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Extraordinary stability of hemocyanins from *L. polyphemus* and *E. californicum* studied using infrared spectroscopy from 294 to 20 K⁺

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Hemocyanins are large oligomeric respiratory proteins found in many arthropods and molluscs. Here we give infrared spectroscopic evidence of a high stability towards exposure to sub-zero temperatures for hemocyanins from the arthropods *Limulus polyphemus* and *Eurypelma californicum* at different pH values. Small but distinct temperature induced changes of the secondary structure were observed, but a stable core of at least 40% α -helical structure is preserved as identified in the infrared spectra obtained between 294 and 20 K. The structural changes differ in detail somewhat for the two hemocyanins, with overall fewer changes observed in the case of *E. californicum*. Notably, in both cases the overall changes in the α -helical content are found to be fully reversible. The small changes in the secondary structure and reversibility upon cold treatment seem to be a particular property of the two hemocyanins, since it was not observed for myoglobin studied in the same way.

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Introduction

Respiratory proteins facilitate oxygen transport and storage and are thus key elements of oxygen supply in vertebrates and higher invertebrates. Three types can be distinguished based on the structure of the active site: hemoglobins (Hb), where oxygen is bound via an iron-ion to a porphyrin ring; hemerythrins, which contain merely two iron-ions and hemocyanins (Hcs) where oxygen is bound between two copper ions, which are held in the protein by six histidines.¹ Here we focus on hemocyanins (Hcs). These extracellular multi-subunit proteins are found in the hemolymph of several species of molluscs and arthropods.^{2,3} Subunits of arthropod Hcs are arranged in hexamers or multiples of hexamers such as 2×6 mers, 4×6 mers, 6×6 mers and 8×6 mers.^{3,4} Each subunit of a molecular mass around 72–75 KDa carries one active site.^{5–7} To date it has not been clear why some species regulate their oxygen supply based on hexameric Hcs while others developed larger oligomers. Even in a particular species two different variants might exist, which in a few cases have been shown to play a slightly different physiological role.8 A clear correlation of the size of hemocyanins with the living conditions of the host organism seems to be lacking. However, based on the data available so far there seems to be a tendency that larger oligomers display a higher stability with respect to temperature-induced denaturation. A remarkable stability of the oxy-forms of *Limulus polyphemus* and *Eurypelma californicum* Hcs (91 and 92.5 °C respectively) was revealed using differential scanning spectroscopy.^{9,10}

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Eurypelma californicum is a spider found in the deserts of southwest North America, a particularly hostile environment which is characterized by large temperature fluctuations between day and night. E. californicum Hc is a 24-mer complex with a molecular mass of 1.7 MDa including seven types of subunits.^{11,12} Each subunit shows similar oxygen binding behavior despite their unique immunological and physicochemical properties.^{11,13} The horseshoe crab, Limulus polyphemus Hc, lives in the eastern coast of North and Central America in varying habitats depending on the life cycle.¹⁴ The Hc from L. polyphemus, with a molecular mass of 3.5 MDa, is composed of 48 subunits. The crystal structure of the homohexamer of subunit II of L. polyphemus Hc was resolved with a resolution of 2.4 Å.15-17 A relatively high sequence homology of 60% was reported between the Lpol-II subunit from L. polyphemus and the analogous subunit "a" in E. californicum.18

E. californicum Hc was shown to be more stable with regard to chemical denaturation and temperature induced unfolding in the native oligomer than as an isolated subunit.^{9,19} Pavlina *et al.* investigated the stability of the quaternary structure of *L. polyphemus* and *P. vulgaris* Hcs using fluorescence spectroscopy in the presence of denaturing agents.²⁰ They concluded

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that the quaternary structure of the Hcs proteins is stabilized by hydrophilic and polar forces and that the denaturation process consists of two steps: the dissociation of the native molecule into its subunits and the denaturation of the subunits. Stabilization due to oligomer formation was also reported for other arthropod Hcs.^{21,22}

In the study presented here, we investigated the structural changes induced in these Hcs by very low temperatures. Infrared spectroscopy is an established tool for the characterization of the secondary structure of proteins and peptides. Here we monitored the amide I signal at different pH values as a function of temperature from room temperature down to 20 K for Hcs from *E. californicum* and *L. polyphemus*. For comparison, we also studied myoglobin as an example of small monomeric proteins. Using IR-spectroscopy, samples can be studied at high concentrations, which are close to the *in vivo* concentrations (up to 100 mg mL⁻¹). To the best of our knowledge no data for the cold stability of *L. polyphemus* and *E. californicum* Hcs or any other large oligomeric proteins have been reported yet.

