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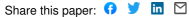
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Advances in the Engineering of Near Infrared Emitting Liquid Crystals and Copolymers, Extended Porous Frameworks, Theranostic Tools and Molecular Junctions Using Tailored Re₆ Cluster Building Blocks.

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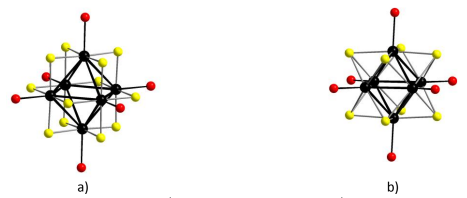
Abstract At the occasion of the fiftieth birthday of the introduction of the term 'metal atom cluster' by F. A. Cotton in inorganic chemistry, it is the good time to make a review on the advances in the engineering of molecular assemblies and nanomaterials based on octahedral Re₆ metal atom clusters. The latter exhibit unique intrinsic structural and physicochemical properties (orthogonal disposition of metallic sites that can be selectively functionalized, photoluminescence, redox, generation of singlet oxygen) that make them relevant building blocks for the structuration at the nanometric scale and functionalization of hybrid organic-inorganic materials and supramolecular frameworks. After synthesis by solid state chemistry techniques at high temperature, inorganic precursors built up on face-capped $[(Re_6Y^i_8)Y^a_6]$ cluster units (Y = chalcogen and/or halogen) can be functionalized via solution chemistry techniques or organic melts to form $[(Re_6Y^i_8)L^a_6]$ (L = CN, OH, various organic ligands...). This work reports advances in the synthesis of $[(Re_6Y^i_8)Y^a_6]$ and $[(Re_6Y^i_8)L^a_6]$ cluster units as well as on their use in the elaboration of supramolecular frameworks, nanoparticles, hybrid nanomaterials (co-polymers and liquid crystals) and active molecular junctions.

 $\textbf{Keywords} \ \text{Rhenium metal atom clusters} \cdot \text{Nanoparticles} \cdot \text{Supramolecular frameworks} \cdot \text{Copolymers and Liquid crystals} \cdot \text{Functional surfaces}$

I. Introduction.

The solid state chemistry of metal atom clusters and in particular the solid state chemistry of octahedral metal atom clusters has been extensively studied since the 60's [1-8]. This chemistry has yielded a wide library of compounds with fascinating crystal structures and properties as superconductivity [9], thermoelectricity [10], intercalation/de-intercalation [11], Mott insulating behaviors [12] with potential applications, for instance, for energy conversion or the design of Random Access Memories (RAM). Besides their physical properties in the solid state, that originate from number of valence electrons per cluster available for metal-metal bonding and to the strength of electronic interactions between cluster units, many solid state compounds and in particular octahedral cluster compounds can be used as precursors to afford discrete soluble molecular species for the structuration and the design of molecular assemblies [13], extended molecular solids [14-25], supported materials and nanomaterials [21, 26, 27] that are of interest in the fields of hydrogen storage, sensors, imaging and displays.

Octahedral nanometer-sized metallic clusters of transition elements are associated with halogen or chalcogen ligands to form edge-bridged $[(M_6L^i{}_{12})L^a{}_6]^{n-}$ and face-capped $[(M_6L^i{}_8)L^a{}_6]^{n-}$ units (a = apical, i = inner). As sketched in Fig. 1, M_6 clusters are bonded to six terminal ligands (L^a) in both kinds of units but they are edge-bridged by twelve ligands in the $[(M_6L^i{}_{12})L^a{}_6]^{n-}$ unit (M = Nb or Ta) and face-capped by eight inner ligands in the $[(M_6L^i{}_8)L^a{}_6]^{n-}$ unit (M = Mo, Mo). The intrinsic properties of M_6 cluster units depend on both the nature of the metal, the number of valence electron and the nature of L^i and L^a ligands.



 $\textbf{Fig. 1} \ \text{Representation of the edge-bridged} \ [(M_6L^i{}_{12})L^a{}_6]^{n-} \ (a) \ \text{and face-capped} \ [(M_6L^i{}_8)L^a{}_6]^{n-} \ (b) \ units$

Both $[(M_6L^i_{12})L^a_6]^{n-}$ and $[(M_6L^i_8)L^a_6]^{n-}$ exhibit a wide absorption window from UV to visible range [28-38] but only the $[(M_6L^i_8)L^a_6]^{n-}$ (M = Re, Lⁱ = S, Se or Te; M = Mo or W, Lⁱ = Cl, Br or I) cluster units exhibit

photoluminescence properties [35, 38-49]. Both units exhibit reversible electron-oxidation processes and magnetic properties when they carry unpaired valence electrons [30, 32, 35, 39, 50-62]. Compared with other metal-based solid state complexes, the added value of $[(M_6L^i_{12})L^a_6]^{n-}$ and $[(M_6L^i_8)L^a_6]^{n-}$ is that the inorganic L^a ligands can be exchanged by other L'a inorganic ones or functional organic ligands [29, 31, 63-68]. This substitution can be complete or partial leading to isomeric $[(M_6L^i_{12})L^a_xL^{'a}_{6-x}]^{n-/+}$ and $[(M_6L^i_8)L^a_xL^{'a}_{6-x}]^{n-/+}$ cluster units. Interestingly, the $(M_6L_8^i)^{x+}$ and $(M_6L_{12}^i)^{x+}$ cores can be viewed as nodes that can be selectively functionalized for their controlled assembling by supramolecular interactions via linker in one, two or three crystallographic directions in extended molecular solid. $[(M_6X_{12}^i)L_6^a]^{n-}$ cluster units (X = halogen and L = H₂O, N₃-, CN⁻ and SCN⁻ groups) [69-72] have been involved in the design of extended frameworks [73, 74], deposited as redox-active moieties on gold electrodes [75] or used, in the case of tantalum, for the structural determination of proteins [76, 77]. It is worth noting that the chemistry of [(M₆Lⁱ₈)L^a₆]ⁿ⁻ face-capped units is much more developed than that of $[(M_6X^i_{12})L^a_6]^{n-}$ edge-bridged ones owing to, probably, a higher chemical stability with higher oxidation potential compared to $[(M_6X_1^i)_2]L^a_6]^{n-}$ cluster units but also because such units exhibit additional unique photoluminescence properties, photocatalytic activity [78, 79] and capability to generate singlet oxygen [62, 80, 81]. Many recent works have been reported on the elaboration and design of Mo₆ cluster-based molecular assemblies, nanomaterials and functional surfaces [13, 20, 21, 27, 82-84]. This review focuses on the recent advances and perspectives on molecular assemblies, nanomaterials and surfaces using rhenium cluster building blocks which is actually a hot topic of research. The idea to prepare solid state compounds and to use them as precursors in the elaboration of nanomaterials is not straightforward and can be decomposed in three important chronologic steps. The first step started with the pioneering work of P. Batail and A. Perrin with the discovery of the solubility of K[(Re₆Seⁱ₅Clⁱ₃)Cl^a₆] in ethanol [85]. This property of Re₆ clusters was subsequently used for the design of charge transfer salts. Then, the second step started with the works of R. H. Holm, J. R. Long, V. E. Fedorov and Y. Sasaki consisting in the functionalization of the Re₆ clusters by donor ligands like CN, phosphine or pyridine derivatives. Indeed, a multiplicity of molecular assemblies and extended supramolecular solids or extended frameworks were synthesized and characterized. The third step started at the beginning of 2000s by the engineering of hybrid molecular compounds and nanomaterials [66, 86-92] using the Re₆ clusters as building blocks or active moieties for applications ranging from biomedicine and bio-imaging to lighting and displays. This field of research is becoming more and more interdisciplinary from the synthesis of solid state compounds, functionalization of clusters, design of multifunctional materials and evaluation of toxicity. Fifty years after the introduction by F. A. Cotton of the term metal atom cluster, this review reports advances of this interdisciplinary work consisting to use Re₆ cluster-based solid state compounds for the engineering of near infrared emitting liquid crystals and copolymers, extended porous frameworks, theranostic tools and molecular junctions. This constitutes a first step before applications in many fields ranging from telecoms, molecular electronics, displays, lighting and medicine.

II. Art of work: from solid state compounds to functional molecular assemblies

The solid state synthesis and excision reactions to form soluble scaffolds for functional molecular assemblies have been largely described earlier [36, 93-96], in the following main results are summarized (Table 1). The synthesis methods of the cluster compounds may be divided into two main groups: the high-temperature synthesis, which is carried out in closed systems (usually, in sealed evacuated tubes), and the synthesis in solution.

II.a. Synthesis of cluster precursors by high temperature and solution routes

The high-temperature reactions are extremely important for the synthesis of the $(Re_6Q^i_8)^{n+}$ cluster cores which hitherto have never been obtained in solution. In the solid state, the cluster core is stabilized by halogen and/or chalcogen to form polymeric frameworks as for instance $Re_6Q_8Br_2$ (Q = S, Se) or Re_6Te_{15} . When alkali metal salts are added to the starting mixture, the cluster core is stabilized by halogen to form discrete charged units as found for instance in the double salts $Cs_4[(Re_6Q^i_8)X^a_6]$ CsX (Q = S, Se; X = Br, I). The preparation method for Re_6 cluster-based compounds is based on the assumption that a phase thermodynamically stable under defined experimental conditions (i.e. temperature, pressure) is obtained from starting precursors taken in the appropriate ratio. Simple substances and binary compounds (for example, metal halides, metal chalcogenides, chalcogenides and halides) are used as starting compounds. For instance, polymeric $Re_6Q_8Br_2$ is prepared from elements, addition of excess of CsX loading results in formation of $Cs_4[(Re_6Q^i_8)X^a_6]$ CsX with discrete clusters. In addition to the ratio of the starting compounds in the reaction mixture, the synthesis temperature is the key point of the procedure. The standard procedure for the synthesis of a cluster chalcohalogenide can be described as follows.

