Abstract: Acid-base properties of cyclodextrins (CDs), persubstituted at C-6 by 3-mercaptopropionic acid, sulaphadex (Suα-CD), subetadex (Suβ-CD) and sugammadex (Suγ-CD, the antidote of neuromuscular blocking steroids) were studied by $^1$H NMR-pH titrations. For each CD, the severe overlap in protonation steps prevented the calculation of macroscopic $pK_a$ values using the standard data fitting model. Considering the full symmetry of polycarboxylate structures, we reduced the number of unknown NMR parameters in the “Q-fitting” or the novel “equidistant macroscopic” evaluation approaches. These models already provided $pK_a$ values, but some of them proved to be physically unrealistic, deceptively suggesting cooperativity in carboxylate protonations. The latter problem could be circumvented by adapting the microscopic site-binding (cluster expansion) model by Borkovec, which applies pairwise interactivity parameters to quantify the mutual basicity-decreasing effect of carboxylate protonations. Surprisingly, only a single averaged interactivity parameter could be calculated reliably besides the carboxylate ‘core’ microconstant for each CD derivative. The specification of protonation isomers hence could not be resolved, but the optimized microscopic basicity parameters could be converted to the following sets of macroscopic $pK_a$ values: 3.84, 4.35, 4.81, 5.31, 5.78, 6.28 for Suα-CD; 3.82, 4.31, 4.73, 5.18, 5.64, 6.06, 6.54 for Suβ-CD and 3.83, 4.28, 4.65, 5.03, 5.43, 5.81, 6.18, 6.64 for Suγ-CD. The pH-dependent charge of these compounds can now be accurately calculated, in support of designing new analytical methods to exploit their charge-dependent molecular recognition such as in cyclodextrin-aided chiral capillary electrophoresis.

Keywords: cyclodextrins; sugammadex; $^1$H NMR pH-titration; protonation microconstant; cluster expansion model; parameter correlation

1. Introduction

Cyclodextrins (CDs) are naturally occurring cyclic oligosaccharides, consisting of six, seven or eight glucopyranose units for α-, β- and γ-CD, respectively. CDs find a wide variety of applications in pharmaceutical sciences [1,2], cosmetic and food industries [3,4], environmental or analytical chemistry [5]. The molecular basis of these applications is the propensity of CDs to encapsulate small apolar organic molecules in their hydrophobic cavity [6–8], where the intrinsic chiral environment also enables enantiospecific molecular recognition, thus the use of CDs as chiral selectors in separation science [9]. In addition to their most common roles as solubilizer or stabilizer excipients for compounds of low aqueous solubility [10], some CDs also gained exquisite applications due to their highly specific interaction with small molecules [11].

Sugammadex (Suγ-CD) is a derivative of γ-CD, persubstituted by 3-mercaptopropionic acid sidechains on its primary hydroxyl groups (see Figure 1; Suγ-CD is in fact the octasodium salt of the polyacid). It is used in the clinical practice as the first selective antidote of steroidal neuromuscular blocking agents rocuronium and vecuronium during surgical...
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were first described twenty years ago [17], complete characterization of their acid-base properties has not yet been reported either experimentally or via computer prediction. In addition to the expanding utilization of Sugammadex (Suγ-CD), there is also a growing interest in its α- and β-CD homologues sualphadex and subetadex [16,17], abbreviated as Suα-CD and Suβ-CD in Figure 1. Besides interaction studies with fentanyl [14], rocuronium bromide [18] and azobenzenes [19], their applicability as chiral selectors was also explored recently [16].

Figure 1. (a) Structures of sualphadex (Suα-CD, n = 1), subetadex (Suβ-CD, n = 2) and sugammadex (Suγ-CD, n = 3); (b) structure of 3,3′-dithiodipropionic acid (DTDPA); (c) structure of monosugammadex (mono-Suγ-CD).

A thorough physicochemical profiling of CD derivatives is essential in order to deepen our understanding and improve the modeling perspectives on their intermolecular interactions upon inclusion complex formation. Although Suα-CD, Suβ-CD and Suγ-CD were first described twenty years ago [17], complete characterization of their acid-base properties has not yet been reported either experimentally or via computer prediction. The potentiometrically derived set of protonation constants of Suβ-CD published by Wenz et al. [20] is incomplete, giving only six of the seven values. There is also a recent example in the literature of pK_a determination for persubstituted CD-derivatives using 13C NMR spectroscopy, although the exploited method only provides a single average pK_a value for each binding site [21]. A full and reliable set of pK_a acid dissociation constants (or equivalently, stepwise log K protonation constants of their conjugate bases) enables the calculation of the net charge of these CD polycarboxylate anions for any arbitrary pH value. The ability to fine-tune the charge of the macromolecular host via solution pH and thus tailoring the selectivity of supramolecular recognition represent advantages of CDs with COOH-bearing side chains over their permanently charged sulfonated or sulfated counterparts [22]. Anionic CDs are used extensively as chiral selectors dissolved in the background electrolyte of capillary electrophoresis methods [23–25] or as mobile phase additives in chiral HPLC [5,26]. Persubstituted and monosubstituted CDs are so-called single isomer CDs, possessing well-defined molecular structures [25,27]. This feature enables the 3D structure of their complexes to be explored via NOE NMR studies or molecular modeling [28–31]. Thus, single isomer CDs offer a better experimental design and reproducibility for chiral separations, which is more difficult to attain with random substituted CD derivatives, with mixtures of CD isomers differing in degrees and positions of functionalization. These advantages generate an increasing demand for single isomer CDs in separation science.
The primary objective of the current study was to determine a reliable set of protonation constants for Suα-CD, Suβ-CD and Suγ-CD. The monobasic mono-Suγ-CD as well as the related bivalent small molecule, 3,3′-dithiodipropionic acid (DTDPA, Figure 1), modeling the isolated substituent side chain of the functionalized CDs were also characterized to investigate the impact of the local molecular environment on the intrinsic basicity of the carboxylate group(s).

