

Scientific paper

Differences in Unfolding Energetics of CcdB Toxins From *V. fischeri* and *E. coli*

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Dedicated to Prof. Dr. Gorazd Vesnaver on the occasion of his 70th birthday

Abstract

Ccd system is a toxin-antitoxin module (operon) located on plasmids and chromosomes of bacteria. CcdB_F encoded by *ccd* operon located on *Escherichia coli* plasmid F and CcdB_{Vfi} encoded by *ccd* operon located on *Vibrio fischeri* chromosome are members of the CcdB family of toxins. Native CcdBs are dimers that bind to gyrase-DNA complexes and inhibit DNA transcription and replication. While thermodynamic stability and unfolding characteristics of the plasmidic CcdB_F in denaturant solutions are reported in detail, the corresponding information on the chromosomal CcdB_{Vfi} is rather scarce. Therefore, we studied urea-induced unfolding of CcdB_{Vfi} at various temperatures and protein concentrations by circular dichroism spectroscopy. Global model analysis of spectroscopic data suggests that CcdB_{Vfi} dimer unfolds to the corresponding monomeric components in a reversible two-state manner. Results reveal that at physiological temperatures CcdB_{Vfi} exhibits lower thermodynamic stability compared to CcdB_F. At high urea concentrations CcdB_{Vfi}, similarly to CcdB_F, retains a significant amount of secondary structure. Differences in thermodynamic parameters of CcdB_{Vfi} and CcdB_F unfolding can reasonably be explained by the differences in their structural features.

Keywords: Toxin-antitoxin module, CcdB, CcdA, protein unfolding, thermodynamics

1. Introduction

Toxin-antitoxin (TA) modules are operons located on plasmids and chromosomes of bacteria and archaea. The *ccd* operon is a TA module encoding the toxin CcdB and the antitoxin CcdA. C-terminal domain of CcdA is intrinsically disordered and vulnerable for proteolytic attack. In the absence of *ccd* operon expression, CcdA is degraded by proteases faster than CcdB and is thus unable to form CcdA:CcdB complexes. This leads to activation of the toxin CcdB which binds to its cellular target, DNA gyrase, and inhibits DNA transcription and replication. When *ccd* operon is expressed CcdB action is inhibited by the formation of the CcdA:CcdB complexes. *ccd* expression is auto-regulated on the level of transcription by binding of the multimeric CcdA:CcdB complex, with CcdA/CcdB molar ratio of about 1:1, to the promoter DNA.^{1–3}

ccd operon located on plasmid F of *Escherichia coli* encodes CcdB_F which has been studied in detail both in terms of structure and thermodynamic stability.^{3–7} On the other hand, very little thermodynamic information is available on CcdBs encoded by bacterial chromosomes. An example of chromosomal CcdB is CcdB_{Vfi} from the marine bacterium *Vibrio fischeri*, that shows 41% sequence similarity to CcdB_F. Both CcdB_F and CcdB_{Vfi} form dimers in the solid state as well as in solution.^{5,7–9} The structures of CcdB_{Vfi} and CcdB_F dimers show similarity of the secondary and tertiary structure (Figure 1). Each CcdB_F and CcdB_{Vfi} monomer consist of a major N-terminal β -sheet, a few smaller β -sheets and a C-terminal α -helix. In contrast to CcdB_{Vfi}, CcdB_F has a notable charge separation due to a large number of positively charged amino acids located on β -sheet and the more negatively charged helix,^{5,8} which may cause differences in thermodynamic stability of CcdB_{Vfi} and CcdB_F.

In this work an attempt was made to characterize urea-induced unfolding of CcdB_{Vfi} by CD spectroscopy. The obtained thermodynamic parameters were compared to the known values reported for CcdB_F.^{4,6} The observed differences in thermodynamic stability of CcdB_{Vfi} and CcdB_F are discussed in terms of structural differences between the two proteins.

2. Experimental

Proteins were prepared and purified as described elsewhere.^{7,8} Solutions of CcdB_{Vfi} were dialyzed against TRIS buffer (0.02 M TRIS, 0.15 M NaCl and 0.001 M EDTA with pH = 7.5). Their concentrations were determined by measuring the absorbance at 280 nm using the absorption coefficients calculated by the method introduced by Gill and von Hippel.¹⁰ CcdB_{Vfi} stock solutions were mixed with concentrated urea solutions to prepare solutions with final urea concentration ranging from 0 to 8 M.

