Force Field Parameterization for the Description of the Interactions between Hydroxypropyl--Cyclodextrin and Proteins

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Cyclodextrins are cyclic oligosaccharides, widely used as drug carriers, solubilizers, and excipients. Among cyclodextrins, the functionalized derivative known as hydroxypropyl-β-cyclodextrin (HP/βCD) offers several advantages due to its unique structural features. Its optimal use in pharmaceutical and medical applications would benefit from a molecular-level understanding of its behavior, as can be offered by molecular dynamics simulations. Here, we propose a set of parameters for all-atom simulations of HP/βCD, based on the ADD force field for sugars developed in our group, and compare it to the original CHARMM36 description. Using Kirkwood–Buff integrals of binary HP/βCD–water mixtures as target experimental data, we show that the ADD-based description results in a considerably improved prediction of HP/βCD self-association and interaction with water. We then use the new set of parameters to characterize the behavior of HP/βCD toward the different amino acids. We observe pronounced interactions of HP/βCD with both polar and nonpolar moieties, with a special preference for the aromatic rings of tyrosine, phenylalanine, and tryptophan. Interestingly, our simulations further highlight a preferential orientation of HP/βCD’s hydrophobic cavity toward the backbone atoms of amino acids, which, coupled with a favorable interaction of HP/βCD with the peptide backbone, suggest a propensity for HP/βCD to denature proteins.

INTRODUCTION

Cyclodextrins (CDs) are cyclic oligosaccharides containing 6 (αCD), 7 (βCD), or 8 (γCD) α-glucopyranose monomers. CDs possess a unique torus-like-shaped structure and are characterized by the presence of a lipophilic cavity and a hydrophilic outer surface. The internal core is surrounded by two rims (primary rim, formed by C6 atoms of the glucopyranose subunits, and secondary rim, consisting of the C2 and C3 atoms).

CDs count several applications as excipients, drug carriers, solubilizers, and adsorption enhancers. They can increase the solubility and bioavailability of hydrophobic drugs by including them within their lipophilic cavity. Because of their distinct features, CDs are present in many marketed drugs, and their field of application is supposed to grow further in the next few years. For instance, CDs have been widely used in formulations for oral, parenteral, nasal, pulmonary, and skin delivery of drugs, and there is widespread interest in their use for delivery to the brain.

Among CDs, βCD has arisen particular interest due to its structural characteristics. For instance, the cavity size of βCD allows the inclusion of aromatic amino acids, such as Phe, Tyr, His, and Trp, and this mechanism is supposed to be at the basis of the effective prevention of protein aggregation observed for this CD and its derivatives. However, βCD is poorly soluble in water (the solubility is only 16 mM at 25 °C), and this makes it unsuitable for parenteral formulations. This problem can nevertheless be mitigated by the addition of hydrophilic derivatizations to the glucopyranose subunit, for instance, by substituting some hydroxyl groups with other moieties. Among the possible derivatizations, hydroxypropyl groups may be linked to the glucopyranose subunit, eventually obtaining the so-called hydroxypropyl-β-cyclodextrin (HP/βCD). HP/βCD is more soluble than its unfunctionalized counterpart (>300 mM at ambient temperature) and displays amphiphilic properties. For instance, it was found to be surface-active, as such mitigating surface-induced aggregation of proteins. HP/βCD is approved as an excipient in parenteral formulations and was authorized by both FDA (in 2010)
and EMA (in 2013) for the treatment of the Niemann–Pick type C (NPC) disease. NPC disease is a progressive neurodegenerative disorder characterized by the intracellular accumulation of cholesterol. The effectiveness of HP/βCD against NPC is related to its action as a cholesterol-scaping compound, capable of transporting cholesterol away from cell bodies. An elevated level of cholesterol is also a risk factor for Alzheimer’s disease, suggesting that HP/βCD may also prove beneficial for its treatment. However, observations on this point are conflicting.

The pharmaceutical and medical applications of HP/βCD would benefit from a better characterization of its behavior at the molecular level, as this could be provided by molecular dynamics simulations. For this purpose, a force field capable of describing with good accuracy HP/βCD–water and HP/βCD–protein interactions is needed. We recently developed a force field for sugars and polyols, named ADD, and showed that it can correctly describe sugar–protein and sugar–water interactions, as well as self-association of sugars. We developed the new parameters in combination with the CHARMM36m force field for proteins, but good compatibility was also observed with other extensively used force fields, such as the AMBER 99SB-ILDN and the OPLS-AA force fields. The ADD parameters represent, therefore, a very promising candidate for the description of the α-glucopyranose subunit of cyclodextrins.