The determination of protonation constants is now widely regarded as a routine task in physical chemistry, although published protonation data for at least hexabasic molecules are rare and often incomplete [32,33]. From the array of experimental methods available [34], potentiometric titration remains the gold standard technique for compounds available in sufficient amount and purity. Protonation constants have been successfully determined by pH-metric titrations for as large symmetric systems as dendrimers [35,36]. Another technique gaining immense popularity is NMR-pH titration, in which the chemical shift of the so-called reporter nuclei is followed as a function of solution pH [37,38]. The recorded sigmoidal (chemical shift vs. pH) titration curves undergo computer evaluation to extract the macroscopic log K or pK_a values [38–41]. Carefully selected NMR reporter nuclei enable even monitoring the protonation of each basic site individually. Site-specific or microscopic protonation constants (log k values) can be derived in favorable cases [37] and distribution curves can be computed for the microspecies, isomers holding the same number of protons, but differing in the site(s) of protonation [42]. Since the number of microspecies (2^n) and microconstants (n2^n−1) escalate rapidly with the growing number of binding sites (n), there is a principal limit for a complete, ab initio resolution of microequilibrium schemes of larger than triprotic molecules [43,44]. Molecular symmetry may reduce the number of independent basicity parameters significantly [43,45,46], since symmetry-equivalent groups possess identical basicities. Choosing one intrinsic microconstant for each site and group pair interactivity parameters as iteration parameters to build up all the remaining microconstants [45,46] represents a more efficient strategy for the computer fitting of NMR titration curves. The site-binding (SB) or cluster expansion model by Borkovec [47–49] succeeded in resolving the complete microspeciation of large symmetric polyprotic molecules such as linear polyamines or dendrimers.

Suα-CD, Suβ-CD and Suγ-CD CDs are C_n symmetric polycarboxylate ligands. We applied 1H NMR-pH titrations for the acid-base profiling of these homo-perfunctionalized CDs. The strong overlap of six, seven or eight consecutive protonation steps resulted in single-sigmoid NMR titration curves, without any intermediate local plateaus. This latter feature aggravated the calculation of reliable protonation constants. Several computational strategies had to be explored to identify the most straightforward one in terms of precision and accuracy of the equilibrium constants.

2. Results
2.1. Acid-Base Characterization of Sualphadex

Figure 2 depicts the pH-dependent series of 1H NMR spectra of Suα-CD. From the signal shapes of the methine protons 1-CH through 5-CH, vicinal coupling constants can be derived, which can in turn converted into dihedral angles and compared to the corresponding reference values of the native α-CD in water. Such an analysis may reveal the distortion of the CD macrocycle due to 6-persubstitution and its changes upon acid-base equilibria, but it may deserve a separate molecular modeling study. Returning to characterization of the acid-base equilibria, the 8-CH_2 protons adjacent to the carboxylate group exhibit the largest change in chemical shift upon protonation, so these nuclei were selected as reporters. At slightly alkaline pH, their triplet signal is characteristic to the A_2B_2 spin system, which is gradually transformed into an AA′B_2 type multiplet upon acidification. Regardless of the actual signal shape, the center of these multiplets can be precisely determined at each degree of titration and the resulting chemical shift values are
plotted against pH in Figure 3a. This experimental $^1$H NMR titration curve was evaluated using the approaches detailed in the following sections.

Figure 2. pH-dependent series of $^1$H NMR spectra of Su$\alpha$-CD, with structure and atom numbering of the compound and assignment of the NMR signals (600 MHz, 298 K, H$_2$O:D$_2$O = 9:1).

Figure 3. (a) The experimental NMR-pH titration curve of Su$\alpha$-CD 8-CH$_2$ protons in squares. The green dashed line is generated by fitting the site-binding model without p$_{ε}$, while the solid red line represents the fits by the ED-macro, Q-fitting or the single-p$_{ε}$ site-binding models. The residuals by the ED-macro or the equivalent Q-fitting models are given in solid red line, while those by the single-p$_{ε}$ microscopic evaluation by dashed blue line above (see detailed description of these calculations in Sections 2.3–2.5); (b) protonation degree functions for individual carboxylates ($f$) and the whole Su$\alpha$-CD molecule ($\bar{p}_H$).
2.2. **The General Macroscopic Evaluation Model**

In the macroscopic description shown in Figure 4a, only the stoichiometry of the H,L species is considered, not their site(s) of protonation. Successive formation of the macrospecies (charges omitted) are characterized by the stepwise \( \log K_i \) or cumulative \( \log \beta_i \) macroconstants, whose relationships can be expressed as follows:

\[
\log \beta_i = \log \left( \frac{[H_iL]}{[L][H^+]^i} \right) = \sum_{j=1}^{i} \log K_j = \sum_{j=1}^{i} pK_{a,n-j+1}
\]  

\( (1) \)

![Figure 4. (a) Macroscopic protonation equilibria of Suα-CD, with protonation and dissociation macroconstants; (b) excerpt of the complete microscopic protonation scheme of Suα-CD.](image)

For each microscopic interaction, the six \( \delta L \) values necessarily become iteration parameters. All the tested software tools including our own R scripts failed to estimate the seven \( \delta L \) and pair interactivity parameters \( \epsilon_{ij} \) are also indicated.