First, a quartz tube is loaded with starting compounds and then evacuated and sealed. If necessary, the tube is cooled down by liquid nitrogen prior to evacuation in order to avoid the loss of volatile components (e.g. Br₂ or I₂). Afterwards, the sealed tube is placed into a furnace and heated at temperature ranging from 500°C to 900°C depending on the chemical system. The reaction time varies from several tens of hours up to several days, in order to complete the reaction. Such high temperature reactions produce exclusively valence electron-precise counts irrespectively to stoichiometry or reaction conditions. The most widely used starting precursors are $Re_6Q_8Br_2$ (Q = S, Se), Re_6Te_{15} , $Cs_4[(Re_6Q_8^i)X^a_6]$ CsX (Q = S, Se; X = Br, I). The subsequent reaction of Re₆Q₈Br₂ and Re₆Te₁₅ with KCN, KOH or KNCS yields hexasubstituted units with the [(Re₆Qi₈)La₆]⁴⁻ formula (L = CN, OH or NCS). Let us note that the reaction of Re₆Se₈Br₂ with molten KOH at 280 °C results in the formation of a (Re₆Seⁱ₄Oⁱ₄)²⁺ cluster core with ordered ligands. Four positions of one face of the Se₄O₄ cube inscribing the Re₆ cluster are occupied exclusively by Se atoms and the four O atoms lie in the opposite face [97]. Cyanocomplexes are formed by reactions of ReS2 (or ReSe2) with KCN leading to the formation of two unusual chalcocyanide compounds, $K_4[(Re_6S^i_8)S^{a-a}_{4/2}(CN)^a_2]$ and $K_4[(Re_6S^i_8)(Se_2)^{a-a}_{2/2}(CN)^a_4]$, that contain bridging ligands S²⁻ and (Se₂)²⁻ together with terminal CN⁻ ligands [98]. A chain polymer Cs₄[(Re₆Sⁱ₈)S^{a-} $^{a}_{2/2}(CN)^{a}_{4}$] which is the next member of a series $M_{4}[(Re_{6}S^{i}_{8})S^{a-a}_{(6-2x)/2}(CN)^{a}_{2x}]$, where M is an alkali metal, was obtained with high yield by high temperature reaction of ReS₂ with KCN in presence of CsCl excess [99]. Four $mixed\ ligand\ octahedral\ cluster\ complexes\ K[(Re_6S_8^i)F^a_3(H_2O)^a_3]\cdot 7H_2O,\ H_3O[(Re_6S_8^i)F^a_3(H_2O)^a_3]\cdot 7H_2O\ and\ number of the complexes o$ $[(Re_6Q^i_8)F^a_2(H_2O)^a_4]\cdot 12H_2O$ (Q = S or Se), have been synthesized using the reaction of corresponding rhenium chalcobromide complexes $K_4[(Re_6S_8^i)Br_6^a] \cdot 2H_2O$ and $Cs_3[(Re_6S_8^i)Br_6^a] \cdot H_2O$ with molten KHF₂ and further crystallization form aqueous solutions under different conditions [100].

II.b. Solution chemistry

The solubility of $[(Re_6Q_8)X_{6}]^{n-}$ cluster units is governed by the charge of the units and the nature of counter cations. $[(Re_6Q_8^i)X_6^a]^{n-}$ (X = halogen) cluster units associated with alkaline earth ions are soluble in alcohol and/or in water. The functionalization by organic moieties requires the solubilization of Re₆ clusters in organic media. Indeed, after dissolution of inorganic precursors in aqueous solutions, the precipitation of $[(Re_6O_8^i)X^a_6]^{n-1}$ units with tetrabutylammonium $((n-C_4H_9)_4N)^+$ organic cations affords $((n-C_4H_9)_4N)_n[(Re_6Q^i_8)X^a_6]$ (n=4 for S, n=4 for S)= 3 for Se) precursors, from which a wide variety of hybrid $[(Re_6Q^i_8)L^a_6]^{2+}$ units based on a $(Re_6Q^i_8)^{2+}$ inorganic cluster core and functional organic moieties utilized as apical ligands can be prepared (L = N- and P-donor ligands). Indeed, the more straightforward functionalization method is the direct reaction of $(TBA)_3[(Re_6Q^i_8)Cl^a_6]$ (TBA = tetrabutyl ammonium) with pyridine and functional pyridine to form bi- and trifunctional species [37, 101]. It has even been demonstrated that monofunctional [(Re₆Qi₈)(Cla₅(4-phenylpyridine)^a³⁻ species is obtained by reaction between (TBA)₄[(Re₆Qⁱ₈)Cl^a₆] and 4-phenyl-pyridine in a MeCN forced by photoirradiation of the solution [102]. In order to obtain higher substituted cluster units, the use of phosphine derivatives is preferred. Usually reactions with phosphine are more complete [66, 86, 87, 94, 103], hexasubstituted $[(Re_6Q_8^i)L^a_6]^{2+}$ functional units were prepared from $[(Re_6Q_8^i)L^a_6]^{3-}$ units (Q = S, Se) after the prior removal of iodine apical ligands from the cluster core by reaction with silver salt in order to facilitate the apical ligand substitution. Partially substituted $[(Re_6Q_8^i)(PEt_3)^a_{6-x}I_x^a]^{(2-x)+}$ units were synthesized by the direct reaction of $[(Re_6Q_8^i)I_6^a]^{3-}$ units with PEt₃, leading to a set of $[(Re_6Q_8^i)(PEt_3)^a_{6-x})I_x^a]^{(2-x)+}$ anionic units with different x values and possible isomeric units. The Re-P bonds being very strong, functional $[(Re_6Q^i_8)(PEt_3)^a_{6-}]$ $_x)L_x^a|^{2+}$ units were obtained by reactions of $[(Re_6Q_8^i)(PEt_3)^a_{6-x}I_x^a]^{(2-x)+}$ with various L in the presence of Ag⁺ in order to remove the apical iodine atoms without destroying the Re-PPh₃ bond. Beyond different types of functional Re₆ supramolecular units, this technique afforded original supramolecular network based on hydrogen-bonding by grafting, for instance, isonicotinamide on the cluster. It turns out that the lower reactivity of the $(Re_6S^i_8)^{2+/3+}$ cluster cores compared to that of $(Re_6Se^i_8)^{2+/3+}$ leads to a smaller set of supramolecular assemblies with sulfur than with selenium. All these derivatives have been widely studied experimentally and theoretically for their spectroscopic and photophysical properties [38-42, 104-108].

II.c. Mixing Solid state and solution chemistries, design of functional Re₆ cluster units containing CN and OH apical ligands: $[(Re_6Q^i_8)(OH)^a_{6-x}(CN)^a_x]^{4-}$

In a similar way to what has been observed with Re–P bonds for phosphine ligands, the Re–CN bond is very stable at room temperature. [(Re $_6Q^i_8$)L $_6^a$] n - (L = CN $^-$, OH $^-$, NCS $^-$, N $_3$ $^-$) cluster units associated with alkaline earth ions are readily soluble in water [48, 49, 109-120]. Thanks to the bidentate nature of CN $^-$ group, the [(Re $_6Q^i_8$)(CN) $_6^a$] n - (Q = S, Se, Te) cluster units have been widely used for the elaboration of low dimensional cluster framework by co-crystallization via CN with transition metal cations. The hydroxo species are interesting

since the OH groups react with acid to form water molecules [49, 117]. They also react with carboxylic acids to form corresponding carboxylate cluster complexes [41, 42, 88, 121]. Indeed, based on the fact that the $[(Re_6Q^i_8)(CN)^a_6]^{4-}$ cluster unit is chemically inert and that $[(Re_6Q^i_8)(OH)^a_6]^{4-}$ contains six OH groups capable to react with an acid, the possibility to block apical sites with CN groups will enable to tailor building blocks with controlled number and positions of reactive OH sites. As stressed above, the solid state reaction of Re₆Q₈Br₂ with KCN or KOH [110, 115, 122] affords hexasubstituted units with the [(Re₆Qi₈)La₆]⁴⁻ formula and solid state $reaction \ of \ [(Re_6S^i_8)(CN)_4S^{a-a}_{2/2}]^{4-} \ with \ KOH \ affords \ \textit{trans-}[(Re_6S^i_8)(CN)^a_4(OH)^a_2]^{4-} \ anionic \ cluster \ unit \ [118].$ Protonation of $[(Re_6Q^i_8)(OH)^a_6]^{4-}$ cluster units leads to the formation of $[(Re_6Q^i_8)(H_2O)^a_2(OH)^a_4]^{2-}$ and [(Re₆Qⁱ₈)(H₂O)^a₄(OH)^a₂]⁰ species with low solubility owing to strong hydrogen bonding between clusters [117]. It is worth noting that compounds based on $[(Re_6Q^i_8)(H_2O)^a_2(OH)^a_4]^{2-}$ units like $K_2[(Re_6S^i_8)(H_2O)^a_2(OH)^a_4] 2H_2O$ exhibit extended 3D hydrogen-bonding networks [117]. Protonation of the [(Re₆Sⁱ₈)(CN)^a₆]⁴⁻ cluster complex was not observed under similar experimental conditions up to pH 1. On the other hand, the $K_4[(Re_6S_8^i)(CN)^a_4(OH)^a_2]$ is very soluble in water at pH > 10.5, a pH increase in range 5.5 < pH < 9 enables the formation of $[(Re_6S_8^i)(CN)^a_4(OH)^a(H_2O)]^{3-}$ and finally $[(Re_6S_8^i)(CN)^a_4(H_2O)^a_2]^{2-}$ at pH < 3.5 [49]. The latter constitute thus useful building blocks for the elaboration of monofunctional and trans-bifunctional molecular assemblies by reaction with functional acids in aqueous solutions. A relevant strategy has been developed to obtain organic solvent soluble [(Re₆Qⁱ₈)L^a₆] clusters in high yield without the use of tetrabutyl ammonium salts as developed in the case of rhenium chalcohalogenides. It consists in the reaction between $[(Re_6Q^i_8)(OH)^a_6]^{4-}$ with an excess of tert-butylpyridine (tbp) in water leading to the precipitation of the neutral $[(Re_6Q^i_8)(tbp)^a_4(OH)^a_2]^0$ cluster units. It turns out that $[(Re_6Q^i_8)(tbp)^a_4(OH)^a_2]^0$ is soluble in organic solvents and that the trans-isomer can be separated from the cis-isomer by two successive recrystallizations in CH₂Cl₂-ether mixtures [41]. The added value of all these species is that they are not air sensitive and that they can be produced in a large scale.

