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# Supramolecular reactions of metallo-architectures: Ag<sub>2</sub>-double-helicate/Zn<sub>4</sub>-grid, Pb<sub>4</sub>-grid/Zn<sub>4</sub>-grid interconversions, and Ag<sub>2</sub>-double-helicate fusion†

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Supramolecular reactions are of importance in many fields. We report herein three examples where complexes of hydrazone-based ligands are involved. A  $Ag_2$ -double-helicate was converted, by treatment with  $Zn(OTf)_2$ , into a  $Zn_4$ -grid (exchange of metal ions and change of the nature of the initial complex). A  $Pb_4$ -grid was converted, upon reaction with  $ZnCl_2$  or  $ZnBr_2$ , into a  $Zn_4$ -grid (exchange of metal ions, but conservation of the nature of the initial complex). The reverse conversions were also achieved. The fusion of a  $Ag_2$ -double-helicate with another  $Ag_2$ -double-helicate was performed (exchange of ligands, but conservation of the nature of the complexes) and resulted in a mixture of three helicates (two homostranded ones and one heterostranded one).

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### Introduction

Like covalent molecules, supramolecular<sup>1</sup> assemblies may participate in various reactions. The understanding of supramolecular reactions is of much interest because they are involved in many areas such as complex chemical systems and networks,<sup>2</sup> adaptive<sup>3</sup> and stimuli-responsive<sup>4</sup> chemical systems, fabrication of nanodevices and materials,<sup>4</sup> biomolecular processes. Thus, in the complexity and diversity of supramolecular chemistry, the reactivity of supramolecules plays a crucial role. It includes the processes:

(a) of (self)assembly (*i.e.* formation of supramolecular architectures through assembly, but also their participation, as subunits, in more complex assemblies), and correlatively, partial or total disassembly;

(b) of partial or total reorganization or exchange (at the supramolecular and, additionally and possibly, at the covalent level), that involves the breaking of several or all of the initial supramolecular connections and formation of new ones;

(c) without breaking or formation of new supramolecular connections (*e.g.* covalent modifications after self-assembly<sup>5</sup>).

Amongst supramolecular architectures, double helices and helicates,<sup>6</sup> as well as grids<sup>7</sup> arouse much interest and work. For example, DNA<sup>8</sup> and the ion channel generated by gramicidine<sup>9</sup> have a double helical structure, and there are double helical complexes that act as molecular machines<sup>10</sup> or catalysts.<sup>11</sup> Gridlike complexes have been studied for their electrochemical and magnetic properties,<sup>7</sup> for their capacity to encapsulate ions<sup>12</sup> or as starting materials for building more complex architectures (*e.g.* a Solomon link<sup>13</sup>), amongst other things. However, supramolecular interconversions of grids and helicates have not, except several examples,<sup>14,15</sup> been much explored.

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With these ideas in mind – and using principles such as the displacement of an equilibrium through precipitation, and the preference of Ag<sup>+</sup> for tetrahedral and of Zn<sup>2+</sup> for octahedral coordination – we designed, as reported herein, three supramolecular reactions<sup>16</sup> of reorganization and exchange (Fig. 1) involving grids and double helicates. They are related through the ligands<sup>17</sup> (which are pyrimidine-bis-hydrazones;<sup>18</sup> Fig. 2)

$^{i)}$ $^{2}$ ${\swarrow}$ ${\swarrow}$ ${\Rightarrow}$ ${\longrightarrow}$ $^{I}$ $^{II}$ $^{II}$ $^{II}$ $^{III}$ $^{IIII}$ $^{IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII$	exchange: metal ion nature changes: yes conformation changes: yes nuclearity changes: yes number of ligands changes: yes
$ii) \qquad $	exchange: metal ion nature changes: no conformation changes: no nuclearity changes: no number of ligands changes: no
$\begin{array}{c} \text{iii} \\ \text{iii} \\ \text{Ag}_2 L^1_{2\text{-DH}} \\ \text{Ag}_2 L^2_{2\text{-DH}} \\ \text{Ag}_2 L^1_{2\text{-DH}} \\ \text{Ag}$	exchange: ligand nature changes: no conformation changes: no nuclearity changes: no number of ligands changes: no

Fig. 1 Stylized representation of the three types of supramolecular reactions reported herein.

