DOI: 10.1021/bi500553f
Temperature Dependence of Backbone Dynamics in Human Ileal Bile Acid-Binding Protein. Implications for the Mechanism of Ligand Binding

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Funding Sources

The work was supported by the Hungarian Research Fund OTKA via K 109035 (O.T.).
ABBREVIATIONS

CPMG, Carr-Purcell-Meiboom-Gill; CRBP, cellular retinol binding protein; FABP, fatty acid binding protein; GCA, glycocholic acid; GCDA, glycochenodeoxycholic acid; ITC, isothermal titration calorimetry; I-BABP, ileal bile acid-binding protein; NMR, nuclear magnetic resonance

KEYWORDS: lipid-binding proteins, enterohepatic circulation, bile salts, conformational exchange, protein dynamics, binding cooperativity, site-selectivity, NMR spectroscopy
ABSTRACT

Human ileal bile acid-binding protein (I-BABP), a member of the family of intracellular lipid binding proteins plays a key role in the cellular trafficking and metabolic regulation of bile salts. The protein has two internal and according to a recent study an additional superficial binding site and binds di- and trihydroxy bile salts with positive cooperativity and a high degree of site-selectivity. Previously, in the apo form, we have identified an extensive network of conformational fluctuations on the ms time scale, which cease upon ligation. Additionally, ligand binding at room temperature was found to be accompanied by a slight rigidification of ps-ns backbone flexibility. In the current study, temperature-dependent $^{15}$N NMR spin relaxation measurements were used to gain more insight into the role of dynamics in human I-BABP – bile salt recognition. According to our analysis, residues sensing a conformational exchange in the apo state can be grouped into two clusters with slightly different exchange rates. The entropy-enthalpy compensation observed for both clusters suggests a disorder-order transition between a ground and a sparsely populated higher energy state in the absence of ligands. Analysis of the faster, ps-ns motion of $^{15}$N-$^{1}$H bond vectors indicates an unusual nonlinear temperature-dependence for both ligation states. Intriguingly, while bile salt binding results in a more uniform response to temperature change throughout the protein, the temperature derivative of the generalized order parameter shows different responses to temperature increase for the two forms of the protein in the investigated temperature range. Analysis of both slow and fast motions in human I-BABP indicates largely different energy landscapes for the apo and holo states suggesting that optimization of binding interactions might be achieved by altering the dynamic behavior of specific segments in the protein.
Human ileal bile acid binding protein (I-BABP), expressed in the absorptive enterocytes of the distal small intestine has a key role in the enterohepatic circulation of bile salts. In addition to aiding the absorption of lipidlike compounds in the human body, bile salts (Figure 1) are also known as signal molecules, which play important roles in the regulation of metabolic processes. In particular, by binding to the nuclear farnesoid X receptor α (FXR) they provide a negative feedback mechanism for their own synthesis thereby contributing to the maintenance of whole-body cholesterol homeostasis. In addition, by the activation of various mitogen-activated protein kinase pathways and the interaction with the G-protein-coupled receptor TGR5, they participate in the regulation of triglyceride, energy, and glucose metabolism.

Within the family of intracellular lipid-binding proteins (iLBPs), a group of small 14-15 kDa polypeptide chains which are known to facilitate the cellular trafficking of lipidlike compounds (e.g. fatty acids, retinoids, bile salts), human I-BABP is a protein with unique properties. Unlike most of the other members of the family, I-BABP has two internal and according to a recent study an additional superficial binding site and depending on the hydroxylation pattern of the bound bile salts exhibits a moderate-to-high level of positive binding cooperativity. In addition, the protein has been found to show a high degree of site-selectivity in its interactions with glycocholate (GCA) and glycochenodeoxycholate (GCDA), the two most abundant bile salts in humans. As a result, while in homotypic complexes di- and trihydroxy bile salts occupy both internal binding sites, in the heterotypic complex of I-BABP:GCDA:GCA, they displace each other and selectively occupy site 1 and site 2, respectively. An analogous site-selectivity for di- and trihydroxy bile salts has been reported for the disulfide-containing polymorph (T91C) of chicken liver bile acid-binding protein (cl-BABP). Intriguingly, while in human I-BABP site-selectivity is thought to arise as a result of
localized enthalpic effects, NMR relaxation measurements in cl-BABP suggests a possible connection with differences in protein flexibility.

Although no X-ray structure of the 14.2 kDa I-BABP is available yet, NMR spectroscopic investigation of the apo and the holo protein has revealed a topology characteristic of iLBPs. The dominant feature of this is a β-clam comprised of two antiparallel five-stranded β-sheets and a helix-turn-helix motif. A cavity of ~1000 Å³ located inside of the β-clam hosts two internal binding sites. Additionally, the existence of a third binding site on the protein exterior was proposed recently based on docking calculations and molecular dynamics simulations. The mechanism of ligand entry and exit is not fully understood in I-BABP. In other members of the iLBP family with the same topology, such as fatty acid binding proteins (FABPs), the helical cap region disordered in the apo state is thought to be the main regulator of ligand association/dissociation. Specifically, according to the dynamic portal hypothesis, ligand binding is accompanied by an ordering of the helical cap and the stabilization of the closed state of the protein. Unlike in FABPs, in human I-BABP the two α-helices are well defined in both ligation states and there is no sign of intense motion, raising the possibility of alternative entry/exit mechanisms as it has been suggested for other analogues.