Spectropolarimetry measurements were performed using the CD spectropolarimeter AVIV 62A DS (Aviv Associates, Lakewood, NJ, USA). Experimental conditions: (1) urea induced unfolding: temperature, $T = 5\text{ }^{\circ}\text{C} - 45\text{ }^{\circ}\text{C}$, wavelength, $\lambda = 222\text{ nm}$; (2) thermally induced unfolding: temperatures $T = 20\text{ }^{\circ}\text{C} - 85\text{ }^{\circ}\text{C}$, $\lambda = 225\text{ nm}$ with step size $1\text{ }^{\circ}\text{C}$; (3) CD spectra: $\lambda = 260 - 210\text{ nm}$ with step size 1 nm . Slit bandwidth was set to 2 nm . Temperature equilibration time was 2 minutes for thermally induced unfolding and

30 seconds for other measurements, signal averaging time was 10 seconds for thermally and urea induced unfolding and 3 seconds for other measurements. CD measurements were conducted for solutions with protein (monomer) concentration of about $3\text{ }\mu\text{M}$ (1 cm cuvette) and protein monomer concentration of about $30\text{ }\mu\text{M}$ (0.1 cm cuvette). The ellipticities, θ , measured at given wavelength corrected for the corresponding contribution of the buffer, were converted to molar ellipticities, $[\theta]$, by dividing with optical path-length, l , in cm and protein concentration, c , in μM .

Protein structures needed for surface area calculations were derived from the PDB files 1X75 (CcdB_F) and 3KU8 (CcdB_{Vfi}). Solvent accessible surface areas (SASA) of native protein structures were estimated using the program Naccess 2.1.1.¹¹ SASA of for the unfolded proteins were calculated as the sum of accessibilities of the protein residues X located in the corresponding Ala-X-Ala tripeptides. SASA calculations were performed using the solvent probe size of 1.40 \AA and Z-slices of 0.05 \AA . Other parameters needed for SASA calculations were taken as the program default values.

3. Results and Discussion

3.1. Thermally and Urea Induced Unfolding of CcdB_{Vfi}

Thermal denaturation monitored by CD spectroscopy reveals very high thermal stability of CcdB_{Vfi}. The

