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Rational (supra)molecular design and catalytic activity of cage-like Cu₄-based phenylsilsesquioxanes†

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An extended (*i.e.*, 19 distinct species) family of cage-like Cu_4 -phenylsilsesquioxanes allowed us to accentuate the general regularities behind their structural organization. Influencing factors, namely the (i) size of external alkali metal ions (from Li to Cs) and (ii) nature of bridging linkers (including the smallest possible ones, like a water molecule) on the self-assembly/supramolecular assembly of such Cu_4 -building blocks have been thoroughly explored. A Cu_4K_4 -based complex has been evaluated as a precatalyst in the oxidation of alkanes (cyclohexane, n-heptane, methylcyclohexane) and alcohols. The experimental evidence that radical species participate in the oxidation of alkanes is provided.

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Introduction

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The ability of metallocomplexes based on silsesquioxane [RSiO_{1.5}]_n ligand matrices to form extraordinary cage molecular geometries (cage-like metallosilsesquioxanes CLMSs) is well documented.¹ Enormous variety of compositions and structures of CLMSs endow them with an impressive list of properties and applications. CLMSs may be applied for the design of molecular magnets² (including examples of spin glass³ and single molecule magnets⁴). Recent results have pointed out the rich photophysics of CLMSs,⁵ with the lanthanide-based family of such complexes being especially attractive, holding promise for energy transfer,⁶ remote temperature sensing,⁶ or thin-film optoelectronics.⁸ For prospective materials, CLMSs were applied for the creation of intumescent objects,⁰ anodes,¹⁰ functional coordination polymers,¹¹ calcined derivatives,¹² and fungicides.¹³ Catalytic properties of

CLMSs attracted considerable attention, ¹⁴ including recent results on the Chan-Lam coupling, 15 formation of bio-inspired ethers, 16 oxidation of volatile organic compounds, 17 and deacetalization/ deketalization-Knoevenagel reactions. 18 Further design of CLMSs as potential catalysts towards reactions of high demand, especially functionalization of hydrocarbons, 19 remains an actual scientific task. Regarding the synthetic approaches to CLMSs, two main tactics exist. The first one indicates the involvement of the prebuilt silsesquioxane ligands (mostly Si7-based cubane triol). 1b,e,g,2d,6,12b,16,18 Alternative approach relies on the *in situ* formation of silsesquioxane ligands starting from the simple reactant, e.g., PhSi(OMe)₃. 1k,2b,c,3-5,7,8 This latter approach (self-assembly of ligand just prior self-assembly of metallocomplex) provides considerable flexibility of CLMS structures. In this paper, we discuss the self-assembly features of CLMSs with Cu₄ derivatives as an example. A record-holding large number of tightly related CLMSs (19 individual island-like complexes or coordination polymers) were examined in terms of the role played by (i) solvating ligands (linkers) and (ii) external ions (of alkali metal) in the molecular/supramolecular structure formation. We report on the catalytic properties of a representative Cu4K4-CLMS in the peroxide oxidation of alkanes with hydrogen peroxide and alcohols with tert-butylhydroperoxide.

Results and discussion

Synthesis and structure

A specific feature of heterometallic CLMSs simultaneously bearing transition and alkali metal ions indicates that transition metal ions are located inside the cage while alkali metal

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ions are located outside the cage. 1f,h Due to these external positions, several examples of CLMS' coordination polymers assembled via alkali metal-enabled contacts have been designed.1k These coordination polymers (Chart 1) reveal either crown ether contacts via alkali metal ions leading to (i) porous MOF (Cu₆K₂-compound²⁰) or (ii) 1D extended (Cu₈Cs₂compound²¹) structures.