Experimental

Sample preparation

The Hc from E. californicum was obtained from the hemolymph by dorsal punctuation of the heart. The sample was diluted immediately 1:2 (v/v) with 0.1 M Tris/HCl pH 7.8, 5 mM CaCl₂, 5 mM MgCl₂ in order to stabilize the protein. The sample was then centrifuged for 30 min to remove blood cells. The hemocvanins were purified from the supernatant by gel filtration (S300 26/60, GE Healthcare) in 0.1 M Tris/HCl pH 7.8, 5 mM CaCl₂, 5 mM MgCl₂ at 4 °C.¹⁹ Hc from *L. polyphemus* was obtained from the hemolymph (obtained as described in Martin et al.²³) by centrifugation at 4 °C: 15 min at 10000g, the resulting supernatant was spun again for 15 min at 10000g and the supernatant from this step spun for 2 h at 300 000g. The pellet was resuspended in 10 mM Tris/HCl, 10 mM CaCl₂, 30 mM MgCl₂ (pH 7.8 at 20 °C) and kept at 4 °C. Purity of the samples was verified by SDS-gel-electrophoresis (Fig. S1, ESI†). The Hcs were studied in 20 mM Tris, 5 mM CaCl₂, 5 mM MgCl₂ (E. californicum, pH 7.0, 7.8, 8.5) and 20 mM Tris, 10 mM CaCl₂, 10 mM MgCl₂, 100 mM NaCl (L. polyphemus, pH 7.0, 7.5, 8.5) with the pH adjusted at 20 °C. Samples were transferred into the respective buffer by ultrafiltration using centrifugal filters (100 kDa cut-off) shortly before the experiment. Sample concentration was 2 mM referring to subunits, corresponding to about 150 mg mL⁻¹. Myoglobin from equine heart was purchased from sigma Aldrich (France). The sample was dissolved in 20 mM Tris, pH 7.5 with a final concentration of 2 mM.

Fourier transform infrared spectroscopy

The mid-infrared (MIR) spectra were obtained using a Vertex 70 FTIR spectrometer (Bruker, Germany) equipped with a liquid nitrogen MCT detector. The spectrometer was purged with dry air to avoid the spectral contribution from humidity.

Typically five spectra of 256 scans were recorded at a resolution of 4 cm⁻¹ in the spectral range from 1800 to 1000 cm⁻¹.

The FTIR-ATR measurements of Hcs were performed at different pH values (7, 7.5/7.8 and 8.5) in the attenuated total reflection mode (ATR Harrick crystal, Diamond Prism). 2 μ L of Hcs protein solution were deposited in order to obtain a film on the diamond crystal. It is noted that after drying the samples on the window the concentration was probably significantly higher than 150 mg mL⁻¹ mentioned above and the final pH was not defined.

Temperature-dependent measurements

For temperature-dependent measurements, 2 μ L of the protein sample were air-dried on a 2 mm thick CaF₂ window and placed in a copper sample holder mounted on the cold finger of a closed cycle Helium-cryostat (Model DE-202 AE, Advanced Research Scientific, Allentown, PA, USA). The cryostat system and the sample holder operate at a pressure of 10⁻⁴ mTorr.

The temperature close to the sample was controlled using a silicon diode (Scientific Instruments Calibration, precision of \pm (0.5 K)). The temperature was regulated from 294 to 20 K with a heating resistor monitored using a digital temperature controller (Model 9700-1-1, Scientific Instruments, West Palm Beach, FL, USA).

Determination of oxygen saturation after freezing

Samples at pH 7.5 and 7.8 were exposed to -80 and -194 °C overnight at concentrations similar to the ones for FTIR measurements. The samples were thawed and then diluted with the corresponding buffer, to final concentrations between 0.25 and 0.8 mg mL⁻¹. The absorption spectrum at ambient oxygen pressure (about 150 Torr) was measured using a Cary 1E (Varian, path length 1 cm) at 20 °C. The oxygenation level is expressed as the ratio of the absorbance of the copper–oxygen band at 340 nm to the aromatic band at 280 nm.

