Sol-gel selection of hybrid G-quadruplex architectures from dynamic supramolecular guanosine libraries[†]‡

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We report an original strategy to transcribe and to fix the self-assembly of G-quadruplex architectures in self-organized nanohybrids. In the first resolution step the G-quartet is pre-amplified in solution in the presence of metal cations from a dynamic pool of ribbon-type or cyclic supramolecular architectures. Then in a second selection sol-gel step the G-quadruplex is irreversibly fixed in a siloxane inorganic network. The interface between the guanosine moiety and the heterosiloxane **3** is based on a *double reversible covalent* iminoboronate connection. This contributes to the high levels of adaptability and correlation in the self-organization of the supramolecular G-quadruplex and the inorganic siloxane systems. Biomimetic-type hybrids can be generated by using this strategy.

1. Introduction

Constitutional dynamic chemistry (CDC)¹ that expresses the adaptive behaviour of molecular and supramolecular systems brings into play the self-evolution of such dynamic systems toward the selection of discrete architectures from mixtures of exchanging components. These concepts may be connected with the simple definition of dynamic multicomponent self-assembly in chemical and biological systems.² The spontaneous formation of higher order superstructures from simple molecular blocks concerns inter-exchanging supramolecular architectures, and often yields discrete devices and materials. By virtue of the reversible interchanges, the components can adapt to internal and external constraints. Kinetic resolution,^{3,4} self-assembly processes followed by covalent modification⁵ or the use of crystallization^{6,7} are useful strategies to control and to drive supramolecular self-organization toward producing functional complex systems. Previous investigations have highlighted the wide potential of resolution strategies controlled in the second step by an irreversible enzymatic³ or crystallization^{6,7} tandem process. The control of supra-molecular self-organization by a phase-change can also be driven by using the sol-gel process,⁶⁻⁸ providing a simple method for synthesis of well-defined hybrid constitutional materials.^{5h} In this context, the supramolecular H-bonded macrocycle of four guanine building blocks, the G-quartet, has been proposed as a powerful scaffold for the construction of synthetic supramolecular devices and materials.8

The G-quartet architecture represents a dynamic supramolecular system in which the G-quartet is reversibly exchanging with linear ribbons (Fig.1).^{5a,b} It is stabilized by cations, templating the eight carbonyl oxygens of two sandwiched G-quartets in the G-quadruplex, the columnar device formed by the vertical stacking of four G-quartets.⁹ It plays a very



Fig. 1 CD spectra at room temperature in CH₃OH of $5(10^{-4} \text{ M})$ (black line), 5 + 0.25 eq. KTf or 5 + 0.25 eq. BaTf₂ (red line) and 5 + 1 eq. KTf or 5 + 1 eq. KTf or 5 + 1 eq. BaTf₂ (blue line).

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important role in biology in particular in nucleic acid tellomers of potential interest to cancer therapy.8 New strategies using reversible polymerization or phase-change-driven selection were successfully used to generate G-quadruplex functional nanostructures.^{10,11} In our group, the guanine building block and the sol-gel chemistry were used to conceive hybrid chiral materials at nanometric and micrometric scales. Our efforts involved the synthesis and the self-assembly of a guanine-siloxane monomer G_{Si} (Fig.1), resulting in the formation of G-ribbons, G-quartet and G-quadruplex supramolecular architectures, dynamically exchanging in solution. The generation of G-quadruplex hybrid materials can be achieved by mixing the Gsi derivative with potassium triflate, where G-quartet superstructures have been amplified. Then the sol-gel selection process has been followed by a second inorganic transcription into inorganic silica replica materials, by calcination.^{5a} Dynamic nanomaterials, characterized by their aptitude to continuously organize at the macroscopic level the generation of their own components with different blocks forming in coupled equilibria, have been used as a straightforward approach for the design of functional G-quartet membranes^{5b} and surfaces.^{11d} Despite such progress, considerable challenges still lie ahead and a more significant one is the dynamic convergence between supramolecular selfassembly of G-quartet architectures and the resolution process (polymerization, phase change). The reversibility of interactions between the G-quartet and polymeric components might be a crucial factor and accordingly, a reversible covalent connection might render the emergence of self-adaptive system mesophases, which may mutually adapt their 3D spatial distribution based on their own structural constitution during the simultaneous formation of mixed self-organized domains.

