

Cite this: *Chem. Sci.*, 2020, 11, 8151

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 28th April 2020  
Accepted 14th July 2020

DOI: 10.1039/d0sc02399j

rsc.li/chemical-science

## The dark side of disulfide-based dynamic combinatorial chemistry†

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During the last two decades, disulfide-based dynamic combinatorial chemistry has been extensively used in the field of molecular recognition to deliver artificial receptors for molecules of biological interest. Commonly, the nature of library members and their relative amounts are provided from HPLC-MS analysis of the libraries, allowing the identification of potential binders for a target (bio)molecule. By re-investigating dynamic combinatorial libraries generated from a simple 2,5-dicarboxy-1,4-dithiophenol building block in water, we herein demonstrated that multiple analytical tools were actually necessary in order to comprehensively describe the libraries in terms of size, stereochemistry, affinity, selectivity, and finally to get a true grasp on the different phenomena at work within dynamic combinatorial systems.

### Introduction

Since its discovery, Dynamic Combinatorial Chemistry (DCC) has grown into a powerful strategy to produce, at a low synthetic cost, self-assembled architectures displaying complex topologies and/or innovative functions.<sup>1</sup> DCC from disulfide-based building blocks has been particularly used in the field of molecular recognition to deliver tailored receptors for target (bio)molecules.<sup>2</sup> DCC simultaneously allows the generation of a library of potential receptors, to screen their affinity for a targeted partner, to detect the best lead, and possibly to prepare this species by optimized scaled-up procedures. DCC relies on the amplification phenomenon, which is the increase in the amount of an assembled library member upon introduction of the targeted partner called the template. Amplification generally transcribes a stable and preferential association between both partners. A bottleneck of the DCC strategy is sometimes the difficulty to accurately quantify the amount of each library member within mixtures, and very frequently the impossibility to selectively extract the best binders from the mixtures. Commonly, only the nature of the members and their relative amounts are provided from HPLC-MS analysis of the libraries. These relative amounts and, more specifically, the amplification factor (*i.e.*, the ratio between the amount of potential binder in

the presence and absence of the template) are then used to estimate the binding constant rather than measure it experimentally through titration between the isolated partners.<sup>3,4</sup>

During the last five years, we extensively re-investigated the dynamic combinatorial libraries generated from the single 2,5-dicarboxy-1,4-dithiophenol monomer **M** upon slow aerobic thiol oxidation and simultaneous fast disulfide exchange at physiological pH in water (Fig. 1A).<sup>5</sup> Such libraries were first described in 2006 by some of us to quantitatively deliver the racemic homochiral tetramer of (pS)<sub>4</sub>/(pR)<sub>4</sub> configuration **M<sub>4</sub>-a** upon incubation with the biogenic polyamine spermine (Fig. 1A and B).<sup>6</sup> This collection of readily accessible constitutionally dynamic analogues of pillararenes, to which **M<sub>4</sub>-a** belongs, was enlarged in 2016 and named dyn[*n*]arenes.<sup>5d</sup> A landmark was the straightforward purification procedure of these new cyclophanes by selective precipitation and separation from the buffer and template. Herein, by extending this procedure to entire libraries instead of single members, we thoroughly investigated the impact of the degree of molecular information (*i.e.*, the number of positive charges) borne by homologous templates (from spermine **T4** to ammonium acetate **T1**, through cadaverine **T3** and butylamine **T2**) on the (stereo)chemical composition of libraries made from monomer **M** (Fig. 1A).

### Results and discussion

When building block **M** undergoes spontaneous disulfide bridge formation and exchange in buffered water, the HPLC-MS analysis of the resulting libraries **L<sub>0</sub>-L<sub>4</sub>** after 24 hours of incubation showed that – regardless of the template (**T1-T4**) used – the racemic homochiral tetramer **M<sub>4</sub>-a** of (pS)<sub>4</sub>/(pR)<sub>4</sub>

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† Electronic supplementary information (ESI) available: Full experimental and computational details. See DOI: 10.1039/d0sc02399j