Owing to the rapid kinetics of proton exchange on the \( ^1H \) NMR chemical shift timescale, the observed methylene peak position \( \delta_{\text{obs}} \) becomes a weighted average of the intrinsic chemical shifts \( \delta_{L} \) of the individual macrospecies [37]:

\[
\delta_{\text{obs}} = \delta_{L} x_{L} + \delta_{HL} x_{HL} + \ldots + \delta_{H^nL} x_{H^nL} = \sum_{i=1}^{n} \delta_{H^nL} x_{H^nL}
\]  

\( (2) \)

The \( x_i \) weighting factors are the pH-dependent molar fractions of the macrospecies, which can be expressed in terms of the \( \beta_i \) formation constants as follows [32]:

\[
\delta_{\text{obs}} = \frac{\delta_{L} + \sum_{i=1}^{n} \delta_{H^nL} \beta_i [H^+]^i}{1 + \sum_{i=1}^{n} \beta_i [H^+]^i} = \frac{\delta_{L} + \sum_{i=1}^{n} \delta_{H^nL} 10^{log \beta_i - i p_{H}}}{1 + \sum_{i=1}^{n} 10^{log \beta_i - i p_{H}}}
\]  

\( (3) \)

In principle, the pH-independent limiting chemical shifts \( \delta_{L} \) and \( \delta_{H^nL} \) can be read off as alkaline and acidic plateaus of the titration curve in Figure 3a. In practice, these are co-iterated parameters with the log \( \beta_i \) (or log \( K_i \)) values. Due to absence of local plateaus, the five intermediates \( \delta_{H^nL} \) values necessarily become iteration parameters. All the tested software tools including our own R scripts failed to estimate the seven \( \delta_{H^nL} \) and the six log \( K_i \) parameters together. The severe overlap of six protonation steps rendered the gold standard macroscopic evaluation unfeasible for the fully-symmetric Suα-CD. In our view, the culprit for this failure was the strong mathematical correlation between the \( \delta_{H^nL} \) and log \( K_i \) parameters. A similar conclusion emerged from the NMR titrations of complexones [46] and a two-step CD-steroid equilibrium system [50].
2.3. The Equidistant Macroscopic Evaluation Model

To avoid the correlation of spectroscopic and equilibrium parameters, the NMR titration curve of Suα-CD was interpreted from a different viewpoint. Due to the C₆₀ symmetry, each carboxylate group experiences the same degree of protonation upon acidification:

\[ f = f_A = f_B = \ldots = f_F. \]

As evident from Figure 3b, the \( f(pH) \) function ranges from 0 to 1 and it is linearly proportional to the observed displacement of the 8-methylene signal,

\[ \delta^\text{obs}(pH) = \delta_L + \Delta \delta \cdot f(pH) \tag{4} \]

where \( \Delta \delta = \delta_{H,L} - \delta_L \) is the protonation increment [37]. On the other hand, a similar protonation degree function can be defined for the whole Suα-CD molecule:

\[ \pi_H(pH) = \sum_{i=1}^{n} i \cdot x_{i,H,L}(pH) = \frac{\sum_{i=1}^{n} i \cdot 10^{\log \beta_i - i \cdot pH}}{1 + \sum_{i=1}^{n} 10^{\log \beta_i - i \cdot pH}} \tag{5} \]

The \( \pi_H \) function was introduced by Bjerrum and plays a central role in the calculation of protonation constants from potentiometric titration curves. Figure 3b demonstrates that \( \pi_H \) ranges from 0 to \( n = 6 \), being the sum of contributions of the six equivalent carboxylates [43],

\[ \pi_H = f_A + f_B + \ldots + f_F = 6f \tag{6} \]

Expressing \( f \) from Equation (4) and substituting it into Equation (6) yields the following formula:

\[ \pi_H(pH) = 6 \frac{\delta^\text{obs}(pH)}{\Delta \delta} - 6 \frac{\delta_L}{\Delta \delta} \tag{7} \]

This equation states that apart from the subtracted, pH-independent term, the experimental NMR titration curve \( \delta^\text{obs}(pH) \) is directly proportional to the \( \pi_H \) function. In other words, the evaluation of potentiometric and NMR titration curves becomes mathematically equivalent for a fully symmetric polyacid. Rearranging Equation (7) for \( \delta^\text{obs} \) and combining it with Equation (5) yields the following NMR fitting function:

\[ \delta^\text{obs} = \delta_L + \Delta \delta \cdot \frac{\sum_{i=1}^{n} i \cdot 10^{\log \beta_i - i \cdot pH}}{1 + \sum_{i=1}^{n} 10^{\log \beta_i - i \cdot pH}} \]

\[ = \delta_L + \Delta \delta \cdot \frac{\sum_{i=1}^{n} \left( \delta_L + i \frac{\Delta \delta}{n} \right) 10^{\log \beta_i - i \cdot pH}}{1 + \sum_{i=1}^{n} 10^{\log \beta_i - i \cdot pH}} \tag{8} \]

A comparison of Equation (8) with the general Equation (3) reveals that for fully symmetric ligands, the intermediate \( \delta_{H,L} \) values increase in strictly equidistant steps from \( \delta_L \) to \( \delta_{H,L} = \delta_L + \Delta \delta \) (see Figure 3a). They cease to be iteration parameters, reducing the parameter space considerably.

This equidistant (ED) macroscopic evaluation already converged in both Microcal Origin and our R-script, yielding statistically sound values for the optimized parameters:

\[ \delta_L = 2.4638 \text{ (standard deviation: 0.0008 ppm), } \delta_{H,L} = 2.7064 \text{ (0.0008 ppm) and the log } K \]

protonation constants collected in Table 1 (see Table S1 for more details). This approach succeeded in diminishing the correlation among equilibrium and NMR parameters, see Table S2. At the first sight, the resulting log \( K \) values seemed to be physically realistic. Nevertheless, a subtle problem with log \( K_5 \) was traced by the monotonicity test of micro-constants detailed in the following chapter. We note that potentiometric titration would also be prone to yield biased macroconstants for persubstituted CDs, since the fitting of model Equations (5) and (8) is equivalent from the mathematical statistical viewpoint.
Table 1. Macroscopic and averaged microscopic protonation constants of the studied cyclodextrin derivatives, with their standard deviations in parentheses. The ED macro and the Q-fitting evaluation approaches gave physically unrealistic values for some microconstants (suggesting cooperative proton uptake), the affected macro- and microconstants are marked by question marks. The site-binding microscopic evaluation yielded the correct protonation constants, whose errors were assessed by Monte Carlo simulations.

