Crystallization of Hemoglobins II and III of the Symbiont-Harboring Clam Lucina pectinata

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Crystallization of Hemoglobins II and III of the Symbiont-Harboring Clam *Lucina pectinata*

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Abstract

Diffraction data to 2.7 Å resolution were measured on crystals of the homotetramers of components II and III of the cytoplasmic hemoglobin of the symbiont-harboring clam *Lucina pectinata*. Even though the crystallization conditions are different and the sequence homology of the two hemoglobins is only 63%, the crystals are isomorphous to each other and to the heterotetramer Hb II/III, implying that the residues primarily involved in the intermolecular interactions and responsible for crystal cohesion may be invariant.

Introduction

The symbiont-harboring gills of the clam *Lucina pectinata* which inhabits the sulfide-rich coastal sediments in the Caribbean contain abundant cytoplasmic hemoglobin (Read, 1966; Wittenberg, 1985) which consists of three components (Kraus & Wittenberg, 1990). Two of these components, Hb II and Hb III, combine with oxygen and are unaffected by the presence of sulfide. The third, Hb I, combines with oxygen and in addition reacts with sulfide in the presence of oxygen to form a ferric hemoglobin sulfide. The oxygen-reactive components exhibit moderately high oxygen affinities ($P_{50} = 0.1$–0.2 Torr) and unusually slow rates for oxygen association and dissociation.

Hb II consists of 150 amino acids and has a calculated molecular weight of 17476, including heme and an acetylated N-terminal residue (Hockenhull-Johnson et al., 1991). Hb III consists of 152 amino acids and has a calculated molecular weight of 18068, including heme and an acetylated N-terminal residue (Hockenhull-Johnson et al., 1994). The two hemoglobins share 95 identical residues. These include a Tyr at the tenth position of the B helix, which optical and EPR spectra (Kraus, Wittenberg, Lu & Peisach, 1990) and model-building studies (Hockenhull-Johnson et al., 1991) implicate as the distal ligand to the heme iron. Hemoglobins II and III form hetero- and homomeric complexes (Kraus & Wittenberg, 1990).

As part of our studies involving the determination of the structure of *Lucina* hemoglobin proteins, we report here the crystallization of the homomeric complexes of Hb II and Hb III and the preliminary X-ray diffraction results. A preliminary X-ray diffraction study of the hetero complex of these two hemoglobins has been reported previously (Kemling et al., 1991).

Experimental

Red rectangular-shaped crystals were obtained for both hemoglobins by the hanging-drop method at 295 K. The Hb II crystals were grown under conditions similar to those used for the Hb II/III complex (Kemling et al., 1991), namely, from reservoirs (1 ml)
containing 8% saturated ammonium sulfate and 50 mM Tris–HCl pH 7.0. The drops initially consisted of 2 µl reservoir solution and 2 µl protein solution containing 1.9 mM Hb II, 0.05 mM EDTA and 50 mM phosphate buffer pH 7.5. The crystals of Hb III were obtained under different conditions, from reservoirs (1 ml) containing 15% PEG 8000 and 500 mM lithium sulfate. The drops initially consisted of 2 µl reservoir solution and 2 µl protein solution containing 1.1 mM Hb III, 0.05 mM EDTA and 50 mM phosphate buffer pH 7.5. These conditions were established by sparse-matrix sampling (Jancarik & Kim, 1991) using Crystal Screen (Hampton Research) and could not be further optimized by varying the precipitant concentration. The Hb II crystals are generally larger than those of Hb III.

