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# Thermodynamically driven self-assembly of pyridinearene to hexameric capsules†

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Pyridinearene macrocycles have previously shown unique hostguest properties in their capsular dimers including endo complexation of neutral molecules and exo complexation of anions. Here, we demonstrate for the first time the formation of hydrogen bonded hexamer of tetraisobutyl-octahydroxypyridinearene in all three states of matter - gas phase, solution and solid-state. Cationic tris(bipyridine)ruthenium(II) template was found to stabilize the hexamer in gas phase, whereas solvent molecules do this in condensed phases. In solution, the capsular hexamer was found to be the thermodynamically favoured self-assembly product and transition from dimer to hexamer occurred in course of time. The crystal structure of hexamer revealed 24 N-H···O direct intermolecular hydrogen bonds between the six pyridinearene macrocycles without any bridging solvent molecules. Hydrogen bond patterns correlate well with DFT computed structures. Thus, all structural chemistry methods (IM-MS, DOSY NMR, DFT, X-ray crystallography) support the same structure of the hexameric capsule that has a diameter of ca. 3 nm and volume of 1160 Å3.

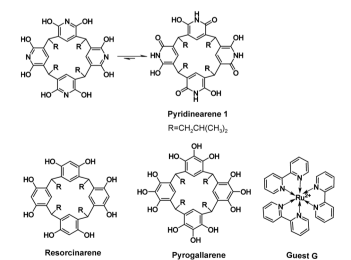
long known for their capability to spontaneously self-assemble into large hexameric capsules.<sup>2</sup> Interestingly, the hydrogen bonding network in resorcinarene hexamers typically involve bridging solvent molecules, which might even be responsible for their unique catalytic activity.<sup>4</sup> Both resorcinarenes and pyrogallarenes have demonstrated their ability to encapsulate various guest molecules in the voids of their capsular dimeric and hexameric assemblies.<sup>2,3</sup>

Pyridinearene<sup>5</sup> (1 in Scheme 1) is closely related to resorcinarene and pyrogallarene. Despite their obvious structural similarities, pyridinearenes display different binding properties due to amide–iminol tautomerism and electronic properties arising from the pyridine ring.<sup>6</sup> From the two tautomeric forms, dihydroxy and hydroxy-oxo, only the latter has been observed in crystal structures.<sup>6</sup> Recently, conformational analysis for the tautomers also revealed that the hydroxy-oxo

#### Introduction

Capsular supramolecular structures are tempting species, as they enable encapsulation of guest molecules and their voids may act as confined spaces suitable for nanocatalysis.<sup>1</sup> Resorcinarenes and pyrogallarenes (see Scheme 1), have been

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Scheme 1 Structures of tetraisobutyl-pyridinearene 1, tris(bipyridine) ruthenium(III) guest (G), resorcinarene and pyrogallarene. For pyridinearene two tautomeric forms are shown, but equilibrium is greatly favoring hydroxy-oxo tautomer.

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tautomer (Scheme 1) is  $\sim$ 150 kJ mol $^{-1}$  lower in energy than the dihydroxy tautomer.

The cavity of pyridinearene was earlier considered electron deficient and suitable to bind anionic guests. <sup>8,9</sup> Recently, instead of encapsulation of anions, the pyridinearene dimers were shown to actually favour encapsulation of neutral guests of matching size, and hence maximizing dispersion interactions. In fact, pyridinearenes bind anions, but outside the cavity, in *exo* position between the lower rim alkyl chains *via* CH···anion interactions. <sup>6</sup> Although the cavity of pyridinearenes is most suitable for neutral guests, also anion-driven encapsulation of cations inside the pyridinearene dimers was recently demonstrated. <sup>7</sup>

In one of the earliest reports concerning the self-assembly of pyridinearene, two species were observed in chloroform solution which were initially erroneously assigned as a mixture of monomers and dimeric assemblies. Cohen *et al.* later proved these as dimeric and hexameric species using diffusion ordered (DOSY) NMR spectroscopy experiments. In here, we report a detailed study of pyridinearene hexamer formation and structure in solution (NMR, DOSY NMR), in solid-state (X-ray crystallography) and in the gas phase (IM-MS, ESI-QTOF MS, DFT calculations).

