Induced CD of iron(II) clathrochelates: sensing of the structural and conformational alterations of serum albumins

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An ability of inherently achiral macrobicyclic metal complexes iron(II) clathrochelates to acquire an induced CD (ICD) output in the visible spectral range upon interaction with bovine serum albumin (BSA) was recently discovered. In the present work, the CD-reporting properties of iron(II) clathrochelates to proteins and the thermodynamic parameters of their binding to albumins are evaluated. It is shown that iron(II) clathrochelates functionalized by six ribbed carboxyphenylsulfide groups are able to discriminate between serum albumins of relative structure (here human and bovine albumins) by giving distinct ICD spectra. Besides, by the variation of the shape and intensity of CD bands, these cage metal complexes reflect the pH-triggered alterations of the tertiary structure of albumins. The constitutional isomerism (ortho-, meta- or para-isomers) of terminal carboxyphenylsulfide groups of iron(II) clathrochelates strongly affects both the character of their ICD output upon binding with proteins and the parameters of the formed guest–host associates. Using isothermal titration calorimetry, it was determined that cage metal complexes bearing meta- and ortho-isomers of carboxyphenylsulfide groups possess higher association constants (K_a ≈ 2 × 10^5 M^{-1}) and clathrochelate-to-BSA binding ratios (n = 2) than the para-isomer (K_a ≈ 5 × 10^3 M^{-1}, n = 1). The iron(II) clathrochelates are suggested to be potential molecular three-dimensional scaffolds for the design of CD-sensitive reporters able to recognize specific elements of protein surfaces.

1. Introduction

Circular dichroism (CD) spectroscopy is a powerful spectral technique for studies of protein–ligand interactions. Biological macromolecules exhibiting inherent chirality are able to induce asymmetry of achiral organic and coordination compounds and, therefore, may cause an appearance of a signal in their CD spectra. These induced CD (ICD) bands are very sensitive to the arrangement of the binding sites of hosting biomolecules. As a result, the chirality of small molecules gained upon their binding to proteins may reflect both the structural alterations and the conformation transitions of proteins (to be observed via changes in the ICD spectra). In the present work we demonstrate that hexafunctionalized iron(II) clathrochelates (that are inherently achiral) give a CD output in the visible range upon their supramolecular binding to serum albumins (HSA and BSA). Since the reported clathrochelates are able to reflect alterations of albumin structure and conformational transitions by their ICD outputs, we suggest that these cage metal complexes are prospective three-dimensional molecular scaffolds for the design of probes giving chiroptical output for use in studies of proteins.
binding to proteins may reflect both the structural alterations and the conformation transitions of proteins (to be observed via changes in the ICD spectra). Thus molecules giving an ICD response upon interaction with proteins are studied as potential reporters with the ability to discriminate structural variants of proteins connected with various diseases via reflection of their structural differences.

Serum albumin is a main transport protein of blood plasma, and its certain structural variations may act as markers of various pathological processes, such as albumin glycation in diabetes,6,7 oxidation in liver disease,6,8 N-terminal modifications in ischemia/reperfusion,6 and allostery modifications by tumor metabolites in cancer.9

Various organic compounds and metal complexes have been reported to be ICD probes due to their binding to proteins and sensitivity to their structural alterations.10,11 In particular, an adenosine derivative that senses the hydrophobic cavity of bovine serum albumin (BSA) was reported,12 as well as a bilirubin molecule that is able to signal the conformational alternations of human serum albumin (HSA) caused by a variation of pH.13 Diazepam and dansylglycine were reported to be site-specific ligands for albumin binding site 2 (vide infra).14 cis-Parinaric and retinoic acids were used as ligands for the hydrophobic interior cavity of β-lactoglobulin.2,15 A derivative of cobalt porphyrin was proposed to serve as a chiral probe for a few globular proteins, such as albumin, lysozyme and insulin.16 A well-known organic dye, Congo Red, was used in ICD studies as an indicator of the alterations in the structure of albumin regarding its binding sites and the formation of amyloid aggregates.17

A wide range of cage complexes with an encapsulated metal ion, known as clathrochelates, have been prepared and characterized to date.18–20 Metal complexes of this type, which are formed by rigid quasiaromatic polyazomethine macropoly cyclic ligands, are metal-centered compounds because their caging (encapsulating) ligands do not exist in the metal-free form due to thermodynamic reasons.19 Iron(II) clathrochelates exhibit unique chemical and biological activities, and allow easy chemical modification of their macropoly cyclic frameworks as molecular scaffolds. Data on their biological activity as submicromolecular inhibitors in the transcription systems of T7 RNA and Taq DNA polymerases,21–25 and as antifibrillogenic agents,26–28 together with axially coordinated phthalo cyanines,29 were recently highlighted. The ability of these cage metal complexes to form supramolecular assemblies with serum albumins has been established using steady state and time-resolved fluorescent spectroscopy methods.30–32