Data for Hb II were measured on a Siemens imaging proportional counter employing a three-axis camera data-acquisition system operating on a Rigaku rotating-anode X-ray generator (Cu target, 1.5418 Å wavelength) at 40 kV 70 mA, and using a Supper graphite monochromator; crystal dimensions: 0.60 × 0.15 × 0.15 mm, 2θ = 18.0°, crystal-to-detector distance = 21.0 cm, 10 589 reflections with I > σ to 2.7 Å resolution, Rsym (unweighted absolute R factor based on intensity) = 6.4%. The same system with the upgrade of a four-circle goniostat was used for Hb III; crystal dimensions: 0.35 × 0.20 × 0.10 mm, 2θ = 18.0°, crystal-to-detector distance = 16.0 cm, 9243 reflections with I > σ to 2.7 Å, Rsym = 12.5%. The difference in the Rsym values between the Hb II and Hb III crystals is probably due to their difference in size.

Results and discussion

The crystals of Hb II and Hb III are tetragonal, and systematic absences along the axes suggest that the space group is P42212 for both. They are isomorphous to each other and to the Hb II/III complex (Kemling et al., 1991). The cell constants are a = 76.5, c = 152.6 Å for Hb II, and a = 76.8, c = 153.6 Å for Hb III. The asymmetric unit probably consists of two monomers. In this case, the volume to molecular weight ratios of 3.20 for Hb II and 3.13 for Hb III are within the limits found by Matthews (1968). The observation of a dimer in the asymmetric unit suggests that the two dimers in the tetramer are related by a crystallographic twofold axis as is the case in the homotetrameric hemoglobin from Urechis caupo (Kolatkar et al., 1992).

The isomorphism of the crystals of Hb II and Hb III to each other and to the Hb II/III complex is worth noting for two reasons. First, Hb III grows under dramatically different conditions from Hb II and Hb II/III. Second, Hb II and Hb III have only 63% sequence homology and this does not necessarily imply isomorphous structures. Furthermore, it is known that vertebrate or invertebrate globins share similar tertiary structures, the myoglobin fold (Lesk & Chothia, 1980; Bashford, Chothia & Lesk, 1987). The sequence homology of Hb II and III suggests similar quaternary structures for the heterotetramer and the two homotetramers; this is confirmed by the isomorphism of the crystals. Presumably, some of the invariant residues are those involved in the subunit association. Similarly, the residues involved in the intermolecular interactions that are responsible for crystal cohesion are also likely to be invariant.

A comparison of the amino-acid sequences of the eight helices in Hb II and Hb III gives the following homology figures: 75% A, 75% B, 71% C, 29% D, 80% E, 82% F, 44% G and 50% H. The low homology of helices G and H between the two globins suggests that a quaternary structure similar to that of vertebrate hemoglobin is unlikely. In the latter, the G and H helices are in the interior of the molecule and are involved in subunit interactions. We note that other invertebrate hemoglobins differ in their quaternary structures with respect to each other and to vertebrate hemoglobins. Thus, in the dimeric and tetrameric hemoglobins from the blood clam Scapharca inaequivalvis (Royer, Love & Fenderson, 1985; Royer, Hendrickson & Chiancone, 1989), the E and F helices are involved in subunit interactions. In the homotetrameric hemoglobin from Urechis caupo (Kolatkar et al., 1992) the region of contact for subunits related by the molecular twofold axis is the A/B turn and E helix while the G/H turn and D helix interact at the other subunit interface.

It is noteworthy that optimal crystallization of Hb II occurred at approximately 0.34 M ammonium sulfate, which is rather low for protein crystallization. The solubility of a protein is primarily a function of its surface, which is usually covered more or less uniformly by polar groups that attract an excess of ions of opposite charge. At low ionic strength this screen of ions reduces the electrostatic free energy of the protein and consequently, the solubility of the protein increases. At high ionic strength, protein solubility decreases because the salt interacts preferentially with the water and increases its surface tension, which in turn increases the free energy of the solvent cavity that accommodates the protein (Creighton, 1993). This latter effect also explains why hydrophobic interactions are strengthened at all ionic strengths. The result we observe with Hb II suggests that rather than having a uniform distribution of polar groups on its surface, Hb II has one or more hydrophobic patches whose aggregation is favored by moderate increases in salt concentration.
We are in the process of trying to solve these structures by the molecular-replacement method.

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