

Probing topology of supramolecular complexes between cyclodextrins and alkali metals by ion mobility-mass spectrometry

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- 1 Probing topology of supramolecular complexes between cy-
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- 3 etry
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ABSTRACT

In this study, the size and shape of supramolecular assemblies between cyclo-oligosaccharides and proton, ammonium or a series of alkali metals by electrospray coupled to trapped ion mobility-mass spectrometry (ESI-TIMS) have investigated. Native cyclodextrins (CD) were selected as models, and collision cross section (CCS) values were deducted for the main positive singly and doubly charged species. Experimental CCS values were in good agreement with those obtained from molecular modeling. Due to the high mobility resolving power and resolution, it was possible to highlight the presence of various conformers. Also, TIMS allowed to discriminate and estimate the content of various orientations from non-covalent nanotubes-based CD, involving secondary/secondary rim hydroxyl groups (head-to head), primary/secondary rim (head-to-tail) hydroxyl groups or primary/primary rim (tail-to-tail) hydroxyl groups interactions. Such results pave the way for a better knowledge of the topology of cyclo-oligosaccharides based supramolecular complexes, demonstrating that TIMS can be a particularly attractive molecular descriptor.

- Keywords: Cyclo-oligosaccharides; Cyclodextrins; Supramolecular assembly; Conformers; Ion-
- 26 mobility; Mass spectrometry.

1. INTRODUCTION

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Starch is among the most abundant polysaccharides present in the plant kingdom. It is composed of numerous glucose (Glc) units predominantly linked by α –(1 \rightarrow 4) glycosidic bonds and occasionally branched by α –(1 \rightarrow 6) moderately long chains.(Tester et al., 2004) The combined action of enzymes such as α -amylase and cyclodextrin glycosyltransferase, has attracted major interest from industry due to their unique capacity to catalyse the formation of α –(1 \rightarrow 4) malto-oligosaccharides as linear (maltodextrins) or cyclic ones (cyclodextrins; CDs) all with possible branching in α –(1 \rightarrow 6).(Ao et al., 2007; French et al., 1965; Taniguchi & Honnda, 2009; Terada et al., 1997) On the basis of the number of residues, commercially available CDs are torus-shaped cyclic oligosaccharides composed of 6, 7 and 8 α -(1 \rightarrow 4) linked D-glucopyranose units for α -CD, β -CD, and γ -CD, respectively. (Szejtli, 1998) CDs form a hydrophobic cavity able to encapsulate several organic molecules, (Kfoury et al., 2018; Song et al., 2009) and exhibit peripherally hydrophilic features forming intra- and inter-molecular networks thanks to hydrogen bonds. (Loftsson et al., 2005) Such particular properties is abundantly exploited to form various inclusion complexes in many industrial and research fields like food, flavours, cosmetics, drug delivery, catalyst or chiral selectors amongst other.(Davis & Brewster, 2004; Singh et al., 2002; Szejtli, 1998; Szente & Szemán, 2013) Also, the non-bonding electron pairs of the glycosidic oxygen bridge directed towards the inside of the cavity lead to a high electron density which lends CDs some Lewis base character.(Saenger, 1980; Szejtli, 1998; Wenz, 1994) Moreover, the 6-hydroxyl group of the primary rim can constitute very good attachment sites for various metals.(S. Angelova et al.,

2017; S. E. Angelova et al., 2017; Dossmann et al., 2021; Przybylski et al., 2015) One of the central concepts of molecular recognition is the issue of "selectivity", i.e. the preferential binding of one guest over another by the host. In this sense, one of the elusive structural characteristics when studying CD complexation is how the cavity shape changes upon complexation. Hence, the determination of effective complexation, as well as size and shape of host with and without any guest at a molecular level are mandatory conditions. In a common way, such piece of information was commonly accessed using NMR based diffusion experiments, (Schneider et al., 1998) X-ray diffraction analysis (Steiner & Koellner, 1994) and molecular modeling.(Quevedo & Zoppi, 2018) Nonetheless, the introduction of a new technology, named ion mobility coupled to mass spectrometry (IM-MS), offers a promising complementary tool to reach aforementioned objectives. IM-MS have successfully been applied to the gas-phase ion separation of a large range of (bio)molecules based isomers such as peptides/proteins, oligonucleotides, lipids, glycans, synthetic polymers or complexes. (Ben-Nissan & Sharon, 2018; Butcher et al., 2018; Charles et al., 2020; Dodds & Baker, 2019; Gray et al., 2016; Kalenius et al., 2019; Li et al., 2020; Zheng et al., 2018) Unfortunately, its potentiality in supramolecular assemblies investigations remains still underestimated. (Kalenius et al., 2019; Polewski et al., 2021) Such discrimination based on structural differences can be characterized by determination of their collision cross section (CCS) values in a given gas. Of special interest, IM-MS has been previously applied to CDs analysis using different instrumental configurations such as matrixassisted laser desorption or electrospray coupled to drift-tube ion mobility spectrometry (ESI-DTIMS). (Liu & Clemmer, 1997; Klein et al., 2018; Fenn & McLean, 2011; S. Lee et al., 1997)

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ESI with Traveling-Wave Ion Mobility Spectrometry (ESI-TWIMS) was used to study negatively charged mono- to trimeric β -CD(Berland et al., 2014), or positively charged complexes like amino- β -CD/sesamins, (Sugahara et al., 2015) α - and β -CD with o-, m- and p-coumaric acids, (Kralj et al., 2009) with piperine/curcumin, (Nag et al., 2018) or with amino acids. (S.-S. Lee et al., 2018; Chen et al., 2018). Unfortunately, the aforementioned studies with complexes involving at least two molecules reported a quite low/medium mobility resolution. A high resolution IM-MS prototype using serpentine ultra-long path was described for α -CD with bile acids analysis in negative and positive mode .(Chouinard et al., 2018) Very recently, a high resolution IM-MS instrument named trapped ion mobility spectrometry (TIMS) was commercially introduced. (Michelmann et al., 2015) TIMS operates at low electric field, preventing ion heating and exhibiting very high resolution. (Jeanne Dit Fouque & Fernandez-Lima, 2019; Ridgeway et al., 2018, 2019) Nevertheless, its application to carbohydrates analysis is still in infancy, although it was successfully applied for the analysis of glycosaminoglycan, (Wei et al., 2019) permethylated lacto-N-tetrasaccharides, (Pu et al., 2016) or separation of 13 isomeric trisaccharides. (Przybylski & Bonnet, 2021) In the study herein, we hypothesize that TIMS can be a useful analytical tool to both probe the influence of alkali metals on cyclo-oligosaccharides conformational modification (size/shape) in complexes, as well as to identify the sub-units orientation in supramolecular hierarchical assemblies and estimate the relative content of these last ones.