The internal binding cavity of I-BABP is unusual in the sense that it contains a large number of hydrophilic side chains that are involved in extensive networks of salt bridges and hydrogen bonds. In fact, NMR spectroscopic and mutagenesis studies have identified two hydrogen-bonding networks as a likely way of energetic communication between the two binding sites. More recently, in an NMR relaxation study of the protein backbone, we have detected a µs-ms fluctuation in the unligated protein affecting several β-strands and the C-D loop, a motion which ceases upon ligation. Our NMR dynamic measurements with previously
obtained kinetic data suggest that conformational fluctuations have an important role in bile salt-human I-BABP recognition. A conformational transition on a similar time scale has been proposed earlier for several other members of the iLBP family, including the intestinal FABP, cellular retinol binding proteins (CRBP I and II), and cl-BABP, indicating that it might be a general way of mediating ligand binding in the protein family.

To improve our understanding of the role of dynamics in human I-BABP-bile salt recognition, we report here a characterization of the temperature dependence of slow (microsecond to millisecond) and fast (picosecond to nanosecond) backbone motions in the protein through the use of 15N NMR spin-relaxation analysis. Atomic scale dynamic parameters obtained from the NMR measurements are discussed in the context of previously obtained macroscopic thermodynamic and kinetic data.

MATERIALS AND METHODS

Sample Preparation. The methods used for the expression and purification of 15N, 13C/15N, and 2H/15N-labeled human I-BABP used in the experiments are detailed elsewhere. Protein was dialyzed into a buffer containing 20 mM potassium phosphate, 50 mM KCl, and 0.05 % NaN3 at pH =6.3. Protein concentration was 1 mM in all experiments. In the case of the holo sample, protein was complexed with an equimolar mixture of GCA and GCDA at a molar ratio of I-BABP:GCDA:GCA=1.0:1.5:1.5, ensuring that over 99.9% of the protein was in its bound state.

NMR Data Collection. Multidimensional NMR experiments were carried out at 283, 287, 291, 298, and 313 K on 600 MHz Varian NMR SYSTEM™ spectrometer equipped with a 5-mm indirect detection triple 1H13C15N resonance z-axis gradient probe. The backbone resonance assignment of apo human I-BABP and that of the heterotypic doubly-ligated complex
at 298 K has been published earlier.\textsuperscript{18} To obtain the amide $^1$H and $^{15}$N assignment at the newly investigated temperatures, the 298 K assignment has been transferred to the corresponding spectra and confirmed by 3D gradient enhanced HNCA\textsuperscript{25-27} and/or CBCACONH\textsuperscript{28} experiments performed on uniformly [U-$^{13}$C/$^{15}$N]-enriched human I-BABP. Spectral processing, computer-assisted spin-system analysis, and resonance assignment was carried out using Felix 2004 (Accelrys, Inc.). The $^{15}$N $T_1$, $T_2$, and steady-state $\{^1$H$\}$.\textsuperscript{15}$N$ NOE measurements\textsuperscript{29-31} were collected on U-[$^{15}$N]-enriched apo and holo human I-BABP at 283, 291, 298, and 313 K. Backbone amide $^{15}$N $T_1$ values were measured from two series of eight spectra (24 transients, interscan delay of 1.5 s) at each temperature with the following relaxation delay times $T = 20, 100, 190, 290, 390, 530, 670, \text{and} 830$ ms, and $T = 20, 50, 100, 170, 240, 340, 480, \text{and} 630$ ms. Amide $^{15}$N $T_2$ values were obtained similarly: $T = 10, 30, 50, 70, 110, 150, \text{and} 190$ ms, and $T = 10, 30, 50, 90, 130, 150, \text{and} 170$ ms. Steady-state $\{^1$H$\}$.\textsuperscript{15}$N$ NOE values were obtained in triplicate (32 transients each) by recording spectra with and without (blank) the use of $^1$H saturation applied during the last 5 s of a 7 s delay between successive transients. Presaturation was achieved with the use of 120º $^1$H pulses applied every 5 ms.\textsuperscript{32} The RF field strength of the $^1$H hard pulse was 11.6 kHz.

Relaxation dispersion data were obtained on [80\% $^2$H, 99\% $^{15}$N]-labeled protein at 283, 287, and 291 K, using a relaxation compensated Carr-Purcell-Meiboom-Gill (CPMG) dispersion experiment performed in a constant time manner.\textsuperscript{33-34} The constant time delay ($T_{CP}$) was set to 40 ms. Spectra were collected as a series of 20 two-dimensional data sets with CPMG field strengths ($\nu_{CPMG}$) of 25, 50, 74, 99, 123, 147, 172, 195, 219, 242, 289, 335, 380, 425, 469, 556, 641, 764, and 883 Hz. A reference spectrum was obtained by omitting the CPMG period in the pulse sequence.\textsuperscript{35} Spectra (3 s interscan delay, 24 transients) were acquired in duplicate.
Model-Free Analysis. Spectral densities were calculated from the $^{15}$N $T_1$, $T_2$, and steady-state $\{^1\text{H}\}$-$^{15}$N NOE relaxation parameters according to Abragam. Amide N-H bond lengths were assumed to be 1.02 Å and the $^{15}$N chemical shift anisotropy was estimated as -172 ppm during the calculation. To characterize the spatial restriction of the $^{15}$N-$^1$H bond vector on the picosecond to nanosecond time scale, the NMR relaxation data were analyzed within the extended model-free formalism. Motional parameters have been determined using FAST-Modelfree (Facile Analysis and Statistical Testing for Modelfree), interfaced with Modelfree 4.2 as detailed elsewhere. To minimize the inconsistencies with a continuous temperature dependence of the order parameter arising from different model selection at different temperatures, if at least at two of the four investigated temperatures FAST-Modelfree chose a more complicated model with two or three parameters, the same model was imposed onto the amide at the two remaining temperatures as well. An initial estimate of the rotational diffusion tensor was calculated from the filtered $T_1/T_2$ ratios using the programs r2r1_diffusion (http://www.palmer.hs.columbia.edu/software/r2r1_diffusion.html) and pdbinertia (http://biochemistry.hs.columbia.edu/labs/palmer/software/pdbinertia.html). The criteria for inclusion of residues in the diffusion tensor estimate relied on the method by Kay et al. Coordinates for the apo and holo form were obtained from PDB files 1O1U and 2MM3, respectively.

