


Fig. 7 Views of the Silverton-type $\{UMo_{12}\}$ -based structures. (a) $[(VO)_2(H_2O)_4(UMo_{12}O_{42})]^{4-}; \quad (b) \quad [(UO_2)_2(H_2O)_3(UMo_{12}O_{42})]^{4-}; \quad (c) \quad [Th]$ $(H_2O)_3(UMO_{12}O_{42})]^{4-}$. Color codes: Mo, blue; U^{IV} , light green; U^{VI} , yellow; Th^{IV}, violet; V: turquoise.

were incorporated into the structure by the coordination of three terminal oxygen atoms of $\{UMO_{12}\}$ with a similar formula of $\{M_a(H_2O)_b(UMO_{12}O_{42})\}^{c-}$ (a=1 or 2; b=0, 3, 4, 6, 12; c=2 or 4). However, most of them are 0D structures, only Th^{4+} and UO_2^{2+} incorporated compounds showed extended structures. $[Th(H_2O)_3(UMO_{12}O_{42})]^{4-}$ is a 1D chain built from Th^{4+} and $(UMO_{12}O_{42})^{8-}$, alternately. $[(UO_2)_2(H_2O)_3(UMO_{12}O_{42})]^{4-}$ represented a rare inorganic (4,4)-connected 2D network if $\{UMO_{12}\}$ clusters were regarded as nodes.

2.1.4 Hybrid structures. Loosely, there are only four cases that are constructed from more than one basic building block to date. The first was one of the three similar triangular sandwich structures reported by Pope in 2001, i.e., [(UO₂)₃(H₂O)₅ ture based on a Keggin-{AsW₉} and a Dawson-like {(AsW₉)₂(W₂)} unit, which was in situ transformed from $Na_{27}[NaAs_4W_{40}O_{140}]\cdot 60H_2O$ (Fig. 8a). The other two compounds were classified as cryptand structures. The second case of a hybrid structure was reported by Pope in 2002.³⁴ The reaction of Na₉[AsW₉O₃₃]·27H₂O and UO₂(NO₃)₂·6H₂O in an aqueous solution resulted in the formation $(NH_4)_{17}[(UO_2)_3(H_2O)_4As_3W_{26}O_{94}]\cdot 16H_2O$ and part of Keggin-{AsW₉} in situ transformed into Dawson-like {As₂W₁₇} (Fig. 8d). This {As₂W₁₇} unit is not the classical monovacant Dawson-{As₂W₁₇} and is built from two rare Keggin-{AsW₈} units and one bridging $\{WO_6\}$. As a result, one $\{AsW_9\}$ and one $\{As_2W_{17}\}$ sandwiched three uranyl to form a triangle configuration. The third one is an iso-heteropolytungstate, [(H₃SbW₁₇O₅₉) $U^{IV}(HW_5O_{18})]^{11-}$, which was constructed from a monovacant Lindqvist- $\{W_5\}$ and a monovacant Dawson- $\{Sb^{III}W_{17}\}$. 61 Significantly, this compound was synthesized from simple materials and is the first case of an in situ synthesized U-POM as far as we know.

Recently, our group and Zhang's group reported a new uranium-containing selenotungstate that consisted of one Anderson-{SeW₆} unit and three Keggin-{SeW₉} units (Fig. 9).⁶²

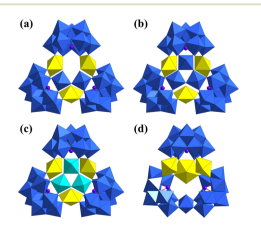


Fig. 8 Views of (a) $[(UO_2)_3(H_2O)_5As_3W_{29}O_{104}]^{19-}$, (b) $[(UO_2)_3(H_2O)_6As_3W_{30}O_{105}]^{15-}$, (c) $[(UO_2)_3(H_2O)_6As_3W_{29}V^{IV}O_{105}]^{17-}$, and (d) $[(UO_2)_3(H_2O)_4As_3W_{26}O_{94}]^{17-}$. Color codes are the same as in Fig. 1.

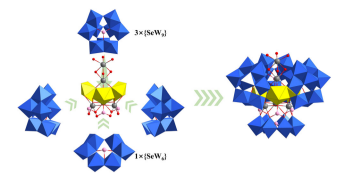


Fig. 9 View of $[(UO_2)_3(SeO_3)_3Na_5(H_2O)_6(SeW_6O_{21})(SeW_9O_{33})_3]^{21-}$. Color codes are the same as in Fig. 6.

In this hybrid polyanion, $[(UO_2)_3(SeO_3)_3Na_5(H_2O)_6(SeW_6O_{21})(SeW_9O_{33})_3]^{21-}$, a rare $\{Se_3U_3Na_5\}$ cluster was surrounded by one Anderson- $\{SeW_6\}$ and three Keggin- $\{SeW_9\}$ units from four directions to form a bouquet-like structure. This Anderson- $\{SeW_6\}$ is different from traditional Anderson units and can be viewed as a derivative of Keggin- $\{SeW_{12}\}$ removing the top and the bottom $\{W_3\}$ units. This compound was also a rare case of an *in situ* synthesized U-POM.

2.1.5 Cryptand structures. Compared with lacunary POMs, cryptand POMs show more rigid coordination sites. Meanwhile, the assembly ability of cryptand POMs is also limited by their stability and the sizes of open sites. The Preyssler-type $[P_5W_{30}O_{110}]^{15-}$ ($\{P_5W_{30}\}$) is a typical cryptand POM and is also one of the earliest models used to study the reactions between POMs and actinides. In 1993, Pope used $\{P_5W_{30}\}\$ to capture various transition metal ions. ⁷² The U^{4+} ion was successfully assembled in the central cavity of {P₅W₃₀} by ion exchange under high-temperature reflux conditions (140-180 °C). Three years later, Pope reported a similar ammonium salt with a formula of (NH₄)₁₁[U(H₂O) $P_5W_{30}O_{110}$]·12 H_2O (Fig. 10).⁶³ Single-crystal data revealed the positional disorder of U⁴⁺ within the cavity. The U⁴⁺ ion and an additional H_2O are located on the C_5 axis of $\{P_5W_{30}\}$. Compared with the U4+ ion, the central H2O was closer to the central position.

Another typical cryptand POM is $\left[As_4W_{40}O_{140}\right]^{28-}$ $\left(\left\{As_4W_{40}\right\}\right)^{.73}$ $\left\{As_4W_{40}\right\}$ exhibits two kinds and five in total coordination sites, theoretically. These two kinds of coordination sites are called S1 (the only one positioned at the center

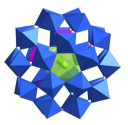


Fig. 10 View of $[U(H_2O)P_5W_{30}O_{110}]^{11-}.$ Color codes are the same as in Fig. 1.

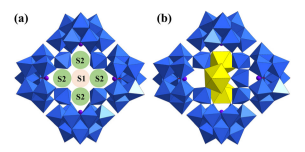


Fig. 11 Views of (a) two kinds of coordination sites S1 and S2 in ${As_4W_{40}}$ and (b) ${[Na(UO_2)_3(OH)(H_2O)_6As_4W_{40}O_{140}(WO)]^{18-}}$. Color codes are the same as in Fig. 1.

of {As₄W₄₀}) and S2 (doubled at two positions between two opposite {AsW₉} sub-units) (Fig. 11a). The first {As₄W₄₀}-based structure was one of the five cases of U-hetero-POMs reported by Pope in 2002 (Fig. 11b).34 The tetramer [Na(UO2)3(OH) $(H_2O)_6As_4W_{40}O_{140}(WO)]^{18-}$ was synthesized from UCl₄ and Na₂₇[NaAs₄W₄₀O₁₄₀]·60H₂O and crystallized from solution at room temperature. However, only three uranyl were assembled in the open sites of {As₄W₄₀}. U1 and U2 at S2 in this structure were both coordinated by two unsaturated oxygen sites from the $\{AsW_9\}$ part of $\{As_4W_{40}\}$, two single H_2O , and a μ_2 -OH. But U3 exhibited positional disorder with a 5-coordinated W atom at another two S2 sites.

It is interesting to note that the three {As₄W₄₀}-based U-hetero-POMs were also synthesized using UCl4 and $Na_{27}[NaAs_4W_{40}O_{140}]\cdot 60H_2O$ by Duval in 2018 (Fig. 12).⁶⁴ By changing the amounts of UCl₄ or {As₄W₄₀}, three similar tetramers were obtained from a similar HCOONa/HCOOH buffer at room temperature, namely $[\{U_2(\mu_2\text{-OH})_3(\mu_2\text{-O})\}\text{As}_4W_{40}O_{140}]^{25-}$, $[\{U_2(\mu_2\text{-O})(HCOO)_3\}As_4W_{40}O_{140}]^{25-}, \text{ and } [\{U_4(\mu_2\text{-O})_2(HCOO)_4\}$ $As_4W_{40}O_{140}]^{20-}$. But the U^{4+} ions in $[\{U_2(\mu_2-OH)_3(\mu_2-O)\}]$ As₄W₄₀O₁₄₀]²⁵⁻ were all positionally disordered with 0.5 occupancy for each S2 position. In $[\{U_2(\mu_2\text{-O})(HCOO)_3\}$ $As_4W_{40}O_{140}$]²⁵⁻, two U⁴⁺ ions were connected by three μ_2 formate and a μ_2 -O at two S2 positions. One of the {AsW₉} subunits was also transformed into a different configuration. The four U^{4+} ions in $[\{U_4(\mu_2-O)_2(HCOO)_4\}As_4W_{40}O_{140}]^{20-}$ were all located at S2 sites and an additional Na⁺ was also assembled at the S1 site.

Besides, the above-mentioned $[(UO_2)_3(H_2O)_6As_3W_{30}O_{105}]^{15-}$ and [(UO₂)₃(H₂O)₆As₃W₂₉V^{IV}O₁₀₅]¹⁷⁻ could also be viewed as

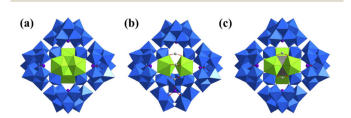


Fig. 12 Views of (a) $[\{U_2(\mu_2-OH)_3(\mu_2-O)\}As_4W_{40}O_{140}]^{25-}$, (b) $[\{U_2(\mu_2-O)\}As_4W_{40}O_{140}]^{25-}$ $(HCOO)_3$ $As_4W_{40}O_{140}$ C1, and (c) $[\{U_4(\mu_2-O)_2(HCOO)_4\}As_4W_{40}O_{140}]^{20-1}$. Color codes are the same as in Fig. 1.

cryptand-based structures (Fig. 8b and c). 60 Three {AsW₉} were linked by {W₃} and {VW₂} to form new {As₃W₃₀} and {As₃W₂₉V} building units with a triangular configuration. Three uranyl ions were coordinated by the oxygen atoms of the open sites. It was notable that the three atoms in {VW2} exhibited positional disorder and the occupancies of V and W were 1/3 and 2/3 in total, respectively. The other cryptand-based structures were the four {P₈W₄₈}-based structures as mentioned in section 2.1.2.