|            | Suα-CD | Suβ-CD | Suγ-CD |
|------------|--------|--------|--------|
|            | ED or Q | SB Micro | ED or Q | Lit. [20] | ED or Q | SB Micro |
| $\log K_1$ | 6.43 (0.07) | 6.28 (0.02) | 6.60 (0.03) | 6.54 (0.02) | 6.49 (1) | 6.68 (0.08) | 6.64 (0.02) |
| $\log K_2$ | 5.79 (0.09) | 5.78 (0.02) | 6.18 (0.04) | 6.06 (0.02) | 5.69 (1) | 6.17 (0.12) | 6.18 (0.02) |
| $\log K_3$ | 5.26 (0.11) | 5.31 (0.02) | 5.27 (0.08) | 5.64 (0.02) | 5.25 (1) | 5.88 (0.16) | 5.81 (0.02) |
| $\log K_4$ | 4.70 (0.14) | 4.81 (0.02) | 5.53 (0.06) | 5.18 (0.01) | 4.75 (1) | 5.28 (0.22) | 5.43 (0.01) |
| $\log K_5$ | **4.53** (0.11) | 4.35 (0.02) | 4.54 (0.06) | 4.73 (0.02) | 4.31 (1) | 5.14 (0.22) | 5.03 (0.01) |
| $\log K_6$ | 3.77 (0.07) | 3.84 (0.02) | **4.46** (0.05) | 4.31 (0.02) | 3.78 (1) | 4.59 (0.17) | 4.65 (0.02) |
| $\log K_7$ | - | - | 3.69 (0.03) | 3.82 (0.02) | <3 | **4.34** (0.12) | 4.28 (0.02) |
| $\log K_8$ | - | - | - | - | - | 3.78 (0.07) | 3.83 (0.02) |

2.4. The Q-Fitting Model

In the microscopic protonation scheme in Figure 4b, all protonation isomers of the $H_iL$ macrospecies ($i = 1$ to 5) are considered as distinct entities. The hexabasic Suα-CD ligand has $2^6 = 64$ protonation microspecies, whose formation are interconnected by $6 \times 2^5 = 192$ microconstants $k_x^y$, whose upper index $x$ identifies the protonating, the lower index $y$ (if any) the already protonated carboxylate(s). Fortunately, molecular symmetry significantly reduces the number of constitutionally different microspecies. For instance, the following isomers protonated at two adjacent carboxylate groups are indistinguishable: $AB \equiv BC \equiv CD \equiv DE \equiv EF \equiv FA$. Thus, the multiplicity of microspecies $AB$ becomes six. Considering all such symmetry-based simplifications, a reduced microscopic scheme with only 13 different microspecies can be constructed (see Figure S26) and certain microconstants coincide, e.g., $k_A^B = k_B^A$ or $k_{AE}^C = k_{EA}^C$, etc. A micro-formation constant denoted by $\kappa$ [43,45] is defined for each microspecies, which reads, e.g., for microspecies $ABD$ as follows:

$$\kappa_{ABD} = \frac{[ABD]}{[L][H^+]} = k_A^B k_B^D k_A^D$$

In contrast to the stepwise $k$ microconstants, the 12 $\kappa$-values (with $\kappa_1$ being 1 by definition) constitute a non-redundant set of parameters for the complete description of the protonation network [43].

The protonation degree function of carboxylate $A$ is the concentration sum of all the microspecies protonated at this site, divided by the total ligand concentration,

$$f_A = \frac{[A] + [AB] + [AC] + \ldots + [ABC] + [ABD] + \ldots + [ABCDEF]}{[L] + [HL] + [H_2L] + [H_3L] + [H_4L] + [H_5L] + [H_6L]}$$
Rewriting this equation in terms of equilibrium constants yields the following ratio of 6-degree polynomials,

\[ f_A(pH) = \frac{Q_1[H^+] + Q_2[H^+]^2 + Q_3[H^+]^3 + Q_4[H^+]^4 + Q_5[H^+]^5 + Q_6[H^+]^6}{1 + \beta_1[H^+] + \beta_2[H^+]^2 + \beta_3[H^+]^3 + \beta_4[H^+]^4 + \beta_5[H^+]^5 + \beta_6[H^+]^6} \]  

(11)

where both the \( Q_i \) and \( \beta_i \) coefficients can be expressed as linear combinations of the 12 unknown \( k_i \) micro-formation constants:

\[ Q_1 = \kappa_A = 1\beta_1/6 \]  

(12)

\[ Q_2 = 2\kappa_{AB} + 2\kappa_{AC} + \kappa_{AD} = 2\beta_2/6 \]  

(13)

\[ Q_3 = 3\kappa_{ABC} + \kappa_{ACE} + 6\kappa_{ABD} = 3\beta_3/6 \]  

(14)

\[ Q_4 = 4\kappa_{ABCD} + 4\kappa_{ABCE} + 2\kappa_{ABDE} = 4\beta_4/6 \]  

(15)

\[ Q_5 = 5\kappa_{ABCDE} = 5\beta_5/6 \]  

(16)

\[ Q_6 = 6\kappa_{ABCDEF} = 6\beta_6/6 \]  

(17)

It was demonstrated earlier \[43–45\] that fitting of an Equation (11) type ratio of polynomials yields unequivocal results only for the \( Q_i \) coefficients. Any attempt to compute the constituent \( k_i \) constants led to overparametrization of the regression for more than triprotic molecules. Fitting the combination of Equations (4) and (11) to the experimental dataset of Su\( \alpha\)-CD yielded an excellent fit (Figure 3a) and statistically sound results for the iterated log \( Q_{1–6} \), \( \delta_L \) and \( \Delta \delta \) parameters (see Tables S3 and S4). The log \( Q_{1–6} \) coefficients were then used to back-calculate the log \( \beta_i \) values by Equations (12)–(17), the differences of which yielded the stepwise log \( K_i \) constants listed in Table 1. The resulting macroconstants, quality of fit and even the residual distributions coincide for the ED-macro and Q-fitting models, the two models are mathematically isomorphous.