Herein we report an original strategy to transcribe and to fix the supramolecular self-assembly of the dynamic constitutional G-quartet architectures, in self-organized nanohybrids.¹² In the first resolution step the G-quartet is pre-amplified in the presence of metal cations from a dynamic pool of oligomeric ribbon-type or cyclic supramolecular architectures (Scheme 1). Then in a second sol-gel step the polycondensation reactions of the inorganic alkoxysilane network take place around the tubular twisted superstructure of the G-quadruplex. It should be noted that the dynamic G-quadruplex architectures are irreversibly fixed in a covalently bonded siloxane network and the structural constitution of the G-quadruplex is transcribed in the hybrid material. These results provides new insights into the basic features that control the convergence of *supramolecular selforganization toward functional hybrid materials*.

2. Experimental

2.1. Materials and methods

All reagents were obtained from Aldrich and used without further purification. All organic solutions were routinely dried by using sodium sulfate (Na₂SO₄). ¹H, ¹³C and ²⁹Si NMR spectra were recorded on an ARX 300 MHz Bruker spectrometer in d4-MeOD using the residual solvent peak as a reference. ²⁹Si MAS NMR was performed on a Bruker ASX 400 spectrometer. ESI-MS studies were performed in the positive and negative ion mode using a quadrupole mass spectrometer (Micromass, Platform II). Samples were continuously introduced into the mass spectrometer through a Waters 616 HPLC pump (60 °C; extraction cone voltage: Vc = 30 V). X-ray powder diffraction measurements were performed with Cu-Ka radiation at 20 °C using a Philips X'Pert Diffractometer equipped with an Xcelerator detector. Thermogravimetric analysis (Hi-Res TGA 2950, TA Instruments, Nitrogen, 50-600 °C at 10 °C min-1) and differential scanning calorimetry (DSC 2920 Modulated, TA Instruments) were used to evaluate the thermal stability of the hybrid materials A-E. CD spectra (220-400 nm) were collected on a MOS-450/ AF-Biologic CD spectrometer with a rate of 0.5 nm/step, using



Scheme 1 Cation-template resolution of a dynamic supramolecular guanine system in which the G-quartet is reversibly exchanging with linear ribbons followed by a secondary irreversible sol–gel selection of G-quadruplex hybrid materials.^{5a}

10 mm quartz cells, at room temperature. The measurements were carried out on MeOH solutions (10⁻⁴ M of 3) and on mixtures of 3/4 (1/1, mol/mol) briefly ultrasonicated with various amounts of K⁺ and Ba²⁺ ions. The SEM micrographs were obtained with high resolution transmission electron microscopy (HRTEM) JEOL 2010 FEG apparatus, working with an accelerating voltage of 200 kV and a point resolution of 2.0 Å.

2.2. Synthesis of 2-((3-(triethoxysilyl)propylimino)methyl) phenylboronic acid, 3

3-(triethoxysilyl)propan-1-amine 2 (0.8 mL; 3.3 mmol) was added to a chloroformic solution (20 mL) of 2-formyl-phenylboronic acid, 1 (0.50 g; 3.3 mmol). The mixture was refluxed for 6 h. The solvent was removed under vacuum to give 3 as a yellow oil. (74%) ¹H-NMR (MeOD, 300 MHz): δ 0.55 (t, 2H); 1.04–1.17 (m, 9H); 1.99 (m, 4H); 3.33 (t, 2H); 3.85 (dd, 6H); 4.55 (s, 2H); 7.37 (m, 1H); 7.57 (t, 2H); 7.64 (d, 1H); 8.71 (s, 1H). ¹³C-NMR (MeOD, 300 MHz): δ 156.7, 153.7, 150.7, 136.2, 136.1, 129.7; 126.8; 116.7; 106.5; 103.0; 88.3; 87.7; 86.5; 84.5; 84.3; 83.1; 82.3; 80.8; 61.6.

2.3. General procedure for synthesis of $A_{K}^{+}-D_{K}^{+}$, $A_{Ba}^{2+}-D_{Ba}^{2+}$ and the reference E hybrid materials

In a typical run, 3 mL of methanolic solution of 3 (2.5×10^{-3} M) was mixed with 1 eq. of guanosine 4, 0.25 or 1 eq. of templating metal salt (KTf or BaTf₂) and 3 eq. of benzylamine as catalyst. Whereas hybrids A_K⁺, C_K⁺, A_{Ba}²⁺, C_{Ba}²⁺ were prepared by a solgel process in the presence of 6 eq. of deionized water the hybrids $\mathbf{B_K}^+$, $\mathbf{D_K}^+$, $\mathbf{B_{Ba}}^{2+}$, $\mathbf{D_{Ba}}^{2+}$ were prepared by using only the air humidity (Table 1). The solutions were briefly stirred and allowed to react at room temperature under static conditions for 1 week. Then samples were slowly evaporated at room temperature to give yellow powders. A reference sol-gel sample E was prepared in the absence of ionic salts, under the same conditions.