2.2 U-oxo-clusters

U-oxo-clusters discussed in this review are those structures that could be defined as POMs and uranium and oxygen atoms are the main building units of the structures. Uranium shows unique self-assembly modes in aqueous solution, which are distinctly different from traditional POMs built from early TM (W, Mo, etc.). Uranyl ions are always assembled into capsule-like uranyl peroxide clusters or other similar uranyloxo-clusters in the presence of H2O2 or other organic or inorganic ligands, while tetravalent uranium is easy to hydrolyze into U^{IV}-oxo-clusters with various structures. It is worth noting that there are no known U^{IV}-containing peroxide clusters due to rapid oxidation caused by H₂O₂. Burns is the pioneer in the field of actinide oxo-clusters, especially uranyl peroxide clusters, and Nyman also made outstanding contributions in this field. 10,74-76 One finds that the chemistry of uranyl peroxide clusters has already developed into a significant field from structure-driven to deeper research of chemical behaviors and has broadened the scope of POMs. 77-80 The U-oxo-clusters discussed in this section were mainly reported in recent years.

2.2.1 Uranyl peroxide clusters and POM-related uranyl peroxide phosphate clusters. Most of the uranyl peroxide clusters can also be called uranyl peroxide capsules due to their hollow-shell structures and uranyl peroxide rings or crowns. Various organic and inorganic ligands have been used as linkers of the clusters among those uranyl peroxide clusters. The organic ligands are mainly various organic phosphoric acids as well as oxalic acid (ox). Sun summarized the structural chemistry of uranium phosphonates based on the structures of organic phosphonate ligands and the hetero-metals.81 Inorganic ligands include PO₄³⁻, HPO₃²⁻, P₂O₇⁴⁻ (Pp), NO₃⁻, WO_4^{2-} , MoO_4^{2-} , and other anions in addition to O_2^{2-} . Those previously reported uranyl peroxide clusters from {U₁₆} to {U124} showed abundant topologies but could also be divided into two categories.⁷⁶ The first kind of uranyl peroxide clusters built only from uranyl ions, O22-, and OH are the most typical representatives of uranyl peroxide capsules. The biggest case is the highly symmetrical {U₆₀} cluster ([UO₂(O₂) (OH)]₆₀⁶⁰⁻). These kinds of uranyl peroxide capsules could be viewed as the combination of 4/5/6-membered uranyl polygons or bigger secondary building units (SBUs) of $\{(UO_2)_x(O_2)_x\}$ by sharing edges. The other kinds of uranyl peroxide clusters are those clusters in which different $\{(UO_2)_x(O_2)_x\}$ SBUs are linked by various organic and inorganic ligands as mentioned above. In consideration of the concept of polyoxometalates, only those structures with typical clusters are discussed here. These