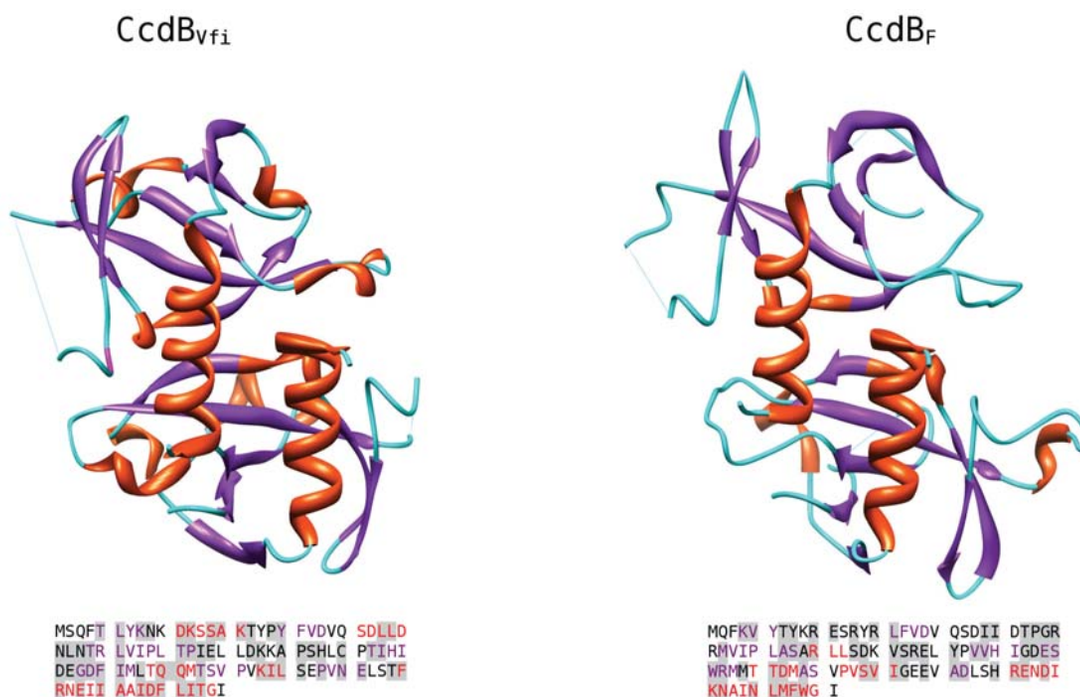


Figure 1: Comparison of crystal structures of CcdB_{Vfi} and CcdB_F dimers. Elements of secondary structure are colored red (α -helix) and violet (β -sheet). The structures were drawn with the program UCSF Chimera¹⁵ from PDB files: 1X75 (CcdB_F) and 3KU8 (CcdB_{Vfi}). Gray marked regions that represent the difference in protein primary sequences (monomers) were obtained using the program SIM.¹⁶

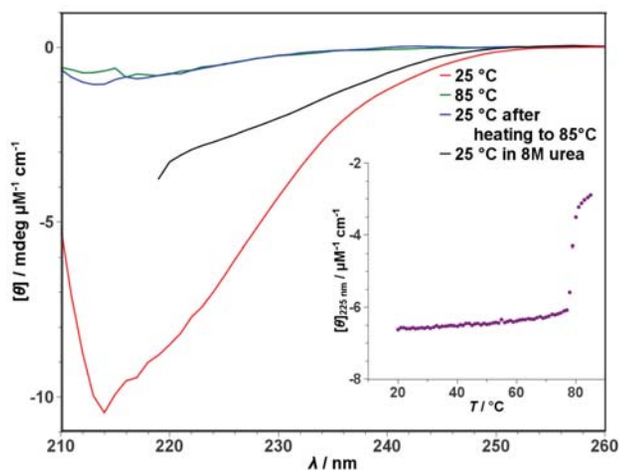


Figure 2: CD spectra and melting curve of CcdB_{Vfi}. Rescan at 25 °C was performed after cooling from 85 °C and 10 min incubation at 25 °C. CD spectrum measured in 8 M urea indicates significant fraction of residual secondary structure of CcdB_{Vfi} in the urea denatured state. Inset: Thermal denaturation followed at 225 nm.

thermally induced denaturation transition is irreversible and occurs at temperatures above 70 °C (Figure 2). Denaturation was also induced by addition of denaturant urea and monitored by CD spectroscopy at various temperatures (Figures 3 and 4). To test the reversibility of the observed urea induced denaturation transition we prepared solutions with fixed protein and different urea concentrations by dilution of (denatured) protein solutions prepared in 8 M urea. Since the extent of recovery of ellipticity is very high (Figure 3) and independent on the protein concentration between 3 and 30 μM we considered the observed urea denaturation to be a reversible process.

3. 2. Urea Induced Unfolding of CcdB_{Vfi} as Two-state Dimer-monomer Transition

Since CcdB_{Vfi} exists as a (native) dimer in both, the solid state and in the solution,^{8,9} we attempted to describe its urea induced unfolding as a reversible two-state process



where N₂ represents the native CcdB_{Vfi} dimer, D the denatured CcdB_{Vfi} monomer and K_(T,u) the apparent equilibrium constant which is a function of temperature (T) and molar urea concentration (u). K_(T,u) is defined as:

$$K_{(T,u)} = \frac{[D]^2}{[N_2]} = \frac{\alpha^2_{(T,u)}}{1 - \alpha_{(T,u)}} 2c \quad (2)$$

where [N₂] and [D] represent equilibrium molar concentrations of N₂ and D. In equation 2 α_(T,u) is the fraction of

CcdB in the denatured state at given T and u defined as α_(T,u) = [D]/c, where c is the total molar concentration of CcdB_{Vfi} monomers. According to the suggested model (equation 1), the measured ellipticity at a given wavelength corrected for the corresponding buffer contribution normalized to c = 1 μM and l = 1 cm, [θ]_(T,u), can be expressed as:

$$[\theta]_{(T,u)} = (1 - \alpha_{(T,u)})[\theta]_{N_2(T,u)} + \alpha_{(T,u)}[\theta]_{D(T,u)} \quad (3)$$

where [θ]_{N₂(T,u)} and [θ]_{D(T,u)} represent the corresponding molar ellipticities of N₂ and D given per mol of CcdB_{Vfi} monomer that can be estimated at any measured T as linear functions of urea concentration u (pre- and post-transitional baselines presented in Figure 3). The measured α_(T,u) (Figure 4) can be expressed as:

$$\alpha_{(T,u)} = \frac{[\theta]_{(T,u)} - [\theta]_{N_2(T,u)}}{[\theta]_{D(T,u)} - [\theta]_{N_2(T,u)}} \quad (4)$$