 $\textbf{Table 1} \ \ \text{Synthetic approaches to the common} \ \ (\text{Re}_6\text{Q}_8)^{2+}\text{-cluster-core-based precursors used for preparation of cluster containing functional materials} \ \ \text{and hybrid materials}$

Complex	Starting materials	Reaction conditions	Reference
$[(Re_6S_8)Cl_6]^{4-}$	1) Re, S, ReCl ₅ and MCl	1) 900 °C, ~116 h	[123, 124]
	(M = Tl or Cs)	2) 850 °C, 7 days	
	2) Re, S, S ₂ Cl ₂ and KCl	050 °G 1001	F2 (1251
$[(Re_6Q_8)X_6]^{4-}$	Re, Q, X_2 and CsX	850 °C, 100 h	[36, 125]
(Q = S, X = Br or I;			
Q = Se, X = I)	(D 0)(OH) 14 G W 14W	Deiling on annual solution 10 min	[115]
$[(Re_6Q_8)X_6]^{4-}$	$[(Re_6Q_8)(OH)_6]^{4-}$, CsX and HX	Boiling an aqueous solution, 10 min	[115]
(Q = S, X = Br;			
Q = Se, X = Cl	F/D - O \W 14-		[26 55]
$[(Re_6Q_8)X_6]^{3-}$	$[(Re_6Q_8)X_6]^{4-}$	oxidation of [(Re ₆ Q ₈ X ₆] ⁴⁻ by SOCl ₂ , SOBr ₂ or H ₂ SO ₄ in an acetonitrile or a CH ₂ Cl ₂	[36, 55]
(Q = S, X = Cl, Br or I;		solution	
Q = Se, X = I)		635 °C, 25 h	[111]
[(Re6Q8)(CN)6]4- (Q = S or Se)	cesium salt of $[(Re_6S_8)Br_6]^{4-}$ or	033 C, 23 II	[111]
	[(Re ₆ Se ₈)I ₆] ⁴ and NaCN	500 °C 24 b	[100]
$[(Re_6S_8)(CN)_6]^4$	Re ₆ Te ₁₅ and KSCN	500 °C, 24 h	[109]
$[(Re_6Q_8)(CN)_6]^{4-}$	Re ₆ S ₈ Br ₂ , Re ₆ Se ₈ Br ₂ or Re ₆ Te ₁₅	600 °C, 48 h	[110, 112,
(Q = S, Se or Te)	and KCN	600 °G 241	122]
$[(Re_6Q_8)(CN)_6]^{4-}$	1) Re ₆ Te ₁₅ , Se and NaCN	600 °C, 24 h	[93]
(Q = Se or Te)	2) Re ₆ Te ₁₅ and NaCN	1) 200 °C 11	[112]
$[(Re_6Q_8)(NCS)_6]^{4-}$	1) [TBA] ₃ [(Re ₆ S ₈)Cl ₆] and	1) 200 °C, 1 h	[113]
(Q = S or Se)	KSCN	2) reflux of a chlorobenzene-DMF (2:1 v/v) solution under Ar, 12 h	
	2) $[(Re_6Se_8)(MeCN)_6]^{2+}$ and	3) reaction in a mechanochemical reactor at	
	SCN	room temperature, 20h	
	3) Cs ₂ [(Re ₆ Se ₈)Br ₄] and KSCN	·	506 041
$[(Re_6S_8)(PEt_3)_xX_{6-x}]^{x-4}$	$[(Re_6S_8)Cl_6]^{4-}$ or $[(Re_6S_8)Br_6]^{3-}$	reflux of a DMF solution under N ₂	[86, 94]
(X = Cl or Br)	and PEt ₃		F1.023
$[(Re_6Se_8)(PEt_3)_xI_{6-x}]^{x-4}$	$[(Re_6Se_8)I_6]^{3-}$ and PEt ₃	reflux of an acetonitrile or a DMF solution under N ₂	[103]
[(Re6Se8)(PEt3)6-	$[(Re_6Se_8)(PEt_3)_xI_{6-x}]^{x-4}$ and Ag^+	reaction in a mixture of CH ₂ Cl ₂ with MeCN,	[96, 103,
$_{x}(\mathbf{L})_{x}]^{2+}$		DMF, DMSO or pyridine, respectively, at	126]
(X = MeCN, DMF,		room temperature	
DMSO or pyridine)			F40 1141
$[(Re_6Q_8)(N_3)_6]^{4-}$	$[(Re_6S_8)Br_6]^{4-}$ or	reaction in an aqueous solution at room	[48, 114]
(Q = S or Se)	[(Re ₆ Se ₈)(OH) ₆] ⁴⁻ and N ₃ ⁻	temperature, 24 h	
$[(Re_6Q_8)(OH)_6]^{4-}$	Re ₆ Q ₈ Br ₂ and KOH or CsOH	$280 ^{\circ}\text{C} \text{ (for Q = S) or } 200\text{-}230 ^{\circ}\text{C (for Q = Se)},$	[115, 116]
(Q = S or Se)		30 min	54.4 - 3
$[(Re_6Q_8)(H_2O)_n(OH)_{6-}]$	$[(Re_6Q_8)(OH)_6]^{4-}$ and H ⁺	reaction in an aqueous solution at room	[117]
n] n =4		temperature	
(Q = S or Se, n = 0-6)	5 (D. 19.) (OTD. 14		[100]
$[(Re_6Q_8)F_{6-n}(H_2O)_n]^{n-4}$	$[(Re_6S_8)(OH)_6]^{4-}$ or	reaction with molten KHF ₂ (270-300 °C, 2 h)	[100]
(Q = S or Se, n = 3 or 4)	$[(Re_6Se_8)(OH)_6]^{3-}$, F ⁻ and H ₂ O	followed by recrystallization from an aqueous solution	
$[(Re_6S_8)(RCOO)_6]^{4-}$	$[(Re_6S_8)(OH)_6]^{4-}$ and $RCOO^-$	reaction in an aqueous solution at room	[42, 121]
$(R = H \text{ or } CH_3)$		temperature	
$[(Re_6Q_8)(TBP)_4(OH)_2]$	$[(Re_6Q_8)(OH)_6]^{4-}$ and TBP	reflux of an aqueous solution, 2 days	[41]
(Q = S or Se, TBP = p			
tert-butylpyridine)			
$[(Re_6S_8)(CN)_4(OH)_2]^{4-}$	$Cs_4[Re_6S_9(CN)_4]$ and KOH	reflux of an aqueous solution, 24 h	[118].
$[(Re_6Se_8)(CN)_4(OH)_2]^{4-}$	$[(Re_6Q_8)(OH)_6]^{4-}$ and CN^-	reflux of an aqueous solution, 2 h	[119]

In the following, we will focus on new perspectives based on recent work dealing on in-situ synthesis of supramolecular compounds using organic melts and on hydroxo, cyano and cyano-hydro complexes as building blocks for the elaboration of supramolecular frameworks, nanoparticles, liquid crystals and functional surfaces.

III. Description of Photophysical Properties

As described above, during the last two decades, a large number of hexarhenium chalcogenide cluster complexes with various apical ligand environments have been synthesized and their structures and properties were evaluated. Following are summarized some important features about electronic structures, redox and luminescence properties.