Iron(II) clathrochelates with mono- and dicarboxyphenylsulfide substitution that are inherently CD-silent have been recently reported to give a strong CD output in the visible spectral range upon their binding to BSA.31 The induction of chirality has been explained by selective fixation of the metal centered framework of a clathrochelate in one of the two (Δ or Λ) trigonally antiprismatic distorted conformations upon host–guest assembly with BSA. It was shown that the intensity of the ICD signal is affected by the constitutional isomerism of the macrobicyclic reporter molecule (in particular, by ortho-, meta- or para-positions of its terminal carboxyl groups).32

The development of supramolecular compounds, particularly metal complexes, for protein surface recognition, and application as potential sensors, affinity tags, tools for protein immobilization etc., is a prospective research and practical area.33 The specificity of recognition is supposed to be achieved via non-covalent interactions between large areas of the host protein surface and multiple binding groups of a guest compound.34 In this light, the discovered ICD sensitivity of clathrochelates to albumins generates an interest in further studies of these macrocyclic metal complexes as potential reporters for proteins.

In the present work, we explore the ability of iron(II) clathrochelates bearing six carboxyphenylsulfide ribbed groups to sense and reflect the structural alterations and conformational changes of tertiary structure of proteins using ICD. For this study, “relative structure” proteins, i.e. human and bovine serum albumins, were used; their conformation transitions were triggered by pH variation. All the constitutional isomers of a hexacarboxyphenylsulfide iron(II) clathrochelate (i.e. ortho-, meta-, or para-positions of its terminal carboxyl group, Scheme 1) were tested as the reporters. The characterization of clathrochelate-to-albumin association was monitored by CD and complemented by protein fluorescence quenching studies. Isothermal titration calorimetry (ITC) was used to determine the thermodynamics of host–guest assembly of these cage metal complexes with BSA. Through the obtained data, the perspective of application of iron(II) clathrochelates as chirooptical probes for the sensing of proteins is explored.

Scheme 1 Chemical drawings of the hexacarboxysulfide iron(II) clathrochelates under study.
2. Materials and methods

2.1. Materials

The clathrochelates were prepared as described elsewhere.\textsuperscript{35} BSA and HSA were obtained commercially (fraction V from Sigma-Aldrich\textsuperscript{36} and fraction V protease free from Fisher Bioreagents\textsuperscript{37}), respectively. 0.05 M Tris–HCl aqueous buffer (pH 7.9), 0.05 M aqueous phosphate buffers (pH 5 and pH 6), and 0.05 M aqueous acetate buffer (pH 3.7) were used as solvents.

2.2. Preparation of the solutions for protein fluorescence quenching studies

Aqueous solutions of BSA and HSA (concentration of 0.2 mg ml\textsuperscript{-1} – 3 \mu M) were prepared by dissolution of their weighed amounts in corresponding buffer. 2 mM stock DMSO solutions of the clathrochelates were prepared, and an aliquot was added to Tris–HCl buffer solution of a chosen protein. Since the amount of added DMSO solution was rather small (up to 2.5% of the total volume), the protein’s concentration remained practically unchanged (0.2 mg ml\textsuperscript{-1}), while that of the clathrochelate changed from 0 to 15 \mu M.

2.3. Fluorescence assay

To study the quenching of albumin fluorescence upon the binding of iron(II) clathrochelates, the fluorescence spectra of the protein itself and after the addition of a DMSO solution of clathrochelate were recorded. Albumin fluorescence was excited at 280 and 295 nm, and the signal was recorded at \( \lambda_{\text{max}} \) of corresponding emission. Fluorescence spectra were measured at 5 nm width of the excitation and emission slits at room temperature on a Cary Eclipse spectrophotometer (Varian) in a 1 cm quartz cuvette. To avoid possible errors caused by differences in the media used for the fluorescent and ITC experiments (Tris–HCl aqueous buffer and sodium acetate aqueous solution, respectively), fluorescence quenching measurements were also performed in sodium acetate aqueous solution. The observed differences were found to be insignificant.