2. MATERIALS AND METHODS

2.1. Chemicals and Materials

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92 α-, β- and γ-Cyclodextrin (α-, β- and γ-CD, respectively) were kindly supplied by Wacker Chimie
 93 S.A.S. (Lyon, France). LiCl, NaCl, KCl, RbCl and CsCl alkali salts were purchased from Sigma-Aldrich
 94 (Saint-Quentin Fallavier, France).

2.2 Solvents

- 96 Methanol used for sample preparation was of MS grade and was purchased from VWR (West 97 Chester, PA, USA). Water was of ultrapure quality (18.2 $M\Omega$).
- **2.3 Samples**
- Stock solutions were made at 1 mM in water and then diluted to 1 μ M in methanol/water (1:1 v/v) with or without salts (0.3 μ M) for further analysis.
- **2.4 TimsTOF™ Experiments.**

We used ESI-timsTOFTM (Bruker Daltonics, Bremen, Germany) operating with oTOF control v5.0 software. The source temperature was hold at 200°C, and the drying and nebulizing gas (N₂) operate at a flow rate of 3 L. min⁻¹ and at a pressure of 0.3 bar. The instrument was calibrated using Tuning Mix G24221 (Agilent Technologies, Les Ulis, France). Applied voltages were +4 kV and -0.5 kV for capillary and endplate offset, respectively. Acquisition was achieved in the m/z 400-4000 range with a centre at m/z 500. TIMS separation depends on the gas flow velocity (v_g), elution voltage ($V_{elution}$), ramp time (t_{ramp}), base voltage (V_{out}) and the electric field (\vec{E}). The reduced mobility, K_0 , can be calculated as follows:

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$$K_0 = \frac{v_g}{\vec{E}} = \frac{A}{(V_{elution} - V_{out})}$$
 (Eq. 1)

The mobility calibration constant A was determined using known reduced mobilities of tuning mix components. The resolving power (R) and resolution (r) are defined as $R=(1/K_0)/w$ and $r=1.18\times[(1/K_0)_2-(1/K_0)_1]/(w_1+w_2)$, where w is the full peak width at half-maximum. To improve separation efficiency, scan rate $(Sr=\Delta V_{\rm ramp}/t_{\rm ramp})$ was tuned thanks to imeXTM technology. For this, t_{ramp} is automatically set as function of manually adjusted ΔV_{ramp} . N₂ was used as buffer gas at funnel temperature (T = 305 K) with v_g set by the pressure difference of 1.69 mbar. A potential of 350 Vpp was applied to radially confine the trapped ion cloud. The measured inverse reduced mobilities were converted into collision cross sections (CCS) using the Mason-Schamp equation:

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$$\Omega = \frac{(18\pi)^{1/2}}{16} \times \frac{q}{(k_B \times T)^{1/2}} \times \left[\frac{1}{m_i} + \frac{1}{m_q}\right]^{1/2} \times \frac{1}{N} \times \frac{1}{K_0} \text{ (Eq. 2)}$$

where q is the ion charge, k_B is the Boltzmann constant, N is the gas number density, m_i is the ion mass, and m_g is the gas molecule mass. TIMS-MS spectra and mobilograms were analysed using Compass Data Analysis 5.1 (Bruker).

2.6 ESI-TIMS-MS analysis of the cyclo-oligosaccharides

Throughout this study, all cyclo-oligosaccharides were analysed in the positive ion mode with or without salt doping. All samples were continuously infused at 3 μ L.min⁻¹ via a 250 mL syringe.

2.7 Theoretical Collision Cross Section Calculations

All initial geometry relaxations were performed using the Merck molecular force field (MMFF94) implemented in Avogadro (v1.95.1). Geometry optimization was finalized using density functional theory (DFT) as previously described (Dossmann et al., 2021) and calculations with NWChem (v7.0). Theoretical CCS calculations were carried out in IMoS (v.1.10)(Larriba & Hogan, 2013) using the average of ten processes by Elastic/Diffuse Hard Sphere Scattering or Trajectory method Diffuse Hard Sphere Scattering.⁴⁶

3. RESULTS AND DISCUSSION

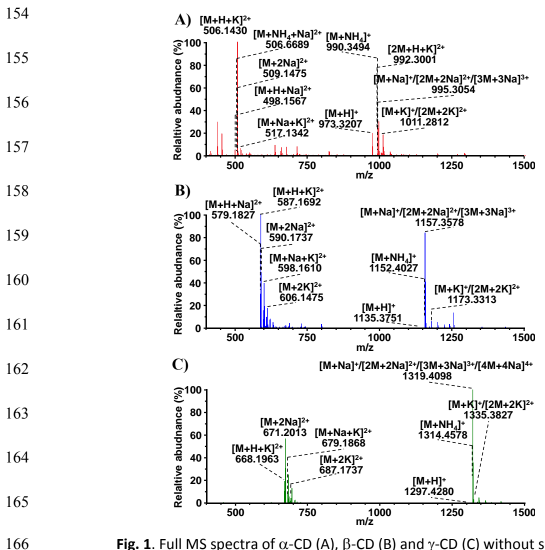
As all neutral carbohydrates molecules, cyclodextrins exhibit intrinsically a strong affinity towards usual alkali metals. (Frański et al., 2005; Metzger et al., 1991; Przybylski et al., 2015; Przybylski & Bonnet, 2013; Reale et al., 2005) We have previously demonstrated that a solution containing an equimolar mixtures of β -CD and LiCl, NaCl and KCl allowed to determine apparent solution selectivity and the relative affinity of various complexes. (Przybylski et al., 2015) However, here we only aimed to study size and conformation of the complexes by IM-MS, and not the abundance. In this sense, as glassware provides enough source of Na⁺ and K⁺, we have directly acquired a first series of spectra with separate α -, β - and γ -CD without any addition. In a second series, α -, β - or γ -CD was mixed with each alkali metals.