We will here test the application of the ADD force field to CDs, comparing the output to previous force fields and available experimental data. In particular, the Kirkwood–Buff integrals will be used as target experimental data for validation because they were found to be an excellent benchmark for force field development. We will focus our attention on HP/βCD because of its unique properties and numerous applications and because of the extensive experimental characterization available for this functionalized CD. Different possible forms of HP/βCD exist, depending on the degree of substitution and position of the derivatization, and we here study the form where the hydroxypropyl group is linked to the O2 atom of the glucose unit (2-HP/βCD) and fully substituted for all of the seven residues. A ball-and-stick representation of the 2-HP/βCD variant studied in this work is shown in Figure 1A, where the primary and secondary rims have been highlighted.

In the following, we will first validate the ADD description for 2-HP/βCD and show how it outperforms previous force fields. We will then study the interaction of 2-HP/βCD with the 20 naturally occurring amino acid side chains and the peptide backbone to gain an understanding of protein–HP/βCD systems at the molecular level.

**MATERIALS AND METHODS**

**Theoretical Background.** The ADD force field for 2-HP/βCD will be validated against experimental quantities related to the Kirkwood–Buff theory of solutions. In this context, a central role is played by the Kirkwood–Buff integrals (KBIs), which are used to describe the solvation behavior of component j around a reference particle i. The radial distribution function (RDF), which describes the variations in component j density as a function of the distance r from component i. A value of $G_j < 1$ indicates exclusion, while $G_j > 1$ indicates accumulation of component j around the reference i.

From now on, we will refer to a system containing water (component 1), HP/βCD (component 3), and, optionally, also an amino acid or peptide (component 2).

For a binary water–HP/βCD mixture (no component 2), the KBIs are related to the composition of the solution (molar density of water $\rho_1$ and HP/βCD $\rho_3$), to the chemical potential of HP/βCD $\mu_3$, and to the partial molar volumes of water ($V_1$) and HP/βCD ($V_3$) as follows:

$$G_{15} = \frac{\xi - \eta V_1 V_3}{\eta}$$  
(2)

$$G_{35} = \frac{\eta V_1 - 1}{\rho_3} + G_{13}$$  
(3)

$$G_{11} = \frac{\eta V_3 - 1}{\rho_1} + G_{13}$$  
(4)

Figure 1. (A) Ball-and-stick representation of 2-HP/βCD. The primary and secondary rims are highlighted in blue and yellow, respectively. (B) Schematic representation of the 2-HP/βCD subunit. The atom types and partial charges used for the simulations are also shown. When different charges are employed for the CHARMM36 and ADD force fields, they are shown with different colors (black for the original CHARMM36 and red for ADD).
Table 1. Lennard-Jones Parameters $\epsilon$ for the Hydroxyl Groups of HP/CD

| force field            | $\epsilon(O)_{\text{glucopyranose}}$ (kJ/mol) | $\epsilon(H)_{\text{glucopyranose}}$ (kJ/mol) | $\epsilon(O)_{\text{hydroxypropyl}}$ (kJ/mol) | $\epsilon(H)_{\text{hydroxypropyl}}$ (kJ/mol) |
|------------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|
| original CHARMM36      | 0.804                                         | 0.192                                         | 0.636                                         | 0.192                                         |
| ADD                    | 0.450                                         | 0.120                                         | 0.450                                         | 0.120                                         |

Table 2. List of the Simulations Performed in This Work

| sim. type (#) | solute (component 2) | HP/CD conc. (mol/L) (component 3) | box size (nm) | $T$ (K) | duration (ns) |
|---------------|-----------------------|---------------------------------|---------------|--------|--------------|
| 1             | 0.061−0.123−0.186     | 8 x 8 x 8                      | 298           | 60     |
| 2             | capped amino acids    | 0.050                          | 8 x 8 x 8     | 300    | 60           |
| 3             | NAG_A                 | 0.050                          | 8 x 8 x 8     | 300    | 100          |

$$
\eta = \frac{RT(\rho^2/\rho_1)}{(\partial h_3/\partial x_3)_{p,T}}
$$

(5)

$$
\xi = R\beta_4 \eta
$$

(6)

$$
\rho = \rho_1 + \rho_3
$$

(7)

where $R$ is the universal gas constant, $T$ and $p$ are temperature and pressure, respectively, $x_i$ are mole fractions, $\rho$ is the total molar density of the solution, and $\beta_4$ is the isothermal compressibility.

