Subunit-directed click coupling via doubly cross-linked hemoglobin efficiently produces readily purified functional bis-tetrameric oxygen carriers†

Serena Singh, Ina S. Dubinsky-Davidchik, Ying Yang and Ronald Kluger*

While cross-linked hemoglobin (Hb) tetramers can deliver oxygen as a supplement to red cells, they also cause unacceptable increases in blood pressure, presumably from their penetration of the linings of blood vessels (endothelia) where the internal hemes bind endogenous nitric oxide (NO). This penetration would lower the local concentration of NO that normally induces vasodilation. Enlarging the effective size of the oxygen-carrying protein by coupling two Hbs can prevent their extravasation. Efficient and selective protein–protein coupling to produce those species has been a significant challenge. Introduction of an azide within a protein provides a directionally-oriented reaction site for utilization of the Cu(I)-catalyzed azide–alkyne cycloaddition (CuAAC) in the protein–protein-coupling process based on solubility-directed sequential addition to a bis-alkyne. However, it is known that Hb with an azide-containing cross-link between α-subunits is unreactive in CuAAC. To direct reaction away from the α-subunits of Hb, a specific fumaryl cross-link is installed exclusively between the most reactive sites on those subunits, thereby blocking the ε-99 lysyl groups and preventing any further reaction. This modification allows installation of an azide-containing cross-link exclusively between lysine–82 ε-amino groups of the β-subunits of Hb. The multiply interconnected sites establish a geometry that permits initial interfacial interaction of the cross-linked Hb-azide with Cu(i) and a bis-alkyne. After coupling, the protein-linked azide product undergoes CuAAC at the remaining alkyne with a second cross-linked Hb-azide, producing a fully functional cross-linked Hb bis-tetramer whose oxygenation and structural properties include cooperativity and oxygen affinity that should be suitable for testing as an alternative to red cells in transfusions.

Chemical modification of human hemoglobin A (Hb), the oxygen-carrying component of our red cells, has been an important approach to the development of materials that are intended to be safe and effective universal sterile alternatives to red cells in circulation. This class of material is known as a hemoglobin-based oxygen carrier (HBOC).1–3 Since acellular Hb is a tetramer that dissociates into rapidly cleared αβ dimers outside the red cell, stabilization of the intact tetrameric state is essential.1,4 Thus, initial efforts to produce a safe and effective HBOC focused on preventing dissociation of the tetramer, most commonly by chemical cross-linking.5,6 However, clinical trials of cross-linked Hb led to observations of induction of increases in blood pressure to unacceptable levels.8–13 Based on the nature of the observed effects, it is likely that the heme proteins are small enough to extravasate through the endothelia that line blood vessels, consuming endogenous nitric oxide. Since nitric oxide is the essential local signal for relaxation of blood vessels, its resulting reduced concentration leads to vasoconstriction that increases blood pressure.14–17

In an important study, Vandegriff and coworkers18 noted that increasing the size of the circulating Hb species by the addition of chains of polyethylene glycol (PEG) is sufficient to prevent vasoactivity in circulation.19 Presumably, the increased size prevents these species from penetrating the endothelia.20 However, the PEGylation imparts high oxygen affinity along with low cooperativity.21 Those materials have been utilized in specific applications22 and are a potential therapeutic directed to very hypoxic regions of circulation. It has not yet been established in a clinical trial whether cooperativity in oxygen binding and release is necessary for a material to be effective. Calculated curves show that oxygen availability is greatest where there is a steep binding curve within the physiologically significant region. Without cooperativity, the HBOC hyperbolic curves predict materials with low affinity will not be loaded with oxygen in the lungs or high affinity materials with only
small amounts of oxygen released in circulation. We reasoned that for producing a more versatile oxygen carrier, the coupling of two Hb tetramers to form a bis-tetramer could yield a product with the necessary size. Although the minimum size needed to avoid extravasation is not known, it is clear from other studies that bis-tetramers are sufficiently large for this purpose.23–26