Electronic structures and redox properties

The electronic structures of $[(M_6L^i_8)L^a_6]$ clusters were studied by several authors using quantum chemical calculations. The molecular diagram of $[(M_6L^i_8)L^a_6]$ units – considering an O_h symmetry – shows a set of twelve metal-metal bonding molecular orbitals $(a_{1g}, t_{1u}, t_{2g}, t_{2u}, and e_g)$. The population of these twelve bonding molecular orbitals (MO) corresponding to metal–metal bonds in a metal cluster Re_6 leads to an optimum number of valence electrons, the so-called valence electron concentration (VEC) of 24. This corresponds to a rhenium ion with a d^4 electron configuration, found in $[(Re_6Q^i_8)Y^a_6]^{4-}$ (Q = S or Se; Y = halogen, OH or CN) with a $(Re_6Q^i_8)^{2+}$ cluster core. One-electron oxidation can be easily performed in solutions of molecular rhenium complexes. This corresponds to the removal of one electron from the HOMO level leading to a VEC = 23. Redox potentials depend on both the nature inner and apical ligands and vary from 0.065 V for $[(Re_6Te^i_8)(CN)^a_6]^{4-}$ to 1.01 V for $[(Re_6Se^i_8)(\mu\text{-dpph})^a_2I^a_4]^{2+}$ moieties [65]. This oxidation provides compounds with paramagnetic cluster. Further oxidation of Re_6 cluster was demonstrated in electrochemical experiments but up to now 22 electrons moieties were not isolated as individual phases.

Luminescence properties

By many examples, it was shown, that complexes of the general formula $[(Re_6Q_8^i)L^a_6]^{4-}$ (Q = S or Se; L = halogen ion, CN-, NCS-, N₃-, OH-/H₂O, anions of carboxylic acids, pyridine and phosphine derivatives, etc.) with the $(Re_6Q_8^i)^{2+}$ core having 24 valence electrons in the solid state and in solutions emit luminescence in visible and near-infrared regions (NIR) upon ultraviolet or blue light excitation with emission lifetimes in the microsecond range [38-42, 44, 48, 49, 81, 86, 88, 101, 102, 113, 117, 119, 121, 127, 128]. The spectroscopic and photophysical properties of these complexes have been extensively studied experimentally (see references above) and by theoretical calculations [34, 38, 42, 48, 104-108, 115, 119, 129]. On the basis of the fairly long emission lifetimes and the insensitive nature of the emission spectrum to the type of solvents employed, it has been proposed that the emitting excited state of the hexarhenium (III) complex is a spin-triplet type and involves orbitals that are primarily localized on the $(Re_6Q_{i_8})^{2+}$ core. In addition to these experimental observations, theoretical studies of the excited state have demonstrated that the lowest-energy unoccupied molecular orbitals (LUMOs) are primarily localized on the $(Re_6Q_{8})^{2+}$ core [38, 42, 44, 101, 108, 130, 131]. Among the known Re₆ clusters, [(Re₆Seⁱ₈)(DMSO)^a₆]²⁺ is the strongest luminophore with the emission quantum yield and lifetime in CH₂Cl₂ being 0.238 and 22.4 μs, respectively [127]. Systematical studies of spectroscopic and photophysical properties (emission maximum wavelengths λ_{em} , quantum yields Φ_{em} and lifetimes τ_{em}) of hexarhenium cluster complexes allowed to reveal some dependencies and make important conclusions, namely:

- 1) as a rule, the $(Re_6Se^i_8)^{2+}$ -based complexes show a stronger emission than those based on $(Re_6S^i_8)^{2+}$ possessing the same apical ligand environment [38, 40, 127];
- 2) influence of the nature of apical ligands:
 - in the series $[(Re_6S^i_8)X^a_6]^{4-}$ $(X = Cl^-, Br^- \text{ or } I^-)$ the emission maximum shifts to a longer wave-length while Φ_{em} and τ_{em} decrease as the terminal halide ions become heavier [39];
 - in CH_2Cl_2 solutions complexes with oxygen- or nitrogen-based organic ligands (DMSO, DMF, pyridine, acetonitrile) as well as the chalcocyanides $[(Re_6Q^i_8)(CN)^a_6]^{4-}$ deliver maximal quantum yields and the longest lifetimes and tend to the higher-energy emission, while clusters having six soft, polarizable ligands, such as iodide or bromide, emit with less intensity, shorter lifetimes, and redder luminescence. Phosphine ligation (PEt₃) also diminishes Φ_{em} [38, 127];
 - in-depth study of the series of complexes with different N-heteroaromatic ligands demonstrated that the emission is effectively quenched as the result of $(Re_6Q^i_8)^{2+}$ -to-ligand charge transfer (MLCT) if the ligand (such as 4,4'-bipyridine or pyrazine) has free nitrogen sites [44, 101];
 - mixed-ligand cyanohydroxo complexes trans-[(Re₆Qi₈)(CN)a₄(OH)a₂]⁴⁻ (Q = S or Se) in aqueous solutions exhibit the emission spectra and photophysical characteristics appreciably different from hexahydroxo clusters [(Re₆Qi₈)(OH)a₆]⁴⁻ and at the same time very similar to the corresponding hexacyano ones [(Re₆Qi₈)(CN)a₆]⁴⁻ [49, 119];

- the acidification of an aqueous solution with $[(Re_6Q^i_8)(OH)^a_6]^{4-}$ (Q=S or Se) ions leads to transformation of the anionic cluster to $[(Re_6Q^i_8)(H_2O)^a_x(OH)^a_{6-x}]^{x-4}$ through the protonation of the apical hydroxo ligands accompanied by red shift of λ_{em} and irregular changes in Φ_{em} and τ_{em} values [117]. Whereas, spectroscopic and photophysical properties of cyanohydroxo complexes trans- $[(Re_6Q^i_8)(CN)^a_4(OH)^a_2]^{4-}$ and hexacyano ones $[(Re_6Q^i_8)(CN)^a_6]^{4-}$ are completely insensitive to pH of their aqueous solutions, demonstrating that the luminescence properties of trans- $[(Re_6Q^i_8)(CN)^a_4(OH)^a_{2-x}(H_2O)^a_x]^{x-4}$ are mainly defined by the CN^- ligands and not appreciably affected by the OH^- or H_2O ligand(s) [49, 119];
- 3) the quantum yield and lifetimes significantly depend on a solvent while the emission maximum remains the same. In particular, a complex in a deaerated aqueous solution shows much lower Φ_{em} and τ_{em} values than in a deaerated organic solvent that can be explained by interaction of water molecules with the inner chalcogenide ligands providing more efficient vibrational decay pathway of the excited state [40];
- 4) the dependences of photophysical properties on both the inner and apical ligands were discussed in terms of the energy gap law, by which the natural logarithm of the nonradiative decay rate constant (k_{nr}) of a cluster is inversely proportional to the emission maximum energy. Since it was found that the emission lifetime of the cluster is determined mainly by the k_{nr} value, the energy gap dependence of k_{nr} demonstrates that the emission lifetime of a cluster can be predicted approximately from the emission energy [38, 40, 113, 127];
- 5) solid-state emission lifetimes and quantum yields usually occupy a similar range at room temperature [38, 42, 44, 48, 88, 113, 119, 121]. Low-temperature luminescence experiments indicate a progressive red shift of the emission maximum upon cooling, with an increase in the excited-state lifetime and the emission quantum yield [38, 44, 89, 113];
- 6) the long lifetimes, large Stokes shifts, and excited-state quenching by O_2 indicate the spin-triplet nature of the luminescent excited state of the cluster complexes, i.e. the change in spin multiplicity is involved in the electronic transitions [38, 39, 86, 89]. Note that to date there is only one described example of a luminescent $(Re_6Te^i_8)^{2+}$ -based complex, namely the photoluminescence of the complex $[(Re_6Te^i_8)(CN)^a_6]^{4-}$ has been studied in acetonitrile and aqueous solutions and it was shown, that the telluride cluster shows much weaker and short-lived emission comparing to $[(Re_6S^i_8)(CN)^a_6]^{4-}$ and $[(Re_6Se^i_8)(CN)^a_6]^{4-}$ [40]. The lack of data on luminescence of telluride hexarhenium cluster complexes with other apical ligands is a consequence of the fact that the tellurocyanide complex is still the unique representative of soluble $(Re_6Te^i_8)^{2+}$ -based complex.

The luminescence properties of Re_6 cluster complexes provided an incentive to establish their potential applications [61, 81, 88, 90, 132]. Synthetic flexibility of the apical ligand environment affords a range of clusters for developing innovative luminescent materials. The quenching of the phosphorescence by O_2 suggests applications to optical sensor technology and singlet oxygen generation that can be used, for example, in the photodynamic therapy of cancer.

IV. Molten salt approach: Toward the one pot synthesis of supramolecular frameworks

It was found that reactions between chalcohalogenide hexarhenium cluster complexes and some molten organic compounds which can coordinate to rhenium lead to substitution of some apical halogen atoms by the organic units [43, 129, 133-136]. Such approach allows to produce cluster complexes with functional organic ligand environment in high yield. The reactions in a molten organic medium may result in two types of cluster complexes: partially substituted neutral complexes [43, 129, 135, 136] or hexasubstituted cluster cations [133, 134]. However, so far there is no evidence on the nature of the driving force at the origin of formation of neutral (partially substituted) or cationic (hexasubstituted) complexes.

The cations $[(Re_6Q^i_8)(3,5\text{-Me}_2PzH)^a_6]^{2+}$ (Fig. 2a) were obtained by the reaction of rhenium thiobromide $[(Re_6S^i_8)Br^a_6]^{4-}$ or selenobromide $[(Re_6S^i_8)Br^a_6]^{3-}$ complex with molten 3,5-dimethylpyrazole (3,5-Me₂PzH) and separated as salts $[(Re_6Q^i_8)(3,5\text{-Me}_2PzH)^a_6]Br_2\cdot 2(3,5\text{-Me}_2PzH)$. In the crystal structures each cluster cation is accompanied by two Br^- anions and additionally by two molecules of 3,5-Me₂PzH. An extensive system of hydrogen bonds joins these ionic fragments and molecules 3,5-Me₂PzH to one another. The interactions of the given type include $N\text{-H}\cdots N$ bonds between coordinated and uncoordinated molecules of 3,5-Me₂PzH; $N\text{-H}\cdots Br$ bonds between coordinated molecules of 3,5-Me₂PzH and Br^- anions and $N\text{-H}\cdots Br$ bonds between uncoordinated molecules of 3,5-Me₂PzH and Br^- anions [134].