Data analysis

In order to obtain the relative contribution of each secondary structure element, the deconvolution of the amide I signal is necessary. Several different procedures have been probed and the following found to be best suited for our data. First the wavenumber of each component was determined from the second derivative of the individual spectrum. Seven Gaussian curves were fitted to each spectrum (Origin 8.5 software) while the identified position was kept constant during the first fitting. Then, a consecutive optimization of amplitudes, band positions and the half-width of the individual bands was performed.^{24,25}

The individual amide I band components were obtained at 1610, 1618, 1633, 1644, 1660, 1679 and 1694 cm⁻¹ and have been assigned to the secondary structure elements as follows: α -helices (1660 cm⁻¹), unordered structures (1644 cm⁻¹), parallel β -sheets (1633 cm⁻¹), anti-parallel β -sheets (two bands in 1618 and 1694 cm⁻¹) and β -turns (1679 cm⁻¹), on the basis of a large body of experimental data.²⁶⁻²⁹ A complete fit of the amide I bands was only possible when adding a band at 1610 cm⁻¹. The position of this signal is not typical for any known amide group

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implicated in common secondary structure elements. Finally, the area of each individual band was used to calculate its relative contribution to the overall area of the measured spectrum between 1729 and 1591 cm⁻¹. The absolute error due to the deconvolution variability and the baseline correction was estimated to be about $\pm 3.5\%$. All spectra shown were normalized on the basis of the area of the amide I band. No smoothing procedures were applied.

Standard deviation

The standard deviation was calculated for each element of the secondary structure following this formula.

$$S = \sqrt{\frac{\sum (x - \bar{x})^2}{n - 1}}$$

 \overline{x} , represent the average of the difference between the two measurements made by the cryostat and the FTIR-ATR averaged over all conditions (pH) and over both species, *x*, the difference between both values: (ATR – cryostat) for any particular condition and species and *n*, the total number of differences.

Results

The cold stability of *L. polyphemus* and *E. californicum* Hcs was monitored using FTIR spectroscopy at pH 7, pH 7.5/7.8 and pH 8.5 at temperatures between 294 K and 20 K (Fig. 1A–D). In the spectral range between 1450 and 1750 cm⁻¹ two main signals dominate the spectra of proteins, amide I and amide II modes.

The amide II band is seen at around 1537 cm⁻¹ (Fig. S2, ESI†). It includes the coupled stretching ν (C=N) and bending δ (N–H) vibrational modes and it is influenced by the tertiary structure. This is in line with previous reports on the low temperature behavior of proteins, a shift of about 9 cm⁻¹ towards higher wavenumbers, reflecting the changes in the hydrogen bonding structure.³⁰ Here, we will focus on the amide I vibrational mode, which involves the contribution of the ν (C=O) stretching vibration of the protein polypeptide chains, specific to the secondary structure of a protein.^{31,32} This band can be used to follow conformational changes induced by an external parameter, such as temperature.^{33,34}

Fig. 1 shows the comparison of the amide I region for both Hcs at 294 K and 20 K and for different pH values (pH 7, 7.5/7.8 and 8.5). The data reveal slight changes in the overall structure. In the case of Hc from *L. polyphemus*, the amide I band centered at 1660 cm⁻¹ adopts essentially the same shape for the three pH values measured (Fig. 1A and C), while a slight change occurs in the case of *E. californicum* Hc (Fig. 1B and D). In order to highlight these changes, difference spectra were calculated by subtracting the absorbance spectra at extreme pH values (7 and 8.5) from the spectrum at the intermediate pH (7.5 or 7.8). The details of the deconvoluted amide I bands are shown in Fig. S3 and S4, ESI.† Each component was assigned to secondary structure elements as follows: α-helices (1659–1661 cm⁻¹), unordered structures (1642–1646 cm⁻¹), parallel β-sheets (1629–1635 cm⁻¹), anti-parallel β-sheets (two bands in 1616–1625 cm⁻¹)



Fig. 1 Absorbance FTIR spectra of *L. polyphemus* and *E. californicum* Hcs at 294 K (A and B) and 20 K (C and D), respectively, at pH 7 in black, pH 7.5/7.8 in red and 8.5 in blue. The difference spectra are represented in violet and in green by subtracting the spectra at pH values (7 and 8.5) from the spectrum at the intermediate pH (7.5/7.8) respectively. FTIR spectra are normalized by the surface of amide I bands.

and 1693–1695 cm⁻¹ regions) and β -turns (1678–1682 cm⁻¹), on the basis of a large body of experimental data.^{26–29}