Knowing $G_{11}$, $G_{13}$, and $G_{13}$, it is then possible to compute other properties. For instance, the derivative of the mole fraction scale activity coefficient $f_{33}$ with the mole fraction $x_3$ can be evaluated as follows:

$$
f_{33} = \left( \frac{\partial \ln f_{33}}{\partial \ln x_3} \right)_{p,T} = -\frac{\rho_3 x_3 (G_{11} + G_{33} - 2G_{13})}{1 + \rho_1 x_3 (G_{11} + G_{33} - 2G_{13})}
$$

(8)

In a ternary solution containing also an amino acid or peptide (component 2), the KBIs can provide interesting information about the HP/CD—peptide interactions. For instance, the difference $\gamma = G_{33} - G_{12}$ quantifies the preferential interaction or exclusion of component 3 from component 2. Specifically, a negative value of $\gamma$ indicates preferential exclusion and vice versa.

Simulation Details. Molecular dynamics simulations were carried out using Gromacs 2018.6.34 Two different force fields were compared for 2-HP/CD. In the first case, the original CHARMM36 force field was used for the glucopyranose subunits, while the hydroxypropyl derivatization was modeled with parameters obtained by analogy, as provided by the CHARMM general force field (CGenFF) program.35 In the second case, charges were adjusted to comply with the ADD force field for carbohydrates.36 In the ADD force field, the partial charges of the hydrogen atom in the hydroxyl group are modified compared to the original CHARMM36 force field (0.33 for H in the ADD description, compared to 0.42 in the original CHARMM36). Because of this, also the other charges need to be modified to preserve the overall neutrality, and the atom types and charges used for the simulations are shown in Figure 1B.

An additional difference of the ADD description compared to the original CHARMM36 resides in the Lennard-Jones parameter $\epsilon$ of the O and H atoms of the hydroxyl group, as shown in Table 1. Specifically, the original CHARMM36 description uses two different combinations of $\epsilon$ values for hydroxyl groups within the glucopyranose subunit or in the hydroxypropyl derivatization. In contrast, the ADD force field employs a single set of $\epsilon$ values for all hydroxyl groups. No difference exists between the two force fields for what concerns the Lennard-Jones parameters $\sigma$.

For all simulations, the CHARMM TIP3P water model was used, while, whenever present, amino acids were described according to the CHARMM36m force field.22 A scheme of all simulations performed, with the corresponding box size, temperature, and duration, is listed in Table 2.

In sim. type 1, three different concentrations of HP/CD were considered in the range of 61−186 mM. These simulations were carried out with the objective to extract selected properties of binary water−HP/CD solutions (density, Kirkwood−Buff integrals, derivative of activity coefficient) to be then compared with experimental data. A temperature of 298 K was selected for these simulations to allow a direct comparison with the experimental data used for validation.

For sim. type 2, all of the 20 naturally occurring amino acids, in their capped form (i.e., acetylated N-terminus and amidated C-terminus), were simulated in 50 mM HP/CD. Also, the N-acetyl glycaminide series (NAG_A) was simulated (sim. type 3 in Table 2). NAG_A corresponds to a series of molecules with a varying number $x$ of glycine residues linked by a peptide bond and whose termini are blocked by an acetyl (N-terminus) and an amide (C-terminus) moiety. The number of internal glycine units has been varied from 1 (NAG_A) to 6 (NAG_A). For simulations 2 and 3, the box was cubic with an $\approx 8$ nm side length and included 25 amino acid/NAG_A molecules. For charged residues, Na$^+$ or Cl$^-$ ions were added to reach neutrality.

In all cases, the cutoff radius for both Coulombic (calculated using the PME method37) and Lennard-Jones interactions was set to 1.2 nm, and periodic boundary conditions were used. Each box was first energy-minimized with the steepest descent algorithm and then equilibrated for 1 ns at 1 bar and 298 K (sim. 1) or 300 K (sim. 2−3) in the NPT ensemble, using Berendsen pressure (3 ps relaxation time) and temperature (0.5 ps relaxation time) coupling.38 The simulations were then run at the same temperature used for equilibration and at 1 bar in the NPT ensemble, controlling temperature and pressure with the Nosé−Hoover thermostat39,40 (0.5 ps relaxation time) and Parrinello−Rahman barostat41 (3 ps relaxation time), respectively. A 2 fs time step was used, and configurations were saved every 2 ps. The Lincs algorithm was employed for constraining all bonds,42 while the SETTLE algorithm was used to keep the water molecules rigid.35 The last 50 ns (for sim. type 1) or 40 ns (for sim. types 2 and 3) were used for the analyses.