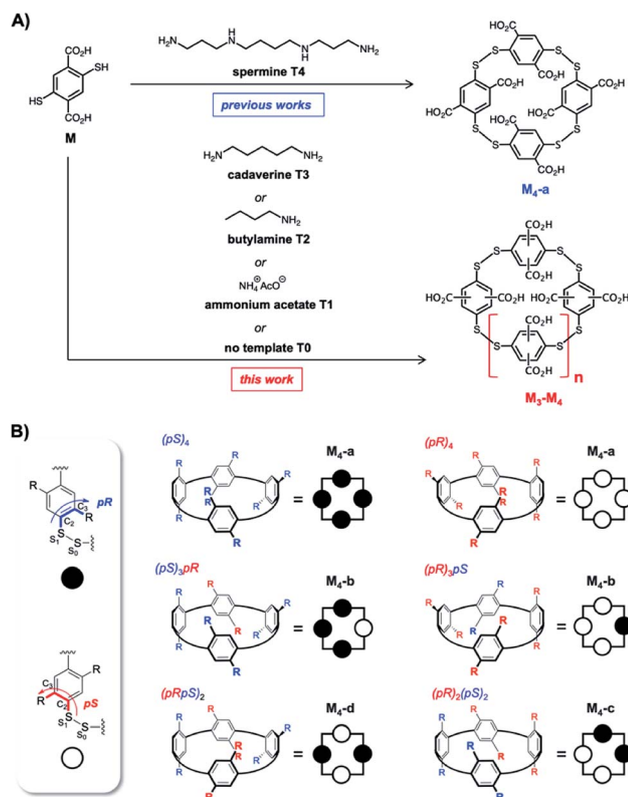


Fig. 1 Previously reported synthesis of homochiral dyn[4]arene  $M_4$ -a using spermine as the template, and current investigations involving truncated analogues of spermine (A), and dyn[4]arene's virtual configurational landscape arising from the rotation of phenyl units yielding hosts  $M_4$ -a to  $M_4$ -d (B).

configuration was the main species produced (in blue in Fig. 2A). In comparison, HPLC analysis of the reference untemplated library  $L_0$  showed almost no detectable peak after one day. Comparison between the peak area corresponding to  $(pS)_4/(pR)_4$   $M_4$ -a in the various templated libraries  $L_1$ - $L_4$  revealed that, although this library member appeared at first sight to be the main or even sole species obtained, its true conversion ranged from 23% to 100% (Fig. 2A). The 100% reference was attributed to the peak area obtained with spermine **T4** (*i.e.*, corresponding to 100% of the disulfide-based material), as this template was previously shown by NMR monitoring to quantitatively deliver the homochiral dyn[4]arene  $M_4$ -a. The fluctuation in the total amount of disulfide-based material within the various libraries  $L_0$ - $L_4$  suggested that a number of species failed to be detected by HPLC, and that complementary analytical tools would be required to properly and fully assess the template effects and provide a clear compositional view of the libraries.

After template and buffer removal by selective precipitation of the library members upon neutralization of their anionic charges using trifluoroacetic acid, the (stereo)chemical composition of the corresponding static template-free libraries  $L_0'$ - $L_4'$  was assessed by ESI-MS and NMR spectroscopy analyses at pH = 7.4 (Fig. 2C). The former revealed that the unidentified species were either dyn[3]arenes or dyn[4]arenes (see Fig. S2†).