Below, we present our investigation of the applicability of the alkali metal ions (Li-Na-K-Rb-Cs) for the purposeful aggregation of CLMSs' coordination polymers. As the transition metal sites, copper(II) ions have been chosen. To obtain the corresponding heterometallic compounds, a well-established scheme, including alkaline hydrolysis of PhSi(OMe)₃ with MOH²² was applied. In the second stage, intermediate [PhSi(O)OM]_n phenylsiloxanolate species formed in situ interact with a source of copper centres (CuCl₂). Due to the lowest ionic radii among alkali metals, a high "inter-cage efficacy" in terms of CLMS assembly is hardly expected for lithium. Indeed, the first run of this synthetic scheme (Fig. 1, top), with LiOH picked as a starting alkali reagent, afforded an islandlike compound $\{[(Ph_{12}Si_{12}O_{24})Li_4Cu_4(EtOH)_8]\}\cdot\{[(Ph_{12}Si_{12}O_{24})$ $Li_4Cu_4(EtOH)_6(H_2O)$ (1, 70% yield). Compound 1 (Fig. 1, bottom) adopts a cage-like structural type, with a large twelvemembered cyclic silsesquioxane ligand embracing the central copper-containing core, while lithium ions are located in external positions. In summary, metal centers form two orthogonal linear fragments with the Li···Cu···Li composition (Fig. 2). Two independent cage components in 1 are structu-

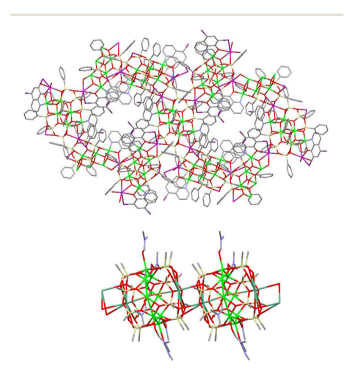
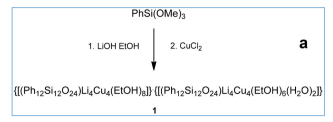


Chart 1 Examples of CLMS-based coordination polymers. Top. Fragment of a porous 3D MOF structure (Cu_6K_2 -silsesquioxane²⁰). Bottom. Dimeric fragment of a 1D extended structure (Cu₈Cs₂compound²¹).



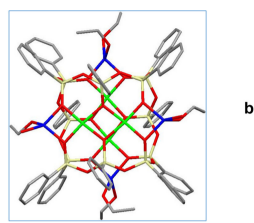


Fig. 1 (a) A general method for the synthesis of Cu₄Li₄-phenylsilsesquioxane 1. (b) The molecular structure of 1. Color code: green Cu; yellow Si; red O; grey C; blue Li.

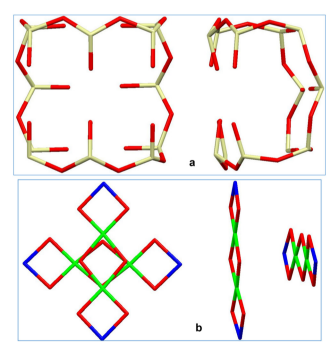


Fig. 2 (a) Two projections of twelve-membered cyclic silsesquioxane ligand in 1. (b) Two projections of the metal-oxo cluster in 1. Color code is the same as in Fig. 1.

rally very similar to each other. In particular, this could be concluded from a comparison of distances between opposite lithium ions in the two independent cages (they are equal to

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8.526(16) Å and 8.515(16) Å, or 8.555(16) Å and 8.565(16) Å, respectively).

In the next stage of our study, we have shifted to sodiumbased compounds (Fig. 3, top). In addition to initial conditions, when ethanol plays a dual role as a solvent for the synthesis and solvating ligand for crystallization, several other solution options have been explored. These included the replacement of ethanol by acetone or by the corresponding solvent pairs (EtOH/MeCN, EtOH/acetone, EtOH/diethyl ether). As a result, corresponding compounds, namely: [(Ph₁₂Si₁₂O₂₄) Na₄Cu₄(EtOH)₆] ·2EtOH (2, 73% yield), [(Ph₁₂Si₁₂O₂₄)Na₄Cu₄ $(Me_2CO)_8$] (3, 69% yield), $\{(Ph_{12}Si_{12}O_{24})Na_4Cu_4(EtOH)_{5,2}\}$ $(MeCN)_2(H_2O)_{0,8}$ $\{(Ph_{12}Si_{12}O_{24})Na_4Cu_4(EtOH)_4(MeCN)_2(H_2O)_2\}$ $(4, 58\% \text{ yield}) [(Ph_{12}Si_{12}O_{24})Na_4Cu_4(Me_2CO)_4(EtOH)_4] (5, 68\%)$ yield), [(Ph₁₂Si₁₂O₂₄)Na₄Cu₄(Et₂O)₄(EtOH)₂] (6, 62% yield; 6a, 60% yield) have been successfully isolated. All compounds 2-6 (Fig. 3, bottom) belong to the structural type 1, with a Si₁₂-silsesquioxane ligand and the Cu₄Na₄ core. To our surprise, we noticed that the replacement of lithium ions in 1 by sodium ions in 2-6 caused a significant structural distortion. Namely, one could observe a bent Na···Cu···Cu···Na fragment in 3, 5-6a