#### Results and discussion

In the gas phase, by ESI-QTOF mass spectrometry, 1 typically ionizes well through deprotonation or anion complex formation (with PF<sub>6</sub>-, BF<sub>4</sub>- or I-). Despite many attempts, no traces of hexamer was detected in the negative mode. Similar hexameric capsules based on resorcinarene or pyrogallarene molecules have been earlier detected on positive mode by using a spherical and cationic transition metal complexes<sup>11</sup> as the stabilizing templates.<sup>3</sup> We, therefore, chose tris(bipyridine) ruthenium(II) (see Scheme 1, guest G) as a suitable complementary template. The cavity size for pyridinearene assemblies are roughly comparable to the size of corresponding assemblies of resorcinarenes and pyrogallarenes, but the affinity of pyridinearene towards cationic guests is lower. Nevertheless, when a sample of 1 (60 µM) with G (1:6) was sprayed from an acetone/dichloromethane (1:1 v/v) mixture, a host-guest hexameric assembly  $[\mathbf{1}_6 + \mathbf{G}]^{2+}$  was detected in the mass spectrum (Fig. 1). In addition, a pentameric host-guest  $[\mathbf{1}_5 + \mathbf{G}]^{2+}$  and a dimeric  $[\mathbf{1}_2 + \mathbf{H}]^+$  complexes were also observed, but the hexameric assembly was clearly the dominating ion in the spectrum (Fig. 1). Ion  $[\mathbf{1}_5 + \mathbf{G}]^{2+}$  likely originates from the hexamer through fragmentation. Assemblies larger than the hexamer, namely heptamers or octamers were not observed. The abundance of hexameric ion and lack of other aggregates in the spectrum suggest that hexamer could be a hydrogen bonded capsule rather than an unspecific aggregated system or a polymer.‡

To get actual proof whether the studied system is really a capsular assembly, ion mobility mass spectrometry (IM-MS) was utilized to acquire ion mobility arrival time distribution (ATD) and to determine the structure related collision cross

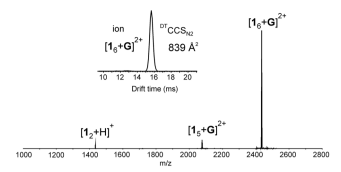


Fig. 1 (+)ESI-MS spectrum of 1 (60  $\mu$ M) and tris(bipyridine)ruthenium(II) (G) with 6:1 ratio measured from acetone/DCM mixture. Inset showing IMS arrival time distribution (ATD) for  $[1_6+G]^{2+}$  ion.

section (CCS) value for the hexamer. IM-MS has already demonstrated its power in structural analysis of different supramolecular systems. 12 In the ATD only a single wellresolved peak was observed. This implies that the corresponding  $[\mathbf{1}_6 + \mathbf{G}]^{2+}$  ion has a relatively rigid well-defined structure. When a drift tube ion mobility (DT-IMMS) instrument employing helium as a drift gas was used, an experimental DTCCS<sub>He</sub> value of 746.4 Å<sup>2</sup> was obtained.§ If a spherical structure is assumed, this corresponds roughly to a diameter of 3.1 nm for the assembly. The theoretical DTM CCSHe was calculated on basis of the DFT calculated structure by using diffuse trajectory method and yielded a DTMCCS<sub>He</sub> of 748.2 Å<sup>2</sup>, <sup>13</sup>¶ which is in good agreement with the experimental value. When more polarizable nitrogen was used as a drift gas, the  $^{DT}CCS_{N2}$  value was determined as 838.7 Å<sup>2</sup>. This roughly corresponds to a sphere with a diameter of 3.3 nm. The theoretical  $^{DTM}CCS_{N2}$  of 928.3  $\mathring{A}^2$  for the  $[\mathbf{1}_6 + \mathbf{G}]^{2+}$  ion indicate slightly bigger difference between experimental and theoretical values when N2 is used as a drift gas. In general, these CCS values are in good agreement with dimensions reported earlier hexameric capsules of pyrogallarenes resorcinarenes, 14,15 and thus, verify that the gas phase structure is indeed a capsular hexameric assembly.