Specific chemical modification of proteins can introduce bioorthogonal functional groups that lead to formation of derivatives with unique and predictable reactivity.16,27–31 Cu(i)- catalyzed azide–alkyne cycloaddition (CuAAC)32,33 has been employed for coupling proteins but the process has been surprisingly inefficient despite the excellent reputation of the process with smaller species.28 In principle, introduction of an azide into a specific site in cross-linked Hb places it on a stabilized location that would allow a reaction to occur with the displayed functional group. Two azide-containing Hb cross-linked tetramers would react with a single bis-alkyne by the displayed functional group. Two azide-containing Hb surprisingly ine

Experimental section

Material and methods

Phase-directed serial cross-linking. The synthesis of the azide-containing linker 2 is described in the ESI.† Trimesyl tris(3,5-dibromosalicylate) (TTDS) was prepared according to the published procedure.28 Bis(3,5-dibromosalicyl) fumarate (DBSF) was purchased from LKT Laboratories, Inc. Highly purified solutions of human hemoglobin A were a gift from Oxygenix Co. Ltd. and are stored sealed in a carbon monoxide atmosphere at 4 °C. The procedure reported by Walder and coworkers6 was utilized to cross-link the α-subunits of native Hb with DBSF at pH 7.4 to give hemoglobin as α99-fumaryl-α99, β2.

Sequentially cross-linked Hb (α99-fumaryl-α99, β82-trimesyl-β82). A solution of α99-fumaryl-α99, β2 cross-linked carbon-monoxide-bound Hb (carbonmonoxyHb, 5.0 mL, 0.05 mM protein) in sodium borate buffer (0.05 M, pH 9.0) was converted to oxygen-bound Hb (oxyHb) by stirring under a stream of oxygen with photo-irradiation at 0 °C for two hours. The solution was then placed under a stream of humidified nitrogen and stirred at 37 °C for two hours to convert the protein to cross-linked deoxygenated Hb (deoxyHb). TTDS (5 eq., 1.3 mg) was added as a solid and the mixture was stirred overnight at 37 °C under the nitrogen atmosphere. A stream of carbon monoxide was passed across the resulting solution (at 0 °C) for 10 min. The mixture was passed through a column containing Sephadex G-25 that had been equilibrated with MOPS buffer (0.1 M, pH 8.0). The protein solution was concentrated through a membrane (30 kDa cutoff) using a centrifuge to accelerate the process (2540g for 20 min). The concentrated protein was stored at 4 °C in a vial sealed and flushed with CO. The composition of the solution was analyzed by reverse phase HPLC using a 330 Å C-4 Vydac column (4.6 mm × 250 mm) with a solvent gradient from 20 to 60% acetonitrile : water with 0.1% trifluoroacetic acid. The eluent was monitored at 220 nm. Observed drifts in retention times may be attributed to solvent evaporation over time (solvents were mixed off-line) and possibly variations in column equilibration time, temperature and degassing. The composition of the protein associated with the peaks was investigated by isolation of the fractions and analysis using electrospray ionization (ESI) high resolution mass spectrometry (AIMS Lab, Department of Chemistry, University of Toronto).