Table 2 Summary of typical uranyl peroxide clusters

Codes	Formula	Shape	SBUs of clusters	Ref.
$\{U_{16}\}$	$[(UO_2)_{16}(O_2)_{24}(OH)_8]^{24-}$	Bowl-like	$\{U_4\}$ quadrangle, $\{U_6\}$ hexagon	82
$\{U_{16}(hedp)_8\}$	$[(UO_2)_{16}(O_2)_{16}(hedp)_8]^{32-}$	Cage	$\{U_4\}$ ring, $\{U_8\}$ double ring	83
$\{U_{16}(hedp)_8P_4\}$	$[(UO_2)_{16}(O_2)_8(OH)_8[(hedp)_8(PO_4)_4]^{28-}$	Cage	$\{U_4\} \text{ ring, } \{(UO_2)_2(OH)_2\}$	83
$\left\{ U_{18}Pp_{2}Pcp_{6}\right\}$	$[(UO_2)_{18}(O_2)_{18}(OH)_2(Pcp)_6(Pp)_2]^{34-}$	Cage	$\{U_8\}$ double ring, $\{U_{10}\}$ ring	84
$\{U_{18}Pp_6\}$	$[(UO_2)_{18}(O_2)_{18}(OH)_4(Pp)_6(H_2O)_4]^{28-}$	Cage	$\{U_8\}$ double ring, $\{U_{10}\}$ triple ring	85
$\{U_{19}(Ppb)_4\}$		Cage	$\{U_8\}$ double ring, $\{U_{11}\}$ triple ring	86
$\{U_{20}\}$	$[(UO_2)_{20}(O_2)_{30}]^{20}$	Cage	$\{U_5\}$ pentagon	87
$\{U_{20R}\}$	$[(UO_2)_{20}(OH)_{16}(O_2)_{28}]^{32-}$	Crown-like	$\{U_5\}$ pentagon	82
$\{U_{20}P_{6}\}$	$[(UO_2)_{20}(O_2)_{27}(HPO_4)_6]^{26-}$	Cage	$\{U_5\}, \{U_8\} \text{ ring}$	88
$\{U_{20}P_{12}\}$	$[(UO_2)_{20}(O_2)_{24}(HPO_4)_6(H_2PO_4)_6]^{26-}$	Cage	$\{U_8\}$ ring	88
$\{U_{20}Pp_{6a}\}$	$[(UO_2)_{20}(O_2)_{24}(Pp)_6]^{32-}$ $[(UO_2)_{20}(O_2)_{24}(Pp)_6]^{332-}$	Cage	$\{U_{10}\}$ triple ring	84
$\{U_{20}Pp_{6b}\}$	$[(UO_2)_{20}(O_2)_{24}(Pp)_6]^{32-}$	Cage	$\{U_5\}, \{U_{10}\} \text{ ring}$	84
$\{U_{20}Pp_{10}\}$	$ \frac{[(UO_2)_{20}(O_2)_{20}(Pp)_{10}]^{40-}}{[(UO_2)_{20}(O_2)_{20}(Pp)_{10}]^{40-}} $	Cage	$\{U_8\}$ double ring, $\{U_{12}\}$ quadruple ring	84
$\{U_{20}Pp_{10a}\}$	$[(UO_2)_{20}(U_2)_{20}(PP)_{10}]$ $[(UO_1)_{20}(O_2)_{20}(PP)_{10}]$	Cage	$\{U_5\}, \{U_{10}\} \text{ ring}$	85
$\{U_{20}(hedp)_{10}\}$	$[(UQ_2)_{20}(Q_2)_{20}[(hedp)_{10}]^{40-}$	Cage	$\{U_5\}, \{(UO_2)_{10}(O_2)_{10}\} \text{ rings}$	83 89
$\{U_{20}(Ppb)_{10}\}$	$(UO_2)_{20}(O_2)_{20}(Ppb)_{10}^{40-}$	Cage	$\{U_5\}, \{U_{10}\} \text{ ring}$	89
$\{U_{20}(Ppb)_{6}\}$	[(UO ₂) ₂₀ (O ₂) ₂₄ (Ppb) ₆] ³²⁻ [(UO ₂) ₂₁ (O ₂) ₂₁ (Pp) ₉ (OH) ₃] ³⁹⁻	Cage	$\{U_{10}\}$ triple ring	85
$\{U_{21}Pp_9\}$	$ [(UO_2)_{21}(P_2)_{21}(P_1)_9(UH_{33}] $ $[(UO_2)_{22}(O_2)_{15}(HPO_3)_{20}(H_2O)_{10}]^{26-} $	Cage	$\{U_4\}$ ring, $\{U_9\}$ triple ring	90
$\{U_{22p}\}$	$ [(UO_2)_{24}(OH)_{24}(O_2)_{24}]^{24} $	Cage Cage	$\{(UO_2)_4(O_2)_4\}, \{UO_2\}$ $\{U_4\}$ quadrangle, $\{U_6\}$ hexagon	91
$\{U_{24}\}$	$ [(UO_2)_{24}(OH)_{26}(O_2)_{12}]^{24-} $	Crown-like	$\{U_6\}$ hexagon	82
$\{U_{24R}\}$	$ [(UO_2)_{24}(O1)_{36}(O_2)_{12}] $ $[(UO_2)_{24}(O_2)_{25}(Pp)_9(OH)_4]^{42-} $	Cage	$\{U_4\}$ ring, $\{U_8\}$ double ring, $\{U_{12}\}$ triple	85
$\left\{ U_{24}Pp_{9}\right\}$	$[(OO_2)_{24}(O_2)_{25}(PP)_9(OH)_4]$	Cage		03
(II Don)	$[(UO_2)_{24}(O_2)_{24}(Pcp)_{10}(H_2O)_4]^{40-}$	Carro	ring $\{U_4\}$, $\{U_8\}$ rings	85
$\{U_{24}Pcp_{10}\}$	$ [(UO_2)_{24}(O_2)_{24}(Pp)_{10}(H_2O)_{4}] $ $[(UO_2)_{24}(O_2)_{24}(Pp)_{12}]^{48-} $	Cage	$\{U_4\}, \{U_8\}$ fligs	84
$\{U_{24}Pp_{12}\}$	$ [(UO_2)_{24}(P_2)_{24}(P_2)_{12}]^{48-} $ $ [(UO_2)_{24}(O_2)_{24}(Pcp)_{12}]^{48-} $	Cage	$\{U_4\}$ ring	84
$\{U_{24}Pcp_{12}\}\ \{U_{24}(hedp)_{12}\}$	$ [(UO_2)_{24}(O_2)_{24}(FCP)_{12}] $ $ [(UO_2)_{24}(O_2)_{24}(hedp)_{12}]^{48-} $	Cage	$\{U_4\}$ ring $\{U_4\}$ ring	83
	$[(UO_2)_{24}(O_2)_{24}(IICup)_{12}]$	Cage		84
$\{U_{26}Pp_6\}$	$ \begin{bmatrix} (UO_2)_{26}(O_2)_{33}(Pp)_6 \end{bmatrix}^{38-} \\ [(UO_2)_{26}(O_2)_{28}(Pp)_{11}]^{48-} \\ \end{bmatrix} $	Cage	$\{U_5\}$ pentagon $\{U_5\}$ ring, $\{(UO_2)_{16}\}$ triple ring	84
$\{ U_{26}Pp_{11} \} $ $\{ U_{26}(Ppb)_{6} \}$	$ [(UO_2)_{26}(O_2)_{28}(Pp)_{11}] $ $ [(UO_2)_{26}(O_2)_{33}(Ppb)_6]^{38-} $	Cage Cage	$\{U_5\}$ pentagon, $\{U_6\}$ hexagon	89
$\{U_{28}\}$	$ [(UO_2)_{28}(O_2)_{33}(IPB)_6] $ $[(UO_2)_{28}(O_2)_{42}]^{28-} $	Cage	$\{U_5\}$ pentagon, $\{U_6\}$ hexagon	91
$\{U_{28a}\}$	$ [(UO_2)_{28}(O_2)_{42}]^{28-} $	Cage	$\{U_4\}$ quadrangle, $\{U_5\}$ pentagon, $\{U_6\}$	92
\O28a\f	$[(OO_2)_{28}(O_2)_{28}(OII)_{28}]$	Cage	hexagon	92
$\{U_{28P}\}$	$[(UO_2)_{28}(O_2)_{20}(HPO_3)_{24}(H_2O)_{12}]^{32-}$	Cage	$\{(UO_2)_4(O_2)_4\}, \{(UO_2)_2(O_2)\}$	90
$\{U_{30}\}$	$[(UO_2)_{30}(O_2)_{36}(OH)_{22}]^{34-}$	Cage	$\{U_4\}$ quadrangle, $\{U_5\}$ pentagon, $\{U_6\}$	92
(55)	2,000(2,000()223	o .	hexagon	
$\{U_{30a}\}$	$[(UO_2)_{30}(O_2)_{30}(OH)_{30}]^{30-}$	Cage	$\{U_4\}$ quadrangle, $\{U_5\}$ pentagon, $\{U_6\}$	93
			hexagon	
$\left\{ U_{30}Pp_{6}\right\}$	$[(UO_2)_{30}(O_2)_{39}(Pp)_6]^{42-}$	Cage	$\{U_6\}$ ring, Bowl-like $\{U_{24}\}$: $\{U_5\}$ penta-	93
			gon, $\{U_6\}$ hexagon	
$\{U_{30}Pp_{10}Ox_{5}\}$	$[(UO_2)_{30}(O_2)_{30}(Pp)_{10}(Ox)_5]^{50-}$	Cage	$\{U_5\}, \{U_{10}\} \text{ rings}$	94
$\left\{ U_{30}Pp_{12}P\right\}$	$[(UO_2)_{30}(O_2)_{30}(Pp)_{12}(PO_4)(H_2O)_5]^{51-}$	Double cage	$\{U_5\}, \{U_{10}\} \text{ rings}$	93
$\{U_{32}\}$	$[(UO_2)_{32}(OH)_{32}(O_2)_{32}]^{32-}$	Cage	$\{U_4\}$ quadrangle, $\{U_5\}$ pentagon, $\{U_6\}$	91
			hexagon	
$\{U_{32R}\}$	$[(UO_2)_{32}(O_2)_{40}(OH)_{24}]^{40-}$	Crown-like	$\{U_5\}$ pentagon, $\{U_6\}$ hexagon	95
$\{U_{32}Pp_{16}\}$	$[(UO_2)_{32}(O_2)_{32}(Pp)_{16}]^{64-}$	Cage	$\{U_4\}$ ring	84
$\{U_{36}\}$	$ (UO_2)_{36}(O_2)_{41}(OH)_{26} ^{30}$	Cage	$\{U_5\}$ pentagon, $\{U_6\}$ hexagon	96
$\{U_{36a}\}$	$[(UO_2)_{36}(O_2)_{36}(OH)_{36}]^{36-}$	Cage	$\{U_4\}$ quadrangle, $\{U_5\}$ pentagon, $\{U_6\}$	92
()	F() (-) 726-	_	hexagon	
$\{U_{36}Ox_6\}$	$[(UO_2)_{36}(O_2)_{48}(OX)_6]^{36-}$	Cage	Bowl-like {U ₁₈ } septuple ring	97
$\left\{ U_{38}Pp_{10}Nt_{4}\right\}$	$[(UO_2)_{38}(O_2)_{40}(Pp)_{10}(NO_3)_4]^{56-}$	Cage	$\{U_8\}$ double ring, $\{U_{11}\}$ triple ring	94
$\{\mathrm{U_{40}}\}$	$[(UO_2)_{40}(OH)_{40}(O_2)_{40}]^{40-}$	Cage	$\{U_4\}$ quadrangle, $\{U_5\}$ pentagon, $\{U_6\}$	98
((1 1))	F() (a.) (1. 1.) 380-		hexagon	
$\{U_{40}(hedp)_{20}\}$	$[(UO_2)_{40}(O_2)_{40}(hedp)_{20}]^{80-}$	Cage	$\{U_4\}$ ring	83
$\{U_{42}\}$	$[(UO_2)_{42}(O_2)_{42}(OH)_{42}]^{42}$	Cage	$\{U_4\}$ quadrangle, $\{U_5\}$ pentagon, $\{U_6\}$	93
(rr - p., .)	[(HO) (O) (OH) (Pu) 148-	0	hexagon	0.0
$\left\{ U_{42}Pp_{3}\right\}$	$[(UO_2)_{42}(O_2)_{42}(OH)_{36}(Pp)_3]^{48-}$	Cage	$\{U_4\}$ quadrangle, $\{U_5\}$ pentagon, $\{U_6\}$	93
(T.T.)	[(HO) (O) 144-	Ellinesi J	hexagon, $\{U_{10}\}$ decagon	0.6
$\{U_{44}\}$	$[(UO_2)_{44}(O_2)_{66}]^{44-}$	Ellipsoid cage	$\{U_5\}$ pentagon, $\{U_6\}$ hexagon	96
$\{U_{44a}\}$	$[(UO_2)_{44}(O_2)_{44}(OH)_{44}]^{44-}$	Dumbbell-like	$\{U_4\}$ quadrangle, $\{U_6\}$ hexagon, $\{U_8\}$	92
(II Des)	$[(HO_1), (O_1), (Pp_1)]^{190-}$	Carra	octagon	0.2
$\{U_{45}Pp_{23}\}$	$ \frac{\left[(\text{UO}_2)_{45} (\text{O}_2)_{44} (\text{Pp})_{23} \right]^{90-} }{\left[(\text{UO}_2)_{50} (\text{OH})_{50} (\text{O}_2)_{50} \right]^{50-} } $	Cage	$\{U_4\}, \{U_5\} \text{ rings}, \{(UO_2)_3(O_2)_2\}$	83
$\{U_{50}\}$	$[(UO_2)_{50}(U\Pi)_{50}(U_2)_{50}]$	Cage	$\{U_5\}$ pentagon, $\{U_6\}$ hexagon	98
$\{U_{50}Ox_{20}\}$	$[(UO_2)_{50}(O_2)_{43}(OH)_4(OX)_{20}]^{30-}$	Cage	$\{U_5\}$ ring, $\{U_8\}$ double ring	99
$\{U_{54}Pp_{27}\}$	$[(UQ_2)_{54}(Q_2)_{54}(Pp)_{27}]^{108-5}$ $[(UQ_2)_{64}(Q_2)_{54}(Pp)_{27}]^{60-5}$	Cage	$\{U_4\}, \{U_5\} \text{ rings}$	85 06
{U ₆₀ }	$[(UO_2)(O_2)(OH)]_{60}^{60-}$	Cage	$\{U_5\}$ pentagon, $\{U_6\}$ hexagon	96 07
$\{U_{60}Ox_{30}\}$	$\begin{bmatrix} (UQ)_{60}(O_2)_{60}(OX)_{30} \end{bmatrix}^{60-} \\ \begin{bmatrix} (UQ)_{60}(O_1)_{60}(OX)_{30} \end{bmatrix}^{128-} \\ \end{bmatrix}$	Cage	$\{U_5\}$ ring	97 83
$\{U_{64}(hedp)_{32}\}$	$[(UO_2)_{64}(O_2)_{64}(hedp)_{32}]^{128-}$	Cage	$\{U_4\}$ ring	03

Table 2 (Contd.)

Codes	Formula	Shape	SBUs of clusters	Ref.
$\{U_{28}U_{40R}\}$ or $\{U_{68}\}$	$\left[(UO_2)_{68} (O_2)_{74} (OH)_{16} (NO_3)_{16} (H_2O)_{16} \right]^{44-}$	$\{U_{28}\}$: cage $\{U_{40R}\}$: saddle-like ring	$\{U_{28}\}$: $\{U_5\}$ pentagon, $\{U_6\}$ hexagon; $\{U_{40R}\}$: $\{U_4\}$ quadrangle, $\{U_5\}$ pentagon	100
$ \begin{cases} U_{120}Ox_{90} \\ U_{124}P_{32} \\ \end{cases} $	$ \begin{array}{l} \left[(UO_2)_{120} (O_2)_{120} (Ox)_{90} \right]^{180-} \\ \left[(UO_2)_{124} (O_2)_{152} (PO_4)_{16} (HPO_4)_8 (H_2PO_4)_8 (OH)_2 (H_2O)_{24} \right]^{132-} \end{array} $	Double-layer cage Five cages combined structure	$\{U_5\}$ rings Bowl-like $\{U_{18}\}$ septuple ring, $\{U_{16}\}$ triple ring, $\{U_5\}$ ring	99 101

Abbreviation: hedp = (1-hydroxyethane-1,1-diyl)bis(phosphonic acid), Pcp = methylenebis(phosphonic acid), Ppb = benzene-1,2-diphosphonic acid.

uranyl peroxide clusters could be classified according to the numbers of uranyl moieties in the cluster and then their topologies. The typically reported uranyl peroxide clusters are summarized in Table 2.⁷⁶ Those uranyl peroxide clusters containing other TM ions are classified in section 2.2.2. Being different from traditional POMs, the counter cations are found to be one of the most important factors influencing the assembly progress, structures, states of aggregation, solubilities of uranyl peroxide clusters, sometimes even more important than pH.6,76,102 The attractiveness of uranyl peroxide clusters is not only their varied topology but also the interesting space chemistry within the inner cavities of the cages. 103,104

In 2017, Burns synthesized the only two uranyl peroxide phosphate clusters with μ-η¹:η² peroxide bridges, $\text{Li}_4\text{Na}_{12}\{(SO_4)\subset \text{Na}_{12}\subset [(UO_2)_{20}(O_2)_{27}(HPO_4)_6]\}$ and Li_xNa_{16-x} $\{(SO_4)\subset Na_{12}\subset [(UO_2)_{20}(O_2)_{24}(HPO_4)_6(H_2PO_4)_6]\}$ 13). Different amounts of Na₂SO₃ and H₃PO₄ and different basic pH values resulted in the two similar but different uranyl peroxide phosphate clusters. The inner cores were both {Na₁₂(SO₄)} clusters, which were further surrounded by different fullerene-like $\{U_{20}\}$ phosphate shells.⁸⁸ At the end of 2018, Burns reported six new uranyl peroxide phosphate clusters, and P₂O₇⁴⁻ and methylenebis(phosphonic acid) ((CH₂) (PO₃H₂)₂, Pcp) were used as bridging ligands (Fig. 14).⁸⁵ The obtained $\{U_{18}Pp_6\}$, $\{U_{20}Pp_{10a}\}$, $\{U_{21}Pp_9\}$, $\{U_{24}Pp_9\}$, $\{U_{24}Pcp_{10}\}$, and {U54Pp27} show different topologies of uranyl peroxide clusters, which were constructed from different SBUs of clusters linked by P2O74- or Pcp ligands. The compositions of these uranyl peroxide phosphate clusters are summarized in Table 3.