The physical validity of the macroconstants was assessed by the monotonicity test of microconstants. For symmetric ligands with \( n \) fully equivalent basic sites, a set of stepwise \( k_i \) microconstants can directly be derived from the macroconstants \[37,43,51\]:

\[ k_i = \frac{i}{n - i + 1} K_i \]  

(18)

The “first” and “last” microconstants calculated in this way coincide with the “true” microconstants of Su\( \alpha\)-CD, \( k_1 = k^A \) and \( k_6 = k^F_{ABCDE} \). However, each remaining intermediate \( k_i \) value becomes an average of all microconstants describing the formation of protonation isomers of \( H_i L \). For this reason, these averaged microconstants are denoted by \( \bar{k}_i \) in Table 1. The majority of calculated \( \bar{k}_i \) values decrease in a monotonic order as expected. However, coordination of the fifth proton to tetraprotonated microspecies seemingly increases the carboxylate basicity from log \( \bar{k}_4 = 4.82 \) to log \( \bar{k}_5 = 4.93 \). Such a positive cooperativity of the carboxylates would require a specific “allosteric” conformational transition of the CD to ease the subsequent protonation step. This type of allosterism is known for proton- or ligand-binding of certain proteins, but it is less conceivable for Su\( \alpha\)-CD. In fact, the outlier nature of \( \bar{k}_5 \) has a root cause in the numerical sensitivity of the evaluation. Consequently, the macroconstants obtained by the ED-macro or Q-fitting approaches are still approximate values.

2.5. The Microscopic Site-Binding Model

Unbiased macroconstants for Su\( \alpha\)-CD could only be deduced by the site-binding model (SB). Its mathematical elaboration has been published by Borkovec \[47–49\]; only the key concept is reiterated here using our notational system.

As a first approximation, the basicity of a given carboxylate is assumed to change negligibly upon successive protonation of the five others. This assumption seems to be
plausible, considering the large covalent and spatial distances among the carboxylates. The formation constants of all conceivable protonation isomers of H$_4$L simply become $k_i = (k^A)^i$ and $k^A$ can be termed as group constant in this statistical case [51]. Plugging these $k$ values into Equations (11)–(17) and using Equation (4) for nonlinear regression yields the results in Tables S5 and S6. However, the quality of fit in Figure 3a is clearly unsatisfactory, indicating that the six carboxylates of Suα-CD do not obey the statistical case of independent protonation.

As carboxylate interactions obviously cannot be neglected, all $k_{ij}$ microconstants are expressed in terms of the ‘intrinsic’ or ‘core’ $k^A$ microconstant of the first protonating carboxylate and the three pair-interactivity parameters $\epsilon_{12}$, $\epsilon_{13}$ and $\epsilon_{14}$ (see Figure 4b). $\epsilon_{12}$ quantifies how protonation of a carboxylate impacts the basicity at the adjacent site and vice versa. $\epsilon_{13}$ and $\epsilon_{14}$ describe the interaction of more distant sites. As already mentioned, negative cooperativity has been experimentally observed for the vast majority of small-molecules ranging from diacids to dendrimers [37,51], thus $\epsilon_{AB} < 1$ or $p\epsilon_{AB} = -\log \epsilon_{AB} > 0$. Taking the microspecies ABD from Figure 4b as an example, its microconstant for the ABD→ABDE protonation step and its $\kappa$ formation constant are expressed in terms of the site-binding model as follows:

$$k_{ABD}^F = k^A \epsilon_{12} \epsilon_{13} \epsilon_{14}$$

$$\kappa_{ABDE} = k^A \epsilon_D \epsilon_{ABD}^E = (k^A)^2 \epsilon_{12} \epsilon_{13} \epsilon_{14}$$

Similar $\kappa$ values are formulated for the remaining twelve non-equivalent microspecies of Suα-CD. The $Q_i$ coefficients in Equations (11)–(17) also become functions of the four “cluster parameters” $k^A$, $\epsilon_{12}$, $\epsilon_{13}$ and $\epsilon_{14}$. The master fitting Equation (4) brings two additional spectroscopic parameters ($\delta_L$ and $\Delta\delta$) to this highly efficient parametrization.

The allowance for merely nearest-neighbor pair interactions via the parameter $\epsilon_{12}$ yielded an excellent fit to the titration profile, see Figure 4a. The optimized $\log k^A = 5.51$ (0.02), $p\epsilon_{12} = 0.44$ (0.02), $\delta_L = 2.4650$ (0.0009) ppm and $\Delta\delta = 0.241$ (0.001) ppm parameters exhibited only moderate correlations (see Tables S7 and S8). For the sake of completeness, the calculation of two interactivity parameters ($p\epsilon_{12}$ and $p\epsilon_{13}$) was subsequently also attempted. The same excellent fit was accompanied here with satisfactory results only for three parameters: $\log k^A = 5.63$ (7), $\delta_L = 2.464$ (0.001) ppm and $\Delta\delta = 0.241$ (0.001) ppm. The optimized $p\epsilon_{12} = 0.28$ and $p\epsilon_{13} = 0.28$ parameters were burdened with disproportionately large standard deviations of 138 and a correlation coefficient of $-1$. The model turned out to be overparametrized with respect to the latter two interactivity parameters. This unexpected failure prompted us to perform extensive tests regarding the robustness of our SB fitting script. To this end, synthetic titration datasets were simulated from known cluster parameters and normally distributed random error added to the chemical shift values generated by Equation (3). We concluded that only one single $p\epsilon$ value is accessible numerically in this symmetric hexaprotonic scenario, any additional interactivity parameter becomes statistically insignificant. Moreover, the successfully optimized $p\epsilon_{12}$ parameter did not exactly coincide with the $p\epsilon_{12}$ value used for the simulation, rather it became a nontrivial combination of the input $p\epsilon_{12}$, $p\epsilon_{13}$ and $p\epsilon_{14}$ values. Due to its averaged nature, the single accessible interactivity parameter will henceforth be denoted by $p\epsilon$. The inability to distinguish between 1 and 2, 1 and 3 or 1 and 4-type carboxylate interactions also implicates the distribution of protonation isomers which cannot be determined for a fully symmetric hexaprotonic ligand. Nevertheless, the $\log k^A = 5.51$ and $p\epsilon = 0.44$ results were combined to $\kappa$ via relationships similar to Equation (19), followed by their conversion to $\kappa_L$, $Q_i$, $\beta_i$, and finally to $\log K_L$ equilibrium constants. The latter values listed in Table 1 represent the first reliable and physically sound protonation macroconstants for Suα-CD. They enable the calculation of speciation curves (in Figure S27) and the average charge of Suα-CD (in Figure S29) as a function of pH.
3. Discussion