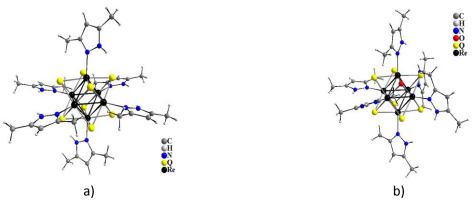
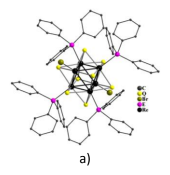
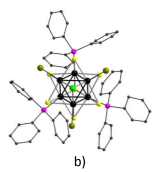


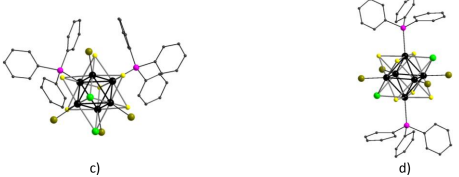
Fig. 2 Structure of cationic cluster complexes $[(Re_6Q^i_8)(3,5-Me_2PzH)^a_6]^{2+}$ (a) and $[(Re_6Q^i_7O^i)(3,5-Me_2PzH)^a_6]^{2+}$ (b)

Two octahedral rhenium cluster complexes $[(Re_6Q^i_7O^i)(3,5-Me2PzH)^a_6]^{2+}$ (Fig. 2b) were obtained also from molten 3,5-Me₂PzH by the reaction with rhenium chalcobromides $[(Re_6Q^i_7Br^i)Br^a_6]^{3-}$ (Q = S, Se) possessing the mixed-ligand cluster core $(Re_6Q^i_7Br^i)^{3+}$. It was found that during the reaction substitution of six apical bromide ligands by 3,5-Me₂PzH molecules accompanied by substitution of the inner μ_3 -Br ligand by oxygen giving cluster cores $(Re_6Q^i_7O^i)^{2+}$. Note that there is no disorder in the present structures: seven corner positions of the Q_7O cube are occupied exclusively by S or Se atoms and one by an O atom. There are two types of Br atoms in the structure: one bromine atom is connected by hydrogen bonds to one cluster whereas the other one is bridged between two cluster cations to form extended chains along the a-axes [133].

A series of partially substituted neutral complexes $[(Re_6Q^i_{8-n}Br^i_n)(EPh_3)^a_{4-n}Br^a_{n+2}]$ (Q = S, n = 0, 1 or 2; Q = Se, n = 0 or 1) were obtained by the reaction between anionic chalcobromide complexes $[(Re_6Q^i_{8-n}Br^i_n)Br^a_6]^{4-n}$ and molten EPh₃ (E = P, As or Sb) [129, 135]. More precisely, reactions of $[(Re_6Q^i_8)Br^a_6]^{3-}$ (Q = S, x = 4; Q = Se, x = 3) and $[(Re_6Q^i_7Br^i)Br^a_6]^{3-}$ and molten EPh₃ resulted in *trans*- $[(Re_6Q^i_8)(EPh_3)^a_4Br^a_2]$ and fac- $[(Re_6Q^i_7Br^i)(EPh_3)^a_3Br^a_3]$, respectively (Fig. 3a,b); $[(Re_6S^i_6Br^i_2)Br^a_6]^{2-}$ and molten PPh₃ produced a mixture *cis*- $[(Re_6S^i_6Br^i_2)(PPh_3)^a_2Br^a_4]$ and trans- $[(Re_6S^i_6Br^i_2)(PPh_3)^a_2Br^a_4]$ (Fig. 3c,d), that were separated by column chromatography. The interesting feature of these structures is the ligand ordering in mixed chalcobromide cluster cores $(Re_6Q^i_7Br^i)^{3+}$ and $(Re_6S^i_6Br^i_2)^{4+}$. Note that the neutral complexes with mixed-ligand apical ligand environment were obtained in reactions with a large excess of EPh₃. It was expected that the reactions result in the substitution of all terminal bromide ligands in the starting cluster compounds by EPh₃ ligands similarly to reactions with an excess of molten 3,5-dimethylpyrazole described above. However, complexes obtained in the reactions with EPh₃ have only four, three, or even two terminal EPh₃ ligands depending on the composition (and consequently, the charge) of cluster core. Such results were explained by the preferred formation of neutral molecular compounds in this system. No remarkable interactions were found between the molecular complexes in all these crystal structures of cluster complexes with EPh₃ ligands.







 $\textbf{Fig. 3} \ \ Structure \ \ of \ neutral \ \ cluster \ \ complexes \ \ \textit{trans-}[(Re_6Q^i_8)(EPh_3)^a_4Br^a_2] \ \ (a), \ \textit{fac-}[(Re_6Q^i_7Br^i)(EPh_3)^a_3Br^a_3] \ \ (b), \ \textit{cis-}[(Re_6S^i_6Br^i_2)(PPh_3)^a_2Br^a_4] \ \ (d)$

By the example of reaction with molten pyrazine it was shown that functional Re₆ cluster units can be obtained by a direct interaction of cluster chalcohalogenides with molten organic ligands based on organic ring containing nitrogen atoms [43]. Such one pot synthesis enables an easier preparation of functional units and do not need a preliminary functionalization of the cluster precursor before self-assembling from solutions as shown in the strategies developed by R. H. Holm, Z. Zheng and T. Yoshimura who oxidized parent chalcohalogenide cluster complexes [37, 96, 101, 103]. It turns out that the number of substituted apical ligands depends on the charge of the unit in the starting precursor. The easy formation of the stable CsBr inorganic salt in the organic melt must be the key point of this reaction. The limiting factor of the technique is the thermal stability of organic ligands that can decompose before the exchange occurs. Among organic ligands stable at mild temperatures, bidentate linkers such as, for instance, bipyridine or pyrazine constitute relevant candidates to build coordination polymers with low dimensionalities or supramolecular frameworks. One of the actual hot topics is the research of porous neutral supramolecular frameworks. A possible strategy is the reaction of cluster unit leading to more than two substituted apical sites. It is clear that substitution of one halide ligand by a functional organic moiety can lead to the formation of dimers, while two substituted complexes should lead to the formation of chains. Thus, we can suppose that cluster units with charges of 3- or 4- should lead to the formation of supramolecular frameworks by relevant choice of ligands. In this frame, reactions of [(Re₆Qⁱ₇Brⁱ)Br^a₆]³⁻ with molten pyrazine were investigated. After the optimization procedure, it turned out that reactions at 140 °C for 2 days in sealed glass tubes afforded very interesting supramolecular compounds built up from octahedral rhenium cluster complexes fac-[(Re₆Qi₇Bri)(pyz)^a₃Bra₃] (Q = S, Se) (Fig. 4a). Despite their bidentate nature, each pyrazine is bonded to only one Re₆ cluster. Indeed, this finding evidenced that the idea to link cluster units together via simple bidentate ligands containing aromatic groups is counter balanced by the formation of strong supramolecular interactions (i.e. hydrogen bonding and π - π stacking). But at the end, fortunately the structures of fac-[(Re₆Qⁱ₇Brⁱ)(pyz)^a₃Br^a₃]·xH₂O have porous structures containing hollow tubular channels (Fig. 4b). The (Re₆Qⁱ₇Brⁱ)³⁺ cluster core is built up from a Re₆ cluster lying in a cube of ligands formed by seven Q's and one Br. It is worth noting that, in the structures of starting salts Cs₃[(Re₆Qⁱ₇Brⁱ)Br^a₆]·H₂O the seven Q's and Br are randomly distributed on the eight inner positions of the anionic cluster complex. The presence of only one Br atom among the inner ligands generates the local C_{3V} symmetry for the [(Re₆Qⁱ₇Br)ⁱBr₆]³⁻ unit and does not enable the formation of isomers. Thus, owing to the orientational disorder, the average apparent symmetry of the [(Re₆Qⁱ₇Brⁱ)Br^a₆]³⁻ unit in Cs₃[(Re₆Qⁱ₇Brⁱ)Br^a₆]·H₂O deduced from X-ray single crystal diffraction analysis is close to O_h . In fac-[(Re₆Qi₇Bri)(pyz)a₃Bra₃] xH₂O, the [(Re₆Qi₇Bri)(pyz)a₃Bra₃]⁰ complex is not submitted to an orientational disorder, and the local C_{3V} of the unit symmetry is preserved. The ordering of μ_3 -Br atoms in the fac-[(Re₆Qⁱ₇Brⁱ)(pyz)^a₃Br^a₃]·xH₂O hybrid structure is driven by the selective substitution of three apical bromines by three pyrazine groups. One of the explanations of this selective ligand coordination is the specific distribution of electron density on the rhenium atoms in the starting precursor. Indeed, owing to the metal-to-ligand charge transfers, charge balances, and the stronger Re-Qi bond compared to the Re-Bri one, the three rhenium atoms bonded to four μ_3 -chalcogen ligands have a more positive charge than the three rhenium atoms bonded to three μ_3 -chalcogen and one μ_3 -halogen atom. The result is that the three Re-Br^a bonds for which Re atoms are exclusively bonded to inner μ_3 -S are weaker than the three opposite Re-Br^a ones favoring their substitution by the neutral pyrazine ligands. This affords the fac isomer of a neutral trisubstituted hybrid unit with an ambivalent inorganic/organic character. The self-assembly of the latter building blocks leads to the formation of a unique structure in which the blocks are held together by halogen/chalcogen van der Waals contacts found in pure inorganic solids along with π - π stacking and hydrogen bonds found in organic solids. Each cluster interacts with three adjacent ones through π - π stacking interactions as well as C-H···N bonds between pyrazine rings,