An increase in the clathrochelate concentration led to the simultaneous increase in the optical density at maxima of BSA excitation and/or emission, thus causing a decrease of observed fluorescence intensity because of the so-called “inner filter effect” (IFE), as well as due to the reabsorption process.\textsuperscript{16,37} These effects should be taken into account for the correct interpretation of the experimental data.\textsuperscript{37–39} The correction factor for these effects can be calculated using eqn (1):\textsuperscript{39}

\[
I_{\text{cor}} = I_{\text{obs}} \times 10^{[(D_{\text{ex}}+D_{\text{em}})/2]}
\]  

where \( I_{\text{cor}} \) and \( I_{\text{obs}} \) are the corrected and observed fluorescence intensities, respectively; \( D_{\text{ex}} \) and \( D_{\text{em}} \) are the total optical densities of a solution at the wavelengths of the fluorescence excitation and emission, respectively. All the experimental fluorescent data were corrected using eqn (1).

2.4. UV-vis spectroscopy

The UV-vis absorbance spectra of BSA-clathrochelate assemblies were recorded at room temperature on a Specord M-40 spectrophotometer (Carl Zeiss, Germany). An aliquot of 2 mM stock DMSO solution of the cage complex was added to 0.05 M Tris–HCl aqueous buffer solution, pH 7.9. The concentration of clathrochelate was varied from 0 to 15 \mu M.

2.5. ICD spectroscopy

The CD spectra were recorded on a Jasco J-1500 spectropolarimeter at room temperature in the 300–600 nm range, in 0.1 nm steps with a scan of 3001 points; three scans were averaged for each of the ICD spectra. The data expressed as ellipticity (mdeg) were obtained in mdeg units directly from the instrument. Tris–HCl aqueous buffer (pH 7.9), phosphate aqueous buffers (pH 5 and pH 6) and acetate aqueous buffer (pH 3.7) were used for the preparation of stock solutions of the proteins as well as for working solutions with an albumin-to-clathrochelate 2:1 molar ratio (\( \epsilon_{\text{albumin}} = 8 \times 10^{-5} \text{ mol l}^{-1} \), \( \epsilon_{\text{clt}} = 4 \times 10^{-5} \text{ mol l}^{-1} \)). It should be mentioned that the observed fluorescence quenching is rather low at such a molar ratio. But due to the peculiarities of the experiment (higher sensitivity of the fluorescence method compared to the CD method), absolute concentrations used in CD studies were much higher as compared to the ones used in fluorescence studies (albumin concentration equals \( 8 \times 10^{-5} \text{ mol l}^{-1} \) and \( 3 \times 10^{-6} \text{ mol l}^{-1} \) respectively). Thus, a significantly higher percentage of clathrochelate molecules is bound to albumins at a 2:1 molar ratio for the conditions of the CD experiment, and induced CD was clearly observed.

The clathrochelate binding site was evaluated by displacement of clathrochelate upon addition of site markers: (i) warfarin,\textsuperscript{3,41} and (ii) ibuprofen.\textsuperscript{13} A 50-fold excess (\( 2 \times 10^{-3} \) M) of warfarin or ibuprofen was added to a mixture of protein (\( \epsilon_{\text{albumin}} = 8 \times 10^{-5} \text{ mol l}^{-1} \)) and clathrochelate (\( \epsilon_{\text{clt}} = 4 \times 10^{-5} \text{ mol l}^{-1} \)).

2.6. ITC experiments

A 5 mM solution of clathrochelate in 0.15 M aqueous sodium acetate was added to a 0.5 mM solution of BSA in 0.15 M aqueous sodium acetate thus obtaining a working solution with clathrochelate-to-protein molar ratios from 0 to 3. ITC experiments were carried out on a Nano ITC calorimeter (TA Instruments), equipped with a standard 1.0 ml cell (24K Gold). The stock solution of clathrochelate was added to the cell using a 250 \mu l syringe. The calorimetric experiments were operated by Nano ITC Run v. 2.2.3 software. All the experimental data were evaluated using the NanoAnalyze v. 2.4.1 program package and an independent model was used for their interpretation. In each case, control experiments were performed and the enthalpies of dilution of the components were subtracted from the data for their supramolecular assembly.

3. Results and discussion

3.1. Supramolecular binding of the hexacarboxyl-terminated iron(II) clathrochelates to BSA and HSA

BSA and HSA are heart-shaped proteins, the crystal structures of which are very similar.\textsuperscript{31} The molecule of HSA consists of three helical domains (I, II and III), each comprising two
subdomains (A and B; Fig. 1). The drug binding site 1 (Sudlow’s I) is a binding pocket within a core in the subdomain IIA. Drug binding site 2 (Sudlow’s II) is located in subdomain IIIA.42–44 Despite the similarity of BSA and HSA molecular structures, the proteins possess certain regions with substantial differences in amino acid sequences;45 the BSA molecule was reported to be less conformationally flexible than HSA.46

The type of serum albumin, BSA versus HSA, was shown to affect the association of various drugs/toxins, such as warfarin, ochratoxin A and phenylbutazone,46,47 with site 1, i.e. the stability constants and competing capacity. The structural distinctions between the binding sites of BSA and HSA are suggested to cause variations of the ICD output of clathrochelates upon their association.