The self-assembly of 1 was further studied in solution by <sup>1</sup>H and DOSY NMR (in CDCl<sub>3</sub> at 298 K). In a 20 mM solution of 1, two different species were observed in <sup>1</sup>H NMR spectra shortly after preparing the sample (Fig. 2a): one causing signals at 4.28 and 7.10 ppm (d/d'), and another (h/h') with signals at 4.36 and 7.22 ppm. The ratio of these two (d:h) was 2:1. The diffusion coefficient determined for assembly d was  $0.411 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$ , and  $0.287 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$  for assembly h. These values are in accordance with earlier results from Cohen, 10 and confirm that the signal sets d/d' and h/h' correspond to the dimer and the hexamer of 1. These values are in the expected range when compared with the diffusion coefficients of the corresponding assemblies of resorcinarene and pyrogallarene, which were used as internal standards. Assuming spherical shapes of these assemblies this translates into diameters of 2.0 nm for the dimer and 2.8 nm for the hexamer using the Stokes-Einstein equation (see ESI†). These

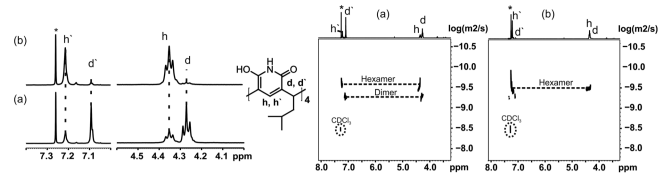


Fig. 2 Left: <sup>1</sup>H NMR of 1 (20 mM in CDCl<sub>3</sub>) (a) measured within 2 hours after sample preparation and (b) same sample after 7 days from sample preparation. Right: DOSY NMR of the same sample (a) measured within 2 hours form sample preparation and (b) after 7 days from sample preparation. Signals d and d' correspond to the dimeric assembly and signals h and h' to the hexameric assembly.

values are also well in line with the observed gas phase structures and determined CCS values. The DOSY NMR experiment in Fig. 2a was measured within 2 hours after sample preparation and shows the dimeric assembly as more abundant (2:1). The same NMR sample was stored and re-measured after seven days showing that the intensity of the signals for the dimer diminished dramatically, while at the same time signals for the hexamer increased and were now dominating (ratio d:h 1:10). This can also be seen in DOSY NMR, where only the signal of hexamer is observed after 7 days from sample preparation (Fig. 2b). This shows that pyridinearene forms indeed both dimeric and hexameric capsules in solution but the dimeric capsule is obviously only the kinetically favoured product while the hexameric capsule is the thermodynamic product, which abundance increases over time.

Single crystals suitable for X-ray diffraction analysis were obtained by slow vapour diffusion of diisopropyl ether into the solution of 1 in chloroform. Block shaped crystals were formed and the hexamer structure (16) in the solid-state could be confirmed by X-ray crystallography (Fig. 3a). The asymmetric unit contains three crystallographically independent molecules of 1, and the resultant self-assembled hexameric structure contains 24 O-H···O and 24 N-H···O hydrogen bonds. As a result of hydroxy-oxo tautomer form, the 6-hydroxyl position in 6-hydroxy-2(1H)-pyridone units manifest four O-H···O circular intramolecular hydrogen bonds to give 1 bowl-like  $C_{4v}$  conformation. The intermolecular N-H···O hydrogen bonding between the pyridone units in 16 provides a compact internal cavity with volume of 1160 Å<sup>3</sup>. This is smaller than corresponding hexameric capsules based on resorcinarenes and

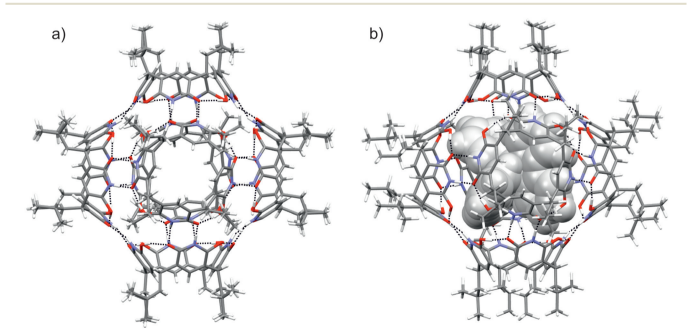


Fig. 3 X-Ray crystal structure of  $\mathbf{1}_6$  and (b) optimized structure of  $[\mathbf{1}_6 + \mathbf{G}]^{2+}$  at the RI-PBE-D3/def2-SVP level of theory in capped stick model. Tris (bipyridine)-ruthenium(II) (G) is shown in grey CPK model in (b). Black dashed lines represent hydrogen bonds. Color code: H = white, C = grey, N = blue, O = red.