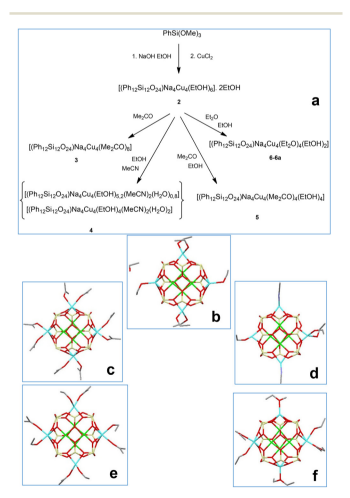
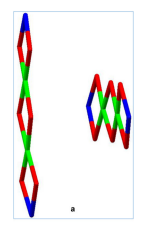


Fig. 3 (a) A general method for the synthesis of Cu₄Na₄-phenylsilsesquioxanes 2-6. (b-f) The molecular structures of 2-6. Color code is the same as for Fig. 1, but cyan Na.



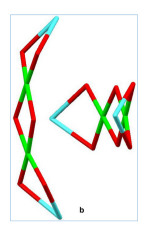


Fig. 4 Comparison of linear (a) and bent (b) metal-oxo clusters in 1 and 3, 5-6a, respectively. Color code is the same as for Fig. 1 and 3.

(Fig. 4) instead of a linear Li···Cu···Cu···Li fragment in 1. Only complex 4 adopts a linear principle similar to 1. Noteworthily, very recently, we have already reported on a specific role played by the lithium ion (along the Li-Na-K series) on the selfassembly of Fe(III)-silsesquioxane/acetylacetonate complexes, ^{19d} and this "Li-feature" is further confirmed here. Another influential aspect altering the structure of cages 2-6 is the nature of the solvating ligands, which could be illustrated by the variation of distances between opposite sodium centers. Namely, Na···Na distances are equal to 8.420(3) and 9.241(4) Å (for 2), 9.246(8) and 8.704(9) Å (for 5), 8.857(4) and 8.396(3) Å (for 6), 8.851(4) and 8.407(4) Å (for 6a). Complexes 6 and 6a represent different conformers and are distinguished only by the conformations of ether ligands bound to sodium ions. Note that acetone-solvated cage 3 possesses the symmetrical structure (both Na...Na distances are equal to 8.600(4) Å), as well as complex 4 with the Na···Na distances of 9.201(9)/9.218 (8) and 9.218(8)/9.213(8) Å (for the two crystallographically independent cages) without such variations.

We suggested that the formation of coordination polymers in the case of all complexes 2-6 was not realized due to screening effects of solvates at sodium centres. To force the formation of desired extended polymeric structures, it would be logical to try to use water since water molecules are characterized by minimum steric limitations among any widely used solvents. In line with these thoughts, we performed the "CuNa-silsesquioxane 2-targeted" syntheses but in the presence of an additional amount of water (Fig. 5, top). As a result, the corresponding water-containing complexes have been isolated, namely, $[(Ph_{12}Si_{12}O_{24})Na_4Cu_4(EtOH)_{1,5}(H_2O)_5]\cdot(EtOH)$ (7, 52% yield) and $[(Ph_{12}Si_{12}O_{24})Na_4Cu_4(H_2O)_4(EtOH)_{3,5}] \cdot \frac{1}{2}(EtOH)$ (8, 58% yield). Both complexes 7-8 represent a type of molecular architecture similar to compounds 2-6. Furthermore, water molecules, due to the lower steric hindrances, give rise to extended 1D chain-like structures (Fig. 5, bottom). Variation of the number of water and ethanol solvate molecules in 6-7 causes a certain distortion in the cage structures (opposite Na···Na distances are equal to 9.343(4) and 8.703(4) Å (for 7),

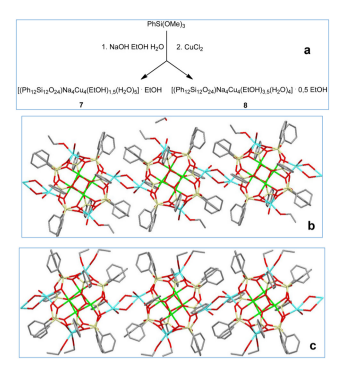


Fig. 5 (a) A general method for the synthesis of Cu₄Na₄-phenylsilsesquioxanes 7-8. 1D supramolecular structures of 7 (b) and 8 (c).