Effects of vacuum on the secondary structure

The data presented in this study were obtained on the basis of experiments of protein samples under vacuum. In order to exclude an effect on the integrity of the protein structure by the vacuum, control experiments with an ATR cell were performed (see the Materials and methods part). Table 1 summarizes the comparison of the spectra obtained at 294 K with those of a film of the same protein on an ATR crystal (Table 1). The content deviation was calculated for each secondary structure element under these conditions and then averaged over both Hcs at three pH values, yielding $f_{\rm ATR}$ - $f_{\rm cryo}$ \pm stddev. This procedure delivered the following values: -4.7 ± 3.1 (α -helix), 3.5 \pm 2.4 (unordered), 1.3 \pm 2.6 (parallel β -sheets), 5.3 \pm 1.3 (anti-parallel β -sheets) and -5.5 ± 2.3 (β -turns) (Table 1). A small increase of contributions from α -helices, β -turns and a decrease of unordered and anti-parallel β-sheet conformations appear to be induced by the vacuum. It seems that the native α -helical conformation is more stable in a vacuum and that the proteins studied here tend to adopt a more organized structure. Indeed, it was previously demonstrated with α-helical

Table 1 Comparison of structural composition of *L. polyphemus* and *E. californicum* Hcs determined by curve-fitting of the amide I band at 294 K, obtained in air (ATR) and under vacuum (cryostat). In the last column statistics of the difference between the values obtained by ATR and cryostat are given

	Hemocyanins	рН 8.5		pH 7.5/7.8		рН 7			
Secondary structure (%)		ATR	Cryostat	ATR	Cryostat	ATR	Cryostat	Average deviation ATR $-$ cryostat \pm stddev.	
α-Helix	L. polyphemus	36	43	36	43	35	43	-4.7 ± 3.1	
	E. californicum	39	42	37	37	36	39		
Unordered	L. polyphemus	29	23	25	23	29	22	3.5 ± 2.4	
	E. californicum	25	23	24	21	24	23		
Parallel β-sheet	L. polyphemus	11	10	13	9	13	9	1.3 ± 2.6	
	E. californicum	11	14	12	12	13	11		
Anti//β-sheet	L. polyphemus	9	4	11	4	8	4	5.3 ± 1.3	
	E. californicum	10	5	11	7	11	4		
β-Turns	L. polyphemus	15	20	15	21	15	22	-5.5 ± 2.3	
	E. californicum	15	16	16	23	16	23		

polyalanine that a native α -helical conformation is stabilized under vacuum conditions.³⁵ On the other hand, different orientations of the proteins on the ATR crystal compared to the samples measured in transmission in the cryostat could also contribute to these changes. Overall, the changes are small enough to be ignored.

In order to compare the stability of hemocyanins with another respiratory protein, an experiment on myoglobin was performed under the same experimental conditions as Hcs (see Fig. 2A). Table 2 shows the contribution of each secondary structure elements obtained on the basis of the deconvolution of the amide I band. The major contribution corresponds to α -helical structures of 64% plus 7% of the unordered structure. This result is in very good agreement with the FTIR analysis reported by Meersman *et al.* which show that the secondary structure consists of 73.5% of the α -helical plus unordered structure.³⁴ The contribution of the extended chain, β -turns and the anti-parallel β -sheet remains stable upon lowering the temperature. Upon cooling the native α -helical structure decreases by (-28%) in favor of the unordered structure (+26%) at 10 K (see Table 2 and Fig. 2B).

Analysis of the effect of pH on the native structures of *L. polyphemus* and *E. californicum* Hcs

In the first series of experiments, the secondary structure of Hcs was probed at different pH values since their oxygen binding affinities are pH dependent.^{36,37} At all pH values probed (pH 7, 7.5/7.8 and 8.5), the proteins are functional at standard temperature.

Fig. 3 shows the relative contribution of each secondary structure element obtained on the basis of the deconvolution. The raw data used to prepare the graphs are summarized in Fig. S3 and S4 and Table S1 (ESI†). In all cases, the major contribution results from the band centered at around 1659–1661 cm⁻¹, reflecting an α -helical structure of 35 to 43%.

The direct comparison of the data of the two species at different pH values reveals that at 294 K the α -helical content is a little higher for the Hc from *L. polyphemus* (differences of 4, 6 and 1% for pH 7, 7.5/7.8 and 8.5, respectively, seen in Fig. 3A and B and Table S1, ESI†); the shift of the later being lower than the calculated error. The parallel β -sheet, anti-parallel β -sheet and β -turn structures are approximately the same in the



Fig. 2 (A) Absorbance FTIR spectra of myoglobin at 294 K in black and 20 K in red. (B) The temperature dependence of the α -helix and unordered structure of myoglobin at pH 7.5. Filled circles: α -Helices; open circles: unordered structure.

structures of *L. polyphemus and E. californicum* Hcs at 294 K and at pH 7, 7.5/7.8. At pH 8.5, the contribution of the β -turn structure decreases for the sample from *E. californicum* at 294 K, explaining the change in the shape of the amide I band seen in Fig. 1. In contrast, the relative contribution of β -turns in the *L. polyphemus* Hc structure remains stable under the same conditions.