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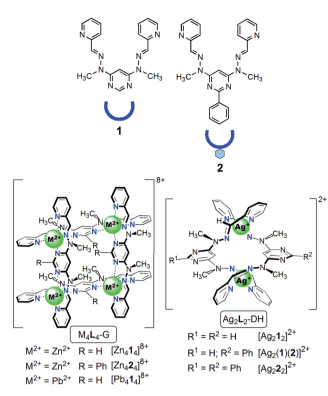
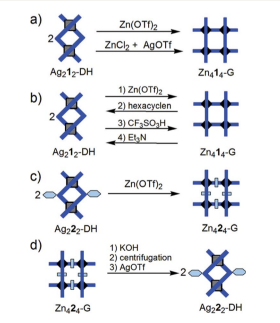


Fig. 2 Structural formulae and stylized representations of ligands 1 and 2, and of grids and double helicates.

that produce the supramolecular complexes, as well as through the nature of complexes, and occur due to the dynamic character of the present metal-ligand connections. These reactions (Fig. 1) can be seen as:



**Fig. 3** Stylized representation of: (a) the conversion of  $Ag_2I_2$ -DH double helicate into the  $Zn_4I_4$ -G grid; (b) the interconversion  $Ag_2I_2$ -DH/Z $n_4I_4$ -G; (c) the conversion of  $Ag_22_2$ -DH double helicate into the  $Zn_42_4$ -G grid; (d) the conversion of  $Zn_42_4$ -G grid into  $Ag_22_2$ -DH double helicate. Charges and stoichiometric coefficients are omitted for simplicity.

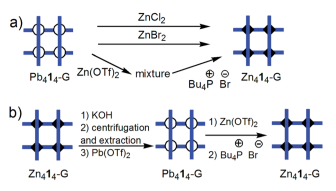


Fig. 4 Stylized representation of: (a) reaction of Pb<sub>4</sub>1<sub>4</sub>-G grid with ZnCl<sub>2</sub>, ZnBr<sub>2</sub>, and Zn(OTf)<sub>2</sub> and Bu<sub>4</sub>P<sup>+</sup>Br<sup>-</sup> (solvent CD<sub>3</sub>CN); (b) Zn<sub>4</sub>1<sub>4</sub>-G/Pb<sub>4</sub>1<sub>4</sub>-G and Pb<sub>4</sub>1<sub>4</sub>-G/Zn<sub>4</sub>1<sub>4</sub>-G conversions. Charges and stoichiometric coefficients are omitted for simplicity.

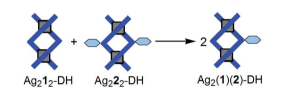


Fig. 5 Stylized representation of the fusion (conproportionation) reaction of double helicates  $Ag_21_2$ -DH and  $Ag_22_2$ -DH with formation of heterostranded species  $Ag_2(1)(2)$ -DH. Charges are omitted for simplicity.

(i) a change of the nature of the supramolecular architecture, from a Ag<sup>+</sup> dinuclear double helicate (DH) into a Zn<sup>2+</sup> tetranuclear grid (G), induced by replacement of Ag<sup>+</sup> by Zn<sup>2+</sup> (Fig. 3). In this reaction, not only the nature of the complex and that of the metal ion change, but also the conformation of the ligand (helical  $\rightarrow$  unfolded), the charge (2<sup>+</sup>  $\rightarrow$  8<sup>+</sup>) and the nuclearity of the complex (2  $\rightarrow$  4) and the number of ligands per complex (2  $\rightarrow$  4). In regard to this last change, this process can be compared with the conversion or the equilibrium between supramolecular dimer and tetramer of bioactive proteins,<sup>19</sup> or between other homo-oligomers<sup>20</sup> with influence on the protein functions.