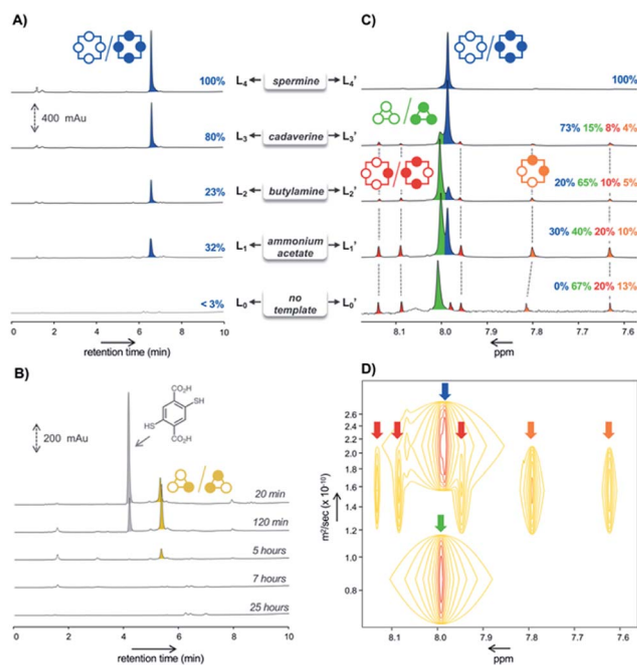


Fig. 2 HPLC analyses of the libraries  $L_1$ - $L_4$  made from monomer **M** (4 mM) in the presence of the templates (A), and  $L_0$  in absence of any template (B);  $^1\text{H}$  NMR analyses of the corresponding libraries  $L_0'$ - $L_4'$  (C), and DOSY NMR analysis of the library  $L_1$  made from monomer **M** in presence of ammonium acetate **T1** as the template (D).

The latter indicated that: (i) library  $L_4'$  generated with spermine as a template expectedly displayed a unique singlet at 7.98 ppm (in blue in Fig. 2C), which was formerly assigned to the homochiral dyn[4]arene  $M_4$ -a of  $(pS)_4/(pR)_4$  configuration,<sup>5d</sup> and that (ii) other templated libraries  $L_1'$ - $L_3'$  also showed the presence of  $(pS)_4/(pR)_4$   $M_4$ -a, along with at least six other visible singlets of varying areas depending on the template used, and (iii) that the homochiral tetramer  $M_4$ -a was indeed absent from the untemplated system  $L_0'$ . According to DOSY NMR experiments, these additional singlets corresponded to species displaying diffusion coefficients 1.6 to 3 times lower than the  $(pS)_4/(pR)_4$  homochiral cyclophane  $M_4$ -a (Fig. 2D). From these observations, it seemed legitimate to conclude that the trimers  $M_3$  and tetramers  $M_4$ , which failed to be detected by HPLC-UV but could be quantified by  $^1\text{H}$  NMR spectroscopy, form (sub)nanoscopic aggregates in solution. This hypothesis was supported by the good agreement between the percentages of homochiral  $M_4$ -a obtained by its integration in the NMR spectra (libraries  $L_i'$ , Fig. 2C) and the relative peak area measured on each of the corresponding HPLC chromatograms (library  $L_i$ , Fig. 2A). It also demonstrated that the composition of the system was unaltered by the precipitation process, *i.e.* that static template-free systems  $L_i'$  were true reflections of the dynamic templated mixtures  $L_i$ .

Among the remaining  $^1\text{H}$  NMR signals, it was possible to unambiguously assign the singlet at 8.01 ppm (in green in Fig. 2C) to the homochiral trimer  $M_3$ -a of  $(pS)_3/(pR)_3$  configuration, as this dyn[3]arene had been previously isolated by us and fully characterized by NMR spectroscopy.<sup>5b</sup> We also ruled out the presence in the libraries of the heterochiral trimer  $M_3$ -



**b** of (pS)<sub>2</sub>(pR)/(pR)<sub>2</sub>(pS) configuration. Indeed, the early stage HPLC monitoring of the untemplated library **L**<sub>0</sub> revealed that **M**<sub>3</sub>-**b** is a kinetic intermediate. While it appeared to be the major (detectable) library members during the first hours of incubation, it rapidly disappeared after a few additional hours (in yellow in Fig. 2B). HPLC-MS confirmed that this species was a trimer, which has to be the heterochiral stereoisomer since the homochiral counterpart **M**<sub>3</sub>-**a** yielded a flat HPLC-UV chromatogram. The remaining unassigned species present in **L**<sub>0</sub>-**L**<sub>3</sub>' could only be heterochiral tetramers **M**<sub>4</sub>-**x** (x = b-d, Fig. 1B), whose potential configuration may either be (pSpR)<sub>2</sub>, (pS)<sub>2</sub>(pR)<sub>2</sub>, or (pS)(pR)<sub>3</sub>/(pR)(pS)<sub>3</sub>. They correspond to C<sub>4</sub>, C<sub>2</sub>, and C<sub>1</sub> symmetries and should consequently display a single, two, and four singlets in <sup>1</sup>H NMR spectroscopy, respectively.