Table 2 Percentages of secondary structure elements of myoglobin from equine heart during freezing experiments at pH 7.5 from 294 to 10 K and heating experiments (in brackets) from 10 to 294 K. The accuracy is estimated to be within $ca. \pm 3.5\%$

Temperature (K)	α-Helices	Unordered	Extended chain	β-turns	Anti-//β-sheet
294	64 (49)	7 (20)	10 (10)	13 (15)	6 (6)
250	59 (49)	14 (22)	10 (11)	13 (14)	4 (4)
200	58 (47)	17 (26)	9 (10)	12 (13)	4 (4)
150	52 (44)	21 (26)	10 (11)	13 (14)	4 (5)
100	50 (45)	21 (24)	11 (11)	13 (14)	5 (6)
80	45 (45)	24(24)	10 (9)	15 (15)	6 (7)
50	43 (43)	26 (26)	12 (12)	14 (14)	5 (5)
40	43 (42)	27 (27)	11 (Ì11)	14(14)	5 (6)
30	43 (42)	28 (28)	11 (11)	14(14)	4 (5)
20	39 (41)	30 (28)	11 (11)	14(14)	6 (6)
10	36 (36)	33 (33)	11 (11)	13 (13)	7 (7)



Fig. 3 Secondary structure elements (α -helix, unordered, parallel β -sheet, anti-parallel β -sheet and β -turns) found on the basis of the deconvolution of the amide I band of *L. polyphemus* and *E. californicum* Hcs at 294 K (A and B) and 20 K (C and D), respectively, and at different pH values: in black at pH 7, grey at pH 7.5/7.8 and white at pH 8.5.

The same pH dependence was then studied at 20 K. A similar behavior was observed for both proteins with respect to changing pH when compared to the data obtained at 294 K. One exception was the unordered structure that decreases for the protein from *L. polyphemus* at 20 K (see Fig. 3C and D and Table S1, ESI[†]).

Secondary structure composition changes reversibly between 20 K and 294 K

In order to investigate the reversibility of the small, but clear structural changes observed under the effect of temperature, difference spectra were calculated for the data obtained for Hc from *L. polyphemus* at pH 8.5 from 294 to 20 K. This revealed consistent temperature dependent changes in the absorption spectrum (Fig. 4). The difference spectra include the sum of all changes induced by cooling/heating.

More precisely, the strong peak at 1663 cm^{-1} decreases, leading to a negative peak in the difference spectra, while the



Fig. 4 Difference FTIR spectra of *L. polyphemus* Hc at pH 8.5. The cooling experiments from 294 to 20 K are represented as full lines and the heating up from 20 to 294 K as dashed lines. For the cooling cycle, FTIR spectra were obtained by subtracting the spectrum at 294 K from the spectra at lower temperatures. For the heating cycle, the spectra at lower temperatures were subtracted from the spectrum at 294 K. FTIR spectra obtained for the cooling and the heating experiments were normalized to the area of the amide I band.

absorbance at 1631 cm⁻¹ increases, producing a positive peak (see full lines in Fig. 4). These structural changes are completely reversible, as seen from the spectra of L. polyphemus Hcs obtained after reheating the sample (dashed lines in Fig. 4 and Table S4, ESI⁺). The signals at 1696 and 1631 cm⁻¹, which include the parallel and antiparallel β -sheets, partially overlap with the decrease in the unordered structure leading to a signal at 1663 cm⁻¹. These structural changes are in agreement with the percentages of parallel β -sheets and unordered structures given by deconvolution (Table S4, ESI⁺). It is important to point out that the shifts observed in the secondary structure for both proteins during cooling correspond to small global changes in the magnitude of 3 to 7% of the corresponding absorbance spectra. For E. californicum Hc we also observed nearly fully reversible difference spectra as a function of temperature at pH 8.5, as seen in Fig. S5 (ESI[†]).

Fig. 5A–C give an overview of the relative contribution of the unordered structure and parallel β -sheets of *L. polyphemus* Hc at all pH values as a function of temperature. It seems that from 294 to 100 K a rather stable secondary structure composition exists, whereas between 50 and 100 K a transition of the secondary structure takes place. This transition involves a gradual decrease of the unordered structure and a simultaneous increase of parallel β -sheets, a tendency that is most pronounced for pH 8.5. For *E. californicum* Hc, however, no transition temperature was observed for any pH values studied (Fig. 5D–F); the structural ensemble was stable during freezing.