To relate the NMR-derived motional parameters to macroscopic thermodynamic data, the order parameter, $S^2$, obtained from the model-free analysis, was used to calculate a conformational entropy value for the angular fluctuation of individual N-H bond vectors according to

$$S_B = k_B \ln \left[ \pi (3 - (1 + 8S)^{1/2}) \right]$$ (1)
in which $S_B$ is the backbone conformational entropy and $k_B$ is the Boltzmann’s constant. The contribution of various sources of entropy change accompanying ligand binding to calorimetrically measurable entropy change is discussed in terms of

$$\Delta S' = \Delta S_{\text{conf}} + \Delta S_{\text{rot-trans}} + \Delta S_{\text{hydr}}$$  \hspace{1cm} (2)

where $\Delta S_{\text{conf}}$ is the change in conformational entropy of the protein and the ligand, $\Delta S_{\text{rot-trans}}$ is the change in the rotational-translational entropy of the system, and $\Delta S_{\text{hydr}}$ is the entropy change due to the changes in the hydration of the interacting partners.

**Relaxation Dispersion Analysis.** Contributions to transverse relaxation rates of conformational exchange were analyzed assuming a two-state exchange process using the all-timescales multiple quantum Carver-Richards-Jones formulation\textsuperscript{48} implemented in GUARDD.\textsuperscript{49}

**RESULTS**

**Temperature Dependence of Slow Conformational Motions.** Representative transverse relaxation dispersions as a function of CPMG field strength as determined for apo human I-BABP at 283 K, 287 K, and 291 K are plotted in Figure 2. Dispersion profiles were first individually fit at each temperature assuming a two-state exchange process between a ground and an excited state as described in Materials and Methods. The average individual exchange constants and excited state populations along with the average values of $R_{\text{ex}}$ and chemical shift differences between the two exchanging states are listed in Table S1-S3 of the Supporting Information. About one third of the assignable ~115 residues have been found to undergo a conformational exchange with a $R_{\text{ex}} > 2$ Hz in the investigated temperature range. Among these, 30 residues showed a measurable $R_{\text{ex}}$ at all three temperatures.
Similarly to measurements conducted previously at 298 K\textsuperscript{18}, residues sensing a millisecond exchange process in the 283-291 K range form an extended dynamic network in apo human I-BABP (Figure 3). Fluctuations with the largest values of $R_{ex}$ (10-25 s\textsuperscript{-1}) are detected in the E-F (T73) and G-H (V91, N96, H98) regions in the C-terminal half of the protein. These four residues are part of the two longest continuous segments (E69-V83 and L90-E102) in the protein exhibiting a $R_{ex}$. Additionally, a near continuous stretch of eleven (H52-K62) and six (K35-V40) residues show evidence of a millisecond motion in the C-D loop throughout beta-strand D as well as in beta-strand B together with a preceding linker to helix-II. While the C-D region and residue K35 showed a measurable $R_{ex}$ at 298 K as well, the slow motion in beta-strand B has not been detected before. Furthermore, evidence of a slow conformational exchange is indicated near the N-terminus on beta-strand A (K5, E7, E11) continuing (N13) in helix-I (D15, E16, F17) as well as toward the C-terminus of the protein in segments of beta-strands I (V109, E110, T113) and J (Y119, R121, V122, K124).

According to the individual fit of the dispersion profiles obtained in the 283-291 K temperature range, backbone amides appear to be clustered into two main groups (Figure 3) indicating the presence of two slightly different exchange processes in the system. Most of the residues in the E-F and G-H regions have been found to undergo an exchange process with values of $k_{ex}$ averaging around 769±59 s\textsuperscript{-1} (283 K), 1088±195 s\textsuperscript{-1} (287 K), and 1401±141 s\textsuperscript{-1} (291 K), whereas the rest of the amides exhibiting a $R_{ex}$ throughout the protein appear to display a slower average $k_{ex}$ of 265±82 s\textsuperscript{-1} (283 K), 658±45 s\textsuperscript{-1} (287 K), and 878±86 s\textsuperscript{-1} (291 K). The similarity of the exchange constants obtained for the amides within each cluster justifies a global fit with a single rate constant for each. As adequate fits of the dispersion profiles were achieved with the assumption of a two-state exchange, two separate A$\leftrightarrow$B and A$\leftrightarrow$C processes have been
considered. Parameters determined from the global fit analysis of the two exchange processes are listed in Table 1. The slightly higher excited state population determined for cluster I indicates a more pronounced presence of the conformational equilibrium involving the C-terminal half of the protein. The temperature dependence of $\ln(k_{BA}/k_{AB})$ and $\ln(k_{CA}/k_{AC})$ has been used to calculate $\Delta H = 7.0 \pm 1.6$ kcal mol$^{-1}$ and $4.2 \pm 0.1$ kcal mol$^{-1}$ as well as $\Delta S = 17.8 \pm 5.4$ cal mol$^{-1}$ K$^{-1}$ and $6.9 \pm 0.3$ cal mol$^{-1}$ K$^{-1}$ for the A to B and A to C transition, respectively, showing a typical entropy-enthalpy compensation characteristic of an order-disorder transition for both clusters. The Gibbs free energies are near 2 kcal mol$^{-1}$ in the investigated temperature range for both transitions. In light of the two clusters of amides observed in the 283-291 K temperature range, the previously collected data at 298 K$^{18}$ was revisited. Although forcing the amides into their corresponding cluster was possible, the resulting values of $k_{ex}$ were nearly the same for both groups ($3260 \pm 560$ s$^{-1}$ and $2900 \pm 400$ s$^{-1}$ for cluster I and II, respectively). The temperature dependence of the forward and reverse rate constants is depicted in Figure 4. Based upon conventional transition state theory, the energy barrier for the A to B and A to C process is estimated to be 19 kcal mol$^{-1}$ and 26 kcal mol$^{-1}$, respectively.