3.1. Protonation of Subetadex and Sugammadex

The NMR-pH titration profiles and the problems of their evaluation were highly similar for the three investigated CD derivatives, so Suβ-CD and Suy-CD are discussed together here. The macroscopic evaluation by Equation (3) failed for the larger CDs as well. The ED macroscopic and Q-fitting approaches yielded excellent fits (see Figures 5a and 6a and Tables S9–S12 and S17–S20), but the resulting log $K$ macroconstants were only partially meaningful. Table 1 reveals two incorrect log $K$ values for Suβ-CD and one for Suy-CD according to monotonicity test of the derived microconstants. We note that the macroconstant set published by Wenz et al. [20] also contains biased values. Henceforth we resorted to applying the site-binding model for Suβ-CD and Suy-CD.

![Figure 5.](image)

**Figure 5.** (a) The experimental NMR-pH titration curve of Suβ-CD 8-CH$_2$ protons in squares. The green dashed line is generated by fitting the site-binding model without $p\varepsilon$, while the solid red line represents the fits by the ED-macro, Q-fitting or the single-$p\varepsilon$ site-binding models. The residuals by the ED-macro or Q-fitting models given in solid red line, while those by the single-$p\varepsilon$ microscopic evaluation by dashed blue line above; (b) the reduced microscopic protonation scheme of Suβ-CD, with multiplicities of the protonation isomers, where white circles indicate deprotonated-, and the yellow circles denote protonated binding sites.

Binding site combinatorics brings increasing complexity to the microscopic protonation schemes of the larger CDs: there are 128 microspecies (20 nonidentical when symmetry is considered, see Figure 5b) for Suβ-CD and 256 microspecies (36 nonidentical in the reduced protonation scheme in Figure 6b) for Suy-CD. Assuming independent carboxylate protonations yielded a bad fit for both CDs, see the dashed lines in Figures 5a and 6a. To allow for the pairwise basicity-modifying interactions of the carboxylates, $p\varepsilon_{12}$, $p\varepsilon_{13}$ and $p\varepsilon_{14}$ interactivity parameters were introduced for Suβ-CD, while an additional $p\varepsilon_{15}$ also emerged for Suy-CD. The excellent quality of solid-line fits displayed in Figures 5a and 6a could be achieved with optimizing at least one $p\varepsilon$ value besides log $k^A$ (Tables S13–S16 and S21–S24). Co-iteration of two or more $p\varepsilon$ values resulted in overparametrization also for these larger CDs, so merely an averaged $p\varepsilon$ value could be extracted with statistical significance. The optimized parameters for Suβ-CD and Suy-CD were converted to the stepwise log $K$ macroconstants listed in Table 1. These latter values are more closely spaced, indicating less basicity-decreasing interactions between carboxylates when compared to the more steeply decreasing, biased values published by Wenz et al. [20]. The uniformity of $p\varepsilon$ values ranging from 0.44 to 0.51 across the three Su-CD homologues corroborate the validity of our results. The macrospecies distribution curves for Suy-CD are shown in Figure 7, while
the same plot for Suβ-CD and the mean charge vs. pH profiles are given as Figures S28 and S29, respectively.

Figure 6. (a) The experimental NMR-pH titration curve of Suγ-CD 8-CH₂ protons in squares. The green dashed line is generated by fitting the site-binding model without pe, while the solid red line represents the fits by the ED-macro, Q-fitting or the single-pe site-binding models. The residuals by the ED-macro or Q-fitting models given in solid red line, while those by the single-pe microscopic evaluation by dashed blue line above; (b) the reduced microscopic protonation scheme of Suγ-CD, with multiplicities of the protonation isomers. The white circles indicate deprotonated-, and the yellow circles denote protonated binding sites.

Figure 7. pH-dependent distribution of protonation microspecies of Suy-CD.

3.2. Basicity Comparison with Related Compounds

NMR-pH titrations were carried out also for the mono-Suy-CD (Figures S30 and S31) and DTDPA model compounds (Figure S32). The standard macroscopic evaluation, Equation (3) for \( n = 1 \) and 2 yielded the protonation macroconstants without any difficulties, due to the significantly fewer iteration parameters. Table S25 lists the microscopic basicity parameters of the symmetric diacid DTDPA, calculated either from the macroconstants by known equations \([37,45,51]\) or by any of the data fitting approaches discussed
in Chapters 2.3–2.5. The log $K = 4.35 \pm 0.01$ value of mono-Suγ-CD agrees well with the basicity of the first protonating carboxylate in DTDPA, log $k^A = 4.33$. The literature value for the latter constant is somewhat lower at 4.16 [52]. One would intuitively expect a similar intrinsic basicity for the thiopropionate side chain in the persubstituted CDs, since the carboxylates on different glucose units are seemingly isolated from one another (Figure 1). However, the log $k^A$ values of 5.51, 5.70 (literature: 5.64 [20]) and 5.73 in Suα-CD, Suβ-CD and Suγ-CD reveal a consistently higher proton affinity for the same side chain. This surprising finding may be explained by the higher overall net charge (electron density) of the persubstituted CDs, and this phenomenon has also been revealed for acrylic acids where the $pK_a$ of the monomeric form shows a 0.9 log $K$ unit lower acidity compared to its polymeric form [53]. The averaged interactivity parameters of carboxylates, $\bar{p}\varepsilon$ ranging from 0.44 to 0.51 in the CDs are also higher than $p\varepsilon = 0.15$ in DTDPA. This latter value is reminiscent of carboxylate interactions ($p\varepsilon \approx 0.1$) in flexible oligopeptides such as oxidized glutathione [45]. The covalently remote COO$^-$ and COOH in monoprotonated DTDPA seem to approach each other by adopting a bent conformation. Earlier literature suggests almost independent carboxylate protonations with $p\varepsilon \approx 0$ for DTDPA [51,52]. Turning back to the fully unprotonated CD polyanions, their side chains are expected to repel one another due to electrostatic reasons. However, this repulsion is presumably tamed upon successive protonation, so the side chain carboxylates on the pendant, flexible side chains may approach each other and even water-molecule-mediated hydrogen-bonding networks might be suspected at some stages of protonation. The “crowding” of COOH and COO$^-$ groups near the narrower rim of the truncated cone of CD enhances basicity-modifying interactions of otherwise distant basic sites and this fact may be reflected in the experimentally found, seemingly elevated $p\varepsilon > 0.4$ values.