In order to assign these remaining unknown species, we performed an NMR titration on the static library of hosts **L**<sub>1</sub>' generated with ammonium acetate as a template (corresponding to the templated library displaying the highest amount of unassigned species) using increasing amounts of cadaverine **T**<sub>3</sub> as a guest molecule (see Fig. S3†). Interestingly, normalizing the chemical shifts' variation for each signal provided a supervised procedure for peak assignment within a complex mixture of homologous species. Not only does it validate the attribution of each C-H proton peak monitored throughout the titration, preventing accidental swapping, but it also allows the gathering of signals corresponding to the same host species on the basis of their evolution profile, even when splitting occurs during titration (Fig. 3A). Upon addition of cadaverine **T**<sub>3</sub>, the first binding event detected involves **M**<sub>4</sub>-**a**. As previously reported, it corresponds to the formation of a pseudorotaxane-type complex between both partners (**T**<sub>3</sub> ⊂ **M**<sub>4</sub>-**a**). This binding event is slow at the NMR time scale: upon guest addition, the singlet corresponding to the free host disappears while the singlet corresponding to the complex appears further downfield. Additional amounts of guest **T**<sub>3</sub> triggered simultaneously all other binding events, which appeared to be fast at the NMR time scale. Three distinct sets of signals could be identified: two singlets belonging to the *meso* (pS)<sub>2</sub>(pR)<sub>2</sub> species (**M**<sub>4</sub>-**c**, in orange in

Fig. 2B), and four singlets belonging to the heterochiral dyn[4]arene of (pS)(pR)<sub>3</sub>/(pR)(pS)<sub>3</sub> configuration (**M**<sub>4</sub>-**b**, in red in Fig. 2B). Among all possible stereoisomers of dyn[4]arene **M**<sub>4</sub>, the *meso* alternate (pSpR)<sub>2</sub> stereoisomer **M**<sub>4</sub>-**d** was missing in every library. This may be imputed to “head-to-head” repulsive coulombic interactions between adjacent aromatic units, which represent up to 4 unfavorable contacts within this *meso* species while only 2 can be found within the macrocycles **M**<sub>4</sub>-**b** and **M**<sub>4</sub>-**c** of (pS)<sub>2</sub>(pR)<sub>2</sub> and (pS)(pR)<sub>3</sub>/(pR)(pS)<sub>3</sub> configuration respectively, and none in the homochiral dyn[4]arene **M**<sub>4</sub>-**a**. To further confirm our assignment, the NMR titration of **L**<sub>2</sub>' was also conducted with **T**<sub>2</sub>, which led to the same conclusions (see Fig. S4†). This thorough compositional and stereochemical analysis dramatically changed the picture of the dynamic libraries **L**<sub>0</sub>-**L**<sub>4</sub> generated from 2,5-dicarboxy-1,4-dithiophenol monomer **M**, a widely used ingredient in disulfide based DCC.<sup>5-7</sup> Relative quantification of the eluted library members by HPLC-UV/MS, which only displayed a minor portion of the system (the “bright side” of the libraries), would in fact lead to the following erroneous conclusions: (i) that templates **T**<sub>1</sub>-**T**<sub>4</sub> stabilized the homochiral tetramer more or less to the same extent through its binding, (ii) that the untemplated system **L**<sub>0</sub> presumably evolved toward the formation of non-eluting polymeric and polydisperse material, and (iii) that the heterochiral tetramers and the homochiral trimer, due to carboxylate-carboxylate contacts and/or ring constraints, were intrinsically too unstable to form and be observed. In fact, this portion of the system (the “dark side” of the libraries), which is not disclosed by conventional monitoring, represents not only most, if not all, the material content, but also most of its constitutional and stereochemical diversity. Unintuitively, this dark portion is stabilized by self-aggregation. From the experimental distribution of these cyclophanes measured in the reference system **L**<sub>0</sub>, and assuming that the limit of detection by <sup>1</sup>H NMR spectroscopy of the homochiral tetramer **M**<sub>4</sub>-**a** is around 1% (with a signal/noise ratio superior to 450),<sup>8</sup> it appeared that members of the dark side of the DCL are around 10 kJ mol<sup>-1</sup> more stable than **M**<sub>4</sub>-**a**. It translates into an interconversion constant  $K_{\text{iso}}^{\text{RPP}}$  in the range of 10 to 10<sup>2</sup> M<sup>-1</sup> (Fig. 5A).<sup>9</sup> This provides an order of magnitude of the counter-driving force that opposes the