3.1.1. Protein fluorescence quenching studies. The effect of clathrochelates on BSA and HSA fluorescence was studied to characterize the BSA/HSA–clathrochelate host–guest assemblies. The Stern–Volmer plots of protein fluorescence quenching intensities versus the clathrochelate-to-protein molar ratio are shown in Fig. 2.

The quenching of proteins’ intrinsic fluorescence observed in the presence of clathrochelate compounds (Fig. 2 and Fig. S1a–i, ESI†) evidenced the formation of their host–guest supramolecular associates. The fluorescence intensity was much more affected by clathrochelate addition in the case of BSA (6.9–7.5-fold quenching, Fig. 2a) than in that of HSA (2–2.4-fold quenching, Fig. 2c). Shifts of the corresponding emission maxima (up to 20 nm to the short-wavelength range for BSA, and up to 9 nm to the long-wavelength range for HSA) were observed (Fig. S1 and S2, ESI†).

Tryptophan (Trp) residues are the brightest fluorophores in albumin, and characteristics of Trp emission (the emission intensity and maximum) are sensitive to the surrounding medium.49,50 Apart from Trp, also tyrosine (Tyr) residues contribute to the intrinsic fluorescence of proteins;51 BSA and HSA contain twenty and eighteen Tyr residues per protein macromolecule, respectively.2,41 However, in comparison to Trp, the fluorescence of Tyr residues is less intense and independent of the local environment.

A weak effect of fluorescence quenching and a small shift of the emission maximum observed upon iron(ii) clathrochelate binding to HSA might be explained by the low accessibility of the single Trp-214 residue that is hidden within a protein globule (subdomain IIA). On the other hand, the BSA macromolecule contains two Trp residues: surface-located Trp-134 (subdomain IB), and pocket-buried Trp-213 (subdomain IIA, Fig. 1).50 The efficiency of BSA fluorescence quenching by
hexacarboxyphenylsulfide iron(II) clathrochelates is similar to that of the ortho- and para-isomers of di-substituted macrobicyclic analogs, but is smaller than that caused by binding of the di-meta-substituted cage complex.31,50

The Stern–Volmer plots for BSA–clathrochelate assemblies (Fig. 2a and b) suggest that ortho- and meta-isomers (complexes 1 and 2, respectively) possess similar fluorescence quenching patterns (with a bend of the corresponding curves at a clathrochelate-to-BSA 3:2 molar ratio), while for para-substituted analog 3 the increase is more gradual (with a bend at a molar ratio of approximately 3:1). For HSA, the quenching patterns for iron(II) clathrochelate isomers are of similar shape (Fig. 2c and Fig. S3, ESI†) with the bend values close to a 1:1 molar ratio. The observed behavior could be explained by “multiple quenching” of two Trp residues in BSA, and “single” quenching of one Trp residue in HSA.

To estimate the effect of the clathrochelate binding on the fluorescence of both Trp and Tyr amino acids, we performed quenching experiments (Fig. 2a–c and Fig. S3, ESI†) upon excitation at 280 nm, where both Trp and Tyr are excited, and at 295 nm where only Trp residues are excited.49 The data clearly show similar quenching patterns upon both used excitation wavelengths for clathrochelate assemblies with BSA or HSA. This indicates that the binding of clathrochelates to albumins quenches mainly Trp emission, and only slightly that of Tyr residues.

Thus we suggest that the quenching of albumin fluorescence is caused by the binding of clathrochelates near Trp-213 and Trp-134 of BSA, or Trp-214 of HSA; the binding most probably occurs at site 1 located in protein subdomain II A.

3.1.3. ICD sensing of the “relative structure” of albumins using clathrochelates. The binding of the hexacarboxyphenylsulfide iron(II) clathrochelates to BSA and HSA was studied by CD spectroscopy. These inherently optically inactive cage complexes (Fig. 3a) were found to gain optical activity upon their binding to both given serum albumins, that resulted in an appearance of intense ICD bands in the 350–600 nm range (Fig. 3a and b). These bands correspond to macrobicyclic UV-vis absorption bands (Fig. S4, ESI†), and are assigned to the metal-to-ligand Fed → Lπ* charge transfer transitions in the quasiaromatic macrobicyclic frameworks.