pyrogallarenes (volumes >1200 Å<sup>3</sup>, see ESI Fig. S4†). 14,15 The DOSY derived hydrodynamic radii (2.80 nm) agrees well with the calculated value obtained from the X-ray crystal structure (2.82 nm). Of special note, the hydroxy-oxo tautomerism renders 1 a rigid cavity and definite directionality for hydrogen bonding. Robust  $C_{4v}$  cavities are seamed together with direct N-H···O intermolecular hydrogen bonding to produce solventfree hexamer 16. Solvent-free hydrogen bonding is also observed in the solid-state pyrogallarene hexamer structures.<sup>14</sup> In the resorcinarene hexamers, due to flexible  $C_{4v}$  cavities and steric reasons solvent molecules typically assist in capsule formation via intermolecular hydrogen bonding interactions, found between two resorcinarene units. 15 Although chloroform is essential to obtain a stable 16 single-crystals its passive role in hydrogen bonding could be observed in the 3-D crystal lattice. The features of pyridinearene lead to unique and complementary alignment of the individual monomers in order to maximize the number of intermolecular hydrogen bonds. In the case of the resorcinarenes, this requires incorporation of solvent molecules as part of the intermolecular hydrogen bond network. Consequently, this causes the differences in the void size for three different hexamers.

Unfortunately, all our attempts to obtain single crystals from  $[\mathbf{1}_6+\mathbf{G}]^{2^+}$  were unsuccessful. Therefore, we decided to carry out the computational analysis for studied systems at the RI-PBE-D3/def2-SVP//RI-PBE-D3/def2-TZVP level of theory in order to investigate the gas phase structure of  $\mathbf{1}_6$  and  $[\mathbf{1}_6+\mathbf{G}]^{2^+}$  and to evaluate their interaction energies in detail.  $^{16-24}$ 

The optimized structures and hydrogen bonding patterns of  $\mathbf{1}_6$  and  $[\mathbf{1}_6 + \mathbf{G}]^{2^+}$  are illustrated in Fig. 3b (see also ESI†). Within  $\mathbf{1}_6$  and  $[\mathbf{1}_6 + \mathbf{G}]^{2^+}$ , each monomer interacts with four neighbouring monomers through intermolecular N–H···O bonds whose distances vary from 1.703 to 1.716 Å and from 1.657 to 1.699 Å for  $\mathbf{1}_6$  and  $[\mathbf{1}_6 + \mathbf{G}]^{2^+}$ , respectively (Table 1). Interestingly, as compared to the dimeric capsule,  $\mathbf{1}_2$ , the N–H···O bonds are slightly shorter in the hexamer. The total number of N–H···O bonds in one hexamer unit is 24. Furthermore, the seam of hydrogen bonds, consisting of four intramolecular O–H···O bonds, of each monomer  $\mathbf{1}$ , is retained during hexamer formation. However, small changes are observed in the O–H···O bonds of  $\mathbf{1}$ ,  $\mathbf{1}_2$ ,  $\mathbf{1}_6$  and  $[\mathbf{1}_6 + \mathbf{G}]^{2^+}$  if compared with each other: the H-bonds are slightly longer in the hexamers and the dimers than in the monomer. The

**Table 1** Calculated O-H···O and N-H···O distances and interaction energies at the RI-PBE-D3/def2-SVP level of theory. RI-PBE-D3/def2-SVP//RI-PBE-D3/def2-TZVP single-point energies in parenthesis

Compound	O–H···O [Å]	N–H···O [Å]	$E_{\rm int}$ [kJ mol <sup>-1</sup> ]
1	1.425-1.432	_	_
$1_2$	1.519-1.525	1.800-1.821	$-280 (-197)^a$ $-942 (-717)^b$
<b>1</b> <sub>6</sub>	1.554-1.560	1.703-1.716	$-942(-717)^{b}$
$\begin{bmatrix} 1_6 \\ [1_6 + \mathbf{G}]^{2+} \end{bmatrix}$	1.528-1.563	1.657-1.699	$-1517(-1205)^{c}$