3.1 Mobility of singly charged monomers

3.1.1 Protonated and ammoniated native cyclodextrins

Cyclodextrins present a lower affinity for H^+/NH_4^+ as compare to usual alkali metals. However, the gas phase basicity of glucose is reported to be 188±3 kcal.mol⁻¹.(Jebber et al., 1996) Hence,

even without any doping, ESI full spectra reveal the presence of such adducts coupled as singly ions at m/z 973.3207/990.3494, 1135.3751/1152.4027 and 1297.4280/1314.4578 for α -, β - and γ -CD, respectively (Fig. 1).



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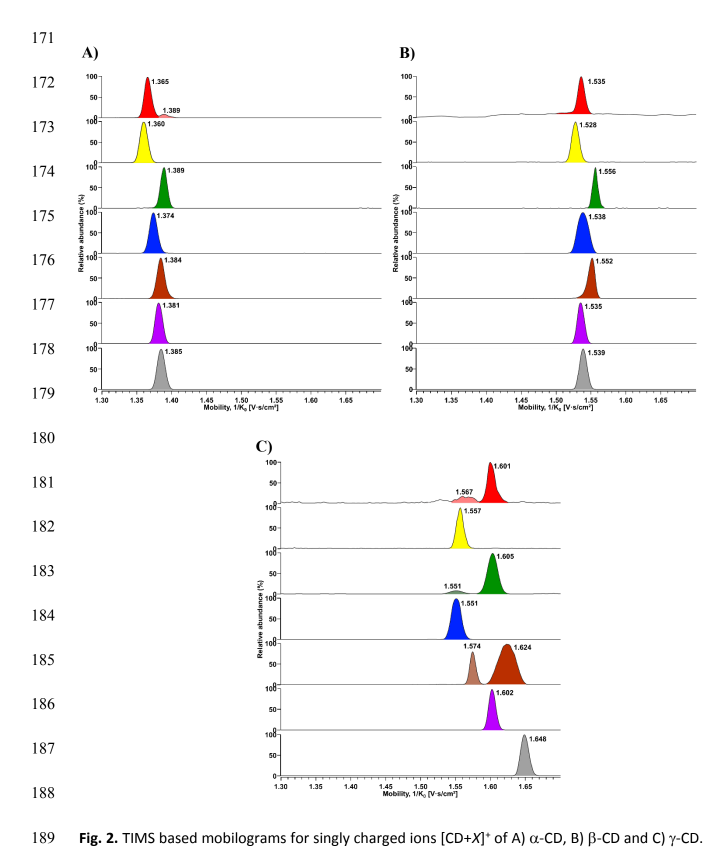
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Fig. 1. Full MS spectra of α -CD (A), β -CD (B) and γ -CD (C) without salt doping.

Examination of the respective extracted mobilograms leads to a couple of prominent/minor mobility peaks for α -CD and γ -CD with 1/K₀ values of 1.365/1.389 and 1.601/1.567 V.s/cm², respectively (Fig. 2).



Color coding for X: H (red), Li (yellow), NH₄⁺ (green), Na (blue), K (brown), Rb (violet) and Cs (grey).

Conversely, β -CD presents a single mobility peak at 1.535 V.s/cm² reflecting a unique stabilized location of the proton. Regarding ammoniated adducts, α -/ β -CD present a single peak with 1/K₀ values equal to 1.389 and 1556 V.s/cm², respectively, while for γ -CD, a couple of values at 1.605/1.551 V.s/cm² were observed (Fig. 2). Such result may be explained by a more pronounced flexibility leading to two distinct complexes [γ -CD+NH₄]⁺.

3.1.2 Native cyclodextrins with alkali metals (Li, Na, K, Rb and Cs)

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Native CDs exhibit lone pairs which greatly favours the attachment and stabilization of alkali metals. Mobility values of singly charged α -CD gradually increase from lithium to sodium (1.360 to 1.374 V.s/cm²), and again with potassium (1.384 V.s/cm²), and then slightly decrease for rubidium (1.381 V.s/cm²) before to be barely higher with cesium (1.385 V.s/cm²) (Fig. 2). As expected, higher mobility values for β -CD than α -CD were measured with all alkali metals. Moreover, the same increasing trend was observed for lithium, sodium and potassium (1.528, 1.538 and 1.552 V.s/cm², respectively). Conversely, lower values than potassium were measured with rubidium and cesium (1.535 and 1.539 V.s/cm², respectively). Regarding γ -CD, except higher values than β -CD, it can be noted a significant difference in the evolution of mobility values that the two previous CDs. If $[\gamma$ -CD+Li]⁺ and $[\gamma$ -CD+Na]⁺ are quasi-similar (1.557 and 1.551 V.s/cm², respectively), at least two distinct peaks at higher mobility values were observed for $[\gamma-CD+K]^+$ (1.624/1.539) V.s/cm², with the latter detected as a twofold lower abundance). Similarly to α -CD and as compared to main $[\gamma-CD+K]^+$ value, mobility peak of $[\gamma-CD+Rb]^+$ is lower (1.602 V.s/cm²) whereas that of $[\gamma-CD+Cs]^+$ is the highest (1.648 V.s/cm²). From these data, and except for $[\gamma-CD+K]^+$, a single mobility peak is systematically obtained for singly charged species, presumably corresponding to a unique and stable complex.