Preparation of cross-linked hemoglobin bis-tetramer. A solution of α99-fumaryl-α99, β2 carbonmonoxyHb (18.4 mL, 0.02 mM) in sodium borate buffer (0.05 M, pH 9.0) was converted to a solution of α99-fumaryl-α99, β2 oxyHb by stirring under a stream of oxygen with photoirradiation at 0 °C for two hours. This solution was placed under a stream of humidified nitrogen and stirred at 37 °C for two hours to produce α99-fumaryl-α99, β2 deoxyHb. The azide-containing cross-linking reagent, 2, in DMSO (8.5 eq., 50 μL, 0.0513 M) was then added. After five hours under the nitrogen atmosphere at 37 °C, the mixture was placed under a stream of carbon monoxide at 0 °C for 10 min, then passed through a Sephadex G-25 column equilibrated with MOPS buffer (0.1 M, pH 8.0). The protein solution was concentrated through a membrane (centrifuge at 2540g for 20 min) and stored at 4 °C. The compositions of the products were analyzed by HPLC equipped with a 330 Å C-4 Vydac reverse-phase column (4.6 mm × 250 mm) and a solvent gradient from 20 to 60% acetonitrile in water spiked with 0.1% trifluoroacetic acid, as previously described. The eluent was monitored at 220 nm. The identities of the peaks were investigated using electrospray ionization mass spectrometry analysis (AIMS Lab, Department of Chemistry, University of Toronto). To the solution containing the modified proteins (1.65 mL, 0.13 mM) in phosphate buffer (0.01 M, pH 7.4) was added alkyne 9 (20 eq., 43 μL, 0.1 M in DMSO), bathophenanthroline ligand 10 (4 eq., 43 μL, 20 mM in DMSO), CuSO4 (2 eq., 22 μL, 20 mM in DMSO) and L-ascorbic acid (40 eq., 86 μL, 100 mM in DMSO). The vial containing this mixture was crimp sealed then flushed with CO and the contents were stirred in the dark.
for 4 h. The resulting solution was passed through a Sephadex G-25 column equilibrated with MOPS buffer (0.1 M, pH 7.2). The collected protein solution was then concentrated through a membrane (30 kDa cutoff) using a centrifuge to accelerate the process (2540g for 20 min). The bis-tetramer was purified from the reagents by passing the mixture through a Sephadex G-100 column equilibrated with Tris-HCl (37.5 mM, pH 7.4) containing magnesium chloride (0.5 M). The first fraction, containing purified bis-tetramer, was concentrated through a membrane (30 kDa cutoff) by centrifugation (2540g for 20 min) and stored at 4 °C in a vial sealed and flushed with CO. The composition of the bis-tetramer containing solution before and after purification was analyzed by HPLC using a Superdex G-200 HR size-exclusion column (10 mm × 300 mm) and a Tris-HCl (37.5 mM, pH 7.4) elution buffer containing magnesium chloride (0.5 M) (see ESI, Fig. 14† for post purification size exclusion HPLC trace). The eluent was monitored at 280 nm. The corresponding reverse-phase HPLC traces are also available as ESI.† The CuAAC coupling reaction was also performed on the deoxyHb version of the doubly cross-linked protein (under a nitrogen atmosphere). The modified carbonmonoxyHb was converted to the modified deoxyHb by stirring under a stream of oxygen with photoirradiation at 0 °C for two hours followed by flushing with a stream of nitrogen for two hours at 37 °C. Post-reaction, the mixture was placed under a stream of carbon monoxide at 0 °C for 10 min then separated from the reagents and characterized as previously described.

SDS-PAGE. The molecular weights were estimated by polyacrylamide gel (12% Tris-HCl) electrophoresis. 2-Dimensional Tris-HCl polyacrylamide gels contained 12% separating gel (pH 8.8) and 5% stacking gel (pH 6.8), each with 10% sodium dodecyl sulfate. Proteins were treated with 2-mercaptoethanol and sodium dodecyl sulfate. Globin chains were denatured by heating at 100 °C for 10 min. Finished gels were stained with Coomassie Brilliant Blue. Further details are in previous papers. The purified bis-tetramer run on the gel was ~90% pure by size-exclusion HPLC analysis.

Analysis of oxygen-binding. The oxygen pressure at half-saturation ($P_{50}$) and Hill’s coefficient of cooperativity at half-saturation ($n_{50}$) were determined using a Hemox Analyzer™ with the sample maintained at 32 °C. These conditions were optimized for laboratory measurements and not as a model for circulatory studies. Hb samples (5 mL, 0.013 M), prepared in phosphate buffer (0.01 M, pH 7.4), were oxygenated prior to analysis by stirring under a stream of oxygen with photoirradiation for 1.5 h at 0 °C. The sample was then contained in a cell connected to the Hemox Analyzer™ for acquisition of the oxygen desaturation curve. The conversion to the deoxy state was achieved by flushing the cell with nitrogen. The data, acquired in triplicate, were fitted to the Adair equation using computation of a best fit by the method of nonlinear least squares (see ESI, Fig. 17 and 18†). Oxygenation properties were also acquired for the native and modified proteins in the presence of inositol hexaphosphate (IHP). To the oxygenated protein sample, 2.3 eq. IHP per heme (7.5 μL of a 0.08 M solution prepared in phosphate buffer (0.01 M, pH 7.4)) was added to give 0.1 mM IHP. The resulting solution was equilibrated in the cell for 2 min prior to deoxygenation. Experiments utilizing half this amount of IHP were also conducted using the same basic protocol.