The chemical shift differences found for individual $^{15}$N spins mapped to the ribbon diagram of the mean coordinates of apo human I-BABP (PDB entry 1O1U$^{12}$) at 287 K are shown in Figure 3B. Similar distribution and amplitude of $\Delta \delta$ have been obtained at 283 K and 291 K. The differences were in general less than 4 ppm, with the largest $\Delta \delta$ values (>1.5 ppm) detected for residues located in the E/F and G/H turns as well as on beta-strand H.

In contrast to the apo protein, in the doubly-ligated state about 95% of the relaxation dispersion profiles remain flat in the investigated 283-291 K temperature range. Representative examples of relaxation dispersion profiles for the two ligation states are depicted in Figure S1 of
the Supplementary Information. Ceasing of conformational motion upon ligand binding is widespread in all regions showing an exchange in the apo state. This agrees well with our previous observation at higher temperature\textsuperscript{18} and indicates that the millisecond conformational motion giving rise to $R_{ex}$ in apo human I-BABP is absent in the doubly-ligated form in the entire 283-298 K temperature range.

**Temperature Dependence of Fast Protein Motions.** Because of their strong effect on entropy, motions on the picosecond to nanosecond timescale have high biological importance.\textsuperscript{50} To characterize the temperature dependence of ps-ns dynamics in the apo and holo forms of human I-BABP, $^{15}$N $T_1$, $T_2$ relaxation and steady-state $\{^1H\}-^{15}$N NOE measurements were performed at 283, 291, 298, and 313 K, and subjected to model-free analysis. For the apo protein, $^{15}$N relaxation parameters could reliably be determined for 117, 115, 112, and 103 backbone amide positions of the 126 nonproline residues at 283, 291, 298, and 313 K, respectively. For the holo state, 115 (283 K), 116 (291 K), 112 (298 K), and 107 (313 K) residues were included in the analysis. Resonances showing severe overlap and those of low intensity were excluded from the study at each temperature. The average values of $^{15}$N $T_1$, $T_2$, and $\{^1H\}-^{15}$N NOE for the two protein states together with the mean values of the generalized order parameters ($S^2$) are given in Table 2. Generalized order parameters were determined for each of the resonances that could be reliably fit to models 1-5. Although at 283 K and 291 K the majority of the amides could be fit by the simplest model-free formalism using $S^2$ alone (model 1), substantially more residues required an apparent conformational exchange term (models 3 and 4) to fit their relaxation parameters than previously observed at 298 K.\textsuperscript{18} For instance, in the apo protein, the number of residues with $R_{ex} > 0$ increased from 11 to 22 to 30 as the temperature was decreased from 298 to 291 to 283 K. A similar tendency was found for the holo protein. In
the apo protein, the position of the amides exhibiting a conformational exchange according to the model-free analysis generally fell into the regions exhibiting a nonflat dispersion profile in CPMG relaxation dispersion experiments. In the holo protein, as dispersion profiles remained almost exclusively flat, no such agreement was found. We should note though that by model-free analysis, only about 20-30% of the affected residues showed a $R_{ex}$ exceeding 2 Hz in both states of the protein and less than 10% could be associated with a $R_{ex} > 5$ Hz. Regarding fast internal motions between 283-298 K, about 20-25% (10-15%) of the assignable ~115 residues required the inclusion of an effective correlation time of $\tau_c$ (models 2, 4, and 5) in the apo (holo) protein and about one third of the affected ones had to be fit by the two-time scale spectral density function (model 5). At 313 K, while the holo form behaved according to the description above, in the apo state a substantially larger number of backbone amides had to be fit with a two- or three-parameter model.

To relate the generalized order parameters determined from the model-free analysis to the density of thermally accessible states and local conformational heat capacities of a specific amide bond vector along the sequence, the temperature dependence of $S^2$ was analyzed. To minimize the inconsistencies with a continuous temperature dependence of $S^2$ and $S_B$ (eq 1) arising from different model selection at different temperatures, if at least at two of the four investigated temperatures FAST Modelfree chose a more complicated model with two or three parameters, the same model was imposed onto the amide at the two remaining temperatures as well. This affected about 30% of the residues in both protein states and resulted in the same motional model across the entire temperature range for nearly 90% of the assignable residues.

Figure 5 shows the temperature dependence of 1-S for a few representative backbone amides located in various parts and various secondary structure elements of the protein in apo
and *holo* human I-BABP. Surprisingly, for the vast majority of the residues, the temperature dependence of 1-S appears to be nonlinear indicating ranges of temperature in which the flexibility of amides decreases rather than increases with increasing temperature. In the case of the *apo* protein, the most general behavior is exemplified by K19 (α-helix) and Y119 (β-sheet). It includes a more or less steady increase of flexibility with increasing temperature in the range of 280-300 K followed by a leveling off or a slight decrease. Intriguingly, the reverse is observed for many of the amides in the *holo* state. Specifically, as the temperature rises from 283 K to 291 K to 298 K, a slight decrease in flexibility is detected in the doubly-ligated form followed by an increase in 1-S above 300 K. While for most of the residues fitting of the temperature dependence of 1-S requires a polynom, for about 15% of the amides it maintains a (near) linearity. Most often this occurs in linker or loop regions (e.g. V65, *apo*; M74, *holo*), but as exemplified in the *apo* state by A31 located in the middle of helix-II, occasionally within well-defined segments as well. There is also a fraction of amides for which in the *apo* form a dramatic increase in flexibility occurs above 300 K. This again is mostly observed in loop regions (e.g. M74), with the exception of two residues in beta-strand E (E68, N70). Upon ligand binding, the sharp increase in 1-S above 300 K disappears in most cases.