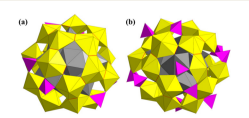


Fig. 13 Views of (a) $\text{Li}_4\text{Na}_{12}\{(SO_4)\subset \text{Na}_{12}\subset [(UO_2)_{20}(O_2)_{27}(\text{HPO}_4)_6]\}$ and (b) $Li_xNa_{16-x}\{(SO_4)\subset Na_{12}\subset [(UO_2)_{20}(O_2)_{24}(HPO_4)_6(H_2PO_4)_6]\}$. Color codes are the same as in Fig. 1.

Burns also used high-temperature drop solution calorimetry to study the enthalpies of formation of several uranyl peroxide clusters and the disassembly and assembly process from $\text{Li}_{24}\text{Na}_{24}\{[(UO_2)(O_2)]_{24}(P_2O_7)_{12}\}\ \text{to}\ \text{Li}_{24}[(UO_2)(O_2)(O_1)]_{24}\ \text{and}\ \text{a}$ series of uranyl peroxide clusters containing {U24Pp12}, $\{U_{26}Pp_6\}, \{U_{28}\}, \{U_{22}(PO_3)_{20}\}, \text{ and } \{U_{28}(PO_3)_{24}\} \text{ with different}$ counter cations. 105,106 Then, the transformation from studtite to different uranyl peroxide clusters was also investigated. The chains of uranyl peroxide polyhedra in studtite, $[(UO_2)(O_2)]$ (H₂O)₂](H₂O)₂, were converted into two kinds of uranyl peroxide clusters under different conditions, i.e., $\{U_{28}\}$ and $\{U_{20}\}$, which was revealed by Raman spectra and electrospray ionization mass spectrometry (ESI-MS). 107 Whereafter, a novel uranyl peroxide cluster, $K_{32}(UO_2)_{19}(O_2)_{26}(OH)_2(Ppb)_4 \cdot 65H_2O$ (Ppb = benzene-1,2-diphosphonic acid), was synthesized with a new "open oyster" structure. Abundant hydrogen bonds, $\pi \cdots \pi$, and $K \cdots \pi$ interactions, and K-O bonds supported the formation of the supramolecular structures. This {U19} cluster also transform into a typical $\{U_{24}\}$ $K_{24}(UO_2)_{24}(O_2)_{24}(OH)_{24} \cdot nH_2O$, with an octanuclear K-cluster within the central cavity. 86 The reaction of Ppb and UO₂(NO₃)₂ in a basic solution could also result in three new organic-inclusters organic hybrid uranyl peroxide $[(UO_2)_{20}]$ $(O_2)_{20}(Ppb)_{10}]^{40-}$, $[(UO_2)_{26}(O_2)_{33}(Ppb)_6]^{38-}$, and $[(UO_2)_{20}(O_2)_{24}(Ppb)_6]^{32-}$. The two $\{U_{20}\}$ clusters both showed similar fullerene topology, while the {U26} cluster exhibited a distorted polyhedron with more than 15 faces. Nyman investigated the influence of three organic solvents (acetone, CH3CN, and EtOH) in the synthesis of different fragments of uranyl peroxide clusters. 79 A new Anderson-like hexameric cluster, {K@ (UO2)6(O2)9(OH)6, was successfully isolated, which was the second stable ligand-free SBU of normal uranyl peroxide clusters (Fig. 15a). The first one, pentameric $\{K@(UO_2)_5(O_2)_5\}$, was also reported by Nyman in 2019 (Fig. 15b). 108 Besides, Forbes also synthesized a hexametric {(UO₂)₆(O₂)₆(HPO₄)₆} in the absence of H₂O₂ (Fig. 15c). 109 An interesting peroxide process occurred with the help of trace levels of benzaldehyde and air. The raw material (aminomethyl)phosphonic acid in situ transformed into HPO₄²⁻ in the solution of methanol and benzyl alcohol with NaOH.

At the end of 2021, LaVerne and co-workers investigated the solid-state transformations of Ca₂[UO₂(O₂)₃]·9H₂O and Li₄[(UO₂)(O₂)₃]·10H₂O under γ-irradiation. ¹¹⁰ Single-crystal and

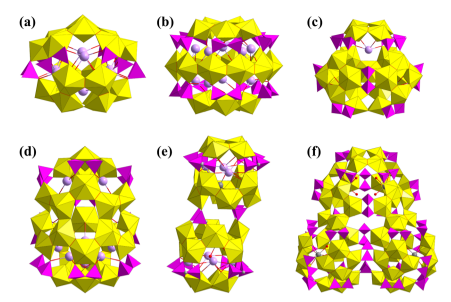


Fig. 14 Views of (a) $\{U_{18}Pp_6\}$, (b) $\{U_{20}Pp_{10a}\}$, (c) $\{U_{21}Pp_9\}$, (d) $\{U_{24}Pp_9\}$, (e) $\{U_{24}Pcp_{10}\}$, and (f) $\{U_{54}Pp_2\}$. Color codes are the same as in Fig. 1.

Table 3 Summary of the structures of $\{U_{18}Pp_6\}$, $\{U_{20}Pp_{10a}\}$, $\{U_{21}Pp_9\}$, $\{U_{24}Pp_9\}$, $\{U_{24}Pp_{10a}\}$, and $\{U_{54}Pp_{27}\}$

Codes	Formula	SBUs of clusters	Linkers
$ \begin{cases} \{U_{18}Pp_6\} \\ \{U_{20}Pp_{10a}\} \\ \{U_{21}Pp_9\} \\ \{U_{24}Pp_9\} \\ \{U_{24}Pcp_{10}\} \\ \{U_{54}Pp_{27}\} \end{cases} $	$\begin{split} & \left[(UO_2)_{18}(O_2)_{18}(OH)_4(P_2O_7)_6(H_2O)_4 \right]^{28-} \\ & \left[(UO_2)_{20}(O_2)_{20}(P_2O_7)_{10} \right]^{40-} \\ & \left[(UO_2)_{21}(O_2)_{21}(P_2O_7)_9(OH)_3 \right]^{39-} \\ & \left[(UO_2)_{24}(O_2)_{25}(P_2O_7)_9(OH)_4 \right]^{42-} \\ & \left[(UO_2)_{24}(O_2)_{24}(O_3PCH_2PO_3)_{10}(H_2O)_4 \right]^{40-} \\ & \left[(UO_2)_{54}(O_2)_{54}(P_2O_7)_{27} \right]^{108-} \end{split}$	$ \begin{aligned} &\{(UO_2)_8(O_2)_8\}, \{(UO_2)_{10}(O_2)_{10}\} \\ &2 \times \{(UO_2)_5(O_2)_5\}, \{(UO_2)_{10}(O_2)_{10}\} \text{ ring} \\ &3 \times \{(UO_2)_4(O_2)_4\}, \{(UO_2)_9(O_2)_9\}, \{(UO_2)_{12}(O_2)_{12}\} \\ &\{(UO_2)_4(O_2)_4\}, \{(UO_2)_8(O_2)_8\}, \{(UO_2)_{12}(O_2)_{12}\} \\ &2 \times \{(UO_2)_4(O_2)_4\}, 2 \times \{(UO_2)_8(O_2)_8\} \text{ ring} \\ &11 \times \{(UO_2)_4(O_2)_4\}, 2 \times \{(UO_2)_5(O_2)_5\} \end{aligned} $	6 P ₂ O ₇ ⁴⁻ 2 × 5 P ₂ O ₇ ⁴⁻ 3 and 6 P ₂ O ₇ ⁴⁻ 9 P ₂ O ₇ ⁴⁻ 2 × 4 and 2 Pcp ⁴⁻ 27 P ₂ O ₇ ⁴⁻

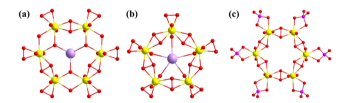


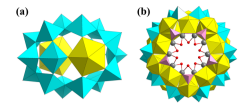
Fig. 15 Views of (a) $\{K@(UO_2)_6(O_2)_9(OH)_6\}$, (b) $\{K@(UO_2)_5(O_2)_5\}$, and (c) $\{(UO_2)_6(O_2)_6(HPO_4)_6\}$. Color codes are the same as in Fig. 1.

powder X-ray diffraction and Raman spectra revealed the different decomposition of peroxide groups into hydroxyl groups, which was influenced by Ca2+ and Li+. For Ca₂[UO₂(O₂)₃]·9H₂O, the products were a mixture of amorphous uranium phase and Ca(OH)₂. The product of Li₄[(UO₂) (O₂)₃]·10H₂O was uranyl oxyhydrate. Very recently, Burns also used γ -ray radiation to promote the conversion of Li₄[UO₂(O₂)₃] into Li₂₄[(UO₂)(O₂)(OH)]₂₄ in aqueous solutions. ¹¹¹ They used ex situ Raman spectroscopy and 18O labeling to track this transformation and found the dependence relationship between Li₄[UO₂(O₂)₃], intermediates of uranyl peroxide clusters, and the radiation dose. These two works show us direct cases to understand the formation mechanism of POMs from monomer to derived species.