3. **Results and discussions**

3.1. Concept, design and synthesis of G-quartet hybrid materials

3.1.1. Design of hybrid molecular component. 2-((3-triethoxysilyl)propylimino)methyl)phenylboronic acid 3 was obtained by refluxing stoichiometric amounts of 2-formyl-phenylboronic acid 1 and 3-(triethoxysilyl)propan-1-amine 2 in CHCl₃ (Scheme 2).

Table 1 The sol-gel synthetic protocol and temperature of 50% mass decomposition, $T_{50\%}$ of the hybrid materials, $A_{K}^{+}-D_{K}^{+}$, $A_{Ba}^{2+}-D_{Ba}^{2+}$

	3 (equiv.)	4 (equiv.)	KTf (equiv.)	BaTf ₂ (equiv.)	H ₂ O (equiv.)g	T _{50%} /°C
A_{K}^{+}	1	1	1	_	_	487
B_{K}^{+}	1	1	1		6	509
C_{K}^{+}	1	1	0.25			467
D_{K}^{+}	1	1	0.25	_	6	450
A_{Ba}^{2+}	1	1	_	1		483
B_{Ba}^{2+}	1	1		1	6	480
C_{Ba}^{2+}	1	1		0.25		464
D_{Ba}^{2+}	1	1		0.25	6	473
E	1	1			_	425

The ¹H, ¹³C NMR and ESI-MS spectra are in agreement with the proposed formula.

3.1.2. Generation of G-quartet architectures in solution. Compounds 1 and guanosine, 4 generate G-quartet architectures by K^+ or Ba^{2+} cation templating, based on three-encoded features:

a) the hybrid materials might be considered to contain superstructures resulting from the combination of cross-linked arrays of G-quartets cross-linked by appending iminoboronate groups connected to the silica matrix, reminiscent of the guanine hybrids recently reported by our group;5a

b) the linkage between the guanosine moiety and the iminoboronic monomer, 3 is based on a double reversible covalent connection. The imino moiety contributes to the stabilisation of the reversible boronate-guanosine ester bond by the formation of a dative N-B bond.13 This then contributes towards dynamic adaptative behaviour on the molecular level in a double "dynamer"-type system:5b it combines reversible supramolecular H-bonding (G-quartets) with the reversible polycondensation (iminoboronate bonds) of components. This might contribute to the high levels of adaptability and correlation in the self-organization of the dynamic supramolecular G-quartets and the rigid inorganic siloxane components;

c) biomimetic guanosine-type hybrids can be generated using this strategy.

The generation of G-quartet architectures can be achieved by using the mixtures of 1 and guanosine 4 (1/1, mol/mol), resulting in the formation of 5 and potassium or barium triflates. Unfortunately such mixtures of 5/KTf or BaTf2 are soluble only in methanol and the ¹H NMR experiments are inadequate to indicate the self-association of the G-quartet due the exchange of NH¹ amide and for NH² amine protons with the solvent.

Next, we used CD spectroscopy to gain evidence that 5 forms supramolecular chiral superstructures in the presence of ions in methanol (Fig. 1). In accordance with the NMR results, a weak Cotton effect was observed in the CD spectrum of a methanolic solution of 5 which can be ascribed to the intrinsic supramolecular chirality, resulting from twisted supra-molecular geometry in which the molecular guanosine components associate and indicating that the G-quadruplex is either unstructured or exchanging very fast in solution (Fig. 1a,b black lines). After addition of 0.25 equiv. of KTf or 0.25 equiv. of BaTf₂ the corresponding CD spectra (Fig. 1a,b red lines) shows a negative Cotton effect with a zero crossing at $\lambda = 320$ nm or $\lambda = 310$ nm, respectively in the region of the iminoboronic chromophores. This effect might be induced by the formation of the G-quartets induced by steric interactions between the chiral sugars groups on the G unit and further inducing asymmetric interactions between iminoboronic pendant chromophores. The observed CD signals in the presence of the cations indicates that chiral aggregates are forming in the presence of metal ions. It is interesting to note that related $oligo(p-phenylene-vinylene)-G^{11a}$ and oligo-thiophene-G^{11b} D4-symmetric octamers, with the appending group attached to the 5'-position of the G ribose, showed a similar negative Cotton effect. Interestingly, increasing the amount of metals ions to 1 equiv. the corresponding CD spectra (Fig. 1a,b blue lines) shows a positive Cotton effect (zero crossing at $\lambda = 314$ nm) for the K⁺ cations and a negative Cotton effect (zero crossing at $\lambda = 330$ nm) for the Ba²⁺ cations corresponding



Scheme 2 The synthesis of iminoboronateguanosine precursor 5 followed by ion-template resolution of G-quartet architectures and sol–gel selection of hybrid materials A_{K}^{+} - D_{K}^{+} , A_{Ba}^{2+} - D_{Ba}^{2+} .

to a helical self assembly of iminoboronic pendant chromophores.¹⁴ Moreover this indirect CD signature of chiral superstructures closely resembles the previously calculated CD spectra by Gray *et al.*¹⁵ of G-quadruplexes whose adjacent G-quartets have opposite polarities.