Fig. 5 Temperature dependence of the unordered and parallel β -sheet structures of *L. polyphemus* Hc (A) at pH 7, (B) pH 7.5 and (C) at pH 8.5 and *E. californicum* Hc (D) at pH 7, (E) pH 7.8 and (F) at pH 8.5. Filled circles: Unordered structures; open circles: parallel β -sheet structure.

Effects of low temperatures on the oxygen binding level

Fig. 6 shows that, in line with the structural integrity of the protein at low temperatures, only very weak changes can be reported when the oxygen saturation level is analyzed after thawing. Analytical ultracentrifugation shows that the oligomeric structure is affected to some extent, with up to 30% of the proteins being dissociated (Table S8, ESI[†]). This is no contradiction, since at ambient oxygen pressure individual subunits are



Fig. 6 Oxygen binding level is not altered by cold treatment. Samples were exposed to -80 and -194 °C overnight. After thawing an appropriate dilution the UV-visible absorption spectra were measured. The oxygenation level is expressed as the ratio of the absorbance of the copper-oxygen band at 340 nm to the aromatic band at 280 nm. Filled circles: *E. californicum* Hc at pH 7.8; open circles: *L. polyphemus* Hc at pH 7.5. The value at 290 K represents the control.

also fully oxygenated for these two Hcs.^{13,38} We can conclude that the oxygen binding capacity is maintained, indicating that the second domain which contains the active site and is mainly α -helical remains intact for both Hcs.

Discussion

In the current study, the structural stability of the Hcs from *L. polyphemus* and *E. californicum* in sub-zero temperatures was investigated. Both proteins were previously shown to be very stable with respect to heat-induced unfolding, with transition temperatures at 92 and 93 °C respectively.^{9,10} In control experiments, we obtained (91 and 93 °C, respectively, see Fig. S6, ESI†).

FTIR spectroscopy showed that upon cooling down to 20 K only small, but largely reversible changes in the secondary structure elements occur. The oxygen binding level is not altered by cold treatment down to -197 °C. Thus, overall both proteins are remarkably stable upon exposure to freezing temperatures. In contrast, freezing of myoglobin and monoclonal antibody 1 (mAb1) in solution was reported to result in protein unfolding with transition temperatures of -12.7 and -23 °C, respectively.^{34,39} In general, cold denaturation unfolding experiments are performed under conditions which circumvent the crystallization of ice in the relevant temperature region (e.g. by choosing destabilizing solvent conditions or high pressure). A possible explanation for the absence of unfolding in our study could then be the employment of thin films with a high protein concentration (about 150 mg mL⁻¹ or even higher, due to drying of the sample to a film), which could lead to a stabilization of the protein structure. We note, however, that the oxygen binding experiments have been performed in solution; thus once back under physiological conditions this putative stabilization is not required any more. Thus it seems that the high stability and activity under extreme conditions is an intrinsic property of the protein. This is supported by the observation that myoglobin exposed to the same treatment recovers some of the α -helical structure having lost at low temperatures, but not all: the level at 294 K is only 49% instead of 64% as observed before cold treatment, thus having irreversibly lost about 15% α-helical structure. The corresponding values for the two Hcs are 1 to 2%.

The active site is mainly α -helical¹⁷ and does not seem to suffer distortions as indicated by the preservation of oxygen binding capacity at least down to –197 °C. This is in line with the observation that even at 20 K the α -helical structure that is observed for all three pH values is stable with values close to those reported on crystal structures (43%, 1NOL.pdb), namely 43–40% at pH 8.5, 43–35% at pH 7.5 /7.8 and 41–37% for pH 7.

A small difference can be noted between the Hcs from the two organisms in terms of stability of the unordered and β -sheets structures. In the case of *E. californicum* Hc, practically no changes in the secondary structure content were found in the whole temperature range, whereas in *L. polyphemus* Hc the amount of unordered and parallel β -sheet structures changes below 50 K, especially at pH 8.5. Studies employing treatment with SDS revealed that Hc from *L. polyphemus* in this case is less stable than the Hc from *E. californicum*.⁴⁰ Together with the result of our study this might indicate that this lowered stability is a general feature.