building a zigzag hexagonal layer of clusters. The cluster layers fit together according to an AA'A sequence. Interactions between layers occur via (*i*) van der Waals contacts between μ_3 -Qⁱ and μ_3 -Brⁱ along the *c* axis; (*ii*) van der Waals contacts between μ_3 -Q and between μ_3 -Br and μ_3 -Br of three adjacent cluster units lying in the same plane but belonging to two successive layers; and (*iii*) N–H··· N, C–H··· Br^a, and C–H··· Brⁱ bonds. Note that the inner chalcogen atoms are not involved in C–H··· Qⁱ nor N–H··· Sⁱ bonds, whereas all bromine ligands are involved in hydrogen bonds. Indeed, the formation of a robust hydrogen network involving μ_3 -Brⁱ must be a driving force that contributes to the Q/Br ordering. The AA'A packing of layers generates large tubular channels parallel to the *c* axis that are arranged according to a honeycomb disposition imposed by the trigonal symmetry of the fac-[(Re₆Qⁱ₇Brⁱ)(pyz)^a₃Br^a₃] cluster unit. The effective aperture of the channel is found roughly equal to 6.10 and 6.43 Å for Q = S and Se, respectively. The total free volume of the unit cell was found to be 43 %, whereas the free volume available for solvent molecules is 29 %. H₂O molecules can easily be adsorbed through C–H···Qⁱ, or O–H···Qⁱ, or O–H···Br^a bonds. They have no effect in the stabilization of the structure and can be removed without collapsing the structures.

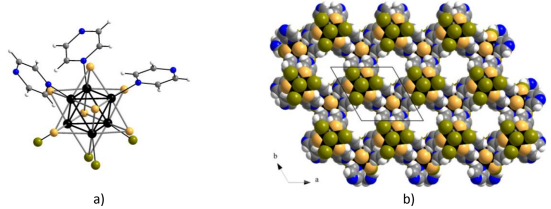


Fig. 4 Structure of neutral cluster complexes fac-[(Re₆Qⁱ₇Brⁱ)(pyz)^a₃Br^a₃] (a) and projection of structure of fac-[(Re₆Qⁱ₇Brⁱ)(pyz)^a₃Br^a₃]·xH₂O along the c direction (b).

This strategy offers now new perspectives in the design of porous framework containing metal atom clusters. The choice of the precursor determines the level of substitution and the choice of ligand determines the type of supramolecular interactions whilst their size should enable to control the aperture of the pores.

V. Hybrid organic-inorganic nanocomposites and functional surfaces

V.a. Hybrid copolymers

Despite visible emission and interesting properties for displays and lighting, the direct use of solid state compounds for applications is limited because of their ceramic nature (brittle, low plasticity). One solution to overcome this drawback is the sintering processing under inert atmosphere however such method drastically limits the size of the targeted devices. Another strategy is the use of hybrid organic-inorganic technologies that associate an organic matrix with photochemically active and reinforcing inorganic moieties. Hybrid materials obtained by this way have enhanced properties and combine the processability of the organic matrix with the intrinsic properties of the inorganic moieties. For luminescent cluster containing hybrid copolymers, it is important to find the best cluster/polymer weight ratio in order to optimize the optical properties without altering significantly the host matrix processability. Recently, luminescent clusters containing co-polymers have demonstrated their high applicative potential in technologies needing deep red emission. For instance, it has been showed that the luminescence of the trivalent Er(III) luminescence used in C-band telecom technologies is sensitized by metal atom clusters [137]. A major difficulty in the elaboration of hybrid polymers, is to avoid the segregation of the organic and inorganic phases. One possible strategy to tackle this problem is to maximize the interactions between both parts of the material by modifying the inorganic moieties with polymerizable ligands and copolymerize them with organic monomers. Two examples have been described up to now in which materials have been obtained by copolymerization of functionalized Re₆ clusters with an organic matrix. The first one is a copolymer resulting from the copolymerization of [(Re₆Seⁱ₈)(PEt₃)^a₅(4-vinylpyridine)^a]²⁺ with styrene [138]. Although a mono functionalized cluster seems the best way to introduce clusters without altering significantly the mechanical properties of the organic host because monofunctionalization does not imply cross linking, the major drawbacks of this technique are that it requests the synthesis of non-stable Re₆ intermediates,

tedious purifications of pentafunctionalized building blocks leading finally to an overall synthetic scheme with poor yield. Moreover, optical properties of such copolymer were not reported. This fact could be eventually explained by the presence of five phosphine ligands around the Re_6 core that have the unfortunate ability to drastically decrease the luminescence ability of the metallic Re_6 core. To prepare new luminescent materials directed towards optical applications, poly-(methyl methacrylate) (PMMA) is a relevant matrix because of its excellent optical properties (i.e., transparency from the near-UV to the near-IR regions), good mechanical and electrical properties, thermal stability, water resistance, and easy shaping. The embedding of Re_6 clusters was performed by a prior functionalization with methacrylate (MA) by reaction of $[(Re_6Q^i_8)(tbp)^a_4(OH)^a_2]^0$ cluster units with methacrylic acid to form $[(Re_6Q^i_8)(tbp)^a_4(MA)^a_2]$.

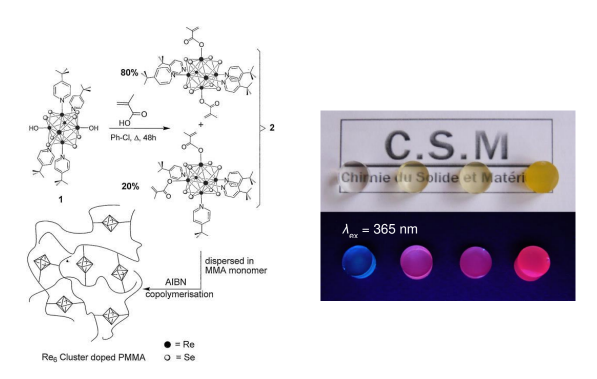


Fig. 5 a) Synthetic pathway for the preparation of the Re₆-doped PMMA hybrid material. b) Digital photographs of the PMMA and Re₆-PMMA pellets obtained by bulk synthesis. Top: under normal daylight; bottom: under UV irradiation at λ_{ex} = 365 nm. The weight percentage of clusters increases from left to right: 0, 0.025, 0.05, and 0.1 wt%, respectively. Reprinted from [88] with permission from John Wiley and Sons.

Copolymers with different cluster content were then prepared to optimize the synergy between the processability and transparency of the organic matrix and the high luminescence of the Re₆ cluster core. Several pellets were obtained by a radical polymerization process for 24 h at 60 °C from a mixture of cluster precursors (from 0.025 to 0.1 wt%), methyl methacrylate and AIBN (azobisisobutyronitrile) as initiator (Fig. 5a). Owing to the low solubility of $[(Re_6Q_8^i)(tbp)^a_4(MA)^a_2]$ in neat MMA, the maximum amount of the cluster in the pellet was limited to 0.1 wt%. Higher cluster content resulting in the loss of pellets transparency. As the Re₆ cluster/MMA ratio was kept very low, the processability of the hybrid copolymer appeared very close to that of neat PMMA with an enhanced thermal stability as demonstrated by TGA analysis. Co-polymers emit bright-red phosphorescence (Fig. 5b) upon excitation anywhere in the absorption band of the cluster (300-550 nm). Luminescence spectrum of copolymers revealed the typical spectra of the cluster core with a broad and structureless band that extended from 600 nm to more than 950 nm, with the maximum wavelength λ_{em} centered at around 710 nm. Absolute emission quantum yield was measured to be about 0.07. It is worth noting that this value is one of the highest photoluminescence quantum yields reported so far for red-NIR luminescent polymers doped with inorganic compounds and does not degrade after several months of ageing. Encapsulation of clusters in this particular polymeric matrix prevents (or minimize) quenching of photoluminescence by oxygen as it is observed in solution. Beyond the possibility to be shaped in bulk form by usual melting processes of polymers technologies, these hybrid materials are soluble in organic solvents which allows their processing into thin films with techniques used for large surfaces preparation like casting, dip coating, or spin coating. Compare to other

red photobleaching proof emitters, Re₆ cluster are not sensitive to their coordination environment as found for rare earths ions and they do not exhibit toxic behavior like quantum dots.

V.b. Liquid crystals

Metal-containing liquid crystals, the so-called metallomesogens, combine the unique properties of anisotropic fluids with the specific properties of metals (e.g. geometry of coordination, optic, electronic, magnetic). Clustomesogens associate mesomorphism with the unique properties of metal atom clusters [139, 140]. The added value of clustomesogens, compared to metallomesogens is that their luminescence properties are not influenced by their supramolecular organization and only poorly by their surrounding ligands. This peculiar point is of interest since known properties can be directly imported in the desired hybrid matrices. The development of such a class of nanomaterials should offer great potential in the design of new electricity-to-light energy conversion systems, optically based sensors, and displays. Two approaches have been developed to obtain clustomesogens: a covalent approach and an ionic one.