3.1.3. Generation of G-quartet architectures in solid state. Having shown that compound 5 equilibrated with K^+ and Ba^{2+} ions forms G-quadruplex superstructures in solution we next evaluated their ability to form higher order superstructures in the solid state, by using a sol-gel resolution process.4ª The generation of *G*-quadruplex hybrid materials $A_{K}^{+}-D_{K}^{+}$ or $A_{Ba}^{2+}-D_{Ba}^{2+}$ can be achieved by mixing guanosine 4 and 2-((3-(triethoxysilyl)propylimino)methyl)phenyl boronic acid, 3 in methanolic solutions with potassium or barium triflates, followed by a solgel process performed at room temperature and using benzylamine as a catalyst (Scheme 2, Table 1). We also carried out the sol-gel process, in the absence of K⁺ or Ba²⁺ cations, resulting in the formation of a G-hybrid material, E. Finally, the organic constituents of these materials have been calcinated to obtain inorganic silica replica architectures induced by the self-organization of G-quadruplex superstructures.

3.2. FTIR and solid state NMR spectroscopic analyses

FTIR and NMR spectroscopic analyses demonstrate the formation of a self-organized organic-inorganic network. The FTIR spectra of the hybrid materials $A_{K}^{+}-D_{K}^{+}$, $A_{Ba}^{2+}-D_{Ba}^{2+}$ or E allowed us to observe the $v_{Si-O-Si}$ broad vibration at 900–1200 cm⁻¹ instead of v_{Si-OEt} vibrations (950, 1070 and 1100 cm⁻¹) which were noticed for the molecular precursor **3**. Clear evidence for H-bonding and metal ions complexation was obtained from FTIR analysis of solids $A_{K}^{+}-D_{K}^{+}$, $A_{Ba}^{2+}-D_{Ba}^{2+}$ and E. Free $v_{C=O} = 727 \text{ cm}^{-1}$ (hybrid E) was downshifted to 1680 cm⁻¹ (hybrids $A_{K}^{+}-D_{K}^{+}$, $A_{Ba}^{2+}-D_{Ba}^{2+}$), indicating the G-quartets formation (Fig. 2). The formation of iminoboronate esters stabilized by a donor N \rightarrow B dative bond is supported the vibration shifts of the -N=C bonds: $v_{C=N} = 1640$ to 1605 cm⁻¹

and the presence of the $v_{B-N} = 814 \text{ cm}^{-1}$ band, mostly predominant in the FTIR spectra of $A_K^+ - D_K^+$, $A_{Ba}^{2+} - D_{Ba}^{2+}$.

²⁹Si MAS NMR spectroscopic experiments are in agreement for both $A_{K}^{+}-D_{K}^{+}$, $A_{Ba}^{2+}-D_{Ba}^{2+}$ and the reference E hybrid materials, with total condensated hybrid materials (the condensation number and the condensation degree are 2.80–2.90 and 93–97% respectively), with major cross linked T³ C–Si(OSi)₃ (70 ppm ~95%) units and minor T¹ C–Si(OSi)(OH)₂ (24%), T² C–Si(OSi)₂(OH) (59 ppm ~5%) units, showing a predominant 3D arrangement. Moreover, that the self assembly processes occur differently in the presence or in the absence of K⁺ templating ions, seems not to influence the hydrolysis-condensation reactions of the alkoxysilane groups during the sol–gel process.

3.4. X-Ray powder diffraction characterization of G-quadruplex hybrid materials

Further valuable insights on the self-organization of $A_{K}^{+}-D_{K}^{+}$ or $A_{Ba}^{2+}-D_{Ba}^{2+}$ and reference E hybrid materials are obtained from the X-ray powder diffraction (XPRD) (Fig. 3 and S2[†]).



Fig. 2 FTIR spectra of G-hybrid material, E and of G-quadruplex hybrid materials A_{K}^{*} and A_{Ba}^{2+} .