9.320(4) and 8.732(4) Å (for 8)). These values correspond well to those observed for 2-6. Moreover, the formation of coordination polymer structures leads to a sharp shortening of intermolecular Na···Na contacts. For 7 and 8, these distances are equal to 3.902(4) and 3.837(4) Å, respectively. Note that the shortest intermolecular Na···Na contact found in the case of 2-6 is equal to 5.971 Å (complex 4).

In the next stage, we concentrated on the syntheses of potassium-containing analogs of complexes 1-8 (Fig. 6, top). A larger radius of potassium ion was expected to better promote the formation of coordination polymers than in the case of sodium-based compounds. As in the case of Na-compounds, several "solvent options" have been explored. These included original (ethanol) synthesis as well as the replacement of ethanol by solvent combinations (DMF/MeOH/EtOH, diethyl ether/MeCN/H2O, or acetone). As a result, three compounds successfully realizing extended structures were isolated $[(Ph_{12}Si_{12}O_{24})K_4Cu_4(EtOH)_4]$ 9 (68% yield), $[(Ph_{12}Si_{12}O_{24})$ $K_4Cu_4(DMF)_2(MeOH)_{1.25}(EtOH)_{0.5}$ **10** (62%) yield), $[(Ph_{12}Si_{12}O_{24})K_4Cu_4(Et_2O)_2(MeCN)_2(H_2O)_2]$ 11 (60% yield) (Fig. 4, center and bottom). The molecular structures of cage complexes 9-11 perfectly correspond to those of compounds 1-7. As in 2, the "ethanol only-solvated" cage 9 reveals identical distances between opposite potassium centers (10.064(3) Å). A replacement of the solvent system causes significant distortion of the cage (the intra-cage K···K distances are equal to 9.857(3) and 10.190(5) Å for 10, or 9.5664(9) and 9.8221(9) Å for 11). At the same time, a change in the solvent systems may provoke a decrease in the dimensionality of the coordination polymer structure (2D in the cases of 9-10, 1D in the case of

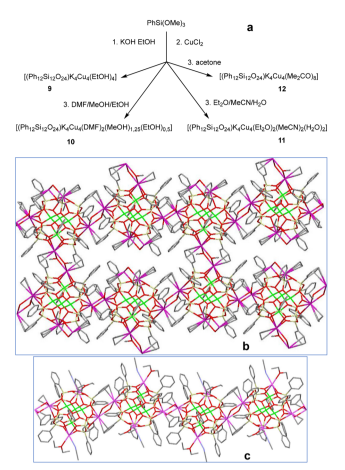


Fig. 6 (a) A general method for the synthesis of Cu₄K₄-phenylsilsesquioxanes 9-12. (b) A simplified (flat layer) representation of 2D "stitch" supramolecular structure of 9. (c) 1D supramolecular structure of 11. Color code is the same as for Fig. 1, but magenta K.

11). It is intriguing that in the cases of 9 and 11, the formation of the coordination polymer is assisted by the dual role of potassium centers. First, they form contacts with linking atoms of the solvating ligands (oxygen atom from the ethanol molecule in the case of 9 and oxygen atom from the water molecule in the case of 11). Second, one could easily identify additional half-sandwich contacts between the potassium center from one cage and a phenyl group (at a silicon atom) from the neighboring cage. The 2D coordination polymer in 9 possesses a non-classical net structure. Indeed, one could see alternating contacts between cages in upper and lower horizontal chains (Fig. 6, center). Namely, odd cages are linked to neighboring ones, forming vertical connections. Moreover, even cages remain unlinked, forming "pore" fragments. This situation is due to different (upward vs. downward) directions in the growth of the coordination polymer structure from any cage in 9, chosen as a starting one (Fig. 7). As a result, it comprises a system of interpenetrating waves (the so-called "stitch" style of 2D coordination polymers). Nevertheless, the resultant 2D coordination polymer in 9 is not porous (Fig. 6, center, shows a cross-section of the extended structure by a plane for clarity).