3.2 Mobility of doubly charged monomers

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The $[\alpha$ -CD+H+Na]²⁺ and $[\alpha$ -CD+H+K]²⁺ ions lead to a close mobility value of 0.798 and 0.791 215 V.s/cm²), respectively. With lithiated complexes ($[\alpha$ -CD+Li+X]²⁺), mobility values were lower at 0.747, 0.763 and 0.759 V.s/cm², for X = Li, Na and K, respectively. Interestingly for both [α -CD+2Na]²⁺ or $[\alpha$ - CD+Na+K]²⁺, two couples of mobility peak were detected at 0.803/0.795 and 218 0.790/0.810 V.s/cm², respectively. Such observation suggests the existence of two distinct ter-219 nary complexes for each ion. None doubly charged species based α -CD involving two potassium 220 or even one/two rubidium and cesium was detected. This is presumably due to higher ionic radii (r_i = 137, 152 and 167 pm for K⁺, Rb⁺ and Cs⁺, respectively)(Haymes, Lide & Bruno, 2016) impairing a good matching, with the coordination sites from this cyclo-oligosaccharide template. As ex-223 pected, the mobility values $[\beta-CD+H+X]^{2+}$ with X=Na and K are higher than for $\alpha-CD$. Nevertheless, no significant differences occurs for the two herein β -CD species with 0.854 and 0.855 225 V.s/cm², respectively. Such results show that $[\beta-CD+H+K]^{2+}$ as compact as $[\beta-CD+H+Na]^{2+}$ suggest-226 ing similar coordination sites and/or flexibility of the CD. Introduction of lithium in ternary complex leads to a single mobility peak at 0.801, 0.814 and 0.830 V.s/cm², for [β-CD+2Li]²⁺, [β-CD+Li+Na]²⁺ and [β -CD+Li+K]²⁺, respectively. Similarly, to α -CD, for adducts with 2Na and 1Na+K, two couples of mobility peaks were observed at 0.852/0.820 and 0.828/0.851 V.s/cm², respectively. Interestingly, the β -CD allows the formation of both $[\beta$ -CD+2X]²⁺ with X = K or Rb exhibiting two distinct mobility values (0.869/0.839 V.s/cm²) for the former and a unique for the latter (0.876 V.s/cm²). Concerning γ -CD, as for β -CD, only a single peak was observed for [γ -CD+2Li]²⁺, 233 $[\gamma-CD+Li+Na]^{2+}$, $[\gamma-CD+Li+K]^{2+}$, $[\gamma-CD+Na+Rb]^{2+}$ and $[\gamma-CD+2Rb]^{2+}$ at 0.826, 0.852, 0.873, 0.900 and 0.878 V.s/cm², respectively. Nonetheless, instead of a unique peak for β -CD, $[\gamma$ -CD+ H+Na]²⁺ and 234

 $[\gamma-CD+K+Rb]^{2+}$, showed two peaks at 0.903/0.879 and 0.892/0.915 V.s/cm², respectively. As regards other doubly charged species, very different behaviours were quoted out for γ -CD compare to both α - and β - ones. Indeed, three distinct peaks were detected for $[\gamma\text{-CD}+X+K]^{2+}$ with X=H or K at 0.873/0.907/0.847 and 0.897/0.855/0.876 respectively. For $[\gamma-CD+Na+X]^{2+}$ with X=Na or K, only one value can be observed at 0.893 and 0.866 V.s/cm², respectively. Conversely to α -/ β -CDs, complexes involving cesium have been successfully detected for γ -CD with only one value for [γ -CD+Na+Cs]²⁺ (0.901 V.s/cm²) and $[\gamma$ -CD+2Cs]²⁺ (0.882 V.s/cm²) and three for $[\gamma$ -CD+K+Cs]²⁺ $(0.876/0.890/0.899 \text{ V.s/cm}^2)$. These last results support the hypothesis that β -CD is the smaller CD able to simultaneously attach two potassium. Also, a minimal number of 7 and 8 glucoses is required to complex two simultaneous potassium and rubidium or cesium based doubly charged species, respectively. Moreover, it was previously observed that the coulombic repulsion between two cations in the ternary complex is reduced for the larger size host. (Wang et al., 2010) Furthermore, two, three and even four mobility peaks can be detected for a given doubly charged ion. That strongly suggests the simultaneous occurrence of several more and less stable complex, which can be estimated by area integration of mobility peaks.

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3.3 Scaling cyclodextrin/metals complexes using the collision cross section as molecular descriptor

After determination of ion mobility values of a given ion and by using Mason-Schamp equation (see experimental section), its collision cross section (CCS) can be deducted. The relationship between structure and CCS has been widely used to offer insights into the structures of gas phase species. More particularly, CCS provide precious pieces of information regarding size and shape of the corresponding molecules. In the case of CDs, we used IM-MS to probe both size difference

informing on location of coordination site including more and less deep insertion in the cavity, and on the shape feature which can be due to the existence of various conformational states during cationization.

3.3.1 Singly charged ions

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Protonated α -, β - and γ -CD have CCS_{N2} equal to 276.8/282.2, 311.1 and 323.8/317.2 \mathring{A}^2 , respectively. Herein values are very close to those reported in the literature (Table 1). (Klein et al., 2018; May et al., 2014) Nonetheless, we noted that herein, α - and γ -CD exhibits two and three distinct mobility peaks respectively, which can be putatively ascribed to different conformational states or protomers. This was obtained thanks to the high resolution of TIMS while unique value was extracted from DTIMS analysis. (Klein et al., 2018) Regarding ammonium adducts, an increase of CCS according to size was observed with 282.1, 315.4 and 324.4/313.9 \mathring{A}^2 for α -, β - and γ -CD, respectively. However, we noted none linearly correlation since we could expect 348.7 Å² for a not deformed γ -CD, but obtained values lower by ~24-30 Å². From these last results, two hypotheses can be formulated either ammonium ion can be more deeply inserted into the γ -CD cavity and/or γ-CD can adopt deformed conformations due to its intrinsic flexibility. In all case, the comparison of any given ion form as a function of CD type shows an expected increase in CCS as a function of increasing molecular weight and/or size (e.g. with [M+Li]+, CSS is 276.1, 309.6 and 315.1 Å^2 for α -, β - and γ -CD, respectively (Table 1 and Fig. S1).

Table 1. Characteristics of singly charged ions from α -, β - and γ -CD under alkali-metal cation complexes. Brackets indicate the relative content. *: large peak, ND: not detected; NC: not calculated.