2.2.2 U-oxo-clusters containing other transition metals.

This kind of U-oxo-cluster is mainly structures incorporating other TM ions. The introduction of a TM into U-oxo-clusters could result in more abundant coordination and connection modes. Burns synthesized a series of U-oxo-cluster phosphates incorporating Mo or W in 2014. 112 In these U-oxo-clusters, different uranyl peroxide clusters were connected by phosphate tetrahedra and Mo- or W-polyhedra to form different cage structures that showed similar corresponding uranyl peroxide clusters topologies, which were named {U18W2P12}, $\{U_{28}W_4P_{12}\},\quad \{U_{28}Mo_4P_{12}\},\quad \{U_{44}Mo_2P_{16}\},\quad \{U_{48}W_6P_{48}\},\quad \text{and}\quad$ $\{U_{50}W_6P_{20}\}$. The number of UO_2^{2+} increased from 18 to 50. Mo- or W-polyhedra all served as linkers at the peaks in place of uranyl. Besides, all these compounds were synthesized in typical acidulous or circumneutral aqueous solutions containing H₃PO₃, H₂O₂, and LiOH. The dissociation of H₃PMo₁₂O₄₀ or H₃PW₁₂O₄₀ provided the corresponding MoO₄²⁻ or WO₄²⁻ ions.

In 2015, Burns reported two uranyl-vanadium oxide clus- $[(UO_2)_2(V_{16}O_{46})]^{8-}$ and $[(UO_2)_{20}(V_2O_7)_{10}(SO_4)_{10}]^{20-}$ (Fig. 16). 113 The {U₂V₁₆} cluster could be viewed as a substitution product of the {V₁₈} cluster with two uranyl ions. The {U20V20} cluster represents a unique nano uranyl-vanadium wheel. Ten uranyl pentagonal bipyramids are connected by



 $[(UO_2)_2(V_{16}O_{46})]^{8-}$ Fig. 16 Views of (h) $[(UO_2)_{20}(V_2O_7)_{10}(SO_4)_{10}]^{20-}$. Color codes: U, yellow; V, turquoise; Na, dark grey; S, pink; O, red.

sharing vertices to form two pentagon-like $\{U_{10}\}$ rings. The two $\{U_{10}\}$ rings are further stabilized by the zigzag $\{V_{20}\}$ ring and SO₄²⁻ to build the unique wheel. Additional symmetric ten halves of Na ions are captured by the cavity of the wheel. One vear later, Burns reported the first uranium-vanadium-phosphorus peroxide cage cluster with a formula of $[(UO_2)_{48}(O_2)_{12}(VO_5)_6(PO_4)_{48}]^{x-}$ ($x \approx 102$), which showed an isostructure to {U₄₈W₆P₄₈} reported in 2014. 114 Subsequently, Burns synthesized an organic-inorganic hybrid uranium-iron cage cluster from a solution containing UO₂(NO₃)₂, H₂O₂, LiOH, 3,3-diphosphono-propanoic acid (C₃P₂), and FeCl₃ (Fig. 17). 115 One C₃P₂ was oxidized to the ligand 2,2-bis(phosphonato)acetate (C₂P₂) to combine with one Fe³⁺. Three Fe³⁺ were connected by $PO_4^{\ 3-}$ and three μ_2 -OH to form a trinuclear cluster {Fe₃(OH)₃(PO₄)(C₂P₂)}. Four uranyl were connected by four μ_2 - O_2^{2-} to form a $\{(UO_2)_4(O_2)_4\}$ cluster. Thus, eight $\{Fe_3(OH)_3(PO_4)(C_2P_2)\}\$ clusters and six $\{(UO_2)_4(O_2)_4\}\$ clusters were connected by the phosphate groups of C2P2 to form the cage cluster $[\text{Li}_8(\text{UO}_2)_{24}(\text{FeOH})_{24}(\text{O}_2)_{24}(\text{PO}_4)_8(\text{C}_2\text{P}_2)_{24}]^{88-}$. This cage could be simplified as a nonstandard quadrangular dodecahedron if Fe trimers and U tetramers were considered as 3and 4-connected nodes, respectively.

In 2019, Nyman investigated the influence of different counter cations on the hydrolysis of U(SO₄)₂ and a series of U^{IV}-oxo-sulfate clusters were obtained based on 6-, 70-, or 84-nuclear U^{IV}-oxo-clusters (Fig. 18). 116,117 Lanthanides from La to Ho (except Pm) resulted in the formation of ten similar clusters of $[U_6(OH)_4(O)_4(SO_4)_{12}]^{12-}$ ($\{U_6\}$). While heavy lanthanides and Y³⁺ with smaller ionic radiuses resulted in a kind of hollow supercluster with more than 84 uranium atoms based on a body-centered-like cubic cluster, $\{[U_6(OH)_4(O)_4]_{14}(SO_4)_x\}$

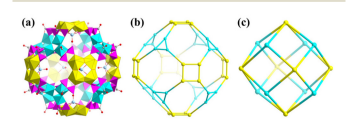


Fig. 17 Views of (a) $[Li_8(UO_2)_{24}(FeOH)_{24}(O_2)_{24}(PO_4)_8(C_2P_2)_{24}]^{88-}$, (b) the simplified diagram, and (c) the simplified quadrangular dodecahedron. Color codes: U, yellow; Fe, turquoise; O, red; P, pink; C, gray; Li, pale blue.

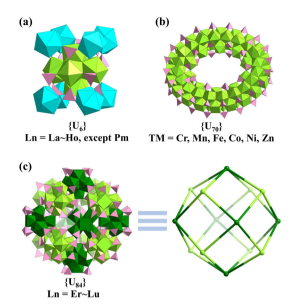


Fig. 18 Views of the (a) $\{U_6\}$, (b) $\{U_{70}\}$, and (c) $\{U_{84}\}$ clusters. Color codes: U^{IV} and {U₆} units, light green and dark green; lanthanides and TM ions, turquoise; S, pink; O, red.

 $(\{U_{84}\}, x = 108 \text{ for Y}, 111 \text{ for Er}, \text{ and } 110 \text{ for Lu})$. The introduction of TM ions (Cr, Mn, Fe, Co, Ni, Zn) could yield wheelshaped clusters of $[U_{70}(OH)_{36}(O)_{64}(SO_4)_{64}]^{12-}$ ($\{U_{70}\}$). Different counter cations and amounts of raw materials also led to different connection modes of these clusters. This series of U-oxo-clusters could also be classified as U^{IV}-oxo-clusters due to the +4 valence of all the uranium atoms.

In 2021, Burns developed two kinds of uranyl vanadates with core-shell structures, which could be classified as U-oxoclusters containing other transition metals (Fig. 19). 118 {U₈V₆}

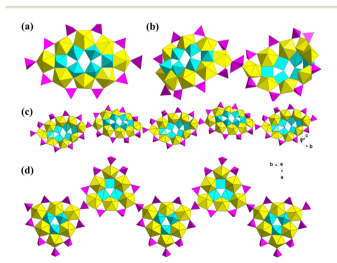


Fig. 19 Views of the (a) monomer, (b) dimer, and (c) 1D chain of {U₈V₆} and (d) 1D chain of $\{U_6V_3\}$. In the $\{U_8V_6\}$ dimer, the $\{PO_4\}$ linker is represented with positional disorder. Carbon atoms for diethyl phosphate are omitted for simplicity. Color codes: U, yellow; V, turquoise; P, magenta; O, red.

and {U₆V₃} could be obtained with different extended structures from an ionic liquid solution containing 1-ethyl-3methylimidazolium diethyl phosphate (EMIm-Et₂PO₄), UO₂(NO₃)₂, NaVO₃, and/or VOSO₄, and Na₂WO₄. The monomer, dimer, and 1D chain of {U8V6} and 1D chain of {U₆V₃} represent a new series of U-oxo-clusters incorporating vanadium atoms.

2.2.3 U^{IV}-oxo-clusters and mixed-valence U-oxo-clusters. Uoxo-clusters built from low valence UIV are usually obtained from the hydrolyzation of U^{IV} salts or the reduction of uranyl under specific conditions. Due to the similarity of some +4 metal ions, U^{IV} could assemble into a hexanuclear U^{IV}cluster $\{U_6O_4(OH)_4\}$ or $\{U_6O_8\}$, 119,120 which was similar to the {Zr₆O₄(OH)₄} cluster in the famous metal-organic framework UiO-66. Various ligands were used to stabilize these U-oxo-clusters, such as carboxylic and N-heterocyclic ligands. Considering the correlation to the theme of this review, those metal-organic compounds with U-oxo-clusters that could be viewed as coordination polymers or metal-organic frameworks are not discussed here and other metal-organic U-oxo-clusters are selectively discussed. 27,121-126

Salmon and co-workers solvothermally synthesized a novel octanuclear U^{IV} -oxo-cluster, $(Hpy)_2[U_8L_4Cl_{10}O_4]\cdot 10py$ $(H_4L =$ N,N'-bis(3-hydroxysalicylidene)-1,2-phenylenediamine) as early as 2004 (Fig. 20). 127 This octanuclear cluster could be viewed as a combination of two parallelogram-like tetranuclear clusters linked by four µ₃-O atoms and further stabilized by four Schiff base ligands, two μ_2 -Cl, and eight terminal Cl ions. In 2016, Mazzanti and co-workers isolated a star-shaped U-oxocluster, among which five uranyl ions and a pentavalent uranium atom were linked by bridging μ_3 -O linkers to form a pentagram configuration (Fig. 21a and b). 128 The photolytic reduction of uranyl resulted in these mixed-valence U-oxo-clusters. Two similar mixed-valence U-oxo-clusters, [U(UO2)5 $(\mu_3 - O)_5 (PhCOO)_5 (Py)_7$ and $[U(UO_2)_5(\mu_3-O)_5(PhCOO)_5(Py)_7]$ [UO₂(PhCOO)₂(Py)₂]·2Py, were synthesized using a photolysis procedure with UO₂(NO₃)₂ in pyridine and the vapor diffusion of hexane into the pyridine reaction mixture. Besides, they also obtained a $\{U_{12}^{IV}U_{4}^{V}\}$ cluster, $\{[K(Py)_{2}]_{2}[K(Py)]_{2}[U_{16}O_{24}(PhCOO)_{24}]\}$ (Py)2], using a similar method (Fig. 21c and d). A similar $\{U_{12}^{IV}U_{4}^{V}\}\ \text{cluster},\ \{[K(MeCN)]_{2}[U_{16}O_{22}(OH)_{2}(PhCOO)_{24}]\}\cdot 4MeCN,$ was also reported by Mazzanti in 2011 (Fig. 21e and f). 129 In this work, they also synthesized two similar {U₁₀} clusters

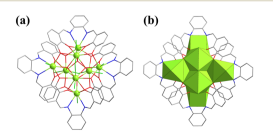


Fig. 20 (a) Ball-and-stick and (b) polyhedral views of $[U_8L_4Cl_{10}O_4]^{2-}$. Color codes: U^{IV}, light green; Cl, bright green; C, gray; O, red; N, dark

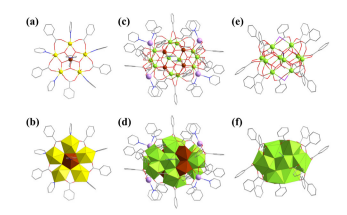


Fig. 21 Views of (a and b) $[U(UO_2)_5(\mu_3-O)_5(PhCOO)_5(Py)_7]$, (c and d) {[K $(Py)_2]_2[K(Py)]_2[U_{16}O_{24}(PhCOO)_{24}(Py)_2]$, (e and f) $[U_{10}O_8(OH)_6(PhCO_2)_{14}I_4]$ (H₂O)₂(MeCN)₂]. Color codes: U^{IV}, green; U^V, brown; U^{VI}, yellow; I, violet. Other color codes are the same as in Fig. 1.