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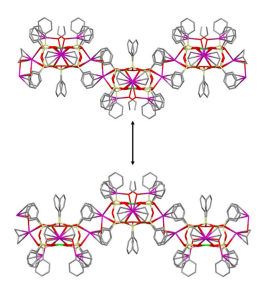


Fig. 7 Two wave-like 1D coordination polymer chains forming a 2D "stitch" structure in 9. Central (shared by both waves) cage is shown by an arrow. Bottom chain is rotated by 90° with respect to the upper one for clarity.

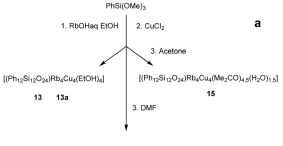
Intermolecular K···K distances for 9 are the same in all directions and equal to 4.378(3) Å.

In turn, compound 10 adopts the classical 2D net structure (Fig. S1†) via oxygen atoms of (i) DMF and (ii) alcohol solvates at potassium centers. Intermolecular K···K distances for 10 are equal to 4.216(3) Å (for DMF-involving contacts) and 4.124(4) Å (for alcohol-involving contacts).

The intercage connectivity of complex 11 (Fig. 6, bottom) is realized only by water-solvated potassium ions. Potassium centers coordinated by ether and acetonitrile molecules cannot form a coordination polymer due to the high steric limitations of ether-solvated molecules. The shortest intermolecular K···K distances are equal to 4.216(3) Å (for 10) and 4.6440(10) Å (for 11).

In line with these thoughts, solvates with even higher screening ability may totally suppress the growth of coordination polymer chains. Indeed, the fourth potassium-based complex, $[(Ph_{12}Si_{12}O_{24}) K_4Cu_4(acetone)_8] \cdot 2(acetone)$ (12, 73%) yield, Fig. S2†) adopts a OD island structure (the intercage K···K distances are all the same and equal to 8.678 Å). In the case of 12, the presence of two acetone solvate molecules at each potassium center effectively prevents any supramolecular assembly.

In the next stage, we concentrated on the syntheses of rubidium-containing compounds (Fig. 8, top). Similar to complexes 7-8 and 9-11, the resultant complexes adopt coordination polymer structures, namely, [(Ph₁₂Si₁₂O₂₄)Rb₄Cu₄ $(EtOH)_4$] (13, 63% yield; 13a, 60% yield), $[(Ph_{12}Si_{12}O_{24})Rb_4Cu_4]$ $(DMF)_4$ (14, 67% yield), and $[(Ph_{12}Si_{12}O_{24})Rb_4Cu_4(Me_2CO)_{4.5}]$ (H₂O)_{1.5}] (15, 71% yield). Furthermore, compounds 13–15 reveal specific distinctions. First of all, these complexes realize purely 2D supramolecular geometries. The nature of solvate molecules responsible for this exact type of supramolecular



 $[(Ph_{12}Si_{12}O_{24})Rb_4Cu_4(DMF)_4]$

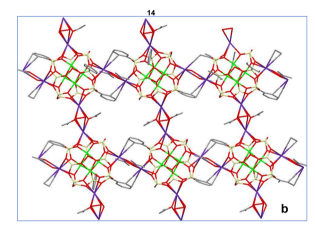


Fig. 8 (a) A general method for the synthesis of Cu₄Rb₄-phenylsilsesquioxanes 13-15. (b) The 2D net supramolecular structure of 15. Color code is the same as for Fig. 1, but purple Rb.

aggregation is intriguing. Namely, three compounds 13-14 belong to a "stitch" 2D structure mentioned above for compound 9. On the other hand, compound 15 shows the classical 2D net structure (Fig. 8, bottom). The intercage connections in 13-15, by analogy to 9-11, are assisted by half sandwich Rb...Ph contacts. The formation of a 2D structure is finalized via additional contacts between rubidium ions and the corresponding linking solvates (two ethanol molecules in the case of 13-13a; two dimethylformamide molecules in the case of 14). In the case of 15, one could see two different linking units: (i) with two acetone molecules and (ii) with three acetone molecules and a half water molecule. As in the case of complexes 1-12, structural parameters of the cage components of 13-15 are sensitive to the nature of the surrounding solvate. Distances between the opposite rubidium centers in the case of DMF-complex 14 are 9.8282(16) Å and 9.9360(17) Å, but 10.0171(5) and 10.3146(5) Å for complex 15 (acetone and water act as solvate molecules therein). The intracage Rb...Rb distances are not the same even for two EtOH4-solvated compounds: 13 (both distances are equal to 10.4168(5) Å) and 13a (10.505(2) and 10.289(3) Å). This indicates that even conformational differences (the tiniest changes in the localization of ethanol molecules) may provide certain distortion of the cage geometry. The same concerns intermolecular Rb···Rb distances (equal to 4.4535(6) Å for 13, while 4.488(2) and 4.421(3) Å for 13a). These distances are 4.3400(5) and 4.4278(5) Å for