Name	m/z		Mass accuracy	1/K ₀ (V.s.cm ⁻²)	CCS _{N2} (Å ²)		
	Experimental	Theoretical	(ppm)	1/K ₀ (V.S.CIII ⁻)	Experimental (n=5)	Theoretical (n=10)	Literature
[α-CD+H] ⁺	973.3207	973.3242	-3.6	1.365 /1.389 (92/8)	276.8±0.6/282.2±0.5	276.8±0.8/281.8±0.7	280.5a;285.2b
[α-CD+Li] ⁺	979.3346	979.3325	2.2	1.360	276.1±0.4	276.3 ± 0.6	
$[\alpha\text{-CD+NH}_4]^+$	990.3494	990.3508	-1.4	1.389	282.1±0.7	282.2 ± 0.8	280.9^{a}
$[\alpha\text{-CD+Na}]^+$	995.3054	995.3062	-0.8	1.374	278.9 ± 0.4	278.8 ± 0.5	278.2a;285.5b
$[\alpha\text{-CD+K}]^+$	1011.2812	1011.2801	1.1	1.384	281.0±0.5	280.9 ± 0.6	280.0a;287.7b
$[\alpha\text{-CD+Rb}]^+$	1057.2269	1057.2282	-1.2	1.381	280.2 ± 0.8	280.2 ± 0.8	
$[\alpha\text{-CD+Cs}]^+$	1105.2208	1105.2218	-0.9	1.385	280.8±0.9	281.2±0.8	
[β-CD+H] ⁺	1135.3751	1135.3770	-1.7	1.535	311.1±0.4	311.5±0.6	312.4a; 301.3/319.6b
[β-CD+Li] ⁺	1141.3843	1141.3853	-0.9	1.528	309.6 ± 0.6	309.9 ± 0.8	
[β-CD+NH ₄] ⁺	1152.4027	1152.4036	-0.8	1.556	315.4±1.1	315.3±0.6	312.5a
[β-CD+Na] ⁺	1157.3578	1157.3590	-1.0	1.538	311.7±0.6	311.8±0.4	309.0°;319.7°
[β-CD+K] ⁺	1173.3313	1173.3329	-1.4	1.552	314.4 ± 0.7	314.6 ± 0.7	308.9 ^a ;320.3 ^b
$[\beta\text{-CD+Rb}]^+$	1219.2796	1219.2810	-1.2	1.535	310.9±0.6	311.2±1.0	
$[\beta\text{-CD+Cs}]^+$	1267.2733	1267.2747	-1.1	1.539	311.5±0.8	311.3±1.1	
[γ-CD+H] ⁺	1297.4280	1297.4299	-1.4	1.601/1.567 (81/19)	323.8±0.8/317.2±0.9	323.6±0.7/317.3±0.9	324.3 ^a ;322.6 ^b
[γ-CD+Li] ⁺	1303.4372	1303.4382	-0.7	1.557	315.1±0.6	314.9 ± 0.5	317.7 ^b
$[\gamma$ -CD+NH ₄] ⁺	1314.4578	1314.4564	1.1	1.605/1.551 (93/7)	324.4/313.9±0.7	324.3±0.9/314.0±1.3	324.3a;
[γ-CD+Na] ⁺	1319.4098	1319.4118	-1.5	1.551	313.9 ± 0.8	313.8±1.0	316.0 ^a ;322.1 ^b
$[\gamma\text{-CD+K}]^+$	1335.3827	1335.3857	-2.3	1.624*/1.574 (77/23)	328.5±0.8/318.8±1.2	328.8±1.1/318.6±1.2	319.3°,324.8°
$[\gamma$ -CD+Rb] ⁺	1381.3364	1381.3383	-1.4	1.602	324.0±1.0	324.0±1.3	327.3 ^b
[γ-CD+Cs] ⁺	1429.3258	1429.3275	-1.2	1.648	333.3±1.1	333.2±1.3	338.2 ^b

^a Klein et al., 2018

^b May et al., 2014

Further examination of complexes involving alkali metals, allows achievement of some observations: i) with α -CD, CCS increase linearly by step of 2.8 and 2.1 Ų from Li† to Na† and from Na† to K†, respectively, and then remains quite constant with Rb† and Cs† (Fig. S1, red star) ii) with β -CD, CCS increase linearly also by step of 2.1 and 2.7 Ų from Li† to Na† and from Na† to K†, respectively, but decrease by \sim 3.5 A² for Rb† and drop only slightly for Cs† (Fig. S1, blue square). iii) with γ -CD, measured CCS portray a completely different behaviour as a slight decrease by \sim 1.2 A² occurred from Li† to Na† and then a drop by 4.9 or 14.6 A² for K† as two peaks were detected. Next, according to previous considered values for K† a variation of -4.5/+5.2 A² occurred for Rb† and a drop by 9.3 A² from this last one to Cs† (Fig. S1, green circle). Also, such a decrease in CCS observed with

CDs and some alkali metals compared to corresponding protonated or ammoniated forms, can be tentatively explained by a greater oxophilicity - stronger coordination of the formers, as previously described for carbohydrates.(Fenn & McLean, 2011) In brief, CCS can be summarized as a consequence of a balance between main factors: size of both cation and CD (including eventual distortions) and more and less deep inclusion of the cation inside cavity (according to coordination features).

3.3.2 Doubly charged ions

Considering the sum of the relative intensities from all alkali metal cations-based CDs observed by MS, the part of 1:2 complexes compared with 1:1 one increased constantly from α -CD to γ -CD. Such trend was similar to that reported elsewhere for an increasing methylation degree on β -CD.(Przybylski et al., 2015) As for singly charged species, any particular ternary complexes the higher the molecular weight and/or size of CD, the higher the CCS (e.g. with [M+2Li]²⁺, 307.6, 328.6 and 337.9 Å² for α -, β - and γ -CD, respectively) (Table S1 and Fig. 3).

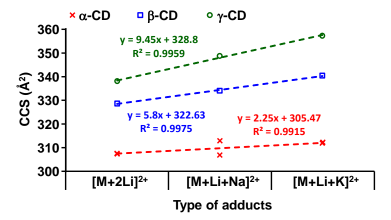


Fig.3. Evolution of CCS as function of doubly charged species $[CD+Li+X]^{2+}$ where X is lithium, so-dium or potassium. α -CD (red cross), β -CD (blue square) and γ -CD (green circle).

Also, interestingly, for a given CD involving both one lithium and in the other hand an additional lithium, sodium or potassium, CCS increase linearly with the size of the associated cation. Examination of the slope value of the resulting curve i.e. 2.25, 5.8 and 9.45 portrays a growing amplitude in CCS drop for α -, β - and γ -CD, respectively (Fig. 3). However, such behaviour according size of the cations was not obvious for all ternary complexes, even if a relative growing linearity of CCS can be deducted for $[\alpha$ -CD or β -CD+Na+X]²⁺ and $[\alpha$ -CD or β -CD+K+X]²⁺ (where X = Li, Na and K) as well as for $[\gamma-CD+K+X]^{2+}$ $[\gamma-CD+Na+X]^{2+}$ (where X=Li, Na, K and Rb) (Fig. S2-S5). We have previously demonstrated by molecular modeling that two cations are stabilized by various coordination sites onto the primary rim of β-CD.(Przybylski et al., 2015) For example, ternary complexes with native β-CD including 2 Li are stabilized each one by three electron donating. Such coordination landscape involves 3 oxygens of the primary rim of n, n+1 and n+3 glucose for one side and n-1, n-2 and n-3 for the second side. (Przybylski et al., 2015) The lower bond length between the three proximal oxygens of doubly lithiated forms highlighted a tighter interaction of the two metals than for corresponding singly charged complexes. Moreover, the coordination of two cations induces a strong distortion of CDs like a spindle with cation localized at each tip. Increasing CCS can be roughly visualized using Van der Waals (WDV) radii for each atom of the molecule, in other words VDW surface through which the molecule might be conceived as interacting with other ones. Examples of VDW surface were given for the three axis of $[\beta-CD+Li+X]^{2+}$ (where X = Li, Na or K) (Fig. 4).