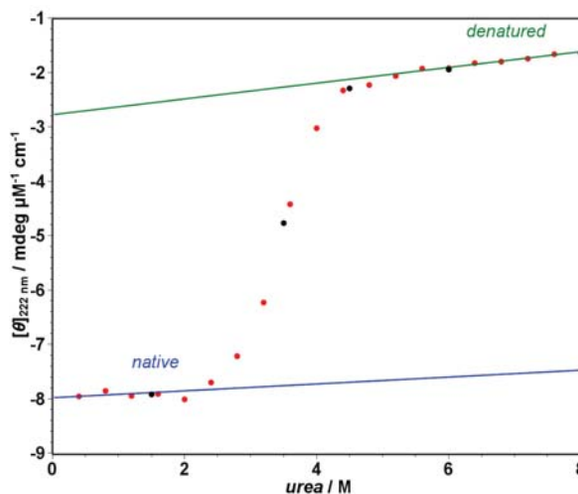


Figure 3: Typical urea-induced CD denaturation curve (red symbols) of CcdB_{Vfi} (molar ellipticity, [θ]_(T,u), versus urea concentration) measured at c = 29.6 μM and 5 °C. Black symbols represent molar ellipticities measured for CcdB_{Vfi} solutions obtained by dilution of CcdB_{Vfi} solutions prepared in 8 M urea (reversibility test). Blue and green line represent the molar ellipticities [θ]_{N₂(T,u)} and [θ]_{D(T,u)} corresponding to the native and denatured state, respectively. Together with the measured molar ellipticities [θ]_(T,u) they were used in calculations of fractions of the protein in the denatured state α_(T,u) (equation 4).

On the other hand, α_(T,u) can be connected to the energetics of unfolding through the two-state transition model (equation 1). According to this model the dependence of the apparent standard Gibbs free energy of unfolding (ΔG^o_(T,u)) on u can be at any T expressed as:

$$\Delta G_{(T,u)}^o = \Delta G_{(T)}^o - m \cdot u \quad (5)$$

where m is an empirical parameter strongly correlated to

the changes of protein accessible surface upon unfolding¹² and assumed to be temperature independent. $\Delta G_{(T)}^{\circ}$ is the standard Gibbs free energy of unfolding in the absence of urea ($u = 0$) that may be expressed in terms of the corresponding standard Gibbs free energy ($\Delta G_{(T_0)}^{\circ}$) and standard enthalpy of unfolding ($\Delta H_{(T_0)}^{\circ}$) at a reference temperature $T_0 = 25$ °C and standard heat capacity of unfolding (ΔC_p°) (assumed to be temperature independent) through the Gibbs-Helmholtz relation (integrated form):

$$\Delta G_{(T)}^{\circ} = T \left\{ \frac{\Delta G_{(T_0)}^{\circ}}{T_0} + \Delta H_{(T_0)}^{\circ} \left[\frac{1}{T} - \frac{1}{T_0} \right] + \Delta C_p^{\circ} \left[1 - \frac{T_0}{T} - \ln \frac{T}{T_0} \right] \right\} \quad (6)$$

It follows from equations 5 and 6 that the model (adjustable) thermodynamic parameters $\Delta G_{(T_0)}^{\circ}$, $\Delta H_{(T_0)}^{\circ}$, ΔC_p° and m define $\Delta G_{(T,u)}^{\circ}$ and also the corresponding $K_{(T,u)}$ ($K_{(T,u)} = \exp(-\Delta G_{(T,u)}^{\circ}/RT)$). Thus, the model function for $\alpha_{(T,u)}$ can be derived from equation 2 as:

$$\alpha_{(T,u)} = \sqrt{\left(\frac{K_{(T,u)}}{4c} \right)^2 + \frac{K_{(T,u)}}{2c} - \frac{K_{(T,u)}}{4c}} \quad (7)$$

its value at any T and u can be calculated for a given set of adjustable parameters and compared to $\alpha_{(T,u)}$ values estimated experimentally from equation 4. The values of adjustable parameters (Table 1) were obtained by global fitting of the model function (equation 7) to the family of CD unfolding curves measured at various temperatures (Figure 4) using the non-linear Levenberg–Marquardt regression procedure. Global fitting results in a good agreement between the model and experimental data. Moreo-