15; while complex 14 reveals the most asymmetrical geometry (Fig. S3†) featuring three different distances (4.5696(13), 4.5348(9), and 4.5486(8) Å).

The final stage of our synthetic studies was aimed at isolating cesium-based compounds. The use of CsOH_{aq} as a source of cesium centers (Fig. 9, top) yields two compounds, namely, $[(Ph_{12}Si_{12}O_{24})Cs_4Cu_4(EtOH)_4]$ (16, 72% yield) [(Ph₁₂Si₁₂O₂₄)Cs₄Cu₄(EtOH)_{4.75}]·EtOH (17, 61% yield). A difference in the number of coordinated and solvated ethanol molecules causes little differences in the resultant cage structures. For example, both intramolecular Cs...Cs distances in 16 are equal to 10.773(2) Å. In the case of 17, these values change to 10.7647(8) and 10.8019(8) Å. To our surprise, much stronger distortions have been found in the supramolecular organizations of these compounds. Complex 16 forms a "stitch"-like 2D coordination polymer analogous to those discussed above for potassium- (9) and rubidium-based (13-14) compounds. The Cs...Ph half sandwich contacts supported by the coordination of each cesium ion by two ethanol solvates act as linking units. In turn, complex 17 represents an intriguing type of a 2D extended structure (Fig. 9, bottom). First of all, the cages in 17 are connected via Cs...Ph contacts and different types of Cs...O contacts. Two cesium ions are coordinated by two ethanol linkers, while two other ones are coordinated only by one ethanol linker. The corresponding intercage Cs···Cs contacts are equal to 4.493(3) and 5.4779(9) Å, respectively. Furthermore, one could see that (in the flat layer projection) the first and the fourth cages in the upper chain are not connected to the ones from the lower chain. This is due to the complicated "long stitch" style of the extended structure realized in 17. Fig. 10 shows differences in the

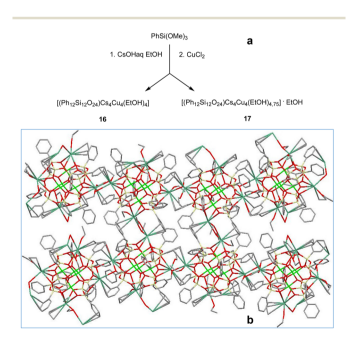


Fig. 9 (a) A general method for the synthesis of Cu₄Cs₄-phenylsilsesquioxanes 16 and 17. (b) Simplified (flat layer) representation of the 2D "long stitch" supramolecular structure of 17. Color code is the same as for Fig. 1, but with dark green Cs.

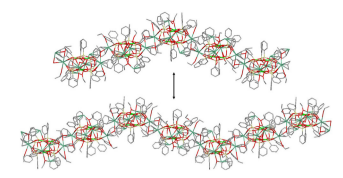


Fig. 10 Two wave-like 1D coordination polymer chains forming the 2D "long stitch" structure in 17. Central (shared by both waves) cage is shown by an arrow. Bottom chain is rotated by 90° with respect to upper one for clarity.

growth of the coordination polymer in two perpendicular directions from one cage chosen as the starting one. One could see that the central cage is on the crest of the wave position in the upper fragment and in the intermediate position in the lower fragment. It is interesting to compare it with the symmetrical style of "stitch" coordination polymer shown in Fig. 7. Due to such a "imbalance" in the central cage position, the 2D packing structure of 17 adopts other periodicity than that in compounds 9, 13-14, and 16. Both wave fragments shown in Fig. 9 are longer than the periodic subunits of the stitch structure (Fig. 7) giving rise to another, "long stitch", style of packing observed in the case of 17.