 $[U^{III}I_3(THF)_4]$ as the using uranium $[U_{10}O_8(OH)_6(PhCO_2)_{14}I_4(H_2O)_2(MeCN)_2]$ and $[U_{10}O_8(OH)_6]$ $(PhCO_2)_{12,79}I_{3,2}(H_2O)_4(MeCN)_4]_2I\cdot 4MeCN$. The average valence of uranium atoms should be +4 according to the bond valence sum (BVS) results and charge balance.

In 2018, Knope and co-workers synthesized a {U₆O₄(OH)₄}-4-hydroxybenzoate, $U_6O_4(OH)_4(4-HB)_{12}(H_2O)_6\cdot 12H_2O$ (4-HB = 4-hydroxybenzoate), and another similar Th-cluster a nitrogen atmosphere. 130 $U_6O_4(OH)_4(4-HB)_{12}$ (H₂O)₆·12H₂O is a typical 0D U^{IV}-oxo-cluster stabilized by ligands. They comprehensively investigated the structural chemistry of these two clusters via various means. Lin and co-workers investigated the influence of counter ions (Na⁺ to Cs⁺) in the assembly of U^{IV}-oxo-sulfate clusters. 131 Na⁺ and Rb⁺ induced the formation of 0D {U₆O₄(OH)₄} clusters with different packing structures, but Cs⁺ resulted in a different trinuclear {U₃(µ₃-O)}based 2D lamellar structure. These UIV-oxo-sulfates were synthesized by the reduction of uranyl in an aqueous solution with Zn amalgam. Nyman also synthesized a $\{U_6\}$ -based cluster $(U(H_2O)_{4.5})_2U_6(OH)_4(O)_4(SO_4)_{10}(H_2O)_9$ using $U(SO_4)_2$ (Fig. 22). ¹³² Each $\{U_6O_4(OH)_4\}$ cluster was coordinated with 12 SO_4^{2-} and connected with four single UIV. Thus, a 4-connected dia-like 3D structure was constructed through the symmetrical extension of $\{U_6O_4(OH)_4\}$ clusters and single U^{IV} atoms.

In addition to the above $\{U_6O_4(OH)_4\}$ cluster, the hydrolyzation of UCl4 could also yield UIV-oxo-clusters with bridging Cl

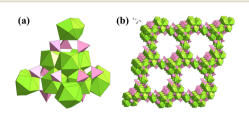


Fig. 22 Views of the (a) monomer and (b) three-dimensional structure of $(U(H_2O)_{4.5})_2U_6(OH)_4(O)_4(SO_4)_{10}(H_2O)_9$. Color codes: U^{IV} , green; S, pink; O, red.

ions. Knope synthesized a series of multinuclear uranium clusters under solvothermal conditions. 133 UCl₄ hydrolyzed into four U-oxo-clusters in a solution of THF with a little water, i.e., $[U_4Cl_{10}O_2(THF)_6(2-FA)_2]\cdot 2(THF), [U_6O_4(OH)_4(H_2O)_3(2-FA)_{12}]\cdot 7$ $[U_6O_4(OH)_4(H_2O)_2(2-FA)_{12}] \cdot 8.76(H_2O)$ $(THF)\cdot (H_2O),$ $[U_{38}Cl_{42}O_{56}(H_2O)_{20}] \cdot m(H_2O) \cdot n(THF)$ (2-FA = 2-furoic acid, THF = tetrahydrofuran). The U^{IV}-oxo-clusters were tetranuclear $[U_4(\mu_3-O)_2Cl_4]$, two hexanuclear $\{U_6O_4(OH)_4\}$, and a mixedvalence $\{U_{38}\}$ cluster (Fig. 23a). A similar $\{U_{38}\}$ cluster was reported by Loiseau with the formula $U_{38}O_{56}Cl_{18}(THF)_8(bz)_{24} \cdot 8THF$ (Hbz = benzoic acid) in 2013 (Fig. 23b). 134 This kind of {U₃₈} cluster could be viewed as a derivative of a face-centered cubic $\{U_{14}\}$ cluster. Six tetranuclear {U₄} clusters combined with the six faces of {U₁₄} to form the {U38} cluster. But it should be noted that there were no additional organic ligands in $[U_{38}Cl_{42}O_{56}(H_2O)_{20}].$ [U₃₈Cl₄₂O₅₆(H₂O)₂₀] was stabilized by bridging Cl ions and water molecules and was a pure inorganic cluster, while [U₃₈O₅₆Cl₁₈(THF)₈(bz)₂₄] was stabilized by the coordination of terminal benzoic acid and THF ligands. Loiseau also synthesized a similar {U38} cluster with the formula $[U_{38}O_{56}Cl_{40}(H_2O)_2(ipa)_{20}]\cdot(ipa)_x$ (ipa = isopropanol) in 2018 (Fig. 23c), ¹³⁵ but the terminal ligands were ionized isopropanol molecules. In 2019, Mazzanti also reported a series of U^{IV}-oxoclusters with multiple U-framework structures, including planiform $[U_6O_4(PhCOO)_{16}(Py)_4],$ fusiform $[U_{13}K_4O_{12}(OH)_4]$ $(PhCOO)_{12}Cl_{14}|Cl_2$ and $[U_{13}K_2O_9(OH)_7(PhCOO)_{12}Cl_{16}]Cl$, heartlike $[U_{16}O_{15}(OH)_8(PhCOO)_{26}(Py)_2],$ oblate spheroidal $[U_{24}O_{30}(OH)_2(PhCOO)_{30}(Py)_4Cl_4]$ cubic and $[U_{38}O_{56}Cl_{18}]$ (PhCOO)₂₂(CH₃CONH₂)₁₀]. Significantly, this {U₃₈} cluster also exhibits a similar structure to the above three $\{U_{38}\}$ clusters (Fig. 23d).

To investigate the kinetic formation of the {U₃₈} cluster, Loiseau studied the evolution of the products of this solvothermal reaction over time and found a new cationic {U₁₂} cluster, $[U_{12}O_7(OH)_4Cl_9(H_2O)_9(bz)_6(gly)_6(ox)_3]^{3+}$ (Hbz = benzoic acid, gly = glycolate, ox = oxalate), with a quadrahedron configuration (Fig. 24a and b). 137 Glycolate and oxalate were in situ

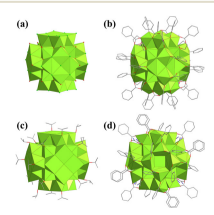


Fig. 23 Views of (a) $[U_{38}Cl_{42}O_{56}(H_2O)_{20}]$, (b) $[U_{38}O_{56}Cl_{18}(THF)_8(bz)_{24}]$, $[U_{38}O_{56}Cl_{40}(H_2O)_2(ipa)_{20}],$ and (d) $[U_{38}O_{56}Cl_{18}(PhCOO)_{22}]$ (CH₃CONH₂)₁₀]. Color codes are the same as in Fig. 20.

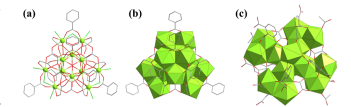
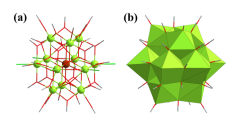


Fig. 24 (a) Ball-and-stick and (b) polyhedral $[U_{12}O_7(OH)_4Cl_9(H_2O)_9(bz)_6(gly)_6(ox)_3]^{3+}$; (c) view of $[U_{14}O_8(OH)_4Cl_8$ $(H_2O)_{16}(1,2,3-Hbtc)_8(ox)_4(ac)_4$]. Color codes are the same as in Fig. 20.

formed from THF and benzoic acid. This $\{U_{12}\}$ cluster could be viewed as a derivative of {U₆O₄(OH)₄} combining three binuclear uranium clusters at three adjacent positions of $\{U_6O_4(OH)_4\}$. All the uranium atoms in $\{U_{12}\}$ are in the +4 valence state. Additional $[(U^{VI}O_2)Cl_4]^{2-}$ balanced the charge. Subsequently, Loiseau also synthesized a {U14} cluster with the formula $[U_{14}O_8(OH)_4Cl_8(H_2O)_{16}(1,2,3-Hbtc)_8(ox)_4(ac)_4]$ (1,2,3- H_3 btc = benzene-1,2,3-tricarboxylic acid) (Fig. 24c). ¹²⁵ This {U14} cluster could also be viewed as a derivative of {U₆O₄(OH)₄}, which combined four groups of binuclear uranium clusters at the four equatorial positions of $\{U_6O_4(OH)_4\}.$

In 2021, Ikeda-Ohno and co-workers synthesized a rare Keggin-type {U₁₃} cluster, [U₁₃(CH₃O)_{35.7}Cl_{2.3}O₈], with mixedvalence U^{IV} and U^V (Fig. 25). This compound was synthesized from a solution of methanol with [U^{IV}Cl(S)-(PEBA)₃] ((S)-PEBA = (S,S)-N,N'-bis(1-phenylethyl)benzamidinate) or $[U^{IV}Cl_2(salen)_2(MeOH)_2]$ (H₂salen = N,N'-bis-(salicylidene)ethylenediamine). Single-crystal structure and BVS results revealed that the valence state of the central uranium atom in $\{U_{13}\}$ was +5 and the other 12 uranium atoms were in the +4 valence state. Ionized MeOH acted as bridging and terminal ligands to stabilize this Keggin framework. Substitutionally disordered Cl ions occupied the equatorial position. This $\{U_{13}\}$ cluster is the first U-based Keggin-type cluster, which broadens the range of

At the beginning of 2021, Wang's group reported a tabular mixed-valence enneanuclear uranium cluster, $[H_3O^+][U^V(U^{VI}O_2)_8(\mu_3-O)_6(PhCOO)_2(Py(CH_2O)_2)_4(DMF)_4]$ (Ph = phenyl, Py = pyridyl, DMF = N,N-dimethylformamide), which was dexterously synthesized from a mixed solution of DMF,



Ball-and-stick and (b) polyhedral views of $[U_{13}(CH_3O)_{35.7}Cl_{2.3}O_8]$. Color codes are the same as in Fig. 21.

