Para-halogenated benzaldehyde molecules included in cyclodextrins: a combined spectroscopic and thermal analysis

Paulo J. A. Ribeiro-Claro,1* António M. Amorim da Costa,2 Manuel L. Vueba,3 M. Eugénia Pina3 and Ana M. Amado2

1 CICECO, Departamento de Química, Universidade de Aveiro, P-3810–193 Aveiro, Portugal
2 Química-Física Molecular, Departamento de Química, Universidade de Coimbra, P-3004-535 Coimbra, Portugal
3 Laboratório de Galêncica e Tecnologia Farmacêutica, Faculdade de Farmácia, Universidade de Coimbra, P-3000–295 Coimbra, Portugal

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The inclusion complexes of 4-X-benzaldehyde guests (X = F, Cl, Br) in α-, β- and γ-cyclodextrins (αCD, βCD and γCD) were prepared and characterized by means of thermogravimetry, differential scanning calorimetry (DSC) and vibrational spectroscopy. The stoichiometry of the inclusion complexes ranges from 2:1 to 1:2, depending on the cavity size of the host. Sensitive vibrational modes such as C=O and ring CH stretching modes were used to monitor the effects of the inclusion process and to assess the preferred inclusion geometry for each host-guest pair. On the whole, the spectral observations suggest that the small αCD cavity imposes important structural restrictions on the guest molecule, while the larger γCD cavity allows ‘liquid-type’ intermolecular contacts. Copyright © 2005 John Wiley & Sons, Ltd.

KEYWORDS: cyclodextrins; benzaldehyde derivatives; hydrogen bonding; thermal analysis

INTRODUCTION

Cyclodextrins (CDs) are known to form inclusion complexes with a wide variety of molecules,1,2 the only requirement being that at least part of the guest can penetrate into the cavity. This ability gives them relevant practical benefits in several areas, namely, in the pharmaceutical, food, cosmetic and agricultural industries.3–14 Many active ingredients, such as flavours and drugs, are protected from degradation and/or evaporation upon encapsulation in the CD cavity. Moreover, CDs can greatly increase the solubility of water-insoluble substances, as well as control the release of encapsulated ingredients, flavours or drugs.

Different analytical techniques have been used to determine whether the product obtained is a true inclusion complex, as well as to analyse the molecular structure and interactions established in the host–guest complex.4,5,15–26 Raman spectroscopy and differential scanning calorimetry (DSC) are among the most accessible experimental techniques for the study of inclusion complexes in the solid state.4–6,18–29

The aim of the present work is to pursue the study and the characterization of the interactions occurring in the α-, β- and γCD inclusion complexes with aromatic guests.18–20 The following benzaldehyde derivatives are considered as guest molecules: 4-fluorobenzaldehyde (4FB), 4-chlorobenzaldehyde (4ClB) and 4-bromobenzaldehyde (4BrB). The guest : host : H2O stoichiometries of the inclusion complexes are inferred from DSC and thermogravimetric (TG) data. Raman spectroscopy is used to characterize the complexes and rationalize the preferences of guest inclusion orientation as well as to establish the host–guest intermolecular interactions. As done previously,18–20 Fourier transform infrared (FTIR) spectroscopy has not been used, as most of the guest bands are either too weak or strongly overlapped by CD bands (even the strong νC=O mode is masked by the H–O–H deformation modes of the structural water molecules).

EXPERIMENTAL

Materials and preparation of inclusion complexes

Cyclodextrins (α-, β- and γCD) were kindly offered by Wacker Chemie GmbH, Germany. The benzaldehyde derivatives (4FB, 4ClB and 4BrB) were obtained commercially (Aldrich) and were used without further purification.

*Correspondence to: Paulo J. A. Ribeiro-Claro, CICECO, Departamento de Química, Universidade de Aveiro, P-3810–193 Aveiro, Portugal. E-mail: pclaro@dq.ua.pt
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The inclusion complexes of the liquid 4FB guest were prepared by mixing the benzaldehyde derivative with an aqueous solution of CD (dropwise addition with strong agitation) at room temperature. In the case of the solid benzaldehyde derivatives (4ClB and 4BrB), because of their low solubility in water, the CD solution was kept under strong agitation in a bath at 50 °C during the guest addition.

In both cases, the process led to the rapid formation of white precipitates, which were filtered and allowed to dry for a few days, at room temperature and ambient relative humidity conditions. After this, the precipitates were ground to a fine powder. The complexes were found to be stable for several months.

**Differential scanning calorimetry (DSC) and thermogravimetric (TG) analysis**

The DSC measurements were carried out using a Shimadzu DSC-50 differential scanning calorimeter with a thermal analyser, Shimadzu TA-50. Samples were heated in sealed aluminium pans under a nitrogen flow of 20 ml/min, using a sealed empty pan as reference, and at a heating rate of 10 °C/min, over the temperature range from 25 to 400 °C.

The thermogravimetric analysis (TGA) measurements were performed on a TGA-50 Shimadzu apparatus. The accurately weighed samples were placed in aluminium pans and sealed. All samples were kept for 5 min at 20 °C, before heating up to 400 °C, at a scanning rate of 2 °C/min.

**FT-Raman spectroscopic experiments**

The room temperature FT-Raman spectra were recorded on a RFS-100 Bruker FT spectrometer using a Nd:YAG laser with the excitation wavelength of 1064 nm. Each spectrum is the average of two repeated measurements of 150 scans each and 2 cm⁻¹ resolution. Samples were sealed in Kimax glass tubes of 0.8-mm inner diameter. Higher-temperature spectra were recorded on a Jobin-Yvon T64000 triple monochromator system (focal length 0.640 m, aperture f/7.5) with a non-intensified charge-coupled device (CCD) detector and an Ar⁺ laser as the excitation source, using a home-made Harney–Miller assembly. An integration time of 5 s and two scans were used.

**Theoretical calculations**

Theoretical calculations were carried out using the Gaussian 98 W program package. Full geometry optimizations and vibrational wavenumber calculations of the guest molecules were performed at the standard B3LYP/6–31G* theory level. The effect of the polarity of the environment on the vibrational wavenumbers of the guest molecules was evaluated using the self-consistent reaction field approach, with the Self-Consistent Isodensity Polarized Continuum Model (SCI-PCM). This approach (SCI-PCM at the B3LYP/6–31G* level) was found to be appropriate to predict the wavenumber dependence on the dielectric media and allowed the comparison with previous studies.

**RESULTS AND DISCUSSION**

**Differential scanning calorimetry (DSC) and thermogravimetric (TG) analysis**

When guest molecules are embedded in CD cavities or in the crystal lattice, their melting, boiling or sublimation points generally shifts to a different temperature. Moreover, as inclusion compounds of CDs are generally prepared in aqueous media, they must be considered as ternary water/CD/guest systems. DSC combined with TGA represents an analytical tool of first choice for an accurate assessment of the inclusion-compound stoichiometry.

The DSC traces of the pure solid guests 4ClB and 4BrB (the DSC curve of pure 4FB was not obtained because of its liquid state) exhibit a sharp endothermic peak centred at 48 and 58 °C, respectively, corresponding to the melting points of those benzaldehyde derivatives (Sigma-Aldrich tabulated values are 47–50 and 55–58 °C, respectively). In each case, two additional broad endothermic peaks are also observed. In the case of 4ClB, the first one is centred at 114 °C, and can be associated with thermal flashing of the sample (Sigma-Aldrich tabulated value is 87 °C), while the second one is centred at 160 °C, and probably indicates thermal degradation of the sample. For 4BrB, these two effects yield the broad peaks centred at 124 °C (Sigma-Aldrich tabulated flashing point value is 108 °C) and 172 °C, respectively. On the other hand, the DSC and TGA traces of α-, β- and γ-CDs are in agreement with those reported in the literature.

Figure 1 shows the DSC and TGA traces of αCD•4BrB inclusion complex. The DSC curves of pure 4BrB and αCD are included for comparison. The thermal curves (DSC and TG) of the remaining CD inclusion complexes are not shown, since they present the same general profile and main features exhibited by the αCD•4BrB complex.

The DSC trace of αCD•4BrB complex displays two endothermic peaks between 20 and 104 °C due to the dehydration of the complex. The occurrence of these two bands indicates the existence of distinctly bound water molecules in the inclusion complex. This stepwise dehydration is evident in the TGA curve and resembles the effect observed for αCD itself. However, the broadening and shifting of the bands indicate differently linked water structures in the two systems (αCD and αCD•4BrB). Moreover, the absence of the two guest peaks centred at 124 and 172 °C confirms the inclusion of the guest (the two peaks are present in the DSC trace of the physical mixture αCD + 4BrB).

The small feature centred at 269 °C is associated with release of the guest from the complex. The mass loss leads to a baseline decline in the TGA curve in the same temperature range (Fig. 1).

On the basis of the combined DSC/TGA, the stoichiometries of the inclusion complexes were achieved by quantification of the different mass losses identified in the TG curves, upon temperature increase. The results are summarized in...
Concerning the water content of the samples, the \( \alpha \)CD inclusion complexes present the lowest number of water molecules per CD (5 to 7). For \( \beta \)CD and \( \gamma \)CD inclusion complexes, the number of water molecules ranges from 9 to 13 (Table 1). These values are in agreement with the reported values for similar inclusion complexes. The X-ray structure of the \( \alpha \)CD-benzaldehyde complex shows the presence of six interstitial water molecules. Thermogravimetric analyses of \( \alpha \)CD\( \cdot \)4-F-phenol, \( \beta \)CD-benzaldehyde, and \( \beta \)CD-anisaldehyde yield nine water molecules per CD. Although there is no information concerning the number of intra-cavity and extra-cavity (interstitial) water molecules, the TG curves clearly show the presence of differently bound water molecules in all systems. For instance, in the TG curve of the \( \alpha \)CD\( \cdot \)4BrB complex (Fig. 1), there are two well-defined steps in the water-loss region, corresponding to the mass of two water molecules (3.31% at ca 50 °C) and three water molecules (4.94% at ca 75 °C).

Vibrational spectra
As in the previous report, emphasis will be given to the C=O stretching mode (vC=O), as it is relatively free from interfering bands of the CD host and is particularly sensitive to the environment around this molecular fragment. Thus, the analysis of this spectral region can yield valuable information concerning the involvement of the C=O groups in intermolecular host–guest and/or guest–guest interactions. The effects of CD inclusion on other spectral regions, namely, those assigned to the ring CC and CH stretching modes (vCC and vCH) and to the CX stretching mode (vCX), are also analysed.

The vC=O region
Figure 2 presents the 1650–1750 cm\(^{-1}\) Raman spectral region of pure 4XB derivatives (X = F, Cl and Br) and their inclusion complexes with \( \alpha \)-, \( \beta \)- and \( \gamma \)CDs. Since 4ClB and 4BrB are solids at room temperature, the spectra of pure guests reported herein are from melted samples, for easier

Table 1. Calculated stoichiometry of the inclusion complexes based on DSC and TGA

<table>
<thead>
<tr>
<th>System</th>
<th>% (Guest)</th>
<th>% (H(_2)O)</th>
<th>Complex stoichiometry</th>
<th>Water molecules per CD</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \alpha )CD( \cdot )4FB</td>
<td>6.79</td>
<td>10.78</td>
<td>( \alpha )CD( \cdot )4FB</td>
<td>7 ( \text{H}_2\text{O} )</td>
</tr>
<tr>
<td>( \alpha )CD( \cdot )4ClB</td>
<td>5.37</td>
<td>9.73</td>
<td>( \alpha )CD( \cdot )4ClB</td>
<td>6 ( \text{H}_2\text{O} )</td>
</tr>
<tr>
<td>( \alpha )CD( \cdot )4BrB</td>
<td>4.51</td>
<td>8.25</td>
<td>( \alpha )CD( \cdot )4BrB</td>
<td>5 ( \text{H}_2\text{O} )</td>
</tr>
<tr>
<td>( \beta )CD( \cdot )4FB</td>
<td>8.93</td>
<td>11.55</td>
<td>( \beta )CD( \cdot )4FB</td>
<td>9 ( \text{H}_2\text{O} )</td>
</tr>
<tr>
<td>( \beta )CD( \cdot )4ClB</td>
<td>8.77</td>
<td>15.25</td>
<td>( \beta )CD( \cdot )4ClB</td>
<td>13 ( \text{H}_2\text{O} )</td>
</tr>
<tr>
<td>( \beta )CD( \cdot )4BrB</td>
<td>6.44</td>
<td>15.00</td>
<td>( \beta )CD( \cdot )4BrB</td>
<td>12 ( \text{H}_2\text{O} )</td>
</tr>
<tr>
<td>( \gamma )CD( \cdot )4FB</td>
<td>14.25</td>
<td>10.25</td>
<td>( \gamma )CD( \cdot )(4FB)( _2 )</td>
<td>10 ( \text{H}_2\text{O} )</td>
</tr>
<tr>
<td>( \gamma )CD( \cdot )4ClB</td>
<td>14.60</td>
<td>9.69</td>
<td>( \gamma )CD( \cdot )(4ClB)( _2 )</td>
<td>9 ( \text{H}_2\text{O} )</td>
</tr>
<tr>
<td>( \gamma )CD( \cdot )4BrB</td>
<td>12.58</td>
<td>13.31</td>
<td>( \gamma )CD( \cdot )(4BrB)( _2 )</td>
<td>13 ( \text{H}_2\text{O} )</td>
</tr>
</tbody>
</table>
The study of para-halogenated benzaldehyde in cyclodextrins reveals significant changes in the carbonyl band envelope upon inclusion. The splitting of the carbonyl band, previously assigned to Fermi resonance or dimerization equilibrium, is observed in the Raman spectra of the pure liquid guests. This splitting is more pronounced in the γCD complexes, where the presence of more than one band is evident, although the intensity ratios differ from those observed in the pure-liquid states. Inclusion in αCD leads to a narrowing of the carbonyl band envelope, while βCD inclusion results in a downward shift of the maximum of the carbonyl band envelope by approximately 4–5 cm⁻¹. In contrast, the γCD inclusion complexes show a distribution of the guest molecules with different inclusion geometries and hydrogen-bond contacts, leading to a decrease in the guest disorder and the assignment of isolated C=O groups.

Figure 2. Raman spectra in the 1650–1750 cm⁻¹ region of the pure 4XB derivatives (X = F, Cl, Br) and of their α-, β- and γCD inclusion complexes.
Figure 3. Raman spectra in the 3000–3150 cm⁻¹ region of the pure 4XB derivatives (X = F, Cl, Br) and of their α-CD, β-CD and γ-CD inclusion complexes.

observed for 4FB and 4ClB complexes, as well as the band at 1706 cm⁻¹ in the 4BrB complex, may be ascribed to the presence of extra-cavity guest molecules.

The νCH region
Figure 3 presents the Raman spectra of the pure 4XB derivatives (liquid) and their α-, β- and γ-CD inclusion complexes in the 3000–3150 cm⁻¹ spectral region, which has been assigned to the aromatic ring νCH modes.

Previous studies of CD complexes with aromatic guests show that the ring νCH modes may be sensitive to the inclusion process. For instance, inclusion of benzaldehyde in the α-CD cavity yields a significant upwards shifting (blue-shift) of the νCH band envelope, which has been correlated with the CH···O=C intermolecular hydrogen bond observed in the X-ray structure of this complex.

As can be seen in Fig. 3, there are two main effects resulting from the inclusion process of the benzaldehyde derivatives reported herein: the narrowing of the band profile (which is particularly evident for the α-CD complexes) and the large downward shifts of the band maxima in the β-CD complexes. In agreement with the observations on the νC=O region, the smallest differences relative to the pure liquid are observed for the γ-CD complexes.

The clear band narrowing observed in the α-CD complexes relative to the pure liquids can be easily related with the structural rigidity imposed by the CD cavity size. Somewhat less straightforward is the interpretation of the 7–10 cm⁻¹ red-shift of the νCH band in the β-CD complexes. The complexes with α-CD and γ-CD present clearly smaller shifts (ca 0–5 cm⁻¹), and this is an evidence of the distinct inclusion geometry in β-CD complexes relative to α-CD and γ-CD.

Other spectral regions
Apart from the νC=O and the aromatic νCH modes, which occur in regions free from CD bands, other guest modes cannot be systematically used to follow the effects of the inclusion process, because of the interference of the host bands. However, in some cases a relevant guest mode can be observed in one of the small windows present in the Raman spectra of CD hosts.

Figure 4 shows the Raman spectra of the pure 4FB and its CD inclusion complexes in the νCF spectral region (1165–1285 cm⁻¹). For pure 4FB, the νCF mode yields a well-defined band at 1227 cm⁻¹, in the vicinity of the νC=C(HO) band (1203 cm⁻¹). The CD inclusion of 4FB leads to an upward shifting of the νCF vibrational mode, which is significantly larger upon α-CD inclusion in relation to the one that occurs for the wider CDs (Δν = +10 cm⁻¹ vs Δν = +3 cm⁻¹). The significant shift of the νCF mode must be related with the different inclusion environment provided by αCD relative to β and γCD, and further supports the proposal of distinct inclusion geometries for these complexes.
Figure 4. Raman spectra in the 1165–1285 cm\(^{-1}\) region of the pure 4FB and of its \(\alpha\)-, \(\beta\)- and \(\gamma\)CD inclusion complexes.

Another interesting case is observed for the 4ClB systems in the region of the ring \(\nu\)CC modes at \(\approx 1600\) cm\(^{-1}\) (\(8a/8b\) pair according to Wilson’s notation for aromatic molecules\(^{45}\)). In the pure guest, the band intensity ratio of the 1596/1588 cm\(^{-1}\) band pair was found to be temperature dependent (results not shown). As shown in Fig. 5, this intensity ratio is also sensitive to the inclusion process. In fact, the inclusion in \(\gamma\)CD mimics the effect of temperature increase, while the inclusion in \(\alpha\)CD acts as in a temperature decrease. Although the temperature dependence of the 1595/1588 cm\(^{-1}\) intensity ratio is not yet fully understood, Fig. 5 stresses the conclusions drawn from the previous spectral regions: the \(\nu\)CC band profile in the \(\gamma\)CD complex is nearly identical to the one of the pure guest, while a large change is observed for the \(\alpha\)CD complex. The effect of \(\beta\)CD inclusion constitutes an intermediate situation, with the two components of the doublet presenting almost equal Raman intensity.