The binding to albumins induces CD bands that are distinct for HSA and BSA, in terms of intensity in the case of clathrochelate meta-isomer 2, and also in the shape for ortho- and para-substituted analogs 1 and 3, respectively (Fig. 3a and b). meta-Isomer 2 shows a positive-sign ICD band with a maximum at 446 nm and higher intensity of the CD signal. The ICD band of para-substituted clathrochelate 3 bound to BSA is less intense, but has a similar shape to that of meta-isomer 2, but with a positive-sign band at 450 nm. In the presence of HSA, the spectrum of para-isomer 3 changes its shape exhibiting a “double peak” negative-sign band with the main maximum near 430 nm. The ICD spectrum of ortho-substituted clathrochelate 1 in the presence of BSA is of low intensity and located in the negative range; with HSA it is much more intense and contains a negative band with a maximum at 447 nm.

The intensities of ICD bands of hexa-carboxyl-terminated iron(II) clathrochelates in the presence of BSA are substantially lower than those of their dicarboxyl-terminated analogs reported earlier.31 Moreover, for all the di- and mono-substituted carboxyphenylsulfide cage metal complexes the shape of ICD bands in the presence of BSA was similar, with an intense negative maximum at 450 nm.31

Hexa-carboxyphenylsulfide iron(II) clathrochelates reveal the ICD sensitivity to related serum albumins, and are capable of discriminating between BSA and HSA hosts, giving different ICD spectra. This spectral discrimination is affected by the constitutional isomerism of the macrobicyclic complexes: it is more pronounced for ortho- and para-substituted cage complexes, 1 and 3, respectively, and substantially less manifested in the case of meta-isomer 2.

The differences in the sequences and thus structures of albumins lead to the alterations of the arrangement of the drug binding sites of BSA and HSA. Hence, variations of ICD outputs of BSA- and HSA-bound hexacarboxy-terminated iron(II) clathrochelates are suggested to be caused by distinct arrangements of BSA- and HSA–clathrochelate assemblies that come from the structural distinctions of BSA and HSA binding sites. The fluorescent data obtained here clearly indicate that the clathrochelate–protein binding is associated with albumin’s drug site 1.
3.2. Sensing of albumin conformational transitions by ICD of clathrochelates

To estimate the effect of albumin tertiary structure alterations on the ICD response given by iron(II) clathrochelates, the protein conformation changes were triggered by the variation of pH (Table 1). The acidic F-form mainly concerns unfolding of domain III, while in the basic B-form an alteration of the conformation of domain I occurs. In the neutral N-form, an interaction between site 1 and site 2 is suggested, while in the B-form, the distance between the corresponding subdomains increases and this interaction disappears.

Changes in ICD spectra of the protein–clathrochelate assemblies with pH variations were studied for both albumins. The corresponding fluorescence quenching experiments were performed only for BSA since its intrinsic emission is more affected by assembling with clathrochelate (see Section 3.1.2).

3.2.1. Changes of albumin fluorescence quenching by clathrochelates upon pH variation. A strong influence of media pH on the quenching of BSA fluorescence by the iron(II) clathrochelates was revealed. The most significant quenching effect (15 to 37-fold) was observed at pH 6, where the N-form predominates, while at pH 3.7 (N→F-transition), the effect was negligible (with only up to 1.8-fold quenching). In a slightly basic medium, i.e. at pH 7.9 (N→B transition), moderate quenching (about 7-fold) occurred (Fig. 4, 5 and Fig. S5, ESI†).

Quenching of BSA intrinsic fluorescence depends on the constitutional isomerism of the iron(II) clathrochelate terminal carboxy groups (Fig. 4, 5 and Fig. S6, ESI†). Maximal variation of quenching versus pH was observed for para-isomer 3 (from 2- to 37-fold, upon the F→N transition and N-forms of BSA, respectively, [Fig. S5, ESI†]). Also, constitutional isomerism affected the quenching patterns of these cage metal complexes (Fig. S5, ESI†).

It has to be mentioned that the pH-dependent experiments were performed in a range of pH that includes the albumin isoelectric point and pK of iron(II) clathrochelates. Both proteins have the same isoelectric point with pI about 4.7 (Fig. 2). pK values for the hexacarboxyphenylsulfide cage metal complexes were estimated by potentiometric titrations, and are in the range 5.5–6.3, depending on constitutional isomerism (data not presented).