Scheme 1 Schematic diagram of the synthesis and structure of $[H_3O^+][U^V(U^{VI}O_2)_8(\mu_3-O)_6(PhCOO)_2(Py(CH_2O)_2)_4(DMF)_4]$. Color codes are the same as in Fig. 21.

methanol, and ethanol with UO2(NO3)2·6H2O, macrocyclic oligoboronate, and benzoic acid (Scheme 1).139 The tetrameric borates in situ transformed into ionized pyridine-2,6-diyldimethanol ligands under solvothermal conditions, which might be the crucial factor for this synthesis. The central uranium ion was in an octahedral geometry and the valence was demonstrated to be +5 by various means. Density functional theory (DFT) and electronic spectroscopy revealed the outstanding semiconductor property of [H₃O⁺][U^V(U^{VI}O₂)₈ $(\mu_3-O)_6(PhCOO)_2(Py(CH_2O)_2)_4(DMF)_4$].

2.3 Other types of U-POMs

In addition to the above types of U-POMs, there are also other structures that are hard to classify as above. Due to the unique bonding law of uranium, fewer U-oxo-clusters are built from typical building units of POMs. Uranyl peroxide clusters and their derivatives are the most common compounds to be seen. There were several Lindqvist-type U-POMs in the early stage. 12,140 In 2001, Clark and co-workers synthesized the first Lindqvist-type uranium cluster with crystal structures, i.e., $[Cp_4(bpy)_2][U_6O_{13}]$ (Cp = 1,2,4-tBu₃C₅H₂, bpy = 2,2-bipyridine), which was stabilized by the coordination of two kinds of aromatic ligands (Fig. 26).141 The average valence of U atoms should be +5 in consideration of charge balance and magnetic susceptibility. Then, Burns reported a non-normal wheel- $\{[W_5O_{21}]_3[(U^{VI}O_2)_2(\mu-O_2)]_3\}^{30-},$ POM, (Fig. 27). 142 This compound was synthesized from a basic solu-

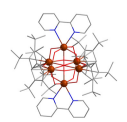


Fig. 26 View of [Cp₄(bpy)₂][U₆O₁₃]. Color codes are the same as in Fig. 21.

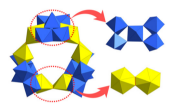


Fig. 27 View of $\{[W_5O_{21}]_3[(U^{VI}O_2)_2(\mu-O_2)]_3\}^{30-}$. Color codes are the same as in Fig. 1.

tion of UO₂(NO₃)₂, H₂O₂, LiOH, NaOH, and H₃PW₁₂O₄₀. Being different from monovacant Lindqvist- $\{W_5\}$, the $\{W_5\}$ units in $\{[W_5O_{21}]_3[(U^{VI}O_2)_2(\mu-O_2)]_3\}^{30-}$ were in a planar configuration and the central W atom was in a tetragonal pyramid geometry. Six uranyl groups were connected by bridging-O₂²⁻ two by two. Three {W₅} and three binuclear uranium clusters were linked alternately to form a triangular arrangement.

Although a series of Silverton-type U-containing polyoxomolybdates (U-POMos) were reported very early, there were still no other types of U-POMos until Nyman reported a U-POMo, i.e., $5Na[UNa(Mo_6P_4O_{31}H_7)_2]\cdot (H_2O)_n$. But there are no typical U-O-Mo bonds in this compound; this is one of the features of traditional POMs. The U4+ cations are linked by Andersonlike $\{Mo_6\}$ moieties through the connection of PO_4^{3-} . This is also the first U-POMo, except for the Silverton-type structures mentioned above.

Furthermore, Nyman also synthesized two Lindqvist-type U-containing polyoxoniobates (U-PONbs), i.e., $[((UO_2)$ $(H_2O)_3Nb_{46}(UO_2)_2O_{136}H_8(H_2O)_4]^{24-}$ and $[(Nb_7O_{22}H_2)_4]$ $(UO_2)_7(H_2O)_6]^{22-}$; these are also currently the only two U-PONbs (Fig. 28). 144 However, there is significant disorder of uranyl ions in both structures. The structure of [((UO2) $(H_2O)_3Nb_{46}(UO_2)_2O_{136}H_8(H_2O)_4]^{24-}$ could be divided into two parts, namely three ordered uranyl ions as linkers and two groups of {Nb₇} triquetrous trimers linked by {NbO₂} or disordered {UO₂}. For the other U-PONbs, two of the three kinds of uranyl ions exhibit positional disorder around a center of symmetry in $[(Nb_7O_{22}H_2)_4(UO_2)_7(H_2O)_6]^{22-}$. The occupancies

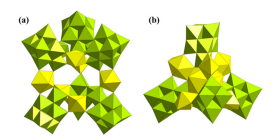


Fig. 28 Views of (a) $[((UO_2)(H_2O))_3Nb_{46}(UO_2)_2O_{136}H_8(H_2O)_4]^{24-}$ and (b) $[(Nb_7O_{22}H_2)_4(UO_2)_7(H_2O)_6]^{22-}$. Color codes are the same as in Fig. 1; Nb,

of U2 and U3 are 0.5 and 0.25, respectively. Each {Nb₇} cluster is connected with two U1, one U2, and three U3 to form a symmetric tetrahedral sandwich structure. These two U-PONbs were synthesized by a similar vapour diffusion method from EtOH diffusing into a solution containing TMA₆[Nb₁₀O₂₈] and Li₄[UO₂(O₂)₃]. The utilization of these two raw materials instead of the normal K salt of {Nb₁₀} and UO₂(NO₃)₂ may be the significant factor for the construction of the only two U-PONbs.

Besides, there is a type of uranium hydroxide cluster outside the above classification, which might be more appropriately called inorganic uranium mineral compounds, but their single-crystal structures are rarely reported. 13 Recently, a new uranium oxide hydrate with a 3D framework was obtained by Zhang and co-workers. $[U(H_2O)_2]\{[(UO_2)_{10}O_{10}(OH)_2][(UO_4)]\}$ (H2O)2] was hydrothermally synthesized from schoepite at high temperature (240 °C) (Fig. 29). 145 Single-crystal structure, BVS, and other characterization methods demonstrated the presence of mixed-valence U^{IV} and U^{VI}.

Characterization, properties, and applications

Early research on U-POMs may be limited by research conditions and only spectroscopy and electrochemistry investigations were carried out for most of the U-POMs. 146 X-ray single-crystal diffraction came into service at the end of the 20th century. 30,63,140 In addition to X-ray single-crystal diffraction, basic spectroscopy investigations were carried out for most of the U-POMs, including FT-IR, Raman, UV/vis, elec-

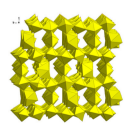


Fig. 29 View of $[U(H_2O)_2]\{[(UO_2)_{10}O_{10}(OH)_2][(UO_4)(H_2O)_2]\}$. Color codes are the same as in Fig. 1.

tronic spectra and so on.65-67 Nuclear magnetic resonance (NMR) spectroscopy of characteristic elements (P, W, etc.) was also utilized to study the structure and solution behaviors of some U-POMs. 30,34,60 Magnetic properties were investigated for some U-POMs due to the paramagnetic nature of U^{IV}/ U^{v.68,133,139,141} Magnetic susceptibility tests could also verify the valence of uranium atoms. Other normal characterization methods such as TGA and PXRD were also investigated for several U-POMs. However, it should be noted that, maybe due to the strong absorption effects in X-ray diffraction, most of the reported PXRD results for U-POMs did not show diffraction peaks as clearly as those of other POMs. 31,32 In recent years, small-angle X-ray scattering (SAXS) was used to investigate POMs in solution, especially uranyl peroxide clusters. 147 The sizes of POMs in solution could be revealed by the SAXS results and thus the free states of POMs in solution could be indicated. SAXS was also used to study the solution assembly and behavior of uranyl peroxide clusters. 148 Burns also used neutron diffraction to study the crystal structures of $Na_{44}K_6[(UO_2)_{24}(O_2)_{24}(P_2O_7)_{12}][IO_3]_2 \cdot 140H_2O$ and $\{U_{60}\}$, which revealed abundant structural information beyond that obtained from X-ray diffraction. 149,150 In addition to BVS and X-ray photoelectron spectroscopy (XPS), X-ray absorption near edge structure (XANES) spectroscopy could also help to determine the valence of uranium atoms. 139 Besides, ESI-MS is a powerful way to study species in solution and understand the assembly of complicated molecules. Burns used Raman spectroscopy and ESI-MS to study the dissolution of UO₂ in a basic solution containing XOH (X = Li, Na, K) and H₂O₂. This process was found to be linearly dependent on the concentrations of alkali, which could rival the usage of boiling HNO₃. Several uranyl peroxide clusters were formed in this process. This work could help to control the dissolution and separation of UO2 in solution. Finally, density functional theory (DFT) calculations could also help people to learn the structure of several U-POMs. 139,142,152 Perhaps the vacuum conditions of the scanning electron microscope (SEM) and transmission electron microscope (TEM) make solvent-rich U-POM crystals decompose easily; SEM and TEM technology are occasionally used to observe U-POMs. 132,133,137,153-155

In terms of properties and applications, only a few investigations were carried out in addition to magnetic studies. Firstly, early research revealed that the study of U-POWs could help people to dispose of radioactive elements because POWs exhibited outstanding thermal stability and radioresistance. Those obtained U-POWs may benefit the development of the immobilization, enrichment, and separation radionuclides. 30,146,156,157 The residue of U-POWs after hightemperature calcination could result in uranium-containing tungsten bronzes, which may provide a direction for handling radioactive materials. 158

Secondly, due to the luminescent nature of uranyl ions, U-POMs may also exhibit good luminescence properties, but there were no studies on the luminescence properties of U-POMs until recently. 41,49,62 In 2021, Kong's group systematically studied the temperature-dependent photoluminescence

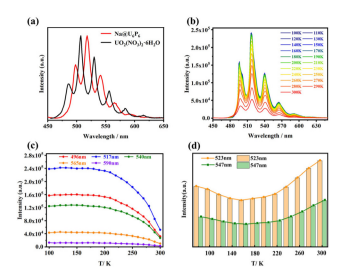


Fig. 30 (a) Solid-state photoluminescence emission spectra of Na@U₆P₆ and UO₂(NO₃)₂·6H₂O (λ_{ex} = 441 nm, normalized intensity); (b) emission map of spectra recorded from 100 to 300 K; variation trend of emission peaks depends on temperature for (c) Na@U6P6 and (d) U3. Reprinted with permission from ref. 41 and 62. Copyright 2021 American Chemical Society and 2022 Elsevier, respectively.