Circular dichroism (CD) spectroscopy. Scans were acquired in triplicate using a Jasco J-710 CD spectrophotometer. CarbonmonoxyHb samples (5 μM heme) in phosphate buffer (0.01 M, pH 7.4) were prepared for far UV (200–260 nm) measurements. CarbonmonoxyHb samples (50 μM heme) in phosphate buffer (0.01 M, pH 7.4) were prepared for UV-vis (245–470 nm) spectral measurements.

Thermal denaturation. A headspace vial containing the sample of carbonmonoxyHb (2.0 mL, 0.02 mM in phosphate buffer (0.01 M, pH 7.4)) was crimp-sealed then flushed with CO. The sample was stirred in the vial, which was immersed in a heated silicone oil bath for 30 min, then cooled in an ice bath for 10 min. The resulting heterogeneous solution, containing denatured protein as a precipitate, was concentrated through a membrane (30 kDa cutoff) by centrifugation (14 000g for 15 min). The supernatant was collected and the final Hb concentration was determined from measuring absorbance at 540 nm.

Turbidity. The samples of native and modified carbonmonoxyHb (2 mL, 0.02 mM in 0.1 M, pH 7.4 phosphate buffer) were contained in quartz cuvettes. Absorbance at 700 nm was determined at 1.0 min intervals with a double beam spectrophotometer (GBC Cintra 40). The temperature was ramped at a rate of 5 °C min⁻¹ from 25 °C to 40 °C, then ramped at 1 °C min⁻¹ from 40 °C to 90 °C.

Results

The initial stabilizing link across the α-subunits was achieved using bis(3,5-dibromosalicyl) fumarate in the presence of inositol hexaphosphate. To test the efficiency of a subsequent reaction at the β-subunits, the α-α cross-linked product was
treated with trimesoyl tris(3,5-dibromosalicylate) (TTDS)\textsuperscript{36}. The product was analyzed by reverse-phase HPLC. The masses of the eluted species were identified by ESI-MS to consist almost entirely of doubly cross-linked Hb (Fig. 1). Based on the pattern of the tryptic digest of the product,\textsuperscript{37} we deduced the sites at which the cross-link is connected to the protein (α99-fumaryl-α99, β82-trimesyl-β82).

To prepare the azide-containing doubly cross-linked protein for CuAAC coupling, we produced reagents containing azides (1 and 2, Fig. 2) that are analogues of TTDS. Compound 2 was prepared by the multi-step synthesis described in Scheme 1. NBS-bromination of the BOC-protected amine produces the benzylic functionalized product 3. Azide 4 was then obtained from 3 by nucleophilic substitution. Deprotection of 4 gives

---

**Scheme 1** Synthesis of the azide-containing linker 2.
amine 5, which is combined with an activated diester to produce amide 6. Hydrolysis of the esters in 6 gives diacid 7. EDC-promoted coupling of the diacid and tert-butyl 3,5-dibromosalicylic acid efficiently produces 8. Finally, cleavage of the tert-butyl esters with TFA generates 2.