Figure 6 depicts the value of S² mapped on the backbone trace for *apo* and *holo* human I-BABP at the four investigated temperatures as obtained using the imposed model selection protocol as described above. The mean values of S² at each temperature are listed in Table 2. In general, the imposed model selection protocol resulted in a slightly smaller mean S² than fast model-free but the differences between the complexed and uncomplexed states are nearly the same using both protocols. According to the mean values of S², temperature change in the investigated region of 283-313 K has a somewhat larger effect on the backbone flexibility of *apo*
I-BABP than that of the complexed form, suggesting that in overall, bile salt binding is accompanied by a reduction in the thermally accessible states for the protein backbone. According to both the mean values of $S^2$ (Table 2) and individual order parameters along the amino acid sequence (Figure 7), ligand binding has a temperature dependent effect on the amplitude of ps-ns motion of backbone amides. This is most clearly seen between the 283 K and the 298 K data. While at 283 K the $S^2$ of most of the individual backbone amides in the *apo* state is exceeding the values of $S^2$ detected in the *holo* form, at 298 K it is the other way around. At 313 K, a slight decrease in $S^2$ occurs upon ligation throughout the protein with the exception of specific regions, where bile salt binding is accompanied by a dramatic decrease in flexibility. The outliers shown in Figure 6 and 7 are located in loop regions or at the termini of secondary structure elements. The largest number of amides with $S^2 < 0.75$ occurs at 313 K in the *apo* state.

Grouping of the temperature dependence of 1-S by secondary structure elements reveals further details about the ligation-induced changes in backbone flexibility. The variation in $d(1-S)/dT$ within each $\alpha$ element is depicted in Figure 8 for *apo* and *holo* human I-BABP at three different temperatures as determined by fitting of 1-S *vs.* T and calculation of the corresponding slope at each specific temperature for individual backbone amides. The box chart form used in Figure 8 allows a simultaneous representation of the median value, the variability, as well as the asymmetry of $d(1-S)/dT$ values in a given structural element. As depicted in Figure 8, at low temperature (near 283 K), while in the *apo* form of the protein the slope of 1-S *vs.* T is positive, upon ligation it changes its sign throughout the protein. For some of the regions ($\beta_A, \beta_B, \beta_C, \beta_D, \beta_i, \beta_l$), the absolute value of the slope becomes smaller indicating a less steep temperature dependence. This can be interpreted as a reduction in the thermally accessible states and a
reduced local heat capacity for the given segment. It is worth noting that in some segments of
the protein while the slope changes its sign upon bile salt binding, in absolute value it remains
the same (α-I, α-II, βF, βG, βH). Moreover, there is one beta-strand (βE) in which according to the
mean value of the slope, the temperature dependence becomes steeper upon ligand binding. It is
also noticeable that as a result of complex formation, the spread of d(1-S)/dT within secondary
structure elements becomes smaller throughout the entire sequence near 283 K, indicating a more
uniform response to temperature in the presence of ligands. At higher temperatures, in particular
near 313 K, d(1-S)/dT becomes smaller and negative for most regions in the apo protein with the
exception of α-II and βE. For these two segments, the mean value of the slope stays positive and
fairly large accompanied by a large variability in the values of d(1-S)/dT among individual
backbone amides. Upon ligation, their spread becomes smaller and comparable to the rest of the
structural elements.

**DISCUSSION**

For a thorough understanding of protein function and the ability to modulate binding
interactions, the intimate relation of protein structure and dynamics needs to be considered.
Human I-BABP is a small but challenging system with two binding sites exhibiting positive
cooperativity and site-selectivity in its interactions with structurally diverse bile salts. To
improve our understanding of the role of flexibility in human I-BABP function, we used 15N
NMR spin relaxation techniques to characterize the temperature dependence of internal motions
occurring in the free and doubly-ligated forms on the μs-ms and ps-ns timescales.

The two clusters of residues found to undergo a millisecond time scale conformational
fluctuation with slightly different exchange rates involve extensive regions of apo human
I-BABP and provide a dynamic connection between distant sites. The ‘faster’ cluster comprises two main continuous segments of residues in the C-terminal half of the protein, whereas the ‘slower’ cluster involves the helical region, the proximate C-D loop, and two beta-strands in the N-terminal half (Figure 3). As approaching room temperature, the exchange rate of the ‘slow’ cluster catches up with the ‘faster’ one (Figure 4) merging into a single network of fluctuation with a $k_{ex}$ of $\sim 2000\text{--}3000\text{ s}^{-1}$. This matches the time scale of an initial unimolecular kinetic step in the binding scheme proposed previously in a stopped-flow kinetic analysis of human I-BABP – bile salt interaction.$^{24}$