Fig. 3 High angles XPRD patterns of the compounds 5, 5 + 0.25 equiv. of M^+ , 5 + 1 equiv. of M^+ ($M^+ = K^+$, Ba^{+2}) and of the hybrid materials a) $A_{K^+}-D_{K^+}$ and b) $A_{Ba}^{2+}-D_{Ba}^{2+}$.

The XRPD patterns of the *G*-quadruplex hybrids $A_{K}^{+}-D_{K}^{+}$ and $A_{Ba}^{2+}-D_{Ba}^{2+}$ present well-resolved peaks (Fig. 3). It is noted that the crystallinity of the Ba²⁺-templated materials $A_{Ba}^{2+}-D_{Ba}^{2+}$ is high compared with the K⁺-templated analogs, as previously observed¹⁶ in view of the signal peaks, indicative of long-range structured material.15,16 This feature was not observed for the XRPD patterns of the non-templated G-hybrid material, E (Fig. 3, blue, patterns). The metal ion templated materials A_{K}^{+} - $\mathbf{D_{K}}^{+}$ and $\mathbf{A_{Ba}}^{2+}$ - $\mathbf{D_{Ba}}^{2+}$ present a peak at $2\theta = 26.8$ (d = 3.3 Å), representative for the π - π stacking distance between two planar G-quartets.^{5b,11e} This peak at $2\theta = 26.7^{\circ}$ (d = 3.3 Å), is strongly amplified by simple addition of metal ion templates before the sol-gel process (Fig. 3, the black and red patterns are compared to the blue one) by simply mixing 5 with metal ions. Further if we proceed with the sol-gel results in the formation of polymerized hybrids. We observed that the peak at $2\theta = 26.7^{\circ}$ (d = 3.3 Å) is more intense for C_{K}^{+} and D_{K}^{+} or C_{Ba}^{2+} and D_{Ba}^{2+} materials prepared in the presence of stoichiometric amounts of metal ions.

3.5. Thermal and morphological studies

To study the stability of the hybrid materials under the influence of the G-quadruplex superstructure, we explored the thermal



Fig. 4 Scanning electron microscopy images of B_{Ba}^{2+} hybrid material after the sol–gel step (bottom) and after the calcination step (lower).

decomposition of A_{K}^{+} - D_{K}^{+} or A_{Ba}^{2+} - D_{Ba}^{2+} and reference E hybrid materials. Table 1 shows the $T_{50\%}$ temperature for which 50% weight decomposes. The formation G-quadruplex structures induce a higher $T_{50\%}$ for $A_{K}^{+}-D_{K}^{+}$ and $A_{Ba}^{2+}-D_{Ba}^{2+}$ materials than reference material E. Scanning electron microscopy (SEM) micrographs reveal that the hybrid materials $A_{K}^{+}-D_{K}^{+}$ and $A_{Ba}^{2+}-D_{Ba}^{2+}$ (Fig. 4) have micrometric plate-like or nanorod morphologies. Moreover, while the classical hybrids are compact dense solids with a granular morphology, the constitutional hybrids A_i and B_i consist of crystalline rods of micrometric dimension. It is interesting to note that similar superior longrange organisation at micrometric length have been previously observed.¹⁶ This phenomenon is certainly dependent on the superior emergence of collective domains adapting along the G-quadruplex reversible interfaces. This may increase the size of self-organized domains, increasing the dimension of resulted microdomains of such constitutional hybrids.¹⁶

4. Conclusions

In conclusion, the results presented here reveal an original strategy to transcribe and to fix supramolecular G-quadruplex architectures in self-organized constitutional hybrids. In particular, the use of a dynamic iminoboronate interface between the organic (supramolecular) and inorganic siloxane networks represents a useful strategy for improving their compatibility. Dynamic self-assembly of supramolecular systems prepared under thermodynamic control, may in principle be connected to a kinetically controlled sol-gel process in order to extract and select the interpenetrated self-organized networks under a specific set of experimental conditions. Such "dynamic convergence" between supramolecular self-assembly and an inorganic sol-gel process which can synergistically communicate, leads to higher self-organized hybrid materials with increased micrometric scales. Sol-gel selection of constitutional architectures from dynamic combinatorial libraries toward systems materials, as highly interconnected networks of reactional and constitutional materials, should expand the fundamental understanding of nanoscale structures and properties as it relates to creating products and manufacturing processes. More generally, applying such consideration to hybrid materials, leads to the definition of constitutional hybrid materials, in which organic (supramolecular)/inorganic domains are reversibly connected.¹⁷ Considering the simplicity of this strategy, possible applications of the synthesis to more complex architectures might be very effective, producing novel expressions of complex matter.

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