Catalytic studies

Oxidation of alkanes. Complex 12, chosen as a representative example, has been studied as a precatalyst in the oxidation of saturated hydrocarbons (Fig. 11 and S7-S11†). The oxidation of cyclohexane with hydrogen peroxide, according to gas chromatography, results in the formation of a mixture of cyclohexanol and cyclohexanone. The ratio of cyclohexanone to cyclohexanol is about 1.2 (0.033 M/0.026 M after 60 min). At the same time, the addition of triphenylphosphine to the reaction mixture according to the method proposed previously by Shul'pin²⁷ gives a much higher concentration of cyclohexanol and a sharp decrease in the amount of cyclohexanone, as shown in Fig. 11. The ratio of cyclohexanone to cyclohexanol after adding triphenylphosphine is about 0.07 (0.008 M/0.120 M after 60 min). The effect of adding triphenylphosphine to the reaction mixture can be explained by the fact that alkyl hydroperoxide is the main product of the oxidation reaction. In the absence of phosphine, it decomposes to form the respective ketone and alcohol in approximately equal amounts. Triphenylphosphine reduces all the alkyl hydroperoxide to alcohol, so one could see a sharp increase in the amount of alcohol on the chromatogram (see Fig. 11B). In many cases, the sum of alcohol and ketone before the addition of triphenylphosphine is much less than that after the addition of PPh3. For example, the sum of ketone and alcohol amounts after 60 minutes is 0.06 M before the addition but

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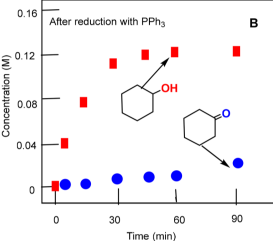


Fig. 11 Accumulation of cyclohexanol and cyclohexanone in the course of the oxidation of cyclohexane (0.46 M, 0.125 ml) with hydrogen peroxide (2.0 M, 50% aqueous, 0.32 ml) catalyzed by compound **12** (1 \times 10⁻³ M, 5.2 mg) in the presence of HNO₃ (0.02 M, 0.02 ml of the stock solution of 65% aqueous HNO₃ in MeCN) in MeCN (up to 2.5 ml) at 60 °C. Concentrations of cyclohexanone and cyclohexanol were determined before adding PPh₃ (graph A) and after the reduction of the aliquots with solid PPh3 (graph B).

increases to 0.128 M after the addition of PPh_3 . The total yield of alcohol and ketone in the oxidation of cyclohexane for a 1-hour reaction is 28%, and TON is 129. To enable the oxidation of alkanes, nitric acid must be added to the reaction mixture. When carrying out reactions in the absence of nitric acid, only the decomposition of hydrogen peroxide (catalase activity) occurred, and the yield of the oxidation reaction products was about 1.4% in 120 minutes (Table S1†).

We also determined the selectivity parameters for the oxidation of certain alkanes (n-heptane and methylcyclohexane) by hydrogen peroxide catalyzed by complex 12. For n-heptane, after two hours, the following concentrations (M) of alcohol isomers were obtained (after reduction with triphenylphosphine): C (1) 0.003 M; C (2) 0.007 M; C (3) 0.008 M; C (4) 0.004 M (yield 7%). These values gave a series of selectivity values: C (1): C (2): C (3): C (4) = 1.0:4.4:4.4:5.0. In the case

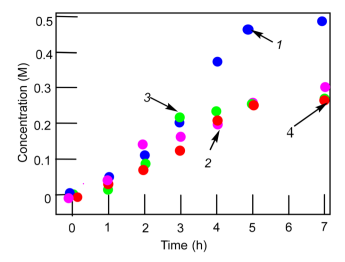


Fig. 12 Accumulation of acetophenone (curve 1, blue, yield 96%, TON = 960, TOF = 61), cyclohexanone (curve 2, pink, yield 57%, TON = 290, TOF = 41), 2-heptanone (curve 3, green, yield 50%, TON = 250, TOF = 36) and 2-hexanone (curve 4, red, yield 47%, TON = 240, TOF = 34) in the oxidation of 1-phenylethanol (0.5 M, 0.16 ml), cyclohexanol (0.5 M, 0.12 ml), 2-heptanol (0.5 M, 0.14 ml) and 2-hexanol (0.5 M, 0.12 ml) respectively, with tert-butyl hydroperoxide (1.5 M, 0.36 ml, 70% aqueous solution) catalysed by complex 12 (1 \times 10⁻³ M) in the presence of HNO₃ (0.02 M) at 60 °C in acetonitrile.