We evaluated the outcome of the reactions of cross-linkers 1 and 2 with α,ω-fumaryl cross-linked Hb. HPLC analysis of the products of the reaction of α99-fumaryl-ω99, β2 Hb with 5.5 eq. of 1 indicated that approximately 50% of the β subunits are cross-linked by the reaction, with small amounts of by-products resulting from other reactions (see ESI, Fig. 11†). This indicates that reactions of 1 lack the necessary selectivity for an efficient process. The addition of 16 eq. of 1 promoted cross-linking of the majority of the β-subunits but half of this cross-linking occurred at an alternative set of sites within the β-subunits (ESI, Fig. 12†). Those azides did not participate in CuAAC. In contrast, the addition of 15 eq. of azide 2 produces the doubly cross-linked product (α99-fumaryl-ω99, β82-azido-β82 Hb, Scheme 2) as the major product with only a minor impurity, most likely an over-modified doubly cross-linked species (Fig. 3). Reaction with 8.5 eq. of 2 produced a defined mixture of 50% of the doubly cross-linked protein and 50% of the unmodified αω-cross-linked species (Fig. 4). Reagent 2, which contains a bromine atom ortho to the central scaffold, replicates the bulk of TTDS, which is a likely source of its regioselectivity. This cross-linker was then utilized to produce a defined mixture of 50% of the doubly cross-linked protein and 50% of the unmodified αω-cross-linked species (Fig. 3). Reaction with 8.5 eq. of azide 2. Peaks: heme (10 min), β subunits (30 min), ββ cross-linked subunits (56 min) and αω cross-linked subunits (73 min).

A basic requirement for successful implementation of CuAAC bis-alkyne protein–protein coupling is that there be only one azide-containing reactant present initially. Therefore, we utilized the material that is 50% β-modified (Fig. 4). We combined the protein-containing solution in phosphate buffer (0.02 M, pH 7.4) with 20 eq. of bis-alkyne 9, 4 eq. of bathophenanthroline ligand 10, 2 eq. of CuSO4 and 40 eq. L-ascorbic acid. This mixture was stirred for 4 h at room temperature in an atmosphere of carbon monoxide according to the previously optimized procedure (Scheme 3). Denaturation of α99-fumaryl-ω99, β2 is a side-effect of CuAAC (see ESI, Fig. 13†) because Hb species with non-cross-linked β-subunits are particularly vulnerable to denaturation by Cu species.28,40

Analysis of the product solution by size-exclusion HPLC at high salt concentrations (to induce dissociation of subunits41) reveals that the desired 128 kDa bis-tetramer species is formed from the azido-doubly cross-linked material (Fig. 5). The bis-tetramer in solution was isolated by passing the mixture through a G-100 Sephadex column equilibrated with tris buffer (37.5 mM, pH 7.4) containing 0.5 M magnesium chloride. From SDS-PAGE the 64 kDa band due to the four covalently linked beta-subunits of the bis-tetramer can be identified (see ESI, Fig. 15 and 16†).

The yield of the CuAAC with the azide-containing modified Hb and the bis-alkyne is not improved with further increases in the amounts of the reagents, adjustment of protein concentration, increases in reaction time or increases in temperature (up to 40 °C). We observed no improvements from additional variations of the relative amount of the various reaction components. Utilizing different ligands, copper species and bis-alkynes decreased the relative yield.

In an important and unexpected contrast, the yield from CuAAC increases from 17% to 32% by utilizing deoxyHb (Fig. 5). Addition of inositol hexaphosphate (IHP) or sodium chloride lowers the yield, which is likely to be due to the anionic species blocking the central cationic channel where the required reactions occur. The yield from CuAAC on the modified COHb also improved marginally at pH 6, consistent with a change to a more reactive conformation.42

---

**Scheme 2.** Addition of azide 2 to α99-fumaryl-ω99, β2 in sodium borate buffer (0.05 M, pH 9.0) to yield α99-fumaryl-ω99, β82-azido-β82 by selective modification of the β-lys-82 residues.

**Fig. 3** Reverse phase HPLC from cross-linking α99-fumaryl-ω99, β2 with 15 eq. of azide 2. Peaks: heme (10 min), ββ cross-linked subunits (45 min), over-modified ββ cross-linked subunits (52 min) and αω cross-linked subunits (62 min).