(ii) a substitution<sup>21</sup> (metal ion exchange or transmetallation), in a sole operation, of the four  $Pb^{2+}$  ions of a grid-like<sup>22</sup> complex by  $Zn^{2+}$  ions (Fig. 4);

(iii) a fusion (conproportionation)<sup>23</sup> between two  $Ag^+$  double helicates<sup>24</sup> (Fig. 5).

While in case (ii) the equilibrium is shifted towards the  $Zn^{2+}$  grid through the precipitation of  $Pb^{2+}$  as its halides (chloride and bromide), in cases (i) and (iii), the conversions can be done without precipitation.

## Results and discussion

(i) The conversion  $Ag_2L_2$ -DH  $\rightarrow Zn_4L_4$ -G (L = 1, 2) through transmetallation is a dramatic reorganization of the nature of the metallo-supramolecular architecture induced by the replacement of Ag<sup>+</sup> by Zn<sup>2+</sup> (Fig. 3a and c): 2 Ag\_2L\_2-DH + 4 Zn<sup>2+</sup>  $\rightarrow Zn_4L_4$ -G + 4 Ag<sup>+</sup>. Ag<sup>+</sup> prefers a tetrahedral coordination geometry which is, in the case of ligands 1 and 2, achieved from 2 two-Nsp<sup>2</sup>-atom bidentate pyridine-hydrazone sites. In this way,  $Ag^+$  induces the formation of double helicates with ligands 1 and 2.  $Zn^{2+}$  prefers an octahedral coordination environment that results from 2 three-Nsp<sup>2</sup>-atom tridentate sites of type pyridine-hydrazone-pyrimidine, thus generating a grid.

Reaction of 1 equiv. of  $Ag_2L_2$ -DH<sup>15c</sup> with 2 equiv. of  $Zn(OTf)_2$ (OTf<sup>-</sup> = CF<sub>3</sub>SO<sup>3-</sup>) produces – without the need to precipitate Ag<sup>+</sup> as a halide – the corresponding grid  $Zn_4L_4$ -G<sup>15b,22a</sup> (solvent: CD<sub>3</sub>NO<sub>2</sub> with 6–14% CD<sub>3</sub>CN; ESI, pp. S9–S11†). Where ZnCl<sub>2</sub> is used in the reaction with Ag<sub>2</sub>1<sub>2</sub>-DH, two equivalents of AgOTf per equiv. of DH are required according to the equation (ESI p. S8†):

$$2Ag_2I_2-DH + 4ZnCl_2 + 4Ag^+ \rightarrow Zn_4I_4-G + 8AgCl_2$$

On treatment of the double helicate  $Ag_22_2$ -DH in  $CD_3NO_2$ with 2 equiv. of Zn(OTf)2 - added as a solution in a small volume of  $CD_3CN$ , or as a solid – the grid  $Zn_42_4$ -G was obtained. When the double helicate  $Ag_2 1_2$ -DH in  $CD_3NO_2$  was treated with 2 equiv. of Zn(OTf)<sub>2</sub>, added as a solution in a small volume of  $CD_3CN$  (about 6–14% of the  $CD_3NO_2$  volume), the grid  $Zn_41_4$ -G was obtained. When  $Zn(OTf)_2$  was added as a solid, without  $CD_3CN$ , was obtained a mixture without the  $Zn_41_4$ -G grid; addition of a small volume of CH<sub>3</sub>CN (about 6-14% of the  $CD_3NO_2$  volume) to this mixture produced the expected grid Zn<sub>4</sub>1<sub>4</sub>-G. A possible explanation could be that, in the case of the reaction  $Ag_2 \mathbf{1}_2$ -DH  $\rightarrow$  Zn<sub>4</sub> $\mathbf{1}_4$ -G, the CH<sub>3</sub>CN acts as a coordinating species for the Ag<sup>+</sup> ions and so contributes to the displacement of the equilibrium from the double helix towards the grid. The grid  $Zn_42_4$ -G should be – due to the  $\pi$ -stacking aromatic interaction between a phenyl ring and the two ligands between which that phenyl is located within the grid - more stable than the grid  $Zn_4 I_4$ -G. This stability may be sufficient to make possible the formation of the grid Zn<sub>4</sub>2<sub>4</sub>-G from the corresponding double helicate without, unlike in the case of the grid  $Zn_41_4$ -G, the assistance of  $CH_3CN$ .