 $Na_{17}\{Na@[(SbW_9O_{33})_2(UO_2)_6(PO_3OH)_6]\}\cdot xH_2O$ properties (Na@U₆P₆) for the first time (Fig. 30a-c). All Na@U₆P₆ displayed a similar emission peak shape to that of UO₂(NO₃)₂·6H₂O. The solid-state photoluminescence quantum yield of Na@U6P6 is 33%. When the temperature increases from 100 to 300 K, $Na@U_6P_6$ shows a relatively stable emission from 100 to 200 K and then the emission intensity sharply decreases from 200 to 300 K. The dominant non-radiative transitions should be responsible for this phenomenon at high temperatures. Whereafter, our group and Zhang's group also studied the temperature-dependent properties of $K_{1.5}Na_{13.25}[(UO_2)_3]$ $(SeO_3)_3Na_5(H_2O)_6(SeW_6O_{21})(SeW_9O_{33})_3$] $\cdot 26H_2O (U_3)_6^{-62}$ While the solid-state photoluminescence quantum yield of U₃ is 72%. What is noteworthy is that Na@U₆P₆ and U₃ show different emission trends depending on temperature. U₃ undergoes a decrease and then increases with increasing temperature, which could result from changes in non-radiative decay and radiative decay 30d). The temperature-dependent luminescence properties of Na@U₆P₆ and U₃ demonstrate that U-POMs have great potential as luminescent materials.

Thirdly, most U-hetero-POMs should retain the catalytic characteristics of POMs and uranium compounds. Thus, U-hetero-POMs could be used as bifunctional Lewis acid-base catalysts in organic synthesis. However, there were also no studies on the catalytic activities of U-hetero-POMs until recently. Our group synthesized six U-POWs to catalyze the synthesis of sulfonyl pyrazoles and/or 3H-benzo[b][1,4]diazepines, $Na_{11}H(H_2O)_{31}[Na(UO_2)(PW_9O_{34})]_2 \cdot 7H_2O$ $(UPW_9-1),$ $Na_{12}(H_2O)_{29}[Na(UO_2)(PW_9O_{34})]_2 \cdot 7H_2O$ (UPW₉-2), $Na_{10.5}H_{3.5}$ $(H_2O)_{36}[Na(UO_2)(\alpha-SiW_9O_{34})]_2\cdot 2.5H_2O$ (USiW₉-1), $Na_{14}(H_2O)_{36}$ $[Na(UO_2)(\alpha-SiW_9O_{34})]_2\cdot 4H_2O$ $(USiW_9-2),$ $(Se_2U_2)Na_3[H_{19}]$ $(UO_2)_2(\mu_2-O)(Se_2W_{14}O_{52})_2$ $\cdot 41H_2O$ (U_2) , and $(NH_4)_{10}[H_4(SeO)_2]$

Scheme 2 Summary of the synthesis of sulfonyl pyrazoles catalyzed by UPW9-1, USiW9-2, and U2.

 $(UO_2)_2(H_2O)_2(H_2Se_2W_{14}O_{52})(Se_2W_{14}O_{52})]\cdot 66H_2O$ (Se₂U₂), which demonstrated the catalytic activities of U-POWs (Schemes 2 and 3).31,32 These catalytic systems showed advantages such as benign conditions, excellent yields, good compatibility, and atom efficiency.

Fourthly, U-oxo-clusters including uranyl peroxide clusters and other similar POMs are a broad category of compounds with a high density of uranium atoms. Therefore, the study of U-oxo-clusters may benefit the development of nuclear fuel cycle technologies and mixed oxide (MOX) nuclear fuel. 159-163

Scheme 3 Substrate scope for the synthesis of benzodiazepines catalyzed by USiW₉-2. Reaction conditions: diamines 1 (1 mmol), 2 (1 mmol), catalyst (0.5 mol%), DMC (0.1 mL) or solvent-free; isolated yields.

Besides, the ion-rich nature of uranyl peroxide clusters and other similar clusters make them good for ion storage. Yin's group recently synthesized a kind of hybrid nanocomposite for solid-state electrolytes using $\{U_{60}\}$ $(Li_{44}K_{16}[UO_2(O_2)(OH)]_{60})$ and cationic surfactants.¹⁶⁴ The cavity makes capsule-like {U₆₀} a good storage and transportation node for Li⁺. Different surfactants could regulate the microphase-separated structures of the nanocomposites from lamellar to columnar structures. This work represents a new direction for the application of uranyl peroxide clusters.

Finally, Wang's work revealed the n-type semiconductor behavior of $[H_3O^+][U^V(U^{VI}O_2)_8(\mu_3-O)_6(PhCOO)_2(Py(CH_2O)_2)_4$ (DMF)₄]. Due to the unique tabular mixed-valence structure, the central UV could act as an electron donor under X-ray radiation. $[H_3O^+][U^V(U^{VI}O_2)_8(\mu_3-O)_6(PhCOO)_2(Py(CH_2O)_2)_4(DMF)_4]$ could produce a photocurrent, which makes it a potential X-ray detecting semiconductor material. This work shows us the potential of U-oxo-cluster-based semiconductor materials due to the multivalent nature of uranium.

Summary and outlook

In summary, one could be aware that the development of U-POMs is still relatively slow despite the early study of U-POMs beginning in the 1990s. Most of the reported U-hetero-POMs and the two main categories of U-oxo-clusters were structure-driven studies. In contrast, the focus of research on mainstream POMs has developed from structure-driven research to performance-driven research. The current situation for U-POMs is that knowledge of the structural diversities and involved elements is still deficient, and the applications of U-POMs are still less explored even though there are dozens of U-POMs. In terms of numbers, although the number is relatively small, there are still dozens of U-hetero-POMs and uranyl hydroxide clusters. In contrast, U^{IV}-oxo-clusters are sporadically reported and the number is quite low. In terms of synthesis, the U-hetero-POMs are still less explored not only due to the difficulty in controlling the self-assembly process of POMs but also the limitations of the vacancy-directed synthetic strategy. Almost all the U-hetero-POMs are synthesized using polyvacant anionic precursors. There were fewer than ten cases of U-hetero-POMs in situ synthesized until recently. U-oxo-clusters reported in recent years have gradually developed from pure inorganic structures to organic-inorganic hybrid structures. However, the discovery of new topologies of uranyl peroxide clusters has gradually slowed down in recent years, which may be because of the limitations of applications due to the soluble nature of uranyl peroxide clusters and expensive costs of synthesis. Meanwhile, the much rarer U^{IV}-oxo-clusters are always synthesized using organic solvent and some ligands under solvothermal synthesis. But the organic ligands are still limited to organic phosphonic acids for uranyl peroxide clusters and a finite number of ligands for U^{IV}-oxo-clusters even if there are multiple changes to these ligands. Thus, the research prospects of U-POMs are still attractive.

From the viewpoint of synthesis and structures, research on U-POMs still exhibits great possibilities.

- (i) First, most U-hetero-POMs are synthesized by a precursor approach from an aqueous solution at room temperature. There were fewer than ten cases synthesized from simple materials, i.e., in situ methods. Thus, the utilization of in situ and hydrothermal methods may result in more unprecedented POMs with various structures.
- (ii) Second, most of the U-POMs are pure inorganic compounds crystallized from an aqueous solution. Thus, these inorganic U-POMs may exhibit considerable solubility in polar solvents, which limits their application in some fields such as catalysis and the immobilization and enrichment of radionuclides. Referring to other POMs, especially lanthanide-containing POMs, the introduction of organic ligands may make the obtained U-POMs more insoluble and diverse. Interesting properties and applications may also be discovered related to these molecules. The study of organic-inorganic hybrid U-POMs may show more possibilities and expansive prospects.
- (iii) Third, the synthesized U-hetero-POMs for now mainly focus on POWs, but other POMs are rarely involved, such as Mo, V, Nb, and Ta. Thus, it should be recognized that the study of U-POMs lags far behind other POMs, which may be limited by the synthesis method. More synthesis strategies could be applied according to other POMs. More synthesis methodologies should be introduced by referring to the synthesis of metal-organic frameworks. Synthesis strategies such as hydrothermal, solvothermal, ionothermal, microwave, ultrasonic, mechanochemical, and in situ ligand reactions should be applied more, especially for U-oxo-clusters.
- (iv) Fourth, due to the unique bonding nature of uranium atoms, the formation of uranyl peroxide clusters and metalorganic U-oxo-clusters is more common, but typical types of POMs (Keggin, Lindqvist, etc.) are hard to see. Thus, the synthesis of U-oxo-clusters with typical types of POMs still faces challenges and risks. For solvothermally synthesized UIV-oxoclusters, the introduction of more different kinds of ligands into U-oxo-clusters could result in unpredictable changes and may help to broaden the scope of U-POMs.

From the perspective of applications, research on most of the U-POMs was only based on the characterizations of the structures by spectroscopic methods. Only a few studies on U-POMs have addressed applications such as magnetism, catalysis, temperature-dependent fluorescence, and the immobilization and enrichment of radionuclides. Thus, performanceoriented research on U-POMs requires more work. Firstly, only a few cases of magnetism studies of low valent U-POMs revealed their potential as magnetic materials such as singlemolecular magnets. But this research is all superficial and deeper investigations are needed with reference to lanthanidecontaining POMs. Secondly, Kong's and our work demonstrated the U-POMs' luminescence ability. U-POMs may exhibit good potential as luminescent materials such as probes and imaging agents, as well as for application in radioactive medicine. 41,62 Thirdly, our work revealed the thermocatalytic activities of U-POWs for the synthesis of N-heterocyclic com-

pounds, and more catalytic systems should be explored. 30,31 Fourthly, the application in solid-state electrolytes and ion transport for ion-rich uranyl peroxide clusters is also a direction for traditional U-oxo-clusters. 164 Fifthly, for those mixedvalence U-oxo-clusters, the inner charge-transfer property makes them potential semiconductor materials. 139 Thus, semiconductor-related applications may be another remarkable direction for those U-oxo-clusters, such as photoelectric conversion, sensors, devices, and so on. Finally, considering the optical nature of uranium and uranyl compounds and the catalytic advantages of POMs, the application of photocatalysis may be an attractive outlet for U-POMs, especially for those mixed-valence uranium-containing U-POMs. U-POMs may show great potential in the field of photocatalysis, including photocatalytic organic synthesis, photocatalytic degradation of pollutants, photocatalytic hydrogen production, and so on.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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