**Fig. 4** Reverse phase HPLC from cross-linking α99-fumaryl-ω99, β2 with 8.5 eq. of azide 2. Peaks: heme (10 min), β subunits (30 min), ββ cross-linked subunits (56 min) and αω cross-linked subunits (73 min).
The purified bis-tetramer has a relatively low affinity for oxygen ($P_50 = 17.4$ torr, Fig. 6, Table 1). Doubly cross-linked Hb ($\alpha_{99}$-fumaryl-$\alpha_{99}$, $\beta_{82}$-trimesyl-$\beta_{82}$) also has a decreased affinity for oxygen compared to acellular Hb. Both species have moderate levels of cooperativity (as indicated by their Hill coefficients, $n_{50}$). Comparison of the $P_{50}$ values for native Hb, $\alpha_{99}$-fumaryl-$\alpha_{99}$, $\beta_2$ and $\alpha_2$, $\beta_{82}$-trimesyl-$\beta_{82}$ suggests that the presence of the fumaryl cross-link in the $\alpha$-subunits contributes significantly to the decrease in oxygen affinity in the resulting bis-tetramer.

IHP dramatically decreases the oxygen affinity of native Hb as it binds to the site that accommodates 2,3-diphosphoglycerate (DPG), an allosteric effector of Hb within red cells. The decrease of sensitivity also occurs as a consequence of cross-linking between the $\alpha$-subunits, as observed by Vandegriff and coworkers. We observed that cross-linking the $\beta$ subunits decreases the allosteric response to DPG or IHP.

### Table 1  The oxygen binding properties of native Hb and chemically modified species and their sensitivity to allosteric modification by IHP

<table>
<thead>
<tr>
<th>Hb species</th>
<th>$P_{50}$ (torr)</th>
<th>$n_{50}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Native</td>
<td>5.0 ± 0.1</td>
<td>3.0 ± 0.1</td>
</tr>
<tr>
<td>Native + 1.2 eq. IHP</td>
<td>28.4 ± 0.2</td>
<td>1.8 ± 0.1</td>
</tr>
<tr>
<td>Native + 2.3 eq. IHP</td>
<td>40.0 ± 0.4</td>
<td>1.8 ± 0.1</td>
</tr>
<tr>
<td>$\alpha_{99}$-fumaryl-$\alpha_{99}$, $\beta_2$</td>
<td>13.9 ± 0.3</td>
<td>2.6 ± 0.1</td>
</tr>
<tr>
<td>$\alpha_{99}$-fumaryl-$\alpha_{99}$, $\beta_2$ + 2.3 eq. IHP</td>
<td>47.3 ± 0.3</td>
<td>1.9 ± 0.1</td>
</tr>
<tr>
<td>$\alpha_2$, $\beta_{82}$-trimesyl-$\beta_{82}$</td>
<td>4.8 ± 0.2</td>
<td>2.4 ± 0.1</td>
</tr>
<tr>
<td>$\alpha_2$, $\beta_{82}$-trimesyl-$\beta_{82}$ + 1.2 eq. IHP</td>
<td>9.9 ± 0.2</td>
<td>2.2 ± 0.1</td>
</tr>
<tr>
<td>$\alpha_2$, $\beta_{82}$-trimesyl-$\beta_{82}$ + 2.3 eq. IHP</td>
<td>22.9 ± 0.2</td>
<td>2.4 ± 0.1</td>
</tr>
<tr>
<td>$\alpha_{99}$-fumaryl-$\alpha_{99}$, $\beta_{82}$-trimesyl-$\beta_{82}$</td>
<td>20.6 ± 0.3</td>
<td>1.7 ± 0.1</td>
</tr>
<tr>
<td>$\alpha_{99}$-fumaryl-$\alpha_{99}$, $\beta_{82}$-trimesyl-$\beta_{82}$ + 2.3 eq. IHP</td>
<td>23.6 ± 0.4</td>
<td>1.8 ± 0.1</td>
</tr>
<tr>
<td>Bis-tetramer</td>
<td>17.4 ± 0.5</td>
<td>1.5 ± 0.1</td>
</tr>
</tbody>
</table>
(as in α2, β82-trimesyl-β82) and installing multiple cross-links (as in α99-fumaryl-α99, β82-trimesyl-β82) decreases the allosteric effect of IHP (Table 1). This attenuation also occurs where the concentration of IHP decreases from 0.1 mM (2.3 eq./heme) to 0.05 mM (1.2 eq./heme).