The covalent approach is based on the covalent anchoring of promesogenic moieties containing organic ligands onto the clusters which leads to hybrid building blocks able of self-assembling. The covalent approach is well developed in the case of molybdenum. Some preliminary results have been reported in the case of Re₆ clusters. The strategy was to functionalize $[(Re_6Q^i_8)(tbp)^a_4(OH)^a_2]^0$ cluster by two gallate where methoxy groups have been replaced by an octyloxy chains. Luminescence experiments ($\lambda_{ex} = 450$ nm) showed a intense emission spectrum centered at 720 nm. Irradiation at $\lambda_{ex} = 405$ nm of trans-functionalized clusters induced a bright red luminescence. A promising birefringent texture could be observed at 25 °C by polarised optical microscopy. However, compounds decompose around 250 °C before reaching the clearing temperature.

The ionic approach is based on the pairing *via* Coulomb interactions of anionic clusters with promesogenic moieties containing organic cations [61]. Indeed, $[(Re_6Se^i_8)(CN)^a_6]^{n-}$ (n = 3, 4) cluster units were paired with appropriate amphiphilic ammonium cations (Fig. 6). These inorganic units are particularly relevant because, thanks to cyano ligands that prevent any ligands exchange, they are chemically very stable and exhibit a low reversible one electron oxidation potential. The latter enables to switch reversibly from an orange colored form (n = 4, VEC = 24) with red-NIR luminescence properties, to a magnetic green colored species (n = 3, VEC = 23). The oxidation-reduction process can be easily implemented by both electrochemical and chemical oxidations. In this frame $(Kat)_3[(Re_6Se^i_8)(CN)^a_6]$ and $(Kat)_4[(Re_6Se^i_8)(CN)^a_6]$ were obtained by pairing $[(Re_6Se^i_8)(CN)^a_6]^{3-}$ and $[(Re_6Se^i_8)(CN)^a_6]^{4-}$ with dialkyldimethylammonium counter cations (denoted as Kat) bearing cyanobiphenyloxy terminated alkyl chains.

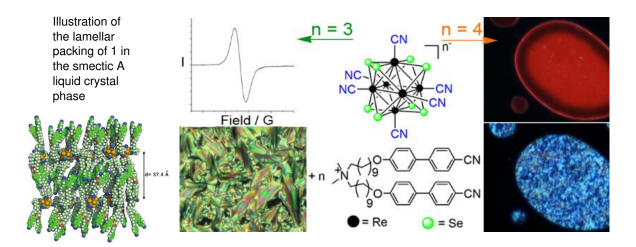


Fig. 6 The pairing of $[(Re_6Se^i_8)(CN)^a_6]^{3-}$ with Kat leads to the formation of smectic liquid crystal phase with a paramagnetic behavior and the $[(Re_6Se^i_8)(CN)^a_6]^{4-}$ with Kat leads to the formation of diamagnetic luminescent phase with liquid crystal properties. Reprinted from [61] with permission from ACS.

The reaction pairing simply consists in a precipitation by addition of a hot ethanol solution of KatBr in a water solution of $[(Re_6Se^i_8)(CN)^a_6]^{n-}$. Both hybrid compounds exhibit a lyotropic behavior with organic solvent such as chloroform: $(Kat)_3[(Re_6Se^i_8)(CN)^a_6]$ shows a texture of nematic type containing broad schlieren features

along with large homeotropic areas, while (Kat)₄[(Re₆Seⁱ₈)(CN)^a₆] exhibits a marbled texture also of nematic type. From the thermotropic liquid crystal point of view, (Kat)₄[(Re₆Seⁱ₈)(CN)^a₆] does not exhibit clear mesomorphic properties but (Kat)₃[(Re₆Seⁱ₈)(CN)^a₆] self-organizes smoothly in a layered phase of Smectic A type between 40°C and 140°C. The intrinsic properties of the molecular inorganic cluster are preserved within the mesomorphic material. It is therefore possible to switch reversibly this LC material from a green-colored magnetic form to a bright red-NIR luminescent one. These compounds, because of their low clearing temperature, can be integrated and annealed directly in devices. Yet that the concept of such class of material is demonstrated, the endless range of possibilities offered by organic chemists to design particular mesogenic cations allowing a fine-tuning of the final material mesomorphic behavior, associated with the versatility of the inorganic clusters (charge, functions) obtained *via* high temperature solid-state synthesis, open new and fascinating perspectives in the design of easy processable functional material containing transition metal clusters.

V.c. Functional surfaces

One successful strategy to design functional devices consists in the immobilization onto surfaces of nanometerscaled building blocks endowed with specific properties. Many approaches can be developed. Indeed, for optical or catalytic properties and even hydrogen storage, metal atom clusters can be immobilized onto a surface like graphene or incorporated in metal organic frameworks. Photocatalytic properties are thus driven by irradiation and depend on the used wavelength. In order to obtain electrically addressable and switchable functional devices (e.g. optical or magnetic arrays, biochemical sensors with electrical detection, or hybrid junctions for charge storage components), active building blocks must be integrated on to conducting surfaces. Many research work has been devoted to the design of molecular memory cells or junctions consisting in the immobilization of redoxactive molecules (e.g., metal-complexed porphyrins, ferrocene, etc.) including metal atom clusters on conducting surfaces. The strategy that has been developed for the anchoring of Mo₆Xi₈La₆ cluster unit on surfaces was based on the synthesis and manipulation of [(Mo₆Xⁱ₈)(CF₃SO₃)^a₆]²⁻ unstable intermediates. The surface grafting consisted in the reaction of [(Mo₆Xⁱ₈)(CF₃SO₃)^a₆]²⁻ with a pyridine-terminated alkyl monolayer-modified silicon semiconducting surface which had been previously prepared from an oxide free, hydrogen-terminated silicon (Si-H). Thanks to the lability of the CF₃SO₃ groups, the cluster was immobilized through a strong covalent bond between the cluster and the nitrogen atom from the pyridine unit endcapping the alkyl chain. After the anchoring 1 step, the modified surface was then dipped into neat pyridine in order to replace the remaining CF₃SO₃ groups by pyridine rings and obtain a stable surface and not reacting with water vapour. For the immobilization of Re₆ clusters and thanks to stability and the reactivity of the [(Re₆Qi₈)(tbp)^a₄(OH)^a₂]⁰ with carboxylic acids, a more straightforward strategy was developed. Indeed, Re₆-functionalized surfaces were obtained by simple acido-basic reaction of [(Re₆Qⁱ₈)(tbp)^a₄(OH)^a₂]⁰ with a Si-H surface modified by an alkyl monolayer terminated by carboxylic groups (Fig. 7). After reaction in refluxing chlorobenzene, carboxylate groups were formed which led to the anchoring of the cluster on the surface through strong covalent Re₆-O bonds. This simple water elimination reaction enabled the formation of a stable organic chain/cluster interface. This grafting technique greatly simplifies the cluster immobilization procedure as compared with that used previously for [(Mo₆Xⁱ₈)L^a₆] cluster units. Interestingly, the surface coverage of the metal cluster could be finely controlled by preparing mixed monolayers containing reactive COOH-terminated alkyl chains diluted with inert alkyl chains with a predefined surface chain ratio. Using mixed dodecyl/undecanoic acid monolayers, the surface coverage of $[(Re_6Q_8^i)(tbp)^a_4(OH)^a_2]^0$ cluster units could be varied from 1×10^{13} to 6×10^{13} cm⁻² [141]. Different surface characterization techniques, such as scanning tunneling microscopy (STM, Fig 7), spectroscopic ellipsometry, and quantitative X-ray photoemission spectroscopy (XPS), were used to demonstrate the covalent immobilization of Re₆ clusters. Furthermore, the cluster integrity within the monolayer was confirmed by its vibrational Raman signature. Charge transport characteristics were evaluated by measuring the current intensity flowing through the functional monolayer upon the application of a voltage between a soft top contact (mercury droplet) and the underlying silicon surface. The as formed Hg/(Re₆Seⁱ₈)-OML-Si(111) molecular junctions (OML = organic chain monolayer) with the highest ($Re_6Se_8^i$) molecular coverage showed current-voltage (I - V) characteristics different from those observed for classical Schottky diodes. The following three main features were observed: (i) a small rectification factor; (ii) a decrease in the conductance G(V) by more than one decade. The observation of the signature of the metal cluster immobilization in G(V) characteristics required a molecular coverage on the order of 4×10^{13} cm⁻² for Re₆-cluster-based junctions.

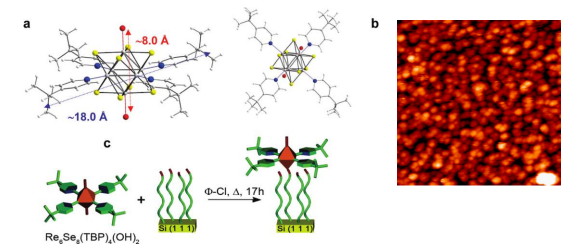


Fig. 7 Left, (a) Representation of the trans-[(Re₆Seⁱ₈)(TBP)^a₄(OH)^a₂] cluster unit. Selenium, rhenium from the Re₆ clusters, and nitrogen atoms are represented as yellow, grey, and blue balls, respectively. (b) Representation of the same units according to a different orientation. (c) Schematic representation of the (Re₆Seⁱ₈)²⁺ cluster immobilization on Si(111) through an acido-basic reaction with carboxylic acid headgroups endcapping an alkyl chain. Right, STM image of a cluster functionalized surface evidencing the homogeneous distribution of clusters onto the surface. Reprinted from [141] with permission from ACS.