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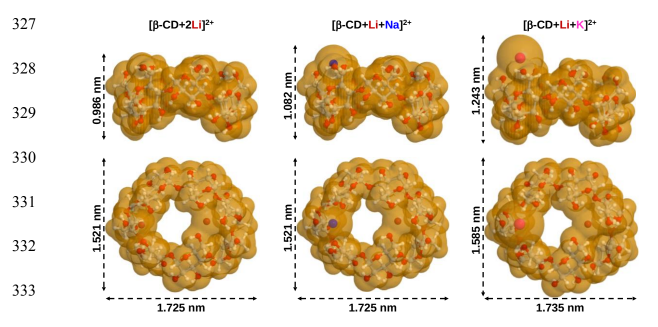


Fig. 4. Van der Waals surface of $[\beta\text{-CD+Li}+X]^{2+}$ (where X = Li, Na or K) from ball and stick model of optimized models of complexes with x,y axis (bottom view) and z-axis side (top view).

Such last examples depict well the possible differences portraying the linear increase of the CCS values where only length on the z axis increases while those of x, y axis remain constant from $[\beta-CD+Li+Li]^{2+}$ to $[\beta-CD+Li+Na]^{2+}$ and then length of the three axis enlarge from $[\beta-CD+Li+Na]^{2+}$ to $[\beta-CD+Li+K]^{2+}$.

3.4 Study of multimeric assemblies of cylodextrins

Literature described that CD can form aggregates with various shapes but based on only 2 types of molecular arrangements: clustered cage or channel.(Saenger et al., 1998) Nevertheless, probability to form disordered assemblies prevails from concentration in the millimolar

range.(Messner et al., 2010) Nevertheless, in our study, we used diluted solution (one micromolar) which must favour formation of ordered assemblies. All the CD have the shape of a basketball net, including a large (head, H) and a small (tail, T) aperture. In tubular arrangements of CDs, they can interact among themselves according to three possible orientations: Head-to-Tail (H $^{\bullet}$ T), Head-to-Head (H $^{\bullet}$ H) or Tail-to-Tail, (T $^{\bullet}$ T).(Staelens et al., 2015) Such organization is the result of the formation of several intermolecular hydrogen bonds constituting the main driving force for holding and stabilizing two or more units together.(Nascimento et al., 2005) As example, we have investigated the ion mobility traces of the singly charged ion at m/z 1967.6199 ascribed to [(α -CD)₂+Na] $^{+}$ and deducted corresponding CCS values (Fig. 5).

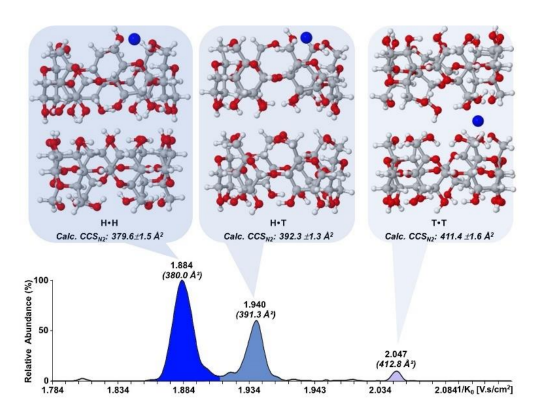


Fig. 5. TIMS based mobilogram for the singly charged dimer of α -CD ([(α -CD)₂+Na]⁺) revealing the three orientations as H•H (bright k blue), H•T (steel blue) and T•T (clear blue).

Examination of the mobilogram revealed two main peaks at 380.0 and 391.3 Å² and one minor one at 412.8 Å². These values are in very good agreement with theoretical CCS calculated for H•H $(379.6\pm1.5 \text{ Å}^2)$, H•T $(392.3\pm1.3 \text{ Å}^2)$, and T•T $(411.4\pm1.6 \text{ Å}^2)$ assemblies, respectively. Such results demonstrated that TIMS can successfully discriminate the three possible orientations of mono sodiated dimers of CDs. Efficiency of separation can be defined with resolving power (R) of a given mobility peak and also with peak-to-peak resolution (r). Here, the values were equal to 109/141/282 and 1.8/5.1 depicting both high R and r. Also, by approximating that equal ionization efficiencies take place and that relative abundance of the overall spectrum could be interpreted as a good image of the solution content, we can estimate content of mono-sodiated species of H•H, H•T and T•T α-CD dimers as 59, 35 and 6%, respectively. Such content is in good agreement with stability order obtained from molecular dynamics (MD) simulations (T•T < H•T < H•H). (Bonnet et al., 2001) Regarding doubly charged species with the two same cations, the nature of the cation leads to contrasted results in terms of both r and R, which rely on their ability to detect and also to discriminate the different orientations as well as their respective content. Results for $[(\alpha-CD)_2+2X]^{2+}$ with X=Li, Na, K, Rb or Cs were given in Fig. 6.

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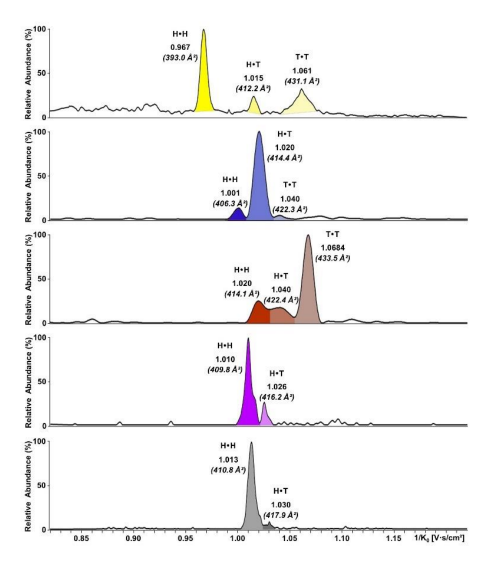


Fig. 6. TIMS based mobilograms for the doubly charged dimer of α -CD ([(α -CD)₂+X]²⁺) revealing the possible orientations (H•H, H•T and T•T). Color coding for X: Li (yellow), Na (blue), K (brown), Rb (violet) and Cs (grey).