CD spectra of both the bis-tetramer and α99-fumaryl-α99, β82-trimesyl-β82 were acquired in the 200–260 nm and 245–470 nm regions. The spectra of both α99-fumaryl-α99, β82-trimesyl-β2 and the bis-tetramer in the far UV region (Fig. 7) are consistent with the secondary structures of the proteins being well-maintained following chemical modification. A notable feature is the enhanced negative ellipticity at 220 nm relative to that for native Hb. This depression is observed where oxyHb is converted to deoxyHb.47 Variations in the vicinity of 285 nm reflect changes in the environment of aromatic residues at the αβ2 interface.47 For α99-fumaryl-α99, β82-trimesyl-β82 and the bis-tetramer, we observe negative ellipticity in this region and a notable red shift (Fig. 8). These features in the CD spectrum are present in the T-state deoxyHb.47 This presence of the T-state is consistent with the reduced oxygen affinity of the modified protein.

Cross-linking of native Hb improves the protein’s thermal stability, making it possible to separate cross-linked species from protein lacking cross-links.48 An additive effect from two cross-links could potentially increase thermal stability,49 creating a route for separation from singly cross-linked material. The minimum temperatures at which native and modified COHb species are fully denatured after 30 min are presented in Table 2. The criterion for indicating the complete denaturation of the protein is the absence of absorbance from the heme at 540 nm as the heme is released from denatured protein. Complete denaturation of a solution of native Hb results from heating a solution for 30 min at 79 °C. Denaturing the singly cross-linked species (α2, β82-trimesyl-β82) requires heating at 95 °C. These results indicate that an additional cross-link does not enhance stability. The degree of heat stability is consistent with heterogeneous bulk solution effects.48,50

We also investigated the thermal stability of the doubly cross-linked species in the presence of oxygen, comparing it to α99-fumaryl-α99 cross-linked Hb.51 Turbidity curves were acquired by slowly increasing the temperatures of solutions of the CO-forms of the modified proteins in solutions that were exposed to air while monitoring absorbance at 700 nm (Fig. 9). Hb in solution does not absorb at 700 nm but the formation of aggregates that occur upon unfolding decrease the transparency of the solutions.52 The doubly cross-linked and singly cross-linked species are essentially indistinguishable in terms of their thermal stability whereas the presence of a single cross-link increases overall stability of the protein relative to native Hb, even where auto-oxidation rates are comparable.

Discussion

It was recently reported that acetylation of amino groups in the α-subunits of Hb direct cross-linking to give β-subunit modification.55 We reasoned that a fumaryl bridge would serve the same function, with the added benefit of providing a defined product that is readily isolated in high yield.6 The precursor to the bis-tetramer would then be doubly cross-linked and could be correlated with earlier reports.53 It was not known whether
this doubly cross-linked hemoglobin could be prepared as efficiently, as the original report of a doubly cross-linked species describes an impure product. In that report, a native deoxygenated Hb (deoxyHb) is reacted with two eq. of trimesoyl tris(3,5-dibromosalicylate) (TTDS) followed by treatment with two eq. of bis(3,5-dibromosalicyl) fumarate (DBSF). The cross-linking of the β-subunits with TTDS is quantitative but the second cross-link from reaction with DBSF forms in lower yield at pH 9.0 (Scheme 4, Path B). At pH 7.4, the reaction is even less complete.

A logical alternative route to doubly cross-linked Hb reverses the order of reactions, introducing a cross-link between the α-subunits followed by reaction of the β-subunits (Scheme 4, Path A). The state of the protein as R or T can control access to reactive groups and thereby control the efficiency of cross-linking. In principle, the materials generated by the two sequences could have different structures and properties as the first cross-link defines the accessible conformational space for the second reaction. In the present study, we find that where the α-subunits are cross-linked ahead of the β-subunits, the doubly cross-linked species is acquired in a higher state of purity than by the original route, with the resulting modified proteins likely to be identical. However, since there are significant differences in the efficiency of the conversion by the two routes, the pathways are not interchangeable. In a formal sense, the results establish that the transition state for the second of the serial cross-linking processes is more closely associated with the reactant than with the product.