4. Materials and Methods

4.1. Materials

4.1.1. Synthesis of Sugammadex-Analogues (General Procedure)

The per-6-iodo-CD was pre-solubilized in dimethyl sulfoxide (DMSO) in a Schott Duran bottle of adequate volume. In order to speed up the process of solubilization, the glass was sonicated at 50 °C. DMSO was added to the reaction vessel under inert atmosphere and 3-mercaptopropionic acid was then poured sequentially. The reaction mixture was cooled down in a water bath ($T \sim 15 \degree C$) and sodium methoxide-methanol solution was slowly added. The reaction mixture turned from a colorless solution to an intense pinkish solution with heat evolution. The DMSO-CD solution was added in one portion to the reaction mixture (Schott Duran is washed with 1 × 30 mL DMSO) causing almost immediate formation of a massive white precipitate. The suspension was stirred at r.t. for 1 h. The reaction mixture was filtered and the solid was extensively washed with methanol until a white solid was obtained. The solid was placed into a drying box and dried until constant weight.

SuαCD: $^1$H NMR (400 MHz, 300 K, D$_2$O) $\delta$(ppm) 5.10 (d, $J = 2.8$ Hz, 6H, H1), 4.13 (bt, $J = 8.5$ Hz, 6H, H5), 3.99 (t, $J = 9.2$ Hz, 6H, H3), 3.67 (dd, $J = 9.8$ Hz, 6H, H2), 3.58 (t, $J = 8.8$ Hz, 6H, H4), 3.29 (m, 6H, H6a), 2.98 (m, 18H, H6b, H7), 2.72 (m, 12H, H8).

$^{13}$C NMR (100 MHz, 300 K, D$_2$O) $\delta$(ppm) 101.2 (C1), 84.6 (C4), 73.1 (C3), 71.8 (C2), 71.2 (C5), 35.4 (C8), 33.8 (C6), 28.3 (C7). The $^{13}$C chemical shifts were read from the DEPT-edited HSQC spectrum.

SuβCD: $^1$H NMR (600 MHz, 300 K, D$_2$O) $\delta$(ppm) 5.18 (d, $J = 3.5$ Hz, 7H, H1), 4.14–3.85 (m, 14H, H3, H5), 3.77–3.51 (m, 14H, H2, H4), 3.33–3.10 (m, 7H, H6a), 3.12–2.75 (m, 18H, H6b, H7), 2.61 (t, $J = 7.3$ Hz, 14H, H8).

$^{13}$C NMR (150 MHz, 300 K, D$_2$O) $\delta$(ppm) 178.5 (C9), 100.7 (C1), 83.1 (C4), 71.8 (C2), 71.2 (C5), 36.6 (C8), 33.4 (C6), 28.8 (C7).

SuγCD: $^1$H NMR (600 MHz, 300 K, D$_2$O) $\delta$(ppm) 5.22 (d, $J = 3.8$ Hz, 8H, H1), 4.01–4.02 (m, 8H, H5), 3.97 (d, $J = 9.4$ Hz, 8H, H3), 3.70 (dd, $J = 9.8$, 3.8 Hz, 8H, H2), 3.61 (t, $J = 9.4$ Hz,
8H, H4), 3.31–3.19 (m, 8H, H6a), 3.01 (dd, J = 13.5, 8.5 Hz, 8H, H6b), 2.97–2.87 (m, 16H, H7), 2.67 (t, J = 7.2 Hz, 16H, H8).

$^{13}$C NMR (150 MHz, 298 K, D$_2$O) $\delta$ (ppm) 178.4 (C9), 100.8 (C1), 82.7 (C4), 72.7 (C2), 72.2 (C3), 71.2 (C5), 36.1 (C8), 33.4 (C6), 28.5 (C7).

4.1.2. Synthesis of Mono-Sugammadex

3-Mercaptopropionic acid (0.025 mol, 2.65 g, 2.2 mL) was added to a viscous DMSO (100 mL) solution of anhydrous 6-mono-O-(p-toluenesulfonyl)-$\gamma$-CD (14.06 g, 0.01 mol) under vigorous stirring and inert atmosphere. The yellowish reaction mixture was cooled down in water bath (T ~ 15 °C) and sodium methoxide-methanol solution (0.050 mol, 10.8 g, 11.4 mL) was slowly added. The reaction mixture turns from a yellowish solution to an intense pinkish solution under heat evolution. The solution was heated at 65 °C for 2 h and a white precipitate formed. The suspension was additionally heated at 65 °C for 3 h. The reaction mixture was cooled down, filtered and the solid was extensively washed with acetone (2 × 100 mL) and methanol (2 × 50 mL) until a white solid was obtained. The solid was placed into a drying box and dried until constant weight (7.5 g).

The structures of the synthetized compounds were confirmed using ESI-MS as well as routine 1- and 2-dimensional NMR techniques (see Figures S1–S25 in Supporting Information).

Mono-Su-$\gamma$CD: $^1$H NMR (600 MHz, D$_2$O, 300 K) $\delta$ (ppm) 5.22–5.04 (m, 8d, J = 3.8 Hz, 8H, H1, H1$'$), 4.08–3.99 (m, 1H, H5$'$), 3.99–3.80 (m, 29H, H3, H3$'$, H5, H5$'$, H6a,b), 3.72–3.63 (m, 8H, H2, H2$'$), 3.59 (m, 8H, H4, H4$'$), 3.14 (dd, J = 13.9, 2.5 Hz, 1H, H6$'$a), 2.93 (t, J = 7.1 Hz, 4H, H7$'$), 2.91–2.85 (m, 1H, H6$'$b), 2.83 (t, J = 7.4 Hz, 2H, H7$''$), 2.60 (t, J = 7.1 Hz, 4H, H8$''$), 2.48 (t, J = 7.4 Hz, 2H, H8$''$).