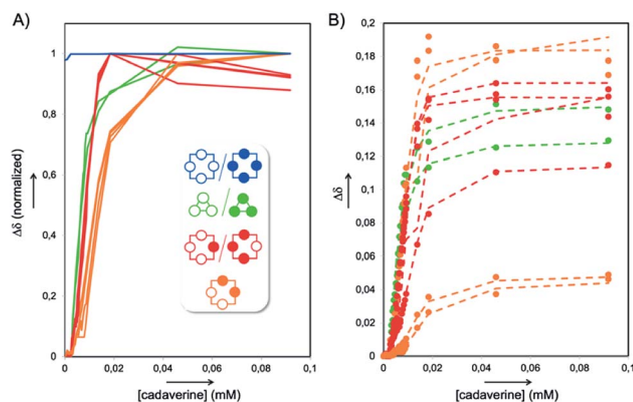


Fig. 3 (A) Normalized chemical shifts' evolution of the various NMR signals from the titration of the static library **L**<sub>1</sub>' with cadaverine **T**<sub>3</sub>, providing full host assignment and (B) corresponding concomitant fitting (dotted lines) of the experimental chemical shifts (circles).

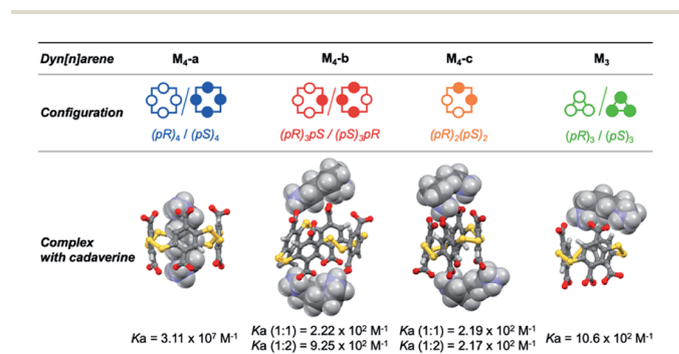


Fig. 4 3D structures of the complexes between cadaverine and dyn[n]arenes **M**<sub>3</sub> and **M**<sub>4</sub> of various configurations obtained by MD simulations, and respective affinities between the partners measured by ITC and NMR titrations.



templated formation of the homochiral tetramer **M<sub>4</sub>-a**, which eventually appeared to be the one of the most unstable architectures within small size cyclo-oligomers. Yet, the fluctuations in the relative amounts of these “invisible” library members (all but the homochiral **M<sub>4</sub>-a**) between the different templated systems **L<sub>1</sub>'–L<sub>4</sub>'** (Fig. 2B) indicated that the production of these species from **M** was guest-dependent, *i.e.* that each of them may display weak yet contrasted affinities toward each guest.

The 1 : 1 binding constant between the homochiral dyn[4] arene **M<sub>4</sub>-a** and cadaverine **T3** was previously determined by ITC to be  $3.11 \times 10^7 \text{ M}^{-1}$  in water at physiological pH.<sup>5a</sup> The apparent binding constants  $K_{\text{bind}}^{\text{app}}$  between **T3** and the dyn[*n*] arenes **M<sub>3</sub>-a**, **M<sub>4</sub>-b** and **M<sub>4</sub>-c** could be extracted from the abovementioned NMR titration by simultaneous curve fitting on every series of downfield displacements for each of the proton NMR signals of the hosts, as the concentration of the guest increased (Fig. 3A and B). To put it simply, the concomitant fitting was performed using homemade MATLAB scripts.<sup>10</sup> The set of equations describing associations and mass balance could be solved using the *fsolve* MATLAB function to deliver the speciation of the system at equilibrium for a defined set of association constants (for the set of equations used, see the ESI†). The *lsqcurvefit* MATLAB function was then used to adjust the association constants by fitting the speciation to the NMR data. This methodology was applied for different binding stoichiometries, and a highly satisfactory average correlation coefficient  $R^2 = 0.991$  was obtained for binary complexes in the case of the homochiral trimer **M<sub>3</sub>-a** and ternary complex in the case of **M<sub>4</sub>-b** and **M<sub>4</sub>-c**.