The results provide a practical outcome: serial modifications of proteins are not commutative: introduction of the first cross-link specifically limits the accessible conformational space of the modified protein compared to that of the native protein. Introduction of the second link occurs within the subset of conformations defined by the initial cross-link. By reversing the order of reaction, the initial product has access to conformations that would not have been available if the other link were already in place. In other words, the initial link excludes the existence of a subset of conformations and defines a new conformational space by restricting structural movement. Failure to cross-link the modified protein completely in the second step (Scheme 4, Path B) is an indication that the deoxy conformational space is not coincident with that of native deoxy Hb. This is consistent with the observations of the structures reported from crystallographic analysis. Interaction of the protein and the reagent within the α-subunits would be reduced. In contrast, Path A (Scheme 4) ensures efficient installation of the second cross-link because the α-α linked fumaryl bis-amide leads to a conformation that is very similar to that of the native protein.

Our results reveal the utility of cross-linking as a probe for detection of otherwise inaccessible conformations, delineating inter-subunit effects and specific residues in dynamic environments. The use of cross-linkers to probe conformational change is especially useful when X-ray crystallography proves to be inadequate because the conformation of the crystallized protein is different from that in solution. Introduction of
two cross-links between subunits in multi-subunit proteins is potentially a general approach to distinguish specific effects of a cross-link as it defines the accessible space for the second link. These general tactics extend to the specific installation of the azido moiety on the cross-link within the β-subunits (Scheme 2). Azides located within the α-subunits of Hb are unreactive and are likely to be within a relatively inaccessible region of the protein. Therefore, the presence of an azide alone is not sufficient to assure its participation as is the case for the reaction of small molecules in solution.

Although azide 2 is more selective in our comparative analysis, its reaction with cross-linked Hb does not go to completion without over-modification (Fig. 3). In order to obtain the bis-tetramer in a pure state, the reaction was stopped after 50% of the protein was modified (Fig. 4). It was this mixture of singly and doubly cross-linked protein that was submitted to the final coupling stage. The non-quantitative CuAAC yield of 17% (Fig. 5) is unexpected since analogous reactions with modified Hb have proceeded efficiently. However, conducting CuAAC on modified Hb in the deoxygen state dramatically increases the yield of the coupled product, suggesting that coupling is controlled to some extent by the accessibility of the azide within the doubly cross-linked system.

To deduce whether this deoxyHb effect is specific to CuAAC, we set up a nucleophilic substitution reaction analogous to that of Yang et al., using the non-hydrolyzed form of α99-fumaryl-α99, β82-trimesyl-β82 as the substrate. We observed that even this substitution occurs more readily when the protein is deoxygenated, confirming that the issue was one of accessibility (see ESI, Fig. 19 vs. 20f). This result illustrates the potential utility of CuAAC as a probe of protein conformation. Furthermore, the switchable nature of the doubly cross-linked framework makes it an interesting candidate for drug conjugation. It would be interesting to know if disulfide bond formation, under the CuAAC conditions competes with the protein-coupling process. So far, we can conclude that a well-defined product results from the coupling step with acceptable efficiency in the overall process.

We find that both the pure bis-tetramer and the doubly cross-linked species (α99-fumaryl-α99, β82-trimesyl-β82) possess low oxygen affinity (Fig. 6, Table 1). Circular dichroism (CD) spectra informed us that the source of this low oxygen affinity is associated with enhanced T-state character (Fig. 7 and 8). The oxygen binding properties of the coupled and uncoupled proteins are slightly different because the two Hb units of the bis-tetramer are close enough to communicate. Our bis-tetramer is the first with an oxygen affinity close to that of Hb within red cells (P50 = 28.0 torr). The bis-tetramer as an HBOC component would not circulate in a DