Hemin and bile pigments are the secondary structure regulators of intrinsically disordered antimicrobial peptides

Abstract: The interaction of protoporphyrin compounds of human origin with the major bee venom component melittin (26 a.a., Z +6) and its hybrid derivative (CM15, 15 a.a., Z +6) were studied by a combination of various spectroscopic methods. Throughout a two-sate, concentration dependent process, hemin and its metabolites (biliverdin, bilirubin, bilirubin ditaurate) increase the parallel β-sheet content of the natively unfolded melittin suggesting the oligomerization of the peptide chains. In contrast, α-helix promoting effect was observed with the also disordered but more cationic CM15. According to fluorescence quenching experiments, the sole Trp residue of melittin is the key player during the binding, in the vicinity of which the first pigment molecule is accommodated presumably making indole-porphyrin π-π stacking interaction. As circular dichroism titration data suggest, the cooperative association of additional ligands subsequently occurs resulting in multimeric complexes with an apparent dissociation constant ranged from 20 to 65 µM. Spectroscopic measurements conducted with the bilirubin catabolite uro- and stercobilin refer to the requirement of intact dipyrromethene moieties for inducing secondary structure transformations. The binding topography of porphyrin rings on a model parallel β-sheet motif was evaluated by VIS absorption spectroscopy and computational modeling showing a slipped-cofacial binding mode responsible for the red shift and hypochromism of the Soret absorption band. Our results may aid to recognize porphyrin-responsive binding motifs of biologically relevant, intrinsically disordered peptides and proteins, where transient conformations play a vital role in their functions.

Keywords: bilirubin; biliverdin; circular dichroism spectroscopy; CM15; fluorescence quenching; hemin; intrinsic disorder; melittin; secondary structure

Introduction

Heme serves as an essential prosthetic group for a wide range of proteins.1,2 Besides the vitally important catalytic and oxygen transfer functions, compelling evidences have been accumulated in recent years which point out that heme, as a regulator molecule, is also involved in numerous cellular and molecular pathways, such as cell proliferation/differentiation, gene transcription/translation and control of ion channel function.3,4 However, the underlying molecular mechanisms by which heme modulates the function of protein targets are poorly understood. From this point of view, the enzymatic degradation products of heme such as biliverdin, the strongly lipophilic bilirubin and its water soluble conjugate have received much less attention. Employing circular dichroism (CD), UV/VIS absorption, and fluorescence spectroscopic techniques, we report herein that hemin and related bile pigments (Scheme 1) can substantially affect the natively unfolded structure of melittin and its hybrid derivative called CM15 (Table 1).

Melittin accounts for major percentage of the crude venom of the honey bee Apis melifera. Its matured form shows multiple pharmacological actions including antibacterial, antiviral, anticancer and anti-inflammatory effects.5,6 According to the primary sequence, the N-terminal part of melittin (1-20) consists mostly of hydrophobic/neural residues whereas the C-terminal (21-26) is enriched in polar, basic side chains (Table 1). This unbalanced charge arrangement renders the peptide chain amphipathic. Melittin contains a lone proline residue that gives rise to a flexible hinge region at the middle of the molecule. Melittin is freely soluble in water and exhibits a monomer-tetramer equilibrium which sensitively depends on experimental conditions. The increase of the ionic strength (~1.5 M NaCl), pH (>8) and/or the peptide concentration (~0.1 mM) of the solution promote the tetramerization and the concomitant helical folding of melittin.13-15 In dilute aqueous environment the extended, monomeric peptide chains are mainly unstructured but also contain two short, somewhat ordered segments (residues 5-9 and 14-20).16 Upon binding to lipid membranes or in membrane mimetic environments, the α-helical content of melittin rises sharply from ~0-20% to ~70%.17-19 CM15 is a synthetic cecropin-melittin hybrid peptide that preserves the antimicrobial activity of the 37-mer cecropin A (from silk moth) but displays decreased hemolysis compared to
Table 2. It consists of 15 residues from the N-terminus of cecropin A (1-7) and melittin (2-9) possessing much higher average charge per residues than its congeners (Table 1).

TABLE 1 Amino acid composition and net charge (Z) at neutral pH of melittin, cecropin A and CM15. Note that positive charge at the N-terminus increases Z by 1 unit. Melittin and CM15 used in this work are amidated at the C-terminus. Residue sequences in melittin and cecropin A are indicated by bold.

![Table 1](image)

**Materials and Methods**

See Supporting Information.

**Results and Discussion**

Pigment binding induced conformational changes in melittin

The far-UV CD spectrum of free melittin measured in pH 7.3 potassium phosphate buffer is dominated by a negative peak centered around 203 nm, together with a weaker positive band below 195 nm and a shoulder above 215 nm (Fig. 1). Regarding the secondary structure, this ellipticity profile refers to the prevalence of the unordered fraction but is also indicative of a significant α-helix and/or β-sheet content. In line with this qualitative assessment, deconvolution of the CD curve showed the dominance of the disordered structure (~45%) and a minor α-helix and β-sheet contribution (Table 2). Upon successive addition of hemin, this CD pattern was converted into a new one. The amplitude of the main negative band gradually diminished and its λ_max shifted to about 209 nm. Concomitantly, the short-wavelength positive peak gained intensity and displayed a red shift up to 195 nm (Fig. 1). Additionally, the bathochromically shifted lowest energy broad band became resolved with appreciable intensity increase. According to the secondary structure analysis, these spectral transformations reflect the four-fold increase of the parallel β-sheet together with the modest rise of the regular α-helix content of the peptide (Table 2). Moreover, the reverse of the relaxed/right-handed twisted sheet ratio within the antiparallel β-sheet subgroups points to some additional conformational redistribution. Upon using biliverdin, bilirubin, and bilirubin ditaurate analogous CD spectral changes were witnessed (Fig. 2, Table 2). Both the visual comparison of the CD curves (Fig. S1) and the secondary structure data show that the parallel β-sheet promoting ability of hemin and bile pigments is close to each other (Table 2). This kind of structural transition of melittin has not been reported previously and is in contrast to the result of membrane interactions, when the low helicity of the peptide rises sharply to ~70%.[1,2,3,4] The development of parallel β-sheet fraction in the pigment loaded form of melittin suggests intermolecular sheet sheet formation, i.e. the ligand-mediated oligomerization of the peptide chains.

The BeStSel algorithm used in this work to predict the secondary structure content discriminates between two kinds of α-helical components.[2,22] The regular, middle part of the helix where all the backbone-backbone H-bonds are formed (H1) and the so called distorted helix (H2) that corresponds to 2-2 residues at both ends of the helix with unsatisfied H-bonding (Table 2). Interestingly, at the beginning of the CD titrations an abrupt, ~50% decline of the H1 fraction was detected which was completed around equimolar pigment-peptide composition (Fig. 3). Subsequent addition of the ligands, however, started the helix content to regain reaching a somewhat higher level at the end of the titration (Table 2). In this respect, hemin was the most effective substance which is also reflected by the emergence of the resolved, 209 nm negative branch of the π-π exciton CD couplet of the right-handed α-helix (Fig. S1).[24]

![Scheme 1](image)

**TABLE 2** Estimation of the secondary structure content of free and ligand-loaded forms of melittin (MLT) and CM15. Far-UV CD spectral data were analysed by using the BeStSel online platform available at bestsel.elt.ehu. BP: bile pigment; PP: peptide; HM: hemin; BV: biliverdin; BR: bilirubin; BRT: bilirubin ditaurate; SB: sterocobilin; UB: urobilin; H1: regular α-helix; H2: distorted α-helix; P): parallel β-sheet; A):1, left-hand twisted β-sheet; A):2, relaxed β-sheet; A):3, right-hand twisted β-sheet.

![Chart](image)
Noticeably, upon melittin association bilirubin ditarurate displays a negative-positive induced CD band pair allied to its VIS absorption band (Fig. S2). Such an exciton CD couplet was reported by several times for the macromolecular complexes of bilirubin and reflects the enantioselective binding of the pigment molecules to the chiral host.31 In line with the so-called exciton chirality rule,32 the CD signal pattern shown in Fig. S2 indicates the slight binding preference of the \( \beta \)-helical (left-handed) conformer (Fig. S3). Besides the main VIS absorption band between 340-550 nm, bilirubin ditarurate also shows light absorption in the UV region (Fig. S4). Accordingly, induced CD signals of ligand origin could also be present here and thus mixed into the far-UV CD spectrum of melittin. This effect may somewhat distort the CD curve and thus the result of the secondary structure analysis.
Pigment binding induced conformational changes in CM15

CM15 is even more disordered than melittin but responds distinctly to porphyrin derivatives (Fig. S5 and S6). Beside the slight increase of the parallel β-sheet fraction, a significant degree of helix content was induced with simultaneous reduction of the unordered fraction (Table 2). For instance, the biliverdin induced qualitative and quantitative CD spectral changes closely match to that obtained with heparin, the polyanionic helicogenic agent used in structural studies of disordered peptides and proteins (Fig. S7).\(^{34,36}\) In relation to bilirubin and its taurine conjugate, hemin and biliverdin were more effective helix promoters. These observations are similar to the impact of heme on the secondary structure of aggregating α-synuclein. In the presence of the pigment, this natively disordered protein adopts α-helical conformation instead of a β-sheet rich structure, the hallmark of pathological protein inclusions.\(^{37}\) Furthermore, the specific α-synuclein binding of porphyrin phthalocyanine tetrasulfonate has also been reported which results in the stabilization of helical structure of the membrane-associated protein.\(^{38}\) This small molecule-mediated structural effect has been attributed to aromatic stacking between the porphyrin ring and Phe/Tyr residues of α-synuclein. Likewise, the more prominent helix-inducing feature of hemin and biliverdin can tentatively be assigned to their aromatic stacking with the Trp (and Phe) side chain of CM15. Presumably due to structural reasons, bilirubins are less suitable for this purpose. The so called ridge-like conformation of these compounds differs sharply from the planar and close planar stereochecmy of hemin and biliverdin, respectively (Fig. S3).\(^{32,35,37}\)

Estimation of melittin binding parameters of hemin and bile pigments

Two approaches were used to calculate the apparent dissociation constants (\(K_d\)) of ligand-melittin complexes. CD curve of the free peptide exhibits a zero crossing point around 194-195 nm (Fig. 1 and 2). Upon successive addition of the folding inducer, the CD signal at this wavelength begins to increase and finally shows saturation at high ligand/melittin molar ratios. On the other hand, spectral position of the zero crossing point is also varied during the titration displaying a gradual bathochromic shift (Fig. 1 and 2). Plotted against the inducer concentration, both of these numerical changes resulted in sigmoidal curves that are indicative of cooperative binding interactions (Fig. 4). Non-linear regression analysis of the data points yielded the \(K_d\) values and the Hill coefficients (\(h\)). Except for hemin, the binding parameters estimated from these different data sets are in close correlation but better fit was obtained by using the zero crossing shift (Table 3). Presumably, aqueous aggregation of the hemin molecules (\textit{vide infra}) may be responsible for the large deviation of the stability constants distorting the CD intensities above the equimolar composition. For a reference, CD titration data obtained with heparan sulfate (not shown) were also analyzed and compared to previously published results.\(^{34}\) The good agreement between the dissociation constants refers to the reliability of our method for the quantitative assessment of small molecule-melittin interactions (Table 3).

<table>
<thead>
<tr>
<th>Ligand</th>
<th>(K_d) ((\mu)M)</th>
<th>(h)</th>
<th>Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemin</td>
<td>65.5 (±1.3)</td>
<td>3.1 (±0.2)</td>
<td>Δastro</td>
</tr>
<tr>
<td>Biliverdin</td>
<td>44.4 (±1.3)</td>
<td>2.4 (±0.2)</td>
<td>Δastro</td>
</tr>
<tr>
<td>Bilirubin</td>
<td>21.0 (±0.5)</td>
<td>2.2 (±0.1)</td>
<td>Δastro</td>
</tr>
<tr>
<td>Bilirubin ditarurate</td>
<td>33.0 (±0.7)</td>
<td>3.4 (±0.2)</td>
<td>Δastro</td>
</tr>
<tr>
<td>Heparan sulfate</td>
<td>10.7 (±0.1)</td>
<td>4.1 (±0.1)</td>
<td>Δastro</td>
</tr>
<tr>
<td>Heparan sulfate</td>
<td>15.8 (±0.4)</td>
<td>3.3 (±0.2)</td>
<td>Δastro</td>
</tr>
<tr>
<td>Heparan sulfate</td>
<td>10.0&quot;</td>
<td>no data</td>
<td>ITC</td>
</tr>
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\(^{4}\) \(K_d\) from isothermal titration calorimetry (ITC).\(^{34}\)

In relation to heparan sulfate, hemin and biliverdin are the weakest melittin binders. The disaccharide unit of heparan sulfate bears ~2-3 negative charges (Fig. S8) which are sterically separated from each other enabling to establish multiple ionic contacts with the cationic residues of melittin. In contrast, the acidic functions of hemin and biliverdin are located on the same side of the rigid porphyrin scaffold that limits conformational adaptation necessary for optimizing the binding interactions (Scheme 1). From this point of view, the structure of bilirubin and its conjugate encode a much larger flexibility which is reflected in their tighter melittin association (Table 3). These molecules can adopt two, nonsuperimposable mirror-image conformations similar to a half-opened book (Fig. S3). By rotation about the central –CH= group, these isoecongener species interconvert rapidly in solution via a succession of non-planar intermediates.\(^{35,40}\) Therefore, bilirubin can accommodate to a variety of macromolecular targets having binding sites of diverse topography.\(^{25,26,41,42}\)

The Hill coefficients indicate the formation of multimeric pigment-melittin complexes where 2-3 ligand molecules are attached to a peptide chain (Table 3). According to earlier reports, bilirubin also binds to some peptides and polypeptides in dimeric and oligomeric forms.\(^{28,43}\)

![FIGURE 3](image3.png)

**FIGURE 3.** Concentration dependent changes of the regular helix content (H1) of 20 μM melittin titrated with hemin and various bile pigments.

**FIGURE 4.** CD titration data of melittin plotted against the ligand concentration in the sample solutions. \(\Delta c_j\): \(\Delta c_j\)residue values calculated at the zero cross-over point of the CD curves. \(\Delta c_j\): the shift in wavelength of the zero cross-over point measured during the titrations. Solid lines are the results of non-linear curve fitting analysis performed by using the “One site - specific binding with Hill slope” equation built in the Graph Pad Prism software (ver. 6.01, San Diego, California, USA). The \(K_d\) values and the Hill coefficients (\(h\)) are listed in Table 3. Data for bilirubin ditarurate are very similar to that of bilirubin and thus are not shown.
Melittin induced changes in the VIS absorption spectrum of porphyrin pigments

Since all ligand molecules used in this study exhibit characteristic absorption band above 300 nm, VIS absorbance curves of the free and peptide-loaded form of the pigments were recorded to gain further insight into the structural features of pigment-melittin complexes (Fig. 5). Upon addition of melittin, the Soret peak of hemin showed a moderate red shift, a broadening of envelope and hypochromism, i.e. an intensity loss.\(^{46-48}\) Contrary to this, the 379 nm band of biliverdin was hardly altered even in the two-fold molar excess of the peptide. It displays a very small hypochromism and a slight band broadening. These spectral differences are attributable to the distinct conformations of hemin and biliverdin: planarity of the hemin molecules allows them to attach close to each other along the peptide chain which favours excitation interaction between the \(\pi-\pi^*\) transitions of the porphyrin chromophores. Depending on the orientation factor of the adjacent rings, the exciton coupling may produce various combinations of red/blue shift and hypo/hyperchromism of the respective absorption band. The bathochromic shift obtained in our experiments suggests the head-to-tail like rather than face-to-face arrangement of the melittin-bound hemin units.\(^{46}\)

In contrast to hemin, steric repulsion between the lactamic carbonyl groups enforces biliverdin to adopt a non-planar, helical geometry.\(^{47}\) It seems from the very similar absorption curves displayed in Fig. 5 that this conformation prevents the spatial proximity of the peptide-bound biliverdin molecules, the prerequisite of intermolecular exciton interaction.

No induced CD signals were observed either with hemin or biliverdin (data not shown) suggesting the lack of stereospecific binding.

The porphyrinic bilirubin behaves as a molecular excitor.\(^{48}\) The \(\pi-\pi^*\) electric dipole transition moments of the intramolecularly H-bonded, unconjugated dipyrrinone moieties are excitonically coupled resulting in a broad absorption band consisting of a longer- and a shorter-wavelength component. The magnitudes as well as \(\lambda_{\text{max}}\) values of such sub-bands are strictly dependent on the relative spatial disposition of the interacting transition moments.\(^{33,35}\) In the case of bilirubin, their amplitudes are nearly equal whereas for the conjugated derivative the longer-wavelength component is more intense (Fig. 5). This indicates that in bilirubin ditalurate the angle of intersection of the two planar dipyrrinone moieties is larger (\(\Theta > 100^\circ\)) than in the unconjugated form (\(\Theta = 100^\circ\)). Bilirubin-melittin interaction provokes a strong hypochromism and band broadening at the red edge of the spectrum (Fig. 5). Intensity loss can also be seen in the absorption spectrum of the conjugate but the envelope broadens toward the opposite, shorter-wavelength direction with the development of two, partially resolved peak centered around \(\sim 406\) and \(\sim 447\) nm (Fig. 5). In both cases, the hypochromism is indicative of the self-association (oligomerization) of the pigment molecules bound to the melittin matrix. During the binding process, however, the ridge-like structure of free bilirubin (\(\Theta = 100^\circ\)) converts into a more extended shape (\(\Theta > 100^\circ\)) whereas opposite change occurs with the ditalurate derivative (\(\Theta < 100^\circ\)) resulting in a more compact conformation.\(^{48}\)

Effect of sterico- and urobilin on the secondary structure of melittin and CM15

These water soluble fecal pigments are the products of bacterial reduction of bilirubin in the gut (Scheme 1).\(^{47}\) Racemic samples were used which thus do not exhibit intrinsic CD activity. Strikingly, either with melittin or CM15, these compounds were unable to trigger conformational transitions witnessed by using hemin and bile pigments and caused only a moderate intensity reduction of the CD values and a small blue shift of the \(\lambda_{\text{CD}}\) (Fig. S9). In proportion to the ligand concentration applied, the largest relative intensity decrease was measured at the beginning of the titration. This implies some initial structure altering effect which, however, is defective in the sense of completing the sort of conformational redistribution observed previously. In line with this presumption, analysis of the CD spectra revealed the initial loss as well as the subsequent partial regaining of the regular helix content, very similar to that obtained with the other pigments (Fig. 3, Table 2). Besides these changes, some increment of the antiparallel but not the parallel \(\beta\)-sheet fraction was observed. Accordingly, these compounds alter modestly the conformation of melittin inducing neither \(\alpha\)-helix nor intermolecular \(\beta\)-sheet. Taking into account that both pigments bear a pair of propionate groups but lack intact, planar dipyrrinone units suggests the potential role of the latter ones in triggering the structural rearrangement seen with hemin and its derivatives. It should finally be noted that mixing of either melittin or CM15 into sterico- and urobilin solutions did not affect their VIS absorption spectra (data not shown).

FIGURE 5 Effect of the addition of melittin on the main VIS absorption band of hemin and bile pigments (10 mM potassium phosphate buffer at pH 7.3, 50 mM Na\(_2\)SO\(_4\), 25 °C).

Sensing of porphyrin binding to melittin via fluorescence quenching method

To obtain additional structural information on pigment-melittin complexes, fluorescence spectroscopy measurements were carried out. We exploited the intrinsic tryptophan (Trp) fluorescence which can sensitively report a binding event. The polarity decrease of the Trp microenvironment leads to a blue shift in the emission spectrum accompanied by intensity fluctuations as shown for peptides embedded into a hydrophobic protein or lipid interior. In other cases, the binding of a partner simply decreases the fluorescence intensity, which we observed for melittin in the presence of hemin-derived compounds. Upon addition of porphyrin agents inducing gross conformational rearrangements detected by CD spectroscopy (Table 2), a significant quenching of the Trp emission was measured (Fig. 6).
Contrary to this, the emission decreasing effect of stercolobilin was quite modest. For this compound, the Stern-Volmer (SV) plot gave rise to a straight line allowing to calculate the quenching constant ($K_Q$) that is equivalent to the equilibrium binding constant in this case. An apparent $K_Q$ of ~150 µM was obtained confirming the low binding affinity of the pigment. The SV plots constructed for bilirubin ditaurate and biliverdin show an upward curvature (Fig. 6B) suggesting the contribution of dynamic quenching besides the static mechanism. However, the linear region displayed between 5-25 µM quencher concentration could be used for $K_Q$ estimation yielding 17 and 27 µM for biliverdin and bilirubin ditaurate, respectively (Fig. 6B).

It is worth to note that at low ligand concentrations (<5 µM), the emission has declined more sharply than at higher pigment loadings (Fig. 6A) suggesting the formation of an initial complex serving as a scaffold for the binding of additional molecules. As such, it is in analogy with the sudden loss of the helical content of melittin measured at the beginning of the CD titrations (Fig. 3). Trp is a typical structure stabilizing side chain in proteins and peptides. In melittin it is the part of the fragment 14-20 shown to be more highly structured than the rest of the sequence.11 According to these mutually supportive CD and fluorescence results, porphyrin-melittin binding can be described by a two-state scenario. It seems that the very first step of the binding process is the accommodation of a pigment molecule close to the indole ring giving rise to a sharp decline in the emission (Fig. 6A) and the simultaneous unfolding the ordered segment (Fig. 3).

During the second phase, cooperative association of additional ligands takes place. Noticeably, the importance of the aromatic residue-porphyrin ring interaction in the binding of heme to amyloid-β peptides has recently been demonstrated, and proposed to be a key point in dismantle of the aggregates.12,13 Finally, it is worth to recall that both CD and fluorescence results refer to the involvement of stereo- and urobilin in the first stage only but they are unable to proceed with forming multimeric complexes what is thought to be required to the conformational conversion of the peptide.

**Conclusions**

Hemin and its metabolites are constantly produced in the human body and their levels show large fluctuations both under physiological and pathological conditions (e.g., neonatal hyperbilirubinemia, malaria, porphyrias, etc.). A plethora of clinical observations have indicated the beneficial effects of mildly elevated serum bilirubin level in reducing the prevalence of a variety of diseases such as diabetes, cardiovascular and even malignant disorders.51-53 Moreover, heme binding to amyloid-β peptides and the altered heme homeostasis in Alzheimer’s disease have also been demonstrated.54-56 There are several lacuna however, in understanding the chemical nature of heme-peptide and bile pigment-peptide interactions which instigated us to study how these compounds affect the secondary structure of intrinsically disordered model peptides. Our work supplies direct spectroscopic evidences on the complex, sequence dependent, significantly different structure modifying ability of hemin and its natural derivatives mediated by hydrophobic (Trp-porphyrin) and most likely ionic (Lys/Arg-proprionate) forces. Being present in the extra- and intravascular space, these porphyrin agents thus can alter the biological activity of melittin and related antimicrobial peptides. In a more broader sense, it could be assumed that these pigments modulate not only the structure of peptidic compounds with external origin, but also endogenous proteins/peptides participating in intrinsic disorder-based interactions. The porphyrin derivates studied here allow a promising opportunity to explore transient conformations of bioactive (poly)peptides to better understand their regulatory functions.
Acknowledgements

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Supporting information

Additional supporting information may be found in the online version of this article at the publisher’s website.

Description: Materials and Methods, Computational studies, Table S1, Figure S1-S10.

REFERENCES AND NOTES