In details, double adduction with lithium allows to easily highlight the presence of the three possible orientations (H $^{\bullet}$ H, H $^{\bullet}$ T and T $^{\bullet}$ T) of α -CD dimer with both R equal to 138, 127 and 71, respectively, and r of 3.2 (H $^{\bullet}$ H to H $^{\bullet}$ T) and 2.0 (H $^{\bullet}$ T to T $^{\bullet}$ T). With two sodium or two potassium, r strongly decreases to 0.8/1.0 and 0.5/0.8 for H $^{\bullet}$ H to H $^{\bullet}$ T/H $^{\bullet}$ T to T $^{\bullet}$ T, respectively. The same trend

was noted regarding R with values for the mobility peak of the three orientations of 80/85/145 and 65/47/92, respectively. Introduction of rubidium lead to detection of only two orientations i.e. $H ilde{ } H ilde{ } H ilde{ } T$ with satisfactory R (141/174) and r (1.2). Such data strongly suggest that the $T ilde{ } T$ dimer of $[(\alpha-CD)_2+2Rb]^{2+}$ cannot be formed and/or stabilized in the gas phase. Similar results were obtained with cesium leading to the two dimers: H•H and H•T with close R (133/138) and r (1.1), and a lesser content of the latter orientation as compared to Rb (9 vs 22%). Interestingly, R and r from $[(\alpha-CD)_2+2Na]^{2+}$ are both lower than $[(\alpha-CD)_2+Na]^+$ (109/141/282 vs 80/85/145 and 0.8/1.0 vs 1.8/5.1). Moreover, difference in CCS of the latter vary from 11.3 Å² and 21.5 Å² for H•H to H•T and H•T to T•T, respectively, while it is constant (8 Å²) for the former. Such results support the hypothesis that the presence of two sodium acts as a more structuring agent than only one, presumably by compacting dimer to minimize natural intrinsic difference resulting from the different orientations. We have demonstrated that for a given CD, the type of adducted cations strongly influences the discrimination ability, but a given cation do not act with the same beneficial/detrimental balance according to the studied CD. For example, the smallest alkali metal studied herein, lithium led to the high-resolution during analysis of $[(\alpha-CD)_2+2Li]^{2+}$ (Fig. 6), while $[(\beta-CD)_2+2Li]^{2+}$ and $[(\gamma-CD)_2+2Li]^{2+}$ only revealed two dimers namely H•H and H•T exhibiting R equal to 124/88 and 160/123, respectively, but also with r of 3.5 and 2.1, respectively (Fig. S6). Conversely, with the largest studied alkali metal, cesium, $[(\alpha-CD)_2+2Cs]^{2+}$ exhibits only H•H and H•T dimers (R: 133/138 and r: 1.1) while the analysis of $[(\beta-CD)_2+2Cs]^{2+}$ unambiguously showed the presence of the three possible orientations H•H, H•T and T•T (R: 286:82/129 and r: 1.4/1.3) respectively (Fig. S7). Finally, a single peak with an important tailing, leading to low R (96), was obtained for $[(\gamma-CD)_2+2Cs]^{2+}$. Such peak was mainly ascribed to H•T by computational methods,

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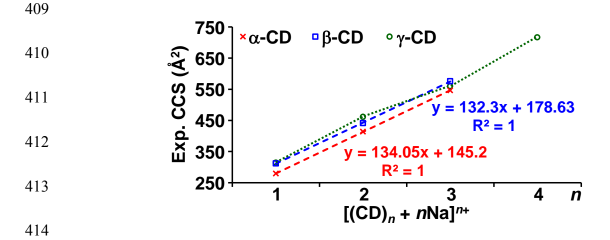
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and the tailing is presumably due to the detection and/or separation difficulty of the other dimers, due to aforementioned high distortion ability of γ -CD and existence of several conformers. To explore the influence of the number (n) of CDs/charge state couple on the formation of small oligomers, we draw experimental CCS as function of $[(CD)_n + nNa]^{n+}$ (Fig. 7).



ular assemblies (Fig. 8A-B).

where n is both the number of CDs, the number of adducted sodium and the charge state. α -CD (red cross), β -CD (blue square) and γ -CD (green circle). In the case of a non-driven CDs aggregation, resulting CCS should be randomly distributed. Conversely, in the case of well-ordered supramolecular assemblies with a tubular topology assimilated to a cylinder or a compact near-spherical nucleation, expected CCS fitting should be near a fully linear or involving a modular exponentiation.(Hanozin et al., 2019; Ruotolo et al., 2008) Herein, we obtain a linear fit for α - and β -CD for 1- to 3-mer portraying both a well-organized into a hierarchical assemblies consecutive to stacked CDs and a structuring role of sodium. Such observations were confirmed by theoretical CCS calculation of single CDs and their supramolec-

Fig. 7. Evolution of the experimental collision cross section as a function of $[(CD)n+nNa]^{n+}$ ions