Analyses of Simulation Results. Kirkwood−Buff Integrals. The Kirkwood−Buff integrals were calculated by taking the average of the running KBIs.
Comparison between experimental (solid black line) and simulation results (black squares, original CHARMM36; red squares, ADD force field). (A) Solution density, (B) HP/βCD–HP/βCD Kirkwood–Buff integral $G_{ij}$, (C) HP/βCD–water Kirkwood–Buff integral $G_{i0}$, and (D) derivative of the mole fraction scale activity coefficient with respect to mole fraction.

RESULTS AND DISCUSSION

Validation of the ADD Force Field for HP/βCD. The first objective of the present work was to compare the original CHARMM36 and the ADD force fields, using target experimental data as reference. The simulations type 1 listed in Table 2 were performed for this purpose. The experimental data considered in this work were the solution mass density, the Kirkwood–Buff integrals $G_{ij}$ and $G_{i0}$, and the derivative of the mole fraction scale activity coefficient with respect to the mole fraction. The solution density and the partial molar volume of HP/βCD ($V_1$, that comes into play in eqs 2 and 4) were obtained from ref 45. Due to the low concentration of HP/βCD considered in this work (the maximum value we simulated was only 0.186 mol/L), we assumed the partial molar volume of water ($V_1 = 18.07$ cm$^3$/mol) and the isothermal compressibility ($\beta_T = 4.52 \times 10^{-10}$ Pa$^{-1}$) to be approximately constant. The molar fraction scale activity coefficient $f_{3w}$ of 2-HP/βCD was obtained from ref 46 and converted to the mole fraction scale using the relation

$$f_{3w} = \frac{\nu}{\nu}$$

where $\nu$ is the number of particles of type $j$ in the system, $V$ is the system volume, $\Delta N_j(r)$ is the excess number of particles $j$ within a sphere of radius $r$ around $i$, and $\delta_j$ is the Kronecker delta.

Amino Acid Inclusion within the HP/βCD Cavity. An amino acid was deemed to be included within the HP/βCD cavity when it was closer than 0.5 nm to the center of mass of HP/βCD. The 0.5 nm cutoff was selected so as to guarantee that only amino acids really included within the cavity, and not simply close to it, were counted. The number of CDs involved in inclusions was counted for each frame, averaged over the total number of frames and subsequently normalized by the total number of cyclodextrins in the system to allow a direct comparison of the different simulations.

Hydrogen Bonding. The number of amino acid–HP/βCD hydrogen bonds was computed. The $\chi$-parameter was evaluated as follows

$$\chi = \frac{\text{number of amino acid–HP/βCD hydrogen bonds}}{\text{total number of amino acid–HP/βCD and amino acid–water hydrogen bonds}}$$

To determine the presence of a hydrogen bond, a geometrical criterion was used, requiring that the distance between the donor and acceptor was less than 0.35 nm and that the angle formed between the hydrogen atom and the line joining the center of masses of donor and acceptor was smaller than 30°.
HPβCD Interacts with Polar and Apolar Side Chains but Is Excluded from Charged Moieties. Simulations type 2 in Table 2 were used to study the interaction of HP/βCD with the different amino acids. In particular, the values of $\gamma = G_{23} - G_{12}$ were extracted from the simulations.

We were particularly interested in studying the interaction of HP/βCD with the different side chains and the protein backbone separately. For this purpose, we followed the approach proposed by Auton et al.\textsuperscript{46,49} and already employed in our previous work.\textsuperscript{21} Briefly, we assumed the existence of additivity for the $\gamma$ values and computed the side-chain contribution $\gamma^\text{sc}$ by subtracting the KBI for glycine $\gamma^\text{gly}$ to the KBI of the specific amino acid $i$ being considered $\gamma_i$.

$$\gamma_i^\text{sc} = \gamma_i - \gamma^\text{gly} \quad (12)$$

Capped amino acids were used for this purpose. As shown in ref 21, the side-chain contribution $\gamma^\text{sc}$ is not influenced by the terminal capping conditions (i.e., zwitterionic vs capped form). This is also in line with previous observations by Nozaki and Tanford,\textsuperscript{40} who showed that the interaction of the side chain of a branched organic compound with the solvent was approximately independent of the interaction of the backbone to which the side chain was attached.