Whether the changes in the structure upon exposure to low temperatures and pH observed for the two proteins relate to differences in the biological function cannot be answered based on the present study. It is interesting to note, however, that *L. polyphemus* Hc is exceptional in the sense that it is one of the few Hcs with an inversed Bohr-effect and regulated by chloride, while Hc from *E. californicum* has a normal Bohr-effect.^{38,41}

Another point that needs to be discussed is the effect of temperature on the pH value in the solution, since the pH value of buffer solutions may depend on temperature. For Tris-HCl buffer, in which both Hcs are solubilized, an increase of ~1.2 units in the temperature range from +25 to -30 °C was reported by Kolhe *et al.*⁴² Estimations of the pH change of Tris-buffer when changing temperature from 20 to -194 °C were identified to correspond to +2.3 units in buffer solution alone, but much less pronounced in the presence of 40 mg mL⁻¹ albumin (+0.1).⁴³ Due to the high protein concentration employed in our study, a similar stabilization of the pH might be expected, although this effect is most likely dependent on the protein type. The structural changes studied here are thus essentially based on the effect of external parameters like temperature or pH, but not on the temperature dependent pH change of the buffer.

Conclusion

In this study we showed that Hcs from two species under conditions of high protein concentration and low water content retained their secondary structure down to 50 K, and their oxygen binding capacity down to 77 K. It seems that Hcs function as their own cryoprotectant under these conditions. Below 50 K the two hemocyanins respond differently to the decreased temperature: *L. polyphemus* Hc showed a re-arrangement of its secondary structure upon decreasing temperature at all pH values, but no structural changes were observed upon increasing the pH; *E. californicum* Hc is characterized by a decrease of the β -turn structure at pH 8.5, a structural change rarely reported in the literature so far.

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References

- S. N. Vinogradov, Comp. Biochem. Physiol., Part B: Biochem. Mol. Biol., 1985, 82, 1–15.
- 2 K. E. Van Holde and K. I. Miller, in *Adv. Protein Chem*, ed. F. M. R. J. T. E. C. B. Anfinsen and S. E. David, Academic Press, 1995, vol. 47, pp. 1–81.

- 3 J. Markl and H. Decker, *Blood and Tissue Oxygen Carriers*, Springer, 1992.
- 4 K. E. v. Holde and K. I. Miller, *Q. Rev. Biophys.*, 1982, 15, 1–129.
- 5 H. Decker, N. Hellmann, E. Jaenicke, B. Lieb, U. Meissner and J. Markl, *Integr. Comp. Biol.*, 2007, 47, 631–644.
- 6 J. Markl, Biol. Bull., 1986, 171, 90-115.
- 7 A. Volbeda and W. G. J. Hol, *J. Mol. Biol.*, 1989, **209**, 249–279.
- 8 A. Kolsch, J. Hornemann, C. Wengenroth and N. Hellmann, *Biochim. Biophys. Acta, Proteins Proteomics*, 2013, 1834, 1853–1859.
- 9 R. Sterner, T. Vogl, H. J. Hinz, F. Penz, R. Hoff, R. Foll and H. Decker, *FEBS Lett.*, 1995, 364, 9–12.
- 10 K. Idakieva, Y. Raynova, F. Meersman and C. Gielens, Comp. Biochem. Physiol., Part B: Biochem. Mol. Biol., 2013, 164, 201–209.
- 11 J. Markl, A. Savel, H. Decker and B. Linzen, *Hoppe-Seyler's Z. Physiol. Chem.*, 1980, **361**, 649–660.
- 12 R. Voit, G. Feldmaier-Fuchs, T. Schweikardt, H. Decker and T. Burmester, *J. Biol. Chem.*, 2000, **275**, 39339–39344.
- H. Decker, J. Markl, R. Loewe and B. Linzen, *Hoppe-Seyler's Z. Physiol. Chem.*, 1979, 360, 1505–1507.
- 14 G. S. Ehlinger and R. A. Tankersley, *Biol. Bull.*, 2004, **206**, 87–94.
- 15 R. Topham, S. Tesh, G. Cole, D. Mercatante, A. Westcott and C. Bonaventura, Arch. Biochem. Biophys., 1998, 352, 103–113.
- 16 K. A. Magnus, B. Hazes, H. Ton-That, C. Bonaventura, J. Bonaventura and W. G. J. Hol, *Proteins: Struct., Funct., Bioinf.*, 1994, **19**, 302–309.
- 17 B. Hazes, K. H. Kalk, W. G. J. Hol, K. A. Magnus, C. Bonaventura, J. Bonaventura and Z. Dauter, *Protein Sci.*, 1993, 2, 597–619.
- 18 A. Buzy, J. Gagnon, J. Lamy, P. Thibault, E. Forest and G. Hudryclergeon, *Eur. J. Biochem.*, 1995, 233, 93–101.
- 19 R. Hubler, B. Fertl, N. Hellmann and H. Decker, *Biochim. Biophys. Acta, Protein Struct. Mol. Enzymol.*, 1998, 1383, 327–339.
- 20 P. Dolashka-Angelova, A. Dolashki, S. Stevanovic, R. Hristova,
 B. Atanasov, P. Nikolov and W. Voelter, *Spectrochim. Acta, Part A*, 2005, 61, 1207–1217.
- 21 A. Dolashki, M. Radkova, E. Todorovska, M. Ivanov, S. Stevanovic, L. Molin, P. Traldi, W. Voelter and P. Dolashka, *Mar. Biotechnol.*, 2015, 17, 743–752.
- 22 P. Dolashka-Angelova, R. Hristova, S. Stoeva and W. Voelter, Spectrochim. Acta, Part A, 1999, 55, 2927–2934.
- 23 A. G. Martin, F. Depoix, M. Stohr, U. Meissner, S. Hagner-Holler, K. Hammouti, T. Burmester, J. Heyd, W. Wriggers and J. Markl, *J. Mol. Biol.*, 2007, 366, 1332–1350.
- 24 W. Yassine, N. Taib, S. Federman, A. Milochau, S. Castano, W. Sbi, C. Manigand, M. Laguerre, B. Desbat, R. Oda and J. Lang, *Biochim. Biophys. Acta, Biomembr.*, 2009, **1788**, 1722–1730.
- 25 S. Castano and B. Desbat, *Biochim. Biophys. Acta, Biomembr.*, 2005, **1715**, 81–95.
- 26 M. Jackson and H. H. Mantsch, *Crit. Rev. Biochem. Mol. Biol.*, 1995, **30**, 95–120.
- 27 J. L. R. Arrondo, A. Muga, J. Castresana and F. M. Goni, *Prog. Biophys. Mol. Biol.*, 1993, **59**, 23–56.