Processes of dissociation of terminal carboxy groups of clathrochelates are considered to affect the albumin–clathrochelate non-covalent interactions, leading to strong quenching of Trp fluorescence at pH 6, as compared to pH 3.7. This also indicates the proximity of the guest clathrochelate to a Trp residue in the formed assembly.

Moreover, the small effect of iron(II) clathrochelates on BSA fluorescence in acidic media could be due to the conformation transition (spatial separation) of the domains III and I (Fig. 2). In the N-form, the bonding of the clathrochelate molecule to site 1 (domain I) could be additionally stabilized by spatial interactions with proximal domain III, thus separation of this domain upon the decrease of pH promotes re-arrangement or dissociation of the clathrochelate–protein associate.

3.2.2. Changes of ICD spectra of clathrochelates with pH variation. ICD spectra were recorded for iron(II) clathrochelates in assemblies with BSA and HSA, in pH range 4–9 (Fig. 6a–c and Fig. S5, ESI†).

The conformation alterations of a host protein molecule triggered by pH variation cause changes of the ICD spectra of the guest clathrochelate; the changes depend on the origin of serum albumin (BSA or HSA).

In associates with HSA, iron(II) clathrochelates change the shape and intensity of their ICD-bands upon pH variation, indicating rearrangements of protein–clathrochelate assemblies. The changes were the most pronounced for meta-isomer 2 (Fig. 6).

Upon pH-triggered conformational modifications of BSA, the intensity and maxima of ICD-bands of meta- and para-clathrochelates were changed, while in the case of ortho-clathrochelate also the shape of the bands was altered (Fig. 6a, b and Fig. S5, ESI†).

In acidic medium (pH 3.7) all iron(II) clathrochelate isomers showed low intensity ICD spectra (up to ΔCD = 2.2 mdeg for
change the shape and intensity of the CD spectra with a variation B-form.

Fig. 6

Fig. 6 pH-Dependent ICD spectra of 

ortho

isomer 1 bound to BSA (a), meta

isomer 2 bound to BSA (b), and meta

isomer 2 bound to HSA (c).

Fig. 6 pH-Dependent ICD spectra of ortho

isomer 1 bound to BSA; together with a very low albumin fluorescence quenching degree (Fig. 4 and 5) this may evidence clathrochelate–albumin assembly dissociation.

In the pH range 5–9 the bands of ICD spectra of clathrochelate–albumin assemblies are the most intense (Fig. 6a–c and Fig. S5, ESI†) and the ability to discriminate between BSA and HSA hosts via the difference of ICD signals is the most pronounced. The variations of the ICD spectra probably reflect rearrangement of the albumin–clathrochelate assembly that may occur upon passing through clathrochelate pK points (at pH 5.5–6.3), and also at more basic pH (7–8) may be caused by change of protein conformation from the N- towards the B-form.

Thus iron(II) clathrochelates in assemblies with albumin change the shape and intensity of the CD spectra with a variation of the pH of the medium, and consequently with protein tertiary structure transitions. This points to the high sensitivity of clathrochelate CD output to the arrangement of the structure of their associates with host proteins. Thus, it allows suggesting these cage metal complexes as potential tools for monitoring of protein conformation changes.

3.3. Competitive binding studies using ibuprofen and warfarin as site specific binders

To evaluate the preferable iron(II) clathrochelate binding site in serum albumins, we carried out experiments of clathrochelate displacement by specific binders of albumin sites 1 and 2, warfarin and ibuprofen, respectively.3,13,40 These site-specific binders show high affinity to albumin, with measured binding constants of about $2.5 \times 10^5 \text{ M}^{-1}$ for warfarin,60 and $2 \times 10^6 \text{ M}^{-1}$ for ibuprofen.61 Upon binding to serum albumin, neither warfarin nor ibuprofen exhibit any ICD spectra in the visible spectral range.7 Therefore, evaluation of the iron(II) clathrochelate preference to certain binding sites is performed by monitoring the changes in the ICD spectra of clathrochelate upon the addition of each of these site-specific binders.

For the competitive binding studies on serum albumin with site-specific ligands, the para-substituted constitutional isomer 3 was used since it shows the binding ratio 1:1 to albumin (see Section 3.4), and certain specificity to only one of the binding sites could be expected. Generally, an addition of low concentrations of site-specific ligands to the clathrochelate–protein assembly did not influence the clathrochelate ICD output. Only the presence of a high excess of the site-specific binders (from 30-fold excess) resulted in changes of the ICD-band intensity (Fig. 7).