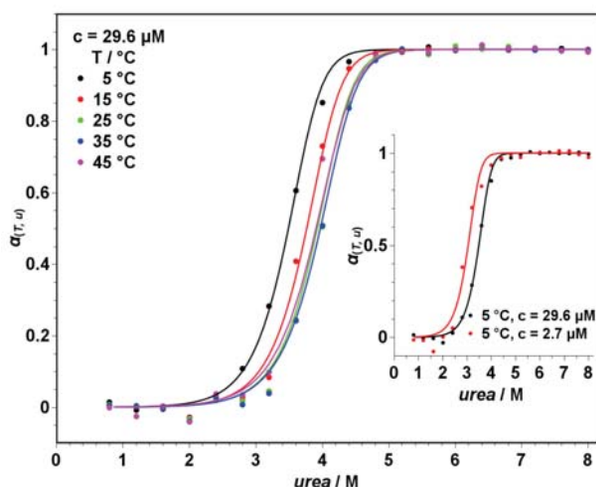


Figure 4: Global model analysis of urea induced denaturation of CcdB_{vfi}. Fraction of the protein in the denatured state $\alpha_{(T,u)}$ as a function of urea concentration determined at various temperatures, T , and protein concentrations, c (inset). The points represent experimental data and the lines represent the best global fit of the model function (equation 7). The inset shows that the transition monitored at low c occurs at lower urea concentration and is well described by the corresponding model function based on the reversible two-state dimer-monomer model (equation 1).

ver, unfolding curves measured at about ten times lower CcdB_{vfi} concentration (Figure 4-inset) show that unfolding at this concentration occurs at lower u suggesting that the transition is not monomolecular. In this light the observation that the model function (equation 7) is able to describe the curves measured at lower CcdB_{vfi} concentration well using the same set of adjustable parameters (Table 1), represents additional support of the proposed model of CcdB_{vfi} dimer denaturation accompanied by dissociation of subunits (equation 1). Therefore, and due to the observed good quality of the global fit we consider the obtained thermodynamic parameters to be reliable and physically sound.

3. 3. Differences in Unfolding Energetics Between CcdB_{vfi} and CcdB_F and Their Structural Interpretation

Thermodynamic profile of CcdB_{vfi} unfolding (Figure 4) was obtained from the best global fit values of $\Delta G_{(T_0)}^{\circ}$, $\Delta H_{(T_0)}^{\circ}$, ΔC_p° (Table 1). They were used to estimate $\Delta G_{(T)}^{\circ}$ (from equation 6), $\Delta H_{(T)}^{\circ}$ from the Kirchhoff's law

$$\Delta H_{(T)}^{\circ} = \Delta H_{(T_0)}^{\circ} + \Delta C_p^{\circ}(T - T_0) \quad (8)$$

and the corresponding entropy contribution, $T\Delta S_{(T)}^{\circ}$, from the Gibbs relation

$$\Delta G_{(T)}^{\circ} = \Delta H_{(T)}^{\circ} - T\Delta S_{(T)}^{\circ} \quad (9)$$

Table 1: Comparison of the thermodynamic parameters of CcdB_{vfi} and CcdB_F unfolding obtained from urea-induced denaturation studies.

	CcdB _{vfi}	CcdB _F ^a	CcdB _F – CcdB _{vfi}
$\Delta G_{(T_0)}^{\circ}$	18 ± 1	21 ± 1	3 ± 1
$\Delta H_{(T_0)}^{\circ}$	3 ± 1	25 ± 4	22 ± 4
$T_0\Delta S_{(T_0)}^{\circ}$	-15 ± 2	4 ± 5	19 ± 5
ΔC_p°	1.0 ± 0.2	2.6 ± 0.3	1.6 ± 0.4
m	3.0 ± 0.1	3 ± 3	0 ± 3

^a Data taken from ref. 6.