Many of the residues implicated in either of the two clusters in apo human I-BABP exhibit a significant chemical shift change upon bile salt binding.$^{18}$ The most affected region appears to be the G-H and E-F regions in both respects. Moreover, in some regions of the protein, particularly those located farther away from the bound bile salts in the holo form, thus not subjected to direct ligand effects, there is a fairly good agreement between dynamic chemical shift differences derived from $^{15}$N relaxation dispersion experiments and backbone nitrogen chemical shift changes observed upon ligand binding (Figure 9). Thus it seems likely that the ms time scale fluctuations detected by the $R_{ex}$ measurements correspond to conformational transitions between a ground and two low-populated excited states exhibiting conformations reminiscent of that of the holo form. A similar correlation has been found by Cogliati and coworkers$^{51}$ for cl-BABP, where an extensive network of conformational fluctuation between a ground and a holo-like excited state conformation has been proposed to be mediating the access of ligands for the binding cavity in the apo form. Interestingly, in other members of the iLBP family, such as FABPs and CRBPs, primarily the helical cap region and the nearby C-D loop have been associated with the regulation of ligand entry by both stopped-flow fluorescence$^{16}$ and
NMR relaxation\textsuperscript{15,17} measurements. Another study of cl-BABP has lead to an extension of the previously proposed ‘dynamic portal hypothesis’ toward a model in which ligands enter through a flexible region consisting of not only the helical cap and the C-D loop but also the E-F region\textsuperscript{19} Yet another investigation of the porcine I-BABP analogue has suggested the existence of a second portal for ligand entry involving the G-H region.\textsuperscript{20} Our temperature-dependent $R_{\text{ex}}$ measurements support a conformational selection model of ligand binding in human I-BABP with the involvement of nearly all of the secondary structure elements providing a dynamic communication network between distant regions. While in FABPs the regions involved in conformational exchange also exhibit an enhanced ps-ns motion\textsuperscript{15}, in BABPs no such correlation between $\mu$s-ms and ps-ns motions is observed. Intriguingly, the temperature dependence of the forward and reverse rate constants associated with the slow conformational fluctuations in apo I-BABP shows an enthalpy-entropy compensation indicative of a disorder-order transition. Accordingly, a more disordered excited state with a population of a few percent must be present in the apo form.

Besides a likely role in mediating ligand entry, the slow motion involving the G-H and E-F regions in wild-type and disulfide-bridge containing cl-BABP analogues has also been associated with the site-selectivity of ligand binding.\textsuperscript{11} Moreover, NMR structural data and molecular dynamics simulations on cl-BABP suggest that conformational flexibility of the G-H region is required for an efficient coupling between the two binding sites.\textsuperscript{52} This view is supported by our stopped-flow kinetic studies of the human analogue indicating a role of a conformational change occurring on the time scale of seconds (thereby being distinct from that of mediating ligand entry) in positive binding cooperativity.\textsuperscript{24} This raises the possibility that a motion mediating ligand entry in the apo protein is transformed into a different timescale-motion
in the ligated form. We note that the recently reported third, superficial binding site in human I-BABP is thought to be involved in an allosteric mechanism of ligand binding and is proposed to have a role in both positive cooperativity and site-selectivity.\textsuperscript{8}

Superimposed onto the slow motions are small-amplitude local fluctuations in proteins occurring at room temperature on the picosecond-to-nanosecond time scale. Their analysis provides insight into the density of energy states thermally accessible for a given protein segment which when studied in a temperature-dependent manner can be related to local conformational heat capacities, in particular when comparing different ligation states. In our study of human I-BABP, we have found an unusual, nonlinear temperature-dependence of backbone amide ps-ns flexibility and conformational entropy (eq 1) corresponding to a temperature-dependent heat capacity in the investigated temperature range. Moreover, according to our analysis, the sign of the slope of $1-S$ vs. $T$ changes upon ligand binding corresponding to different responses to temperature increase in the two forms of the protein. Regarding the variability of the temperature response within secondary structure elements, while at low temperature ($\sim 283$ K) there is a decrease in variability upon ligation for the entire sequence, at high temperature ($\sim 313$ K) it occurs only at specific regions (Figure 8). We note that a similar nonlinear temperature dependence of backbone ps-ns flexibility was recently observed for the human growth hormone\textsuperscript{53} and the glutamine binding protein\textsuperscript{54} based on NMR relaxation data.

The sum of the conformational entropy terms determined for individual residues in the two ligation states by NMR relaxation measurements can be used as an upper estimate\textsuperscript{46-47, 55} of the backbone conformational entropy change associated with ligand binding and can be related to macroscopic thermodynamic parameters. According to a previous calorimetric study by Tochtrop et al.\textsuperscript{9}, the total entropy change accompanying the binding of a 1:1 molar mixture of
GCA and GCDA to human I-BABP at 25 °C and conditions similar to the ones we used in our NMR relaxation study is ~ -2.6 cal/(mol K). As for protein-ligand interactions in general, this calorimetrically measurable entropy change arises from different sources (eq 2). Based on theoretical studies of protein folding\textsuperscript{56-57}, the value of backbone conformational entropy change of $\Delta S_{\text{conf}}^{bb} \sim -89$ cal/(mol K) determined from our NMR relaxation study at 298 K translates into an overall conformational entropy change of $\Delta S_{\text{conf}} \sim -206$ cal/(mol K). By considering rotational-translational entropy changes accompanying ligand binding for protein-ligand interactions of similar size\textsuperscript{58}, $\Delta S_{\text{hydr}}$ of bile salt binding in eq 2 can be approximated as ~254 cal/(mol K). Relying on small molecule thermodynamic transfer data\textsuperscript{59-60}, this corresponds to the release of ~ 200 water molecules, which for a protein with a binding cavity of ~1000 Å\textsuperscript{3}, seems reasonable.\textsuperscript{61} Tochtrop et al.\textsuperscript{9} has also performed ITC experiments to characterize the temperature dependence of the binding interaction for the human I-BABP:GCA complex. Although due to the strong positive cooperativity of ligand binding, the enthalpy change of the first binding step at lower temperatures (15 and 20 °C) has a large ambiguity in their study, the overall trend of the total enthalpy change indicates a nonlinear temperature dependence and similar to our findings a temperature dependent heat capacity in the 10-40 °C temperature range.