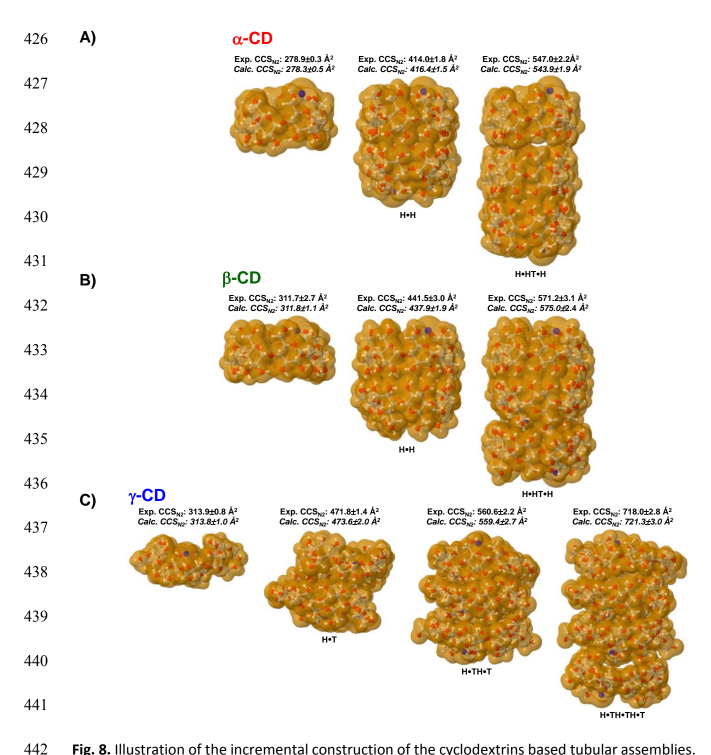


Fig. 8. Illustration of the incremental construction of the cyclodextrins based tubular assemblies. Van der Waals surface from Ball and stick model of optimized models (A) $[(\alpha-CD)_n+nNa]^{n+}$, (B) $[(\beta-CD)_n+nNa]^{n+}$ and (C) $[(\gamma-CD)_n+nNa]^{n+}$ (where n=1,2,3 or 4 for mono-, di-, tri and tetramer). Experimental (n=5) and theoretical (n=10) collision cross section are indicated above structures.

 γ -CD and its short oligomers containing until 4 units show a different behaviour. Firstly, based on a semi-rigid conformation behaviour similar to α - and β -CD, expected CCS of $[\gamma$ -CD+Na]⁺ should be $\approx 348.7 \text{ Å}^2$, but an experimental value of 313.8 \pm 1.0 Å² was obtained. Such discrepancy can be tentatively explained by a distortion of γ -CD. Indeed, it was mentioned in literature that when size of CD increase, probabilities of band flip and kinks occurrence are growing. (Jacob et al., 1999; Saenger, 1980; Steiner & Saenger, 1998) In this sense, MD simulations have previously given evidences that γ -CD exhibits two diametrically opposed glucoses (between n and n+/- 4 unit), which are flipped by ~180°, leading to elliptical and curved structures with a quite narrow, slit-like cavity.(Bonnet et al., 2001, 2002; Suárez & Díaz, 2017) Based on the model from Bonnet et al., (Bonnet et al., 2001, 2002) three glucoses are folded toward the cavity leading to the formation of intramolecular O6-H6 ··O6 by their corresponding primary hydroxyl groups. Such orientation of some groups from the primary rim strongly supports the coordination of the sodium cation (Fig. 8C). Secondary, slope between 2 and 3 CD is lower than that observed between both 1 to 2 and 3 to 4 CDs (Fig. 7). Similar trends were observed for $[(CD)_n + nX]^{n+}$ with X = Li, K and Rb (Fig. S8-S10). Interestingly one exception occurred for $[(\gamma-CD)_n + nCs]^{n+}$ where a linear fit can be calculated from mono to tetramer, presumably to a more structuring role of Cs during nanotubes formation (Fig. S11). As previously observed, the most stable dimer was assigned to H•H assembly whatever CD.(Bonnet et al., 2001) Differently to α - and β -CD, due to the incurved ellipsoidal conformation of γ-CD, such dimer corresponds to superimposed irregular bowls with slightly different orientation of ~90° along z axis close to structure reported by Bonnet et al. (Bonnet et al., 2001) Moreover, such structural arrangement favours a closer contact between each CD and enhances the inter-oligosaccharides van der Waals' stabilization. Furthermore, previous studies

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have demonstrated that favourable elongation of γ -CD based tubular arrangements corresponds to a decrease of the potential energy value of the tubes per CD unit as a function of the number of CDs.(Staelens et al., 2015) The important flexibility of γ -CD leads to a more compressed staked assembly during the trimer formation compared to α - and β -CD where a quite more incremental addition of monomer of semi-rigid structure seems to take place. Such phenomenon can explain why the CCS difference between di- (H•H) and trimer (H•TH•T) is by far lower than from monoto dimer. Then starting for a compressed trimer, an additional γ -CD can be associated leading to an orientated growing of the supramolecular chain by formation of a 4-mer (H•TH•TH•T). Interestingly, slope of the curve between CCS of 3-mer and 4-mer is strictly identical than that between 1-mer to 2-mer suggesting a similar assembly elongation process.

4. CONCLUSIONS

It was unambiguously demonstrated in this work that hyphenation with high resolution TIMS represents a powerful and suitable tool to differentiate various non covalent complexes between α -, β - and γ -CD with both various alkali metals or between their self-assemblies. Indeed, cyclooligosaccharides, as cyclodextrins, are well known to be more rigid than linear or branched ones, suggesting a low probability of several stable attachment sites for the alkali cations. This prompted us to think that detection of two or more mobility peaks, especially for γ -CD, can be rationally ascribed to the presence of different conformers. Indeed, the distinct deducted CCS portrayed well the various size and shape of the CD based complexes, which results as a conse-

quence of a balance between size of both cationized metal (ionic radii) and CD (including eventual distortions) and more and less deep inclusion of the cation inside CD cavity (according to coordination features). Moreover, experimental data have been adequately correlated with theoretical models. In this sense, TIMS can be particularly attractive to be used as a suitable molecular descriptor to enlighten parameters affecting the recognition process, to study the first sphere coordination behaviour as well as rationalization of the catalytic activities based on CDmetal supramolecular assemblies. On the other hand, TIMS was also particularly efficient to highlight the presence and the identity of various orientations of the different constituent CD within hierarchical assemblies' nanotubes as well as to estimate their relative content. According to added metals, it was possible to reveal and discriminate such supramolecular edifices involving secondary/secondary hydroxyl groups (head-to head, H•H), primary/secondary rim (head-to-tail, H•T) hydroxyl groups or primary/primary rim (tail-to-tail, T•T) hydroxyl groups interactions. Such results constitute an important milestone since, at our knowledge, it is the first experimental evidences of the existence of various orientations, which until now have been only investigated by theoretical approaches. The present work paves the way for a better understanding of the topology from cyclo-oligosaccharides based supramolecular complexes, which could of course involve both other metals viz transition, post-transition rare earth or noble one, but also other cyclo-oligosaccharides based self-assemblies with or without any (bio)chemical modifications.

CONFLICTS OF INTEREST

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The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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