Despite the affinity of the site-specific binders being a few orders of magnitude higher than that determined for clathrochelate 3 ($2.5 \times 10^5 \text{ M}^{-1}$ for warfarin, $2 \times 10^6 \text{ M}^{-1}$ for ibuprofen, over $5.3 \times 10^3 \text{ M}^{-1}$ for 3, Table 2), only in the case of a 50-fold excess of ibuprofen over the clathrochelate 3-HSA assembly was a noticeable decrease of the ICD band intensity, up to 2.5-fold, observed (Fig. 7). This indicates that warfarin and ibuprofen displace the iron(II) clathrochelate from its associate with albumin only insignificantly, and thus the competitive binding studies do not provide specific information about the binding site preference.

3.4. Calorimetric study of the supramolecular clathrochelate-to-BSA association

Thermodynamic parameters of supramolecular BSA-to-clathrochelate assemblies were determined using the ITC technique (Fig. 8); the obtained values are summarized in Table 2. The difference between the binding modes of ortho- and meta-isomers and the para-isomer is clearly seen from the shapes of the plots of the binding heat effect versus the clathrochelate-to-BSA molar ratio (Fig. 8), as well as from the values of the corresponding thermodynamic parameters (Table 2). Indeed, $\Delta G$ values for meta- and ortho-isomers of clathrochelate complexes are similar, while that for the para-substituted cage complex is smaller. According to this, the binding constant values $K_D$ are almost the same for the ortho- and meta-isomers 1 and 2.
Initial ICD spectra of the para-isomer 3–HSA assembly and those after addition of a 50-fold excess of albumin site-specific drugs ibuprofen (a) and warfarin (b).

Fig. 8 The experimental (depicted in the points) and fitted (depicted in the solid lines) plots of the binding heat effect versus the BSA-to-clathrochelate molar ratio for the ortho-, meta- and para-substituted hexacarboxyphenyl-terminated iron(II) clathrochelates 1–3 as guests.

Table 2 Thermodynamic parameters of the supramolecular assembly between BSA and iron(II) clathrochelates

| Compound | $\Delta H^a$ (kJ mol$^{-1}$) | $\Delta S^a$ (J mol$^{-1}$ K$^{-1}$) | $\Delta G^a$ (kJ mol$^{-1}$) | $K_a^b$ (10$^3$ mol$^{-1}$) | n $^c$ |
|----------|-----------------|-----------------|-----------------|-----------------|-----------------|
| 1        | -9.1            | 53.2            | -24.9           | 24              | 1.95            |
| 2        | -17.2           | 27.2            | -25.3           | 28              | 1.9             |
| 3        | -23.2           | -6.5            | -19.3           | 5.3             | 0.9             |

$^a$ $\Delta H$, $\Delta S$, and $\Delta G$ are the changes of enthalpy, entropy and Gibbs free energy, respectively, upon the binding of clathrochelates to BSA. $^b$ $K_a$ is a clathrochelate-to-BSA association constant. $^c$ n is the stoichiometry of the interaction, thus also the calculated number of the clathrochelates molecules per BSA macromolecule as a host. The area under each peak was integrated and fit to an independent binding model to determine the binding affinity ($K_a$), enthalpy ($\Delta H$) and stoichiometry ($n$). Since the temperature ($T$) is held constant throughout, the free energy ($\Delta G$) of the binding reaction can be determined by: $\Delta G = -RT\ln K_a$. ITC directly measures $\Delta H$, so the change in entropy ($\Delta S$) can be determined by: $\Delta S = (\Delta H - \Delta G)/T$. The independent model assumes $n$-independent binding sites and close thermodynamic parameters of each site. The binding ratio 2:1 was obtained for ortho- and meta-isomers, thus both isomers bind to BSA with similar $K_a$.

(2.4 × 10$^4$ and 2.8 × 10$^4$ M$^{-1}$, respectively), while for para-functionalized compound 3 this value is 5 times lower (5.3 × 10$^3$ M$^{-1}$). These data are in accordance with the results of the fluorescent quenching studies (Fig. 2a and b), where ortho- and meta-isomers 1 and 2 quench the BSA emission with the same intensity, while para-isomer 3 demonstrates less intense quenching. Thus, the supramolecular binding of the ortho- and meta-substituted iron(II) clathrochelates to BSA is suggested to be more favorable as compared to their para-isomer 3.