- 28 A. Barth and C. Zscherp, Q. Rev. Biophys., 2002, 35, 369-430.
- 29 S. A. Tatulian, in *Lipid–Protein Interactions: Methods and Protocols*, ed. J. H. Kleinschmidt, Springer, New York, 2013, ch. 9, pp. 177–218, DOI: 10.1007/978-1-62703-275-9_9.
- 30 E. S. Manas, Z. Getahun, W. W. Wright, W. F. DeGrado and J. M. Vanderkooi, J. Am. Chem. Soc., 2000, 122, 9883–9890.
- 31 E. Kauffmann, N. C. Darnton, R. H. Austin, C. Batt and K. Gerwert, *Proc. Natl. Acad. Sci. U. S. A.*, 2001, **98**, 6646–6649.
- 32 D. Reinstadler, H. Fabian and D. Naumann, *Proteins: Struct., Funct., Genet.*, 1999, **34**, 303–316.
- 33 B. T. Yesilyurt, C. Gielens and F. Meersman, *FEBS J.*, 2008, 275, 3625–3632.
- 34 F. Meersman, L. Smeller and K. Heremans, *Biophys. J.*, 2002, 82, 2635–2644.
- 35 Y. Levy, J. Jortner and O. M. Becker, Proc. Natl. Acad. Sci. U. S. A., 2001, 98, 2188–2193.
- 36 H. Decker and R. Sterner, J. Mol. Biol., 1990, 211, 281-293.

- 37 B. Hazes, K. A. Magnus, C. Bonaventura, J. Bonaventura,
 Z. Dauter, K. H. Kalk and W. G. J. Hol, *Protein Sci.*, 1993, 2, 597–619.
- 38 M. Brenowitz, C. Bonaventura and J. Bonaventura, Arch. Biochem. Biophys., 1984, 230, 238–249.
- 39 K. L. Lazar, T. W. Patapoff and V. K. Sharma, *mAbs*, 2010, 2, 42–52.
- 40 S. Baird, S. M. Kelly, N. C. Price, E. Jaenicke, C. Meesters, D. Nillius, H. Decker and J. Nairn, *Biochim. Biophys. Acta, Proteins Proteomics*, 2007, 1774, 1380–1394.
- 41 B. Sullivan, J. Bonaventura and C. Bonaventura, *Proc. Natl. Acad. Sci. U. S. A.*, 1974, **71**, 2558–2562.
- 42 P. Kolhe, E. Amend and S. K. Singh, *Biotechnol. Prog.*, 2010, 26, 727–733.
- 43 D. L. Williamssmith, R. C. Bray, M. J. Barber, A. D. Tsopanakis and S. P. Vincent, *Biochem. J.*, 1977, 167, 593–600.