A comparison of $\Delta G_{(T)}^{\circ}$ versus T curves (Figure 5a) for CcdB_{vfi} and CcdB_F indicates that at physiological temperatures the thermodynamic stability of CcdB_F is higher compared to CcdB_{vfi}. On the other hand, the maximum stability for CcdB_F is observed at lower temperature as for CcdB_{vfi}. The obtained ΔC_p° of CcdB_{vfi} unfolding was compared to the corresponding ΔC_p° estimated as a function of no. of amino acid residues for a large set of proteins.¹³ The comparison shows that the measured ΔC_p° represents only 42% of the value expected for the protein of the same size. This suggests that a degree of unfolding of CcdB_{vfi} in concentrated urea solutions is significantly lower than the degree of unfolding

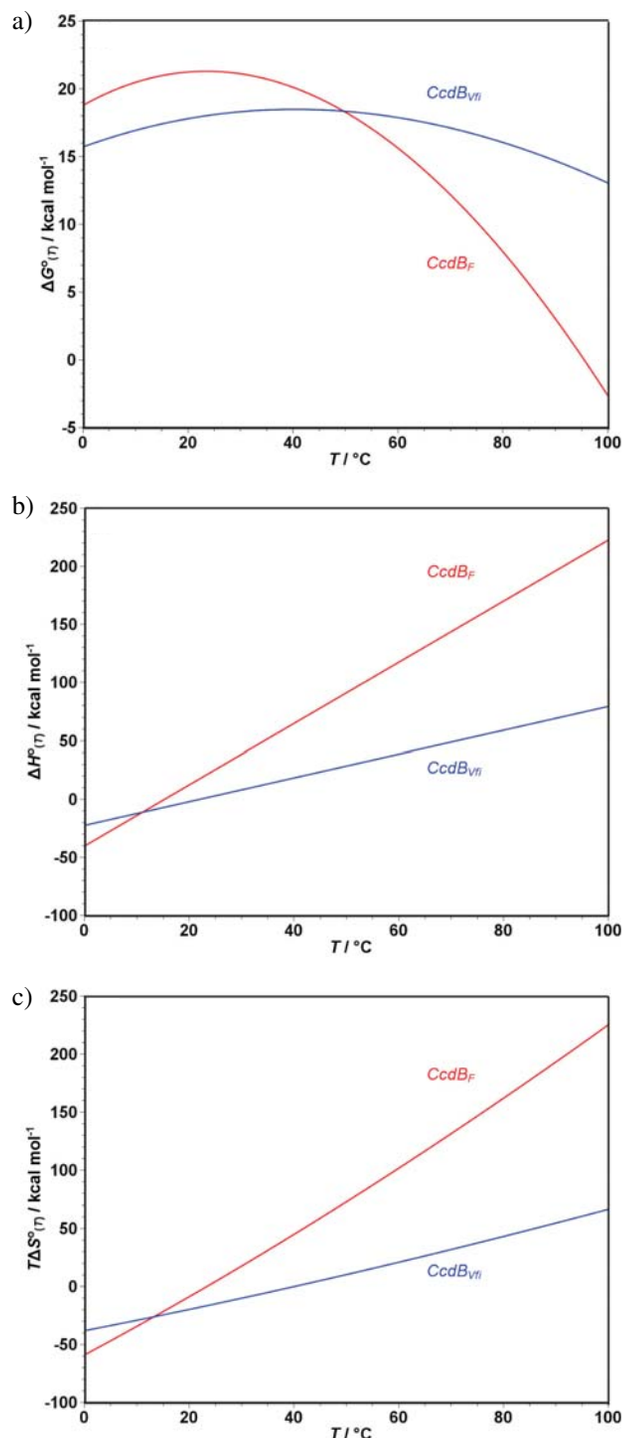


Figure 5. Thermodynamic profile of CcdB_{Vfi} and CcdB_F unfolding. (a) Standard Gibbs free energy, $\Delta G_{(T)}^{\circ}$; (b) enthalpy, $\Delta H_{(T)}^{\circ}$; (c) the corresponding entropy contribution, $T\Delta S_{(T)}^{\circ}$. All thermodynamic quantities extrapolated to urea concentration $u = 0$ are presented as functions of temperature. The profiles were calculated from the best-fit parameters (Table 1) by using equations 6, 8 and 9.

seen in an average protein of the large data set analyzed in ref. 13. The result is in accordance with those of the CD measurements (Figure 2) suggesting that CcdB_{Vfi} in the urea denatured state retains a significant amount of secondary

structure. Similar features of the urea unfolded state were observed also for previously studied CcdB_F.⁶