The values of $\gamma_i^\text{sc}$ obtained in this way are displayed in Figure 3A. On average, HP/βCD was found to be excluded (negative $\gamma_i^\text{sc}$), or only marginally interacting, with charged side chains. This was true for both the original CHARMM36 and ADD force fields and is in line with what observed in Arsiccio et al.,\textsuperscript{21} for sucrose and sorbitol. However, this may be a limitation of the force field, as a mismatch between simulations and experiments was noted for charged moieties.\textsuperscript{21}

The interaction with polar side chains was mostly favorable, with the only exception of asparagine, glutamine, and serine, for which a very small or negative value of $\gamma_i^\text{sc}$ was measured.

The interaction was favorable, and quite intense, also for most apolar side chains, with the only exclusion of alanine, that showed a value of $\gamma_i^\text{sc}$ close to zero. The ability of HP/βCD to interact with both polar and apolar side chains is not surprising considering the amphiphilic nature of this excipient, which was already the subject of both experimental and computational investigation.\textsuperscript{12,14,16}

The interaction of HP/βCD with aromatic side chains (especially tryptophan, phenylalanine, and tyrosine) was particularly pronounced. The preferential interaction of HP/βCD with aromatic groups has been observed experimentally\textsuperscript{9,10} and is well documented in the literature.\textsuperscript{7} The entity of the interaction was substantially different for the two force fields, with the ADD parameters resulting in less pronounced interaction with polar/apolar groups and reduced exclusion from charged side chains. However, it is interesting and important to note that the trend was very similar for both the original CHARMM36 and the ADD force fields, with an almost perfect agreement in the predicted sign of the $\gamma_i^\text{sc}$ values.

**HP/βCD-Backbone Interaction: Evidence of a Denaturing Behavior.** Having characterized the HP/βCD—side-chain interactions, we now set to study the interaction with the backbone. The backbone contribution $\gamma^\text{bb}$ can be computed according to the constant increment method\textsuperscript{51} applied to the NAG$_x$A series (sim. type 3 in Table 2)

$$\gamma = \gamma^\text{eg} + \gamma^\text{bb} x \quad (13)$$

where the value of $\gamma$ for the entire NAG$_x$A is decomposed into the end group contribution ($\gamma^\text{eg}$) and the backbone contribution ($\gamma^\text{bb}$) multiplied by the number of internal glycine units $x$. $\gamma^\text{bb}$ can therefore be obtained by fitting a straight line through the $\gamma$ values as a function of the number of internal glycine units, as shown in Figure 3B.

The $\gamma^\text{bb}$ values obtained are positive for both the original CHARMM36 and the ADD force fields (2.33 and 1.20 nm$^3$, with good coefficients of determination $R^2$ values of 0.84 and 0.94, respectively). This points to a favorable interaction of HP/βCD with the protein backbone. According to Auton et al.,\textsuperscript{46,52} this is indicative of a denaturing character of HP/βCD. The denaturing nature of this cyclodextrin is also in line with previous experimental investigations for immunoglobulin G formulations,\textsuperscript{53} where increasing HP/βCD concentrations were
found to reduce the protein melting temperature. As previously noted for the side-chain contributions already, the original CHARMM36 and ADD force fields differ in the entity of the predicted $\gamma_{bb}$ but not in the sign of the interaction.

**HP/CD Preferentially Orients with Its Secondary Rim toward the Amino Acid Backbone.** We previously mentioned that HP/CD is an amphiphilic molecule, with two rims (Figure 1A) characterized by different properties. Hence, we analyzed the orientation of HP/CD toward the different amino acids.

For this purpose, we computed the radial distribution functions (RDFs) for the atoms of both rims, with respect to the center of mass of the amino acids, and extracted the maximum value of these RDFs (Figure 4). We can observe that, in general, HP/CD mostly oriented with its secondary rim toward the amino acids. The preferential orientation of HP/CD was not very pronounced only for charged amino acids, likely because of the marginal interactions between the CD and the charged moieties as already evidenced in Figure 3A. Again, the overall trend was similar for both the CHARMM36 and ADD force fields, with the ADD description resulting in weaker interactions.