As the full array of spectra corresponding to the titration revealed it, guest complexation with some of the dyn[*n*]arenes also induced a symmetric breaking of the hosts (see Fig. 3 and S3†). As a consequence, some C–H proton signals progressively split into minor (individual) and major (isochronous) peaks (see Fig. S2†). We therefore challenged the robustness of our procedure by fitting not only one series of signals for each species but all of them alternatively, where minor peaks were tested one-by-one while keeping major signals for other species. As mentioned

earlier, this extended fitting allows the detection and self-correction of any assignment error on complex mixtures. Following the same procedure, apparent binding constants displayed individually by every library member contained in **L<sub>2</sub>'** toward **T2** were simultaneously extracted by the same concomitant fitting of the corresponding titration (see Fig. S5†).

From these sets of binding data, it appears that the apparent affinity of cadaverine **T3** for dyn[*n*]arenes of various configurations spans five orders of magnitude (Fig. 4) with a selectivity factor (defined as the ratio of the binding constants) reaching up to  $10^3$  in favour of the template-responsive homochiral host **M<sub>4</sub>-a**. The same general trend was observed for **T2**: the binding constants, one or two orders of magnitude weaker than those displayed by cadaverine **T3**, span four orders of magnitude, with a marked selectivity in favour of **M<sub>4</sub>-a** by at least a factor  $10^3$ .

To confirm the experimental trends, Molecular Dynamics (MD) simulations were performed and allowed to probe the three-dimensional structure of the corresponding complexes. MD simulations of 100 ns at 300 K with constant pressure in a truncated octahedral TIP3P water box were performed with the Amber 12 software package (see the ESI† for full computational details).<sup>11</sup> The most representative structures were extracted after cluster analysis (Fig. 4). Modelling also confirms that the energetically favored association process displayed by the homochiral **M<sub>4</sub>-a** corresponds to molecular recognition through inclusion of the guest into the cavity of the host. Contrary to surface binding, inclusion is accompanied by an enthalpic gain originating from the optimization of the number of electrostatic contacts between partner-borne charges (*i.e.*, the lock-and-key paradigm), as well as from the possible expulsion of high-energy water molecules from the cavity (*i.e.*, the non-classical hydrophobic effect).

This picture of the full network of the affinities and selectivities between hosts and guests within the dynamic system can obviously not be apprehended from the fitted HPLC-UV based apparent amplifications. It underlines the true benefit of analyzing a static system **L<sub>1</sub>'** derived from the parent DCL **L<sub>1</sub>** by NMR spectroscopy. In particular, <sup>1</sup>H and DOSY NMR analysis of the untemplated library **L<sub>0</sub>'** provided the last piece to fully apprehend the various noncovalent interactions and forces simultaneously at work within the dynamic system (Fig. 5A).

As mentioned earlier, the untemplated library gathers self-aggregated cyclophanes (the “dark side”, Fig. 5A), which thereby become substantially more stable than the homochiral cyclotetramer **M<sub>4</sub>-a** (the “bright side”). Guest binding by these species formally required their partial or total disaggregation. This revealed that NMR titration delivers in fact the apparent binding constant  $K_{\text{bind}}^{\text{app}}$ , which is related to the true binding constant  $K_{\text{bind}}$  through the relationship:

$$K_{\text{bind}} = K_{\text{bind}}^{\text{app}} \times K_{\text{agg}}$$

This led to the conclusion that the true selectivity (defined as the ratio of binding constant  $K_{\text{bind}}$ ) displayed by each guest for **M<sub>4</sub>-a** with respect to any member of the dark side of the library

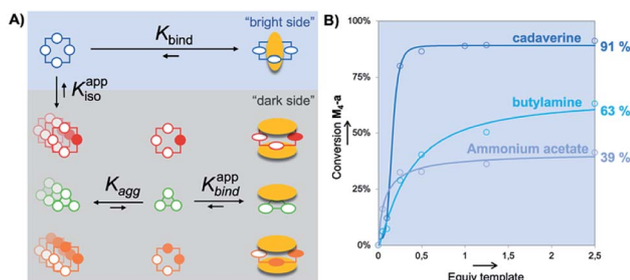


Fig. 5 Merging the bright and the dark side: (A) schematic overview of the supramolecular processes (binding vs. aggregation) operating within the full DCL based on monomer **M** ( $K_{\text{bind}}$ : discrete host–guest binding constant;  $K_{\text{agg}}$ : host self-aggregation constant;  $K_{\text{bind}}^{\text{app}}$ : apparent guest binding constant from aggregated host;  $K_{\text{iso}}^{\text{app}}$ : apparent isomerization constant from initial aggregated state); (B) conversions in **M<sub>4</sub>-a** from quantitative HPLC-UV monitoring with increasing amounts of templates **T1–T3**; x-axis is rescaled by a factor 1/5 for **T1**.