Moreover, from scanning tunneling spectroscopy (STS), it has been found that the HOMO-LUMO energy gap increased from roughly 2 eV to 4 eV after the immobilization step. Therefore, this grafting strategy is particular attractive for controlling the electronic properties of the metal cluster-terminated molecular junctions. Moreover, through a judicious choice of functional ligands substituting the immobilized cluster units, it becomes also possible to develop redox-active interfaces endowed with versatile magnetic, luminescent and catalytic properties.

VI. Toward theranostic tools based on silica nanoparticles

 $[(Re_6Q_8^i)L^a_6]$ clusters units emit in the red and near-infrared (NIR) regions, which constitute a very interesting emitting window for biotechnology since it corresponds to the window of minimum absorption of tissues [142]. Moreover, because they combine the important properties of perfectly reproducible nanosize, photostability, and NIR emission with other properties like the generation of singlet oxygen and do not show acute toxic effects, Re₆ clusters may offer complementary assets to the luminophores nowadays developed for bioimaging and biolabeling technologies. They are mainly based on organic dyes, inorganic quantum dots, or lanthanide based nanocrystals. Several recent results evidence the increasing interest of rhenium atom clusters in theranostic [81, 132, 143, 144]. Beyond interesting labelling properties and thanks to the generation of oxygen singlet, the Re₆ clusters are relevant candidates for photodynamic therapy. Moreover, it has been demonstrated that the [(Re₆Seⁱ₈)I^a₆]³⁻ cluster anionic units in certain conditions selectively increase tumor cell death, leaving nontumoral cells unaffected. Indeed, Re6 clusters may be useful for cancer diagnostics, localization of tumors, and may enable the observation of tumor regression through phosphorescence [132]. Last but not least, the transport and delivering of [(Re₆Qi₈)(OH)a₆]⁴⁻ cluster units can be achieved by sugar-decorated dendritic nanocarriers and the cellular uptake can be considerably improved by the functionalization with an amphiphilic diblock copolymer [89]. Indeed different systems can be imagined and designed and not only for theranostic. In this frame, the versatility of silica nanoparticles is very relevant for the elaboration of multifunctional systems and in particular nanoparticles. Because they can support very different kind of functional nanoobjects, they found numerous applications in photonics, catalysis or biotechnologies [145, 146]. Beyond their non-toxicity, their surface can be functionalized by molecules of interest in biology [147]. Luminescent cluster-based monodispersed nanoparticles prepared by water in oil microemulsion or Stöber processes have been reported [46, 90, 148-151]. Additional properties have been obtained by the inclusion of maghemite at the core of the nanoparticle or by the growth of gold nanoparticles at the surface of silica [152, 153]. Re₆-cluster based silica nanoparticles have been obtained by embedding cationic phosphine-terminated rhenium chalcogenide clusters

[(Re₆Seⁱ₈)(Et₃P)^a₅I^a]I, [(Re₆Sⁱ₈)(Et₃P)^a₅Br^a]Br, [(Re₆Seⁱ₈)(Bu₃P)^a₅I^a]I, and [(Re₆Sⁱ₈)(Bu₃P)^a₅Br^a]Br in silica nanospheres in a one-pot, base-catalyzed hydrolysis in acetonitrile [121]. On the basis of their ability to generate singlet oxygen, they are potential candidates for photodynamic therapy and for other applications. In such nanoparticles, it appears that interactions between cationic cluster units (i.e. [(Re₆Seⁱ₈)(Et₃P)^a₅I^a]⁺, [(Re₆Seⁱ₈)(Et₃P)^a₅I^a]⁺, and [(Re₆Sⁱ₈)(Bu₃P)^a₅Br^a]⁺) and the anionic deprotonated silanol groups occur. Another strategy is the direct synthesis of Re₆ cluster-silica nanoparticles directly from water soluble precursors: K₄[(Re₆Qⁱ₈)(OH)^a₆] (Q = S or Se), Cs₄[(Re₆Sⁱ₈)Br^a₆], and Cs_{1.68}K_{2.32}[(Re₆Sⁱ₈)(CN)^a₄(OH)^a₂] [90]. Indeed, the A₄[(Re₆Qⁱ₈)L^a₆]@SiO₂ nanoparticles were prepared through a microemulsion process adapted from a previously reported synthesis for the encapsulation of hexamolybdenum clusters in silica nanoparticles (Fig. 8).

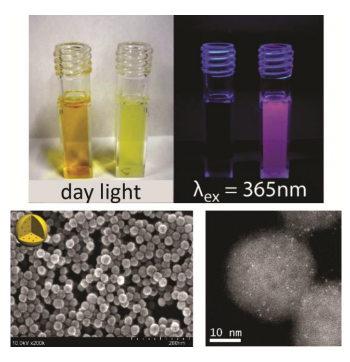


Fig. 8 Top, aqueous solution of $[(Re_6S^i_8)(OH)^a_6]^{4-}$ clusters (a) before and (b) after embedding in silica nanoparticles, under daylight (left picture) and under UV irradiation at 365 nm (right picture). Down-left, FE-SEM pictures of (a) $K_4[(Re_6S^i_8)(OH)^a_6]@SiO_2$ (inset: schematic representation of dispersed clusters inside the nanoparticles). Down-right, ADF-STEM pictures of $K_4[(Re_6S^i_8)(OH)^a_6]@SiO_2$ nanoparticles with discrete cluster units (bright spots). Reprinted from [90] with permission from ACS.

To generate the microemulsion, heptane (oil phase) was first mixed with Brij30 (surfactant), and then an aqueous ammonia solution and an aqueous cluster sol were slowly added to the mixture. The aqueous cluster sol was obtained simply by solubilization of the cluster precursor in water. Once the microemulsion became a clear, stable solution, the silica precursor (tetraethoxysilane, TEOS) was added for the synthesis of the silica shell that occurred directly inside the aqueous nanodroplets already containing the cluster units. After 3 days, the microemulsion was destabilized with ethanol and the nanoparticles were recovered after washing. Resulting nanoparticles could be either redispersed in solution (water or ethanol) or dried in ambient air. The A₄[(Re₆Qi₈)La₆]@SiO₂ nanoparticles showed that they all exhibited a perfectly spherical, monodisperse morphology with an average diameter of 30 nm for the synthesis conditions. The encapsulation of the $[(Re_6Q_8^i)(OH)^a_6]^{4-}$ (Q = Se or S) and $[(Re_6S_8^i)(CN)^a_4(OH)^a_2]^{4-}$ cluster units in the silica matrix resulted in a slight shift of the emission maxima after embedding, but the shape of the emission spectrum remained unchanged (Figure 5a-c). The emission maximum shifting, ranging from 10 to 20 nm, to higher wavelengths between solidstate precursors and the nanoparticles must be due to some perturbations of the molecular orbital diagram of $[(Re_6Q_8)L_6]^{4-}$ units after embedding. One could suspect a slight decrease in the energy gap between excited and fundamental states. Note that the use of $K_4[(Re_6Q_8^i)(OH)^a_6]$ (Q = S or Se) or $Cs_4[(Re_6S_8^i)Br_6^a]$ as starting precursor leads to K₄[(Re₆Qi₈)(OH)^a₆]@SiO₂ and Cs₄[(Re₆Qi₈)(OH)^a₆]@SiO₂ meaning that during the reaction, apical bromine atoms are substituted by OH groups. By analogy with the work on the encapsulation of [(Mo₆Brⁱ₈)Br^a₆]²⁻ units in silica where the substitution of bromine apical bromine by hydroxo groups occur, two kinds of interactions occur with silica. The first one is an interaction by the formation of hydrogen bonds

between OH of the cluster and the silanol groups of the silica and the second one is an interaction by covalent bond between cluster units and the silica matrix. These Re₆ cluster-based nanoparticles have the luminescent properties of the cluster precursor. It means that the luminescent properties of the cluster from the core of the nanoparticles are not quenched by oxygen. Moreover, since the luminescence intensity of nanoparticles decreases until a minimum in the presence of oxygen, it means that the silica is porous and clusters near to the surface of silica are quenched by oxygen and thus generate singlet oxygen. Based on these results and the fact that Re₆ clusters are not significantly toxic and that they exhibit antitumoral properties, Re₆ cluster based silica nanoparticles should be relevant tools for theranostic applications.

VII. Conclusions

Fifty years after the definition of the term "metal atom cluster" by F. A. Cotton, this review showed the step by step evolution of the rhenium metal atom cluster chemistry toward an interdisciplinary field of research from the synthesis of cluster by solid state chemistry, their functionalization by solution chemistry or in organic melts to their use as building block for the elaboration of molecular assemblies, extended molecular frameworks and nanomaterials. It showed in particular all the different strategies that have been used in order to take advantage of their unique properties originating from metal-metal bonding to obtain nanostructured and processable functional materials. We thus believe that the three first steps of the development of metal atom cluster chemistry and in particular the third step dealing with functional materials described in this review will be followed by a fourth step that should be focused on the development and commercialization of devices and nanomaterials built up from transition metal atom clusters in the fields of lighting, display, green chemistry or health. It should not only concern rhenium chemistry but all nanosized transition metal atom clusters.

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