<sup>&</sup>lt;sup>a</sup>  $E_{\text{int}} = E_{\text{dimer}} - 2 \times E_{\text{monomer}}$ . <sup>b</sup>  $E_{\text{hexamer}} - 6 \times E_{\text{monomer}}$ . <sup>c</sup>  $E_{\text{complex}} - (6 \times E_{\text{monomer}} + E_{\text{guest}})$ .

elongation of the O–H···O bonds in the hexamers and dimers most likely arises from the small changes in electron density distribution around/on the oxygen atoms that function as a hydrogen bond acceptors in both, N–H···O and O–H···O, bonds in the hexamers and the dimers. Some electron density is therefore transferred from O–H···O bonds to N–H···O bonds upon the formation of dimers and hexamers. In the monomer, however, these acceptors oxygen atoms participates only in the formation of O–H···O bonds.

Due to the multiple intermolecular N–H···O bonds, 24 in total, the interaction energies of  $\mathbf{1}_6$  and  $[\mathbf{1}_6 + \mathbf{G}]^{2+}$  are very high -717 kJ mol<sup>-1</sup> and -1205 kJ mol<sup>-1</sup>, respectively (Table 1). These energy numbers suggest that the  $[\mathbf{1}_6 + \mathbf{G}]^{2+}$  is  $\sim 500$  kJ mol<sup>-1</sup> more stable than empty hexamer  $\mathbf{1}_6$  and underline the significance of template effect.<sup>3</sup> If the interaction energies of  $\mathbf{1}_6$  and  $[\mathbf{1}_6 + \mathbf{G}]^{2+}$  are compared to the interaction energies of 3 individual dimers  $\mathbf{1}_2$  (3 × -197 kJ mol<sup>-1</sup> = -590 kJ mol<sup>-1</sup>), it is clear that the formation of hexamer is more favourable than the formation of 3 individual dimers from six monomers. Although this very simplified gas phase model does not take into account all forces acting in the solution state, it not only supports the result obtained from MS studies, but is also in agreement with the NMR data: the hexamer  $\mathbf{1}_6$  is thermodynamically more stable than the dimer  $\mathbf{1}_2$  in gas phase and solution state.

### Conclusions

To conclude, the pyridinearene hexamers were structurally characterized in all three states of matter, and hence, for the first time in gas-phase and solid-state. X-ray structure shows that the hexameric assembly is formed via directly linked, intermolecular hydrogen bonds without bridging solvent molecules, which are usually required in the solid-state structures of closely related resorcinarenes. This makes hexamer more compact and decreases the size of empty windows between monomer units, which is a significant feature when applications, especially related to catalysis in confined space, are considered. In chloroform solution the hexamer was found to be the thermodynamically favoured product compared to the corresponding kinetically favoured dimer. All analytical techniques used result in a structure with a diameter of ~3 nm, and support the structural similarity of the pyridinearene hexamer in gas phase, solution and in solid state. As such, it is slightly more compact than the related hexamers of resorcinarenes and pyrogallarenes due to the different intermolecular hydrogen bonding network that implies to unique mutual alignment of the monomers in these assemblies. Interestingly, the use of a cationic, structurally complementary template increased the gas phase stability of the hexamer substantially, which could be rationalized by sophisticated DFT calculations.

Having established the structural properties of these large hydrogen-bonded capsules now paves the avenue to explore its host–guest binding properties arising from its different and unique interactions and geometric features compared to the other similar assemblies.

#### Conflicts of interest

There are no conflicts to declare.

# Acknowledgements

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#### Notes and references

‡Please note, that the template G is not necessary to observe the hexameric assembly in (+)ESI-MS. Without templating guest hexamer can be observed as  $\left[1_6+N_a\right]^+$  even in low (20  $\mu M$ ) concentration, but with relatively lower abundancy (see Fig. S1†).

 $\S^{DT}CCS_{He}$  states for CCS values obtained using drift tube instrument with helium gas, nomenclature recommended in V. Gabelica *et al.*<sup>25</sup>

 $\P^{\rm DTM}{\rm CCS}_{\rm He}$  states for CCS values obtained using diffuse trajectory method with helium gas.