To correlate the observed differences in thermodynamic parameters of CcdB_{Vfi} and CcdB_F (Table 1, Figure 4), with differences in CcdB_{Vfi} and CcdB_F structural features structural characteristics of CcdB_{Vfi} and CcdB_F native (dimeric) and unfolded (monomeric) state are needed. One of those are solvent accessible surface areas (SASA). SASA of native CcdB_{Vfi} and CcdB_F were calculated from crystal structures presented in Figure 1. SASA of the unfolded proteins were approximated as the sum of SASA of the protein residues X located in the corresponding Ala-X-Ala tripeptides (the sum runs over all CcdB_{Vfi} or CcdB_F residues). The values are presented in Table 2.

Table 2: Solvent accessible surface areas (SASA) of CcdB_{Vfi} and CcdB_F

	CcdB _{Vfi}	CcdB _F	CcdB _F – CcdB _{Vfi}
$A_{\text{polar}}(2D) / 10^3 \text{ \AA}^2$	12.09	12.94	0.86
$A_{\text{polar}}(N_2) / 10^3 \text{ \AA}^2$	3.69	3.81	0.12
$\Delta A_{\text{polar}}(N_2 \rightarrow 2D) / 10^3 \text{ \AA}^2$	8.39	9.13	0.74
$A_{\text{non-polar}}(2D) / 10^3 \text{ \AA}^2$	19.33	18.99	–0.33
$A_{\text{non-polar}}(N_2) / 10^3 \text{ \AA}^2$	6.10	5.68	–0.42
$\Delta A_{\text{non-polar}}(N_2 \rightarrow 2D) / 10^3 \text{ \AA}^2$	13.23	13.31	0.09

It can be seen that the changes of polar SASA accompanying CcdB_F unfolding are about 740 \AA^2 higher than for CcdB_{Vfi} unfolding. By contrast, the difference in changes of non-polar SASA of unfolding between CcdB_F and CcdB_{Vfi} is only about 90 \AA^2 . The empirical parameterizations correlating SASA of unfolding to the corresponding thermodynamic parameters^{13,14} suggest that changes of polar SASA have much higher impact on $\Delta H_{(T)}^{\circ}$ while changes of non-polar SASA have much higher impact on ΔC_p° and $\Delta S_{(T)}^{\circ}$. Thus, the observed $\Delta \Delta H_{(T)}^{\circ} > 0$ and smaller magnitudes of $\Delta \Delta C_p^{\circ}$ and $T\Delta \Delta S_{(T)}^{\circ}$ resulting in $\Delta \Delta G_{(T)}^{\circ} > 0$ can be reasonably explained by differences in CcdB_F and CcdB_{Vfi} structural features.

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5. References

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Povzetek

Genetski sistem *ccd* je predstavnik t.i. modulov toksin-antitoksin, ki se nahajajo na plazmidih in kromosomih različnih bakterij. CcdB_F, katerega genetski zapis je vsebovan na plazmidu F *Escherichie coli* in CcdB_{Vfi}, katerega genetski zapis je vsebovan na kromosomu *Vibrio fischeri*, pripadata družini toksinov CcdB. Nativna proteina CcdB_F in CcdB_{Vfi} sta dimera, ki se vežeta na kompleks giraze z DNA in tako inhibirata prepisovanje in podvojevanje DNA. Medtem ko so termodinamska stabilnost in značilnosti razvitja plazmidnega CcdB_F v raztopinah denaturantov dobro poznane, so informacije o omenjenih značilnostih CcdB_{Vfi} zelo redke. Zato smo s spektropolarimetrijo proučevali razvitje CcdB_{Vfi} v raztopinah sečnine pri različnih temperaturah. Globalna modelska analiza spektroskopskih podatkov pokaže, da je denaturacija enostopenjski prehod, pri katerem se dimer CcdB_{Vfi} razvije in hkrati disociira v dva monomera. Rezultati kažejo, da ima pri fizioloških temperaturah CcdB_{Vfi} nižjo termodinamsko stabilnost od CcdB_F. Podobno kot CcdB_F pa pri visokih koncentracijah sečnine ohrani znatno količino sekundarne strukture. Razlike v termodinamskih parametrih razvitja CcdB_{Vfi} and CcdB_F je mogoče smiselno pojasniti z razlikami v njihovih strukturnih značilnostih.