The decrease in the ps-ns backbone flexibility of \textit{apo} and \textit{holo} human I-BABP with increasing temperature as observed in our NMR relaxation study in specific temperature ranges should be caused by intramolecular interactions that increase in strength by increasing temperature. Furthermore, the near concerted response to temperature throughout the entire protein suggests that interactions forming extensive networks between distant protein regions must be responsible for the unusual behavior. Unlike hydrogen bonds, hydrophobic and electrostatic interactions are known to increase in strength with increasing temperature\textsuperscript{62-63} and
our data suggest that they likely have a major contribution to protein stability in human I-BABP. This is supported by a close inspection of the human I-BABP structure revealing a number of possible salt bridges and extensive networks of hydrophobic contacts between different secondary structure elements (Figure 9 and S2).

In conclusion, the analysis of both slow and fast motions in human I-BABP indicates largely different energy landscapes for the two ligation states suggesting that optimization of binding interactions might be achieved by altering the dynamic behavior of specific segments in the protein. Further experiments such as side-chain relaxation data and mutagenesis studies should shed more light on how different time scale motions are channeled into each other upon fine modulation of the identified dynamic interaction networks in the protein.
Table 1. Kinetic and thermodynamic parameters of conformational exchange in *apo* human I-BABP deduced from $^{15}$N backbone relaxation dispersion NMR measurements. Residues were subjected to a global fit analysis in each cluster.

<table>
<thead>
<tr>
<th></th>
<th>283 K</th>
<th>287 K</th>
<th>291 K</th>
</tr>
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<tbody>
<tr>
<td><strong>cluster I</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$k_{ex}$ (s$^{-1}$)</td>
<td>836±59</td>
<td>1049±88</td>
<td>1540±120</td>
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<tr>
<td>$p_b$ (%)</td>
<td>3.1±0.2</td>
<td>3.9±0.2</td>
<td>4.3±0.2</td>
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<tr>
<td>$k_{AB}$ (s$^{-1}$)</td>
<td>26±3</td>
<td>41±4</td>
<td>66±6</td>
</tr>
<tr>
<td>$k_{BA}$ (s$^{-1}$)</td>
<td>810±57</td>
<td>1008±85</td>
<td>1474±115</td>
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<tr>
<td>$\Delta G_{AB}$ (kcal/mole)</td>
<td>1.9±0.1</td>
<td>1.8±0.1</td>
<td>1.8±0.1</td>
</tr>
<tr>
<td>$\Delta H_{AB}$ (kcal/mole)</td>
<td></td>
<td></td>
<td>7.0±1.6</td>
</tr>
<tr>
<td>$\Delta S_{AB}$ (cal/(mole K))</td>
<td></td>
<td></td>
<td>17.8±5.4</td>
</tr>
<tr>
<td><strong>cluster II</strong></td>
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<td></td>
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<td>$k_{ex}$ (s$^{-1}$)</td>
<td>294±40</td>
<td>705±69</td>
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</tr>
<tr>
<td>$p_c$ (%)</td>
<td>1.8±0.2</td>
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<tr>
<td>$k_{AC}$ (s$^{-1}$)</td>
<td>5.3±0.9</td>
<td>14±2</td>
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<td>$k_{CA}$ (s$^{-1}$)</td>
<td>289±39</td>
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<td>$\Delta G_{AC}$ (kcal/mole)</td>
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<td>$\Delta H_{AC}$ (kcal/mole)</td>
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<tr>
<td>$\Delta S_{AC}$ (cal/(mole K))</td>
<td></td>
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<td>6.9±0.3</td>
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Table 2. Summary of $^{15}$N NMR spin-relaxation rates, generalized order parameters, and derived backbone conformation entropy contributions for apo human I-BABP and the heterotypic ternary complex of I-BABP:GCDA:GCA (1.0:1.5:1.5).

<table>
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<td>$^{15}$N R$_1$ (s$^{-1}$)</td>
<td>1.4±0.2</td>
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<td>$^{15}$N R$_2$ (s$^{-1}$)</td>
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<tr>
<td>$^1$H $^{15}$N NOE</td>
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<td>0.79±0.05</td>
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<td>$S^2$, imposed m.s.</td>
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<td>0.87±0.07</td>
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<td>$S_B^{\text{average}}$ (J/K)$^a$</td>
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<td>$S_B$ (cal/(mole·K))$^a$</td>
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<td>-341±128</td>
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<td><strong>holo</strong></td>
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<td></td>
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<td>$^{15}$N R$_1$ (s$^{-1}$)</td>
<td>1.2±0.09</td>
<td>1.5±0.1</td>
<td>1.8±0.2</td>
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<td>$^{15}$N R$_2$ (s$^{-1}$)</td>
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<td>11.6±1.6</td>
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<td>$^1$H $^{15}$N NOE</td>
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<td>$S_B^{\text{average}}$ (J/K)$^a$</td>
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<td>$S_B$ (cal/(mole·K))$^a$</td>
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<td>-270±109</td>
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$^a$Calculated using the imposed model selection described in Materials and Methods. Briefly, if at least at two of the four investigated temperatures FAST-Modelfree chose a more complicated model, the same model was imposed onto the amide at the two remaining temperatures as well.
ASSOCIATED CONTENT

**Supporting Information.** Tables containing the exchange parameters derived from individual fit of the relaxation dispersion curves, generalized order parameters, and contributions to backbone conformational entropy as determined by model-free analysis. Figure showing representative examples of relaxation dispersion profiles for the *apo* and the *holo* protein, figure of salt bridges and hydrophobic contacts in the *holo* protein. This material is available free of charge via the Internet at [http://pubs.acs.org](http://pubs.acs.org).