DOSY NMR was also used to study the conversion  $Ag_2L_2$ -DH  $\rightarrow$   $Zn_4L_4$ -G (L = 1, 2). As expected, the volume of the grid species obtained from double helicates on treatment with  $Zn(OTf)_2$  was found in agreement with that of the grid prepared from the free ligands L and  $Zn(OTf)_2$ .

The reverse conversion  $Zn_4L_4$ -G  $\rightarrow Ag_2L_2$ -DH can be done as follows: after treatment of the grid with KOH, the solvent (CD<sub>3</sub>CN or CD<sub>3</sub>NO<sub>2</sub>) is removed, and the ligand is extracted with CDCl<sub>3</sub> and separated from the solid residue (by centrifugation or filtration); after removal of CDCl<sub>3</sub>, CD<sub>3</sub>NO<sub>2</sub> is added, then AgOTf is added to form the helicate. In order to simplify the procedure, we used ligand **2** and a mixture of CDCl<sub>3</sub> and CD<sub>3</sub>NO<sub>2</sub> where ligand **2**, as well as the corresponding grid and double helicate were soluble. After precipitation of Zn<sup>2+</sup> with KOH, the mixture was centrifuged (the ligand **2** being soluble in the mixture of solvents), and to the recovered liquid phase AgOTf was added to produce the Ag<sub>2</sub>2<sub>2</sub>-DH (ESI, p. S13<sup>†</sup>).

In a pH-dependent system (Fig. 3b), the interconversion between  $Ag_21_2$ -DH and  $Zn_41_4$ -G was achieved as follows (ESI, p. S10†): the grid was generated from the double helicate by reaction with  $Zn^{2+}$ ; then,  $Zn^{2+}$  was complexed with hexacyclen, and the double helicate was regenerated; partial protonation of hexacyclen with TfOH caused release of  $Zn^{2+}$  and formation of the grid (incomplete yield); finally, addition of triethylamine reactivated the hexacyclen that again encapsulated  $Zn^{2+}$  and resulted in the reformation of the double helicate.

(ii) The Pb<sub>4</sub>**1**<sub>4</sub>-G  $\rightarrow$  Zn<sub>4</sub>**1**<sub>4</sub>-G conversion (Fig. 4a) can formally be seen as a substitution of Pb<sup>2+</sup> by Zn<sup>2+</sup> ions, although the real mechanism, involving breaking and formation of supramolecular bonds, must be more complex. Reaction of Pb414-G15b with 4 equiv. of Zn(OTf)2 produces a mixture which no longer contains the grid-like species Pb<sub>4</sub>1<sub>4</sub>-G or Zn<sub>4</sub>1<sub>4</sub>-G (ESI p. S2<sup>†</sup>). This suggests that the affinity of  $Zn^{2+}$  for the ligand, as well as its preference for octahedral coordination are not sufficient to displace the equilibrium towards Zn<sub>4</sub>1<sub>4</sub>-G. We considered that the involvement of Pb<sup>2+</sup> ions in a weakly dissociating or sparingly soluble compound should displace the equilibrium. Indeed, addition of Br<sup>-</sup> (as  $Bu_4P^+Br^-$ ) to the above mixture, or treatment of  $Pb_41_4$ -G with four equivalents of ZnBr2 or ZnCl2 produced - along with the formation of  $PbX_2$  (X = Br, Cl) which precipitates and, doing so, shifts the equilibrium – the expected Zn<sub>4</sub>1<sub>4</sub>-G grid (solvent: CD<sub>3</sub>CN; ESI pp. S3–S4<sup>†</sup>): Pb<sub>4</sub>1<sub>4</sub>-G + 4 ZnX<sub>2</sub>  $\rightarrow$  Zn<sub>4</sub>1<sub>4</sub>-G + 4 PbX<sub>2</sub>.