 $\parallel$  According to Mason–Schamp equation, different gases result in different drift times, and thus to different CCS values. For more detailed explanation see V. Gabelica and E. Marklund.  $^{26}$ 

- D. Fiedler, D. H. Leung, R. G. Bergman and K. N. Raymond, *Acc. Chem. Res.*, 2005, 38, 349; M. D. Pluth, R. G. Bergman and K. N. Raymond, *Acc. Chem. Res.*, 2009, 42, 1650; L. Catti, Q. Zhang and K. Tiefenbacher, *Chem. Eur. J.*, 2016, 22, 9060.
- 2 M. Yamanaka, A. Shivanyuk and J. Rebek, J. Am. Chem. Soc., 2004, 126, 2939; L. Avram and Y. Cohen, J. Am. Chem. Soc., 2002, 124, 15148; L. MacGillivray and J. Atwood, Nature, 1997, 389, 469.
- 3 N. K. Beyeh, M. Kogej, A. Åhman, K. Rissanen and C. A. Schalley, *Angew. Chem., Int. Ed.*, 2006, **45**, 5214.
- 4 Q. Zhang, L. Catti and K. Tiefenbacher, *Acc. Chem. Res.*, 2018, **51**, 2107.
- 5 T. Gerkensmeier, J. Mattay and C. Näther, *Chem. Eur. J.*, 2001, 7, 465.

- 6 A. Kiesilä, L. Kivijärvi, N. K. Beyeh, J. O. Moilanen, M. Groessl, T. Rothe, S. Götz, F. Topić, K. Rissanen, A. Lützen and E. Kalenius, *Angew. Chem., Int. Ed.*, 2017, 56, 10942.
- 7 A. Kiesilä, J. O. Moilanen, A. Kruve, C. A. Schalley, P. Barran and E. Kalenius, Unpublished results.
- 8 A. B. Rozhenko, W. W. Schoeller, M. C. Letzel, B. Decker and J. Mattay, *New J. Chem.*, 2013, 37, 356.
- 9 M. C. Letzel, B. Decker, A. B. Rozhenko, W. W. Schoeller and J. Mattay, J. Am. Chem. Soc., 2004, 126, 9669.
- 10 T. Evan-Salem and Y. Cohen, Chem. Eur. J., 2007, 13, 7659.
- 11 G. Bianchini, A. Scarso, G. La Sorella and G. Strukul, *Chem. Commun.*, 2012, 48, 12082–12084.
- 12 E. Kalenius, M. Groessl and K. Rissanen, *Nat. Chem. Rev.*, 2019, 3, 4.
- 13 C. Larriba and C. J. Hogan, Jr., J. Comput. Phys., 2013, 251, 344; C. Larriba and C. J. Hogan Jr., J. Phys. Chem. A, 2013, 117, 3887.
- 14 T. Gerkensmeier, W. Iwanek, C. Agena, R. Fröhlich, S. Kotila, C. Näther and J. Mattay, Eur. J. Org. Chem., 1999, 2257.
- 15 R. M. Payne and C. L. Oliver, CrystEngComm, 2018, 20, 1919.
- 16 K. Eichkorn, O. Treutler, H. Öhm, M. Häser and R. Ahlrichs, Chem. Phys. Lett., 1995, 242, 652.
- 17 K. Eichkorn, F. Weigend, O. Treutler and R. Ahlrichs, *Theor. Chem. Acc.*, 1997, 97, 119.
- 18 K. Eichkorn, O. Treutler, H. Öhm, M. Häser and R. Ahlrichs, *Chem. Phys. Lett.*, 1995, **240**, 283.
- 19 J. P. Perdew and Y. Wang, Phys. Rev. B: Condens. Matter Mater. Phys., 1992, 45, 13244.
- 20 J. J. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1996, 77, 3865.
- 21 S. Grimme, J. Antony, S. Ehrlich and H. Krieg, *J. Chem. Phys.*, 2010, **132**, 154104.
- 22 S. Grimme, S. Ehrlich and L. Goerigk, *J. Comput. Chem.*, 2011, 32, 1456.
- 23 D. Andrae, U. Haeussermann, M. Dolg, H. Stoll and H. Preuss, *Theor. Chim. Acta*, 1990, 77, 123.
- 24 F. Weigend and R. Ahlrichs, *Phys. Chem. Chem. Phys.*, 2005, 7, 3297.
- 25 V. Gabelica, et al., Mass Spectrom. Rev., 2019, DOI: 10.1002/ mas.21585.
- 26 V. Gabelica and E. Marklund, Curr. Opin. Chem. Biol., 2018, 42, 51.