**Author Contributions**

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

**ACKNOWLEDGMENT**

The authors are indebted to D. P. Cistola for the plasmid of human I-BABP.
**FIGURE CAPTIONS**

**Fig. 1** General structure of physiologically relevant bile salts. The vast majority of the derivatives of cholic (R₁, R₂ = OH), chenodeoxycholic (R₁=OH, R₂=H), and deoxycholic (R₁=H, R₂=OH) acid in the human body exist in a glycine (A= -NHCH₂COOH, 67 %) or taurine (A= -NHCH₂CH₂SO₃H, 23 %) conjugated form.

**Fig. 2** Two-state conformational exchange model for human I-BABP. Transverse relaxation dispersions of the backbone ¹⁵N nuclei of V91 (A) and H52 (B) in apo human I-BABP as a function of CPMG B₁ field strength at temperatures of 283, 287, and 291 K. Solid lines correspond to global two-state exchange models with parameters listed in Table 1 for cluster I (A) and cluster II (B).

**Fig. 3** Contribution to transverse relaxation from conformational exchange as derived from CPMG relaxation dispersion measurements on apo human I-BABP. A) Values of Rₑₓ determined assuming two separate two-state global exchange processes corresponding to cluster I (black) and II (grey) at 283 K (circle), 287 K (triangle), and 291 K (star). B) Backbone ¹⁵N chemical shift modulations observed at 287 K with parameters listed in Table 1. Differences in chemical shifts (Δδᵢ) were normalized with respect to their maximum value (Δδₘₚₙ) and mapped on the ribbon representation of the mean structure of the protein (PDB entry 1O1U₁²) in a pink-to-red (cluster I) and yellow-to-orange (cluster II) gradient. Residues exhibiting a flat dispersion profile or with no available data are colored in grey.
**Fig. 4** Temperature dependence of conformational exchange in apo human I-BABP. Forward (closed) and reverse (open) rate constants of the conformational exchange in cluster I (triangle) and II (circle) obtained from global two-state analysis of relaxation dispersion curves. Linear fit of parameters was obtained using data obtained at 283, 287, and 291 K.

**Fig. 5** Examples of ps-ns backbone flexibility as determined from $^{15}\text{N}$ NMR spin relaxation measurements in apo human I-BABP (open circles/dashed lines) and the heterotypic doubly-ligated complex of human:I-BABP:GCDA:GCA (closed circles/solid lines). The values of 1-S are shown as a function of temperature for residues located in different secondary structure elements: α-helix (K19 and A31), loop (V65 and M74), and β-sheet (N70 and Y119).

**Fig. 6** Temperature dependence of the order parameters for free and doubly-ligated human I-BABP. The values of $S^2$ are mapped onto the backbone trace of the mean coordinates of apo human I-BABP (PDB entry 1O1U$^{12}$) (*top*) and that of the heterotypic I-BABP:GCDA:GCA complex (PDB entry 2MM3$^{14}$) (*bottom*) and color coded in a pink-to-blue gradient. Residues exhibiting a $S^2 < 0.75$ are depicted in darker red. Grey represents proline or overlapping residues or those for which $S^2$ could not be quantified by modelfree-analysis.

**Fig. 7** Temperature dependence of the effect of ligation on backbone amide ps-ns flexibility. Differences of the generalized order parameters determined between the heterotypic doubly-ligated complex of I-BABP:GCDA:GCA (1.0:1.5:1.5) and apo human I-BABP are shown at 283, 291, 298, and 313 K.
**Fig. 8** Temperature response of backbone flexibility by secondary structure elements in *apo* and *holo* human I-BABP. The values of d(1-S)/dT are shown at 286, 298, and 310 K by curve-fitting of 1-S vs. T for each residue for which S^2 could be quantified and calculating the derivative at each specific temperature. The variation in d(1-S)/dT within each secondary structure element is depicted in a box-chart representation. The bar and the dot inside the box indicate the median and the mean value of d(1-S)/dT, respectively, whereas the ends of the box correspond to the upper and lower quartile of the data. The crosses extending from the box indicate minimum and maximum data values within a given secondary structure element. Elements of secondary structure are listed in order of their occurrence in the protein sequence. The ratio of the number of residues used in the analysis relative to the total number of nonproline residues in a given segment is shown at the bottom of the chart at each temperature. The heights of the charts are scaled according to the scale of the vertical axis.

**Fig. 9** Conformational exchange in human I-BABP. A) Correlation between 15N backbone chemical shift differences deduced from relaxation dispersion measurements on *apo* human I-BABP (Δω) and those detected upon binding an equimolar mixture of GCDA and GCA at a molar ratio of 1.0:1.5:1.5 (Δδ). B) Residues exhibiting a linear correlation (R^2=0.98, slope=1.1) between Δω and Δδ are indicated in black on the mean structure of *apo* human I-BABP (PDB entry 1O1U^{12}).

**Fig. 10** Salt bridges (A) and hydrophobic contacts (B) in *apo* human I-BABP.^{12} For clarity, only residues involved in contacts between different secondary structure elements are shown in B). Similarly extensive electrostatic and hydrophobic contacts exist in the *holo* state^{13-14} (Figure S2, Supplementary Information).
REFERENCES


Fig. 5

Fig. 6
Fig. 7
Fig. 8
Temperature Dependence of Backbone Dynamics in Human Ileal Bile Acid-Binding Protein.

Implications for the Mechanism of Ligand Binding

Gergő Horváth, Orsolya Egyed, and Orsolya Toke