The reverse conversion  $Zn_4\mathbf{1}_4$ - $G \rightarrow Pb_4\mathbf{1}_4$ -G grid was achieved in several steps (Fig. 4b). Treatment of  $Zn_4\mathbf{1}_4$ -G (in CD<sub>3</sub>CN) with KOH led to the precipitation of  $Zn^{2+}$  (as  $Zn(OH)_2$  or  $K_2[Zn(OH)_4]$ ), as well as of the free ligand **1**. After removal of CD<sub>3</sub>CN, the free ligand **1** was extracted with CDCl<sub>3</sub> and used further for the preparation of Pb<sub>4</sub>\mathbf{1}\_4-G (see ESI, p. S5<sup>†</sup>).

Thus, in addition to its self-assembly from  $Zn^{2+}$  and a ligand, the same  $Zn^{2+}$  grid,  $Zn_4\mathbf{1}_4$ -G, can be obtained, in reactions (i) and (ii), from a Ag<sup>+</sup> dinuclear double helicate or from a Pb<sup>2+</sup> tetranuclear grid (exchange of metal ions and reorganization of the architectures).

(iii) The fusion (conproportionation) reaction of double helicate  $Ag_21_2$ -DH<sup>15c</sup> with 1 equiv. of  $Ag_22_2$ -DH (Fig. 5) according to the equation

$$Ag_21_2$$
-DH +  $Ag_22_2$ -DH  $\rightarrow 2Ag_2(1)(2)$ -DH

produces a mixture that contains each of the three helicates, namely two homoleptic (homostranded) ones and one heteroleptic (heterostranded) one. Ligands **1** and **2** equally participate to homo- and heteroleptic helicates, and so the observed molar percentages are of approximately 25% for  $Ag_21_2$ -DH, 25% for  $Ag_22_2$ -DH, and 50% for  $Ag_2(1)(2)$ -DH. For characterization of the new compound  $Ag_22_2$ -DH, see ESI pp. S14–S20;† for <sup>1</sup>H, <sup>13</sup>C and DOSY of the mixture of three helicates, see ESI pp. S25–S31.†

For the reactions described above it might appear necessary, in practice, to slightly (2–10%) increase the amounts of reagents with respect to those theoretically calculated.

#### Experimental

For experimental details, see the ESI.†

#### Conclusions

To summarize, three supramolecular reactions were investigated: (i) a  $Ag_2L_2$ -DH double-helicate into  $Zn_4L_4$ -G grid conversion, where the exchange of metal ions changes the nature of the metallo-supramolecular architecture, (ii) a  $Zn_41_4$ -G grid into  $Pb_41_4$ -G grid conversion driven by a halide-induced precipitation and where the nature of the metallo-supramolecular architecture is conserved, and (iii) a double exchange of ligands during the fusion of two double helicates.

The grid/grid and double-helicate/grid conversions were made reversible by precipitation of  $Zn^{2+}$  with KOH and subsequent reaction of the free ligand with  $Ag^+$  or  $Pb^{2+}$ , or, for one DH/G interconversion, in a pH-dependent way.

In perspective, such ligands could be introduced in larger and more complex, suitably decorated, architectures where such supramolecular reactions can act as actuators of various properties (charge, volume, multivalency).

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