Theoretical calculations
Theoretical calculations can be used to assign the vibrational spectra and to predict the wavenumber shifts resulting from both specific and non-specific interactions with the CD cavity (e.g. hydrogen bonding vs changing dielectric environment). Since the CD cavity is known to have lower polarity\(^{46}\) than the region of the rims, the inclusion geometry would affect the \(\nu\)C=O and \(\nu\)CH modes in a predictable way.\(^{33}\)

Wavenumber calculations for the guest molecules in different dielectric media show that an increase of solvent (or environment) polarity leads to a wavenumber decrease (red-shift) of both the \(\nu\)C=O and the \(\nu\)CX modes, while it promotes a blue-shift of the \(\nu\)CH modes (Fig. 6). These results can be used to assist the interpretation of the observed spectral shifts (Figs 2–5).

The results for the \(\nu\)CF mode of 4FB are somewhat straightforward. The inclusion in the wider \(\beta\)- and \(\gamma\)CD yields a much larger upwards shift (+10 cm\(^{-1}\)) than the inclusion in \(\alpha\)CD (+3 cm\(^{-1}\)). These shifts are in agreement with the proposed stoichiometry determined for the complexes: the 2:1 ratio in the \(\alpha\)CD:4FB complex implies that both substituents of the guest are in an intra-cavity environment (low dielectric constant for the C–F group), while in the 1:1

Figure 5. Raman spectra, in the 1550–1650 cm\(^{-1}\) region, of the pure 4ClB derivatives and of its \(\alpha\)-, \(\beta\)- and \(\gamma\)CD inclusion complexes.

Figure 6. Plot of calculated wavenumber shifts (in cm\(^{-1}\)) of CH, CF and C=O stretching modes as a function of the dielectric constant of the medium (SCI-PCM model). The dashed lines are just eye-guides.
and 1:2 ratio of the wider β- and γCD complexes, a deeper inclusion of the guest probably leaves the C–F terminal close to one of the OH-rims.

Concerning the νCH modes, the α 7–10 cm⁻¹ downward shift (red-shift) observed for βCD is consistent with the change from a high dielectric medium (pure guest) to a low dielectric environment (the βCD cavity). This observation suggests deep inclusion geometry of the aromatic ring in the βCD cavity. The small shifts observed for both the αCD and the γCD complexes cannot be so easily interpreted. However, one can assume two reasonable explanations: (i) αCD does not allow the full inclusion of the aromatic ring, leaving the C–H groups in the inter-rims region between CDs and (ii) in γCD, the presence of two guests (and even H₂O molecules) affects significantly the dielectric properties of the host cavity.

The effects of the inclusion process on the νC=O mode seem to be related with specific interactions and not with the dielectric environment. In fact, the observed shifts are generally opposed to those predicted from Fig. 6. For instance, the 1:2 stoichiometry of the αCD complexes indicates an intra-cavity position of the C=O group. The red-shift observed for this mode is not consistent with a low-polarity medium and must result from specific interactions, such as hydrogen bonding with intra-cavity water molecules.

**CONCLUSIONS**

The combined results from TGA, Raman spectroscopy and ab initio calculations allow a consistent description of the inclusion complexes herein studied. The description is in agreement with previous vibrational spectroscopic studies of CD complexes with aromatic guests, and provides further information on the inclusion process in CD hosts.

In the case of the αCD complexes, the structure of the inclusion is determined by the small cavity of the host. The inclusion stoichiometry is 2:1, and the Raman spectra are consistent with a rigid structure, in which both substituent groups are inside the cavities of neighbouring αCD hosts and the phenyl ring lies in the inter-rim region. The presence of differently bound water molecules and the red-shift of the νC=O mode are interpreted in terms of the presence of hydrogen-bonding (C–H ⋯ O) interactions.

The wider cavities of the βCD and γCD hosts allow a deeper inclusion of the guests, with less rigid structures. In the βCD complexes, the results are consistent with a 1:1 inclusion stoichiometry, with full inclusion of the aromatic ring and leaving the substituent groups in the region of the rims. In the case of the γCD complexes, the observed 1:2 inclusion stoichiometry leads to a ‘liquidlike’ distribution and interactions of the guest molecules.

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