was in reality one or two order of magnitudes lower (w.r.t.  $K_{\text{bind}}^{\text{iso}}$  in the range of  $10$  to  $10^2 \text{ M}^{-1}$ ) than the values obtained from the apparent binding constant. Another way to put it is that, within static and dynamic systems  $L_i'$  and  $L_i$ , aggregation competes with binding and thereby reinforces an intrinsically modest supramolecular selectivity of each guest for the series of host molecules (which appears total and completely overestimated by HPLC).

Finally, we addressed the following question: can the quality (or amount) of molecular information contained by a guest be compensated (in terms of template effect on a host) by its absolute amount? Experimentally, we screened the impact (at fixed pH and ionic strength) of increasing amounts of templates **T1–T3** on the absolute conversion (quantified by the peak area of **M<sub>4</sub>-a** measured in HPLC-UV with respect to the area obtained with 0.25 eq. of **T4**) of **M<sub>4</sub>-a** obtained from building block **M** (Fig. S1†). The results, summarized in Fig. 5B, indicate that cadaverine **T3** was in fact the minimal truncated version of spermine, whose lack of molecular information can be compensated by a stoichiometric excess. In other words, almost quantitative amplification of **M<sub>4</sub>-a** could still be induced by equilibrium displacement provided that a strong excess of the  $\alpha,\omega$ -diamine core is used ( $91 \pm 3\%$  vs. 100% with spermine **T4**). In contrast, more truncated versions such as **T1** (AcONH<sub>4</sub>) and **T2** could only afford maximal conversions of respectively  $39 \pm 5$  and  $63 \pm 6\%$  in **M<sub>4</sub>-a**. This was corroborated by non-linear square fitting of the experimental data which, whatever the model used, confirmed that these maximal values were already reached experimentally with respectively 2.5 and 50 equivalents of each template.

## Conclusions

We have shown here that exploring dynamic combinatorial libraries using a single, yet widely used analytical tool can unfortunately lead to a considerable underestimation of the diversity of mixtures in terms of size and stereochemistry, where the identity of the main species can even be misassigned. Quantifying the various host–guest affinities and apprehending the main underpinning supramolecular phenomena (binding, folding, aggregation) at work within a dynamic combinatorial system requires complementary procedures and techniques. In this perspective, we have extended our straightforward protocol of selective template and buffer removal, which herein allowed the preservation of the original composition of the dynamic libraries and provided robust static mixtures eligible for structural analyses and titrations. The propensity of some library members to form aggregates, which affected their detectability by HPLC and participation in the overall thermodynamic equilibrium of libraries, could easily be characterized by routine NMR spectroscopy experiments. Combined with a series of titrations on static mixtures of hosts, this analysis delivered the full array of binding affinities between templates and cyclophanes. It eventually provided the global view of the dark and bright side of this simple DCL, wherein intrinsic selectivity between hosts of varying sizes and configuration is modest but reinforced by self-aggregation processes. These two

antagonistic forces operate both in the present static and dynamic parent mixtures, and presumably in many other water-soluble dynamic combinatorial libraries. Numerical simulations have previously shown that supramolecular interactions between library members could affect both degree and selectivity of the response of the library when a template molecule is added.<sup>12</sup> Herein, we report – for the first time – an experimental illustration of this anticipated scenario. Our final recommendation, to complement HPLC screening by *in situ* and *ex situ* NMR spectroscopy is easy to implement and relies on readily accessible analytical techniques. Such guidelines should allow the community of system chemists to easily apprehend the interactions at work and to generate valuable knowledge on mixtures of increasing complexity.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

This work was supported by the LABEX iMUST (ANR-10-LABX-0064) of the Université de Lyon within the program “Investissements d’Avenir” (ANR-11-IDEX-0007) operated by the French National Research Agency (ANR). We are very grateful to A. Baudouin (CCRMN) for her help with the NMR measurements.

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