After having established that HP/CD preferentially orients with its secondary rim toward the amino acids, we verified whether it was closer to the side chain or the peptide backbone. For this purpose, we calculated the RDFs between the backbone or side-chain atoms of the different amino acids and the center of mass of the HP/CD secondary rim. The maximum values of the RDFs obtained in this way are shown in Figure 5. The trend observed suggests that interactions preferentially occur between the hydrophobic cavity (secondary rim) of the CD and the peptide backbone, in line with the...
favorable HP/βCD—backbone interaction evidenced in Figure 3B. Tryptophan constituted, however, a noticeable exception. In this case, HP/βCD strongly interacted with the aromatic ring of the tryptophan side chain, in line with experimental works suggesting a preferential affinity of CDs towards aromatic groups.\(^7\)\(^-\)\(^12\)

HP/βCD may also form inclusions with amino acids. We calculated the percentage of HP/βCD molecules that, on average, included amino acids within their cavity during the equilibrated trajectories (Figure 6A). We found that the occurrence of inclusions was in agreement with the previously discussed trends of \(\chi^2\) (Figure 3A) and maximum values of the RDFs (Figures 4 and 5). Again, inclusion was negligible for charged residues and more pronounced for polar and apolar ones, especially for the aromatic residues tyrosine, phenylalanine, and tryptophan. The original CHARMM36 and ADD force fields displayed the same overall trend, although the degree of interaction was lower with the ADD description. These same conclusions could be drawn looking at the formation of hydrogen bonds (\(\chi\) parameter; eq 11), as displayed in Figure 6B. This figure suggests that hydrogen bonding contributes to HP/βCD—amino acid interactions.

\section*{CONCLUSIONS}

HP/βCD is emerging as an important agent for drug delivery, as its unique structural characteristics make it particularly well suited to increase the solubility and bioavailability of hydrophobic drugs through encapsulation. In this work, we have developed a new force field for HP/βCD that accurately describes its physical properties. The original CHARMM36 force field was used as the starting point for this investigation, and we have exposed its limitations in reproducing experimental data of binary water—HP/βCD mixtures. We then proposed and validated a new force field based on the ADD description of sugars,\(^7\)\(^-\)\(^11\) which accurately reproduces HP/βCD—HP/βCD and HP/βCD—water interactions.

We have applied the new force field to the characterization of HP/βCD—amino acid interactions, quantified in terms of Kirkwood—Buff integrals (KBIs), and compared our results to the original CHARMM36 description. We have observed that HP/βCD only marginally interacts with charged side chains, while it exhibits strong affinities toward most of the apolar moieties, especially the aromatic ones. For polar side chains, strong interaction was noted with the aromatic ring of tyrosine and the thiol side chain of cysteine. The trends observed for the CHARMM36 and ADD force fields were similar, but CHARMM36 predicted considerably stronger interactions. The poor interaction between charged groups and HP/βCD, observed with both CHARMM36 and ADD, may, however, be a limitation of the force fields, as a mismatch between simulations and experiments has previously been observed for charged moieties in ref 21.

Both the CHARMM36 and ADD force fields evidenced a favorable interaction of HP/βCD with the peptide backbone, which may be indicative of its denaturing behavior. In line with this, we further found that HP/βCD mainly oriented with its secondary rim, i.e., the hydrophobic cavity, toward the peptide backbone. However, the aromatic amino acid tryptophan strongly interacted with HP/βCD also through its side chain.

As cyclodextrins are known for their ability to form complexes with small molecules, we evaluated the degree of amino acid inclusion inside the HP/βCD’s cavity. We observed that the tendency to form inclusion complexes with different amino acids reflected the chemical affinity highlighted by our KBIs. Inclusion was also accompanied by an increased number of hydrogen bonds between the cyclodextrins and the amino acids.

We believe that the improved description of HP/βCD provided in this paper will prove to be useful for the investigation of this promising excipient, enabling the use of computational techniques, such as molecular dynamics, to further clarify its properties. A better understanding of HP/βCD—protein mixtures would be of great interest in the pharmaceutical, biotechnological, and medical fields. In this respect, the characterization of HP/βCD—amino acid interactions described here constitutes the first step toward unraveling this molecule’s behavior toward proteins.
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Notes
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