$^{13}$C NMR (151 MHz, D$_2$O) $\delta$ (ppm) 180.6 (C9$'$, C9$''$), 101.7–101.6 (C1), 101.4 (C1), 83.48 (C4$'$), 80.6–80.3 (C4), 72.9–72.7 (C3, C3$'$), 72.3 (C4), 71.7 (C5), 71.1 (C5$'$), 60.3–60.1 (C6), 37.48 (C8$'$), 36.76 (C8$''$), 34.58 (C7$'$), 32.85 (C6$'$), 29.01 (C7$''$).

The proton and carbon signals marked by * correspond to DTDPA, present as a byproduct formed during the synthesis and already described in the literature [54].

4.1.3. Chemicals

D$_2$O (99.96 atom % deuterium) and acetic acid-d$_4$ (99.9 atom % deuterium) were purchased from Merck KGaA (Darmstadt, Germany). Methanol, sodium hydroxide (1 N solution), hydrochloric acid (1 N solution) and sodium chloride were reagent grade and purchased from Molar Chemicals Ltd. (Halásztelek, Hungary). Standard buffer solutions were purchased from VWR International LLC (Radnor, PA, USA). Water used for the solutions was obtained from a Milli-Q water purification system (Merck Millipore, Burlington, MA, USA).

4.2. Methods

4.2.1. ESI-MS Measurements

Mass spectra were obtained on Bruker ESQUIRE 3000 ES-ion trap instrument with electrospray ionization (ESI) in negative mode. Samples were dissolved in water.

4.2.2. pH Measurements

The pH meter readings were recorded using a Metrohm pH meter, equipped with a Metrohm 6.0234.110 combined glass electrode. A four-point calibration using pH 2.00, pH 4.00, pH 7.00 and pH 11.00 standard buffer solutions were performed directly before pH adjustments. The pH measurements were performed in 25 mL vessels with proper stirring before transferring 600 µL sample solutions into the NMR tubes. Solutions were prepared in the pH range of 1.50–8.70 with 0.30 increments using 0.9 M, 0.5 M and 0.1 M sodium hydroxide and hydrochloric acid solutions in H$_2$O:D$_2$O = 9:1.
4.2.3. NMR Experiments

For structural characterizations of CD derivatives in D$_2$O, a 600 MHz Varian DDR NMR spectrometer equipped with a 5 mm inverse-detection gradient probe or a 400 MHz Varian Mercury Plus spectrometer equipped with a 5mm Varian 400 Automation Triple Resonance Broadband Pulsed Field Gradient probe was used. For NMR-pH titrations, the same instruments were used. The pH-dependent series of $^1$H NMR spectra were recorded in a solvent mixture of H$_2$O:D$_2$O = 9:1 by volume, where the constant ionic strength and the buffer capacity were adjusted using 0.05 M sodium chloride and 0.05 M acetic acid-d$_4$. An appropriate amount of CD was weighed to obtain a 1 mM solution and finally one drop of methanol was added as a chemical shift reference (3.310 ppm). The water resonance was diminished by the dpfgse pulse sequence [55] or presaturation. All the spectra were processed using the MestReNova v9.0.1-13254 (Mestralab Research, S.L., Santiago de Compostela, Spain) software.

4.2.4. Evaluation of Titration Data

The standard macroscopic evaluation of NMR-pH titration curves (Equation (3) in Section 2.2) was performed with the following software tools: Microcal Origin Pro 8 (OriginLab Corp., Northampton, MA, USA), HypNMR2018 [38], OPIUM [39] and home-written R scripts (R version 4.1.2, The R Foundation, Vienna, Austria) using the nls module. The equidistant macroscopic (Equation (8)) and Q-fitting (Equations (4) and (11)) models were fitted both by Origin and our R-scripts to verify their equivalent performance regarding statistical description of the parameter estimates. The microscopic cluster expansion model in Section 2.5 requires summation of >100 variables which is unfeasible in the form of an explicit model function required by commercial data-fitting software like Origin. The enumeration of microspecies was therefore accomplished in a custom-written R script. To assess the robustness of our data-fitting model as well as the error propagation of the optimized log $k$ and p$\varepsilon$ parameters to the derived log $K$ macroconstants, Monte Carlo simulations [56,57] on >1000 replicates were performed in R.

5. Conclusions

Our study demonstrates that the full symmetry equivalence of basic sites in persubstituted CDs generates hitherto unrecognized difficulties when it comes to the determination of macroscopic protonation constants or p$K_a$ values from NMR-pH titration curves. The standard macroscopic evaluation fails to converge, due to strong mathematical correlation between spectroscopic and equilibrium parameters. The Q-fitting or the novel equidistant macroscopic evaluation approaches reduce the dimensionality of parameter space and already yield macroconstant results. Due to symmetry, the latter values can be easily converted to microconstants, which are supposed to decrease monotonously upon proceeding protonation. This simple test is useful to discover systematic error in macroconstant values, thus it is recommended to use more extensively for polybasic compounds. On the other hand, the microscopic site-binding model of Borkovec produces microconstants and interactivity parameters as “building blocks” to guarantee the monotonicity of microconstants, thereby yielding an unbiased set of macroconstants. Surprisingly, we could demonstrate an inherent limitation of this approach in distinguishing group pair interactions according to their covalent distances, which is peculiar to fully-equivalent binding sites. While this problem hindered the resolution of the complete schemes of microscopic protonation equilibria, we publish here the first reliable sets of protonation macroconstants for the pharmacologically relevant sugammadex and its two homologues.

Supplementary Materials: The supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/ijms232214448/s1.
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