The obtained results can be also explained by the peculiarities of the spatial arrangements of the terminal functionalizing carboxyl groups of the clathrochelate guests. In the case of para-isomer 3 of the hexacarboxyphenylsulfide clathrochelate, an external “shell” of the molecule is formed by the “sticking out” terminal polar carboxyl groups, which hampers the ability of this molecule to enter the hydrophobic interactions. In the case of the ortho- and meta-substituted clathrochelate molecules 1 and 2, their terminal carboxyl groups are positioned closer to the cage framework, thus making their inclusion into the hydrophobic cavities of protein macromolecules easier and their supramolecular binding is energetically more favorable. As for the enthalpy effect, we suppose that the para-isomer 3 has a higher local dipole moment of ribbed substituents as compared to those for meta- and ortho-substituted analogs 1 and 2, thus causing an increase in the energy of dipole–dipole interactions.

The number of the iron(II) clathrochelate molecules per BSA macromolecule (n) also depends on their constitutional isomerism. This ratio is estimated as $n = 2$ for the ortho- and meta-isomers 1 and 2 and $n = 1$ for the para-substituted analog 3. This suggests that one additional site on the BSA globule is...
available for the “stable binding” of the ortho- and meta-substituted clathrochelates. This could be explained by the higher number of geometric configurations of the clathrochelate molecule (i.e. the variations of the shape) available due to the rotation of “inherently asymmetric” ortho- and meta-functionalized ribbed substituents compared to the “inherently symmetric” para-functionalized substituents. Higher structural variability allows more flexible tuning of the iron(ii) clathrochelate geometry to the shape of the binding site and promotes more precise adsorption of carboxyphenylsulfide groups to the complimentary binding groups of the protein. The independent binding model suggests that two molecules of each clathrochelate isomer bind to albumin with similar $K_a$. Hence we could consider two “thermodynamically similar” sites on the BSA molecule where ortho- and meta-isomers are able to bind.

Among studied clathrochelates, the para-isomer is the most sensitive to albumin binding site structure distinctions (discriminating HSA and BSA) at neutral and slightly basic pH. This isomer strongly reflects changes that the albumin molecule undergoes when passing from acidic to slightly basic pH, and the quenching of protein fluorescence (i.e. effect on the Trp residue) at slightly basic media is most pronounced for this isomer. The clathrochelate-to-albumin binding ratio of 1:1 (established by ITC) and strong effect on protein fluorescence allow suggesting that the exact binding site for this isomer is Sudlow site 1 of the albumin molecule. Thus, despite the lower binding constant values compared with ortho- and meta-clathrochelate isomers, the para-isomer is considered most suitable as a structure sensitive CD-reporting molecule for albumins.

4. Conclusions
In summary, cage metal complexes iron(ii) clathrochelates have shown potency as molecular three-dimensional scaffolds for the design of CD-sensitive reporters able to recognize specific elements of protein surfaces. These important unconventional properties may lead to their possible future use in biochemistry- and medicine-related fields.

Here we show that hexa-carboxyphenylsulfide iron(ii) clathrochelates discriminate between proteins of similar structure, in this case HSA and BSA, giving distinct ICD spectra. These cage metal complexes sense the variation of the arrangement of this case HSA and BSA, giving distinct ICD spectra. These cage metal complexes discriminate between proteins of similar structure, in the binding of carboxyphenyl substituents to the complimentary binding groups of the protein. The independent binding model suggests that two molecules of each clathrochelate isomer bind to albumin with similar $K_a$. Hence we could consider two “thermodynamically similar” sites on the BSA molecule where ortho- and meta-isomers are able to bind.

Among studied clathrochelates, the para-isomer is the most sensitive to albumin binding site structure distinctions (discriminating HSA and BSA) at neutral and slightly basic pH. This isomer strongly reflects changes that the albumin molecule undergoes when passing from acidic to slightly basic pH, and the quenching of protein fluorescence (i.e. effect on the Trp residue) at slightly basic media is most pronounced for this isomer. The clathrochelate-to-albumin binding ratio of 1:1 (established by ITC) and strong effect on protein fluorescence allow suggesting that the exact binding site for this isomer is Sudlow site 1 of the albumin molecule. Thus, despite the lower binding constant values compared with ortho- and meta-clathrochelate isomers, the para-isomer is considered most suitable as a structure sensitive CD-reporting molecule for albumins.

as the most appropriate structure-sensitive CD-reporter for albumin studies and discrimination.

The thermodynamic parameters of the clathrochelate-to-BSA assemblies were estimated by ITC, which showed that constitutional isomerism determines the characteristics of such assemblies. Thus meta- and ortho-isomers have higher binding constants ($K_a \sim 2 \times 10^4$ M$^{-1}$) and clathrochelate-to-albumin binding ratios ($n = 2$) than the para-isomer ($K_a \sim 5 \times 10^3$ M$^{-1}$, $n = 1$).

Conflicts of interest
There are no conflicts to declare.

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