of methylcyclohexane oxidation, the following selectivity parameters were obtained: $1^{\circ}:2^{\circ}:3^{\circ}=1.0:5.4:14.8$ (yield 15%). The selectivity parameters measured in the oxidation alkanes are close to the parameters typical for the reactions of alkanes with hydroxyl radicals.²⁸ Based on the experimental dependences of the reaction rates on the concentration of the reagents and using previously obtained data for the oxidation reactions with different metallocomplexes, ^{28,29} we can assume that the oxidation reactions of alkanes take place with the participation of hydroxyl species and alkyl hydroperoxides are formed as the main primary products.

Oxidation of alcohols. Complex **12** has also been successfully evaluated in the oxidation of different alcohols with *tert*-butyl hydroperoxide (Fig. 12). In order to quench the oxidation process, the concentrations of products were measured by GC after the reduction of the reaction samples with solid PPh₃. Values of yields, TONs and TOFs are given in the footnote to Fig. 12.

Conclusions

An impressive number (19 distinct compounds) of cage metallosilisesquioxanes have been synthesized using copper(n) ions as core centers and alkali metal ions as peripheral ones. The disposition of alkaline metal cations is mainly determined by their size. First, small lithium cations in 1 coordinate only two oxygen atoms of the siloxanolate ligand and are arranged linearly to the copper cations. Contrarily, large potassium (9–12), rubidium (13, 13a, 14, 15) and cesium (16, 17) cations are arranged in an arcuate manner to the copper cations, binding four oxygen atoms of the siloxanolate ligand. The sodium

cations can also coordinate four oxygen atoms of the siloxanolate ligand; however, their disposition depends on both the cationic size and the different steric factors, for example, the steric volume of the coordinated solvent molecules. As a result, in the investigated cage-like Cu₄Na₄-based phenylsilsesquioxanes, sodium cations can arrange by all three types: (a) only linearly to the copper cations (4), (b) only arcuately to the copper cations (3, 6, 6a), and (c) both linearly and arcuately to the copper cations (2, 5, 7, 8). The molecular structures of all 19 compounds are similar, representing the Cu₄M₄ nuclearity supported by a Si₁₂-based ligand. In turn, the compounds strongly differ from each other in their disposition to form coordination polymers. While the small size of lithium ions predictably resulted in island-like Cu₄Li₄ molecular structures, sodium-containing compounds could form 1D coordination polymers in the case of solvate species with low steric hindrances (i.e., water) interlinking neighboring Cu₄Na₄ cages. In turn, Cu₄K₄ is capable of forming 0D, 1D, or 2D structures depending on the specific screening characteristics of the solvates. Cu₄Rb₄ and Cu₄Cs₄ compounds inevitably form extended (2D) structures. Packing trends of all described 2D coordination polymers (with K, Rb, or Cs ions) are different; along with classical 2D net organization, nontrivial "stitch" structures of interpenetrating waves are also available. Significant amounts of compounds have been isolated using acetone as a key solvating ligand. This supports our recent observations of the high efficiency of acetone surrounding the self-assembly of CLMSs. 20,21,30 A representative acetone-solvated Cu₄K₄-complex revealed a high catalytic activity in the peroxide oxidations of alkanes and alcohols. A strong influence of the nature of peroxide (H₂O₂ vs. TBHP) on catalytic activities towards different types of substrates has been noted. Keeping in mind the evidence that polynuclear compounds are often more active than copper mononuclear derivatives, 19a,23-26,31 further design of copper-based CLMSs, including coordination polymers, is of significant interest, and these investigations are currently ongoing in our team.

The data supporting this article have been included as part of the ESI.†

Crystallographic data for the article have been deposited at the CCDC under 2355320 (1), 2355321 (2), 2355322 (3), 2355323 (4), 2355324 (5), 2355325 (6), 2355326 (6a), 2355327 (7), 2355328 (8), 2355329 (9), 2355330 (10), 2355331 (11), 2355332 (12), 2355333 (13), 2355334 (13a), 2355335 (14), 2355336 (15), 2355337 (16), and 2355338 (17) and can be obtained from https://www.ccdc.cam.ac.uk.†

Conflicts of interest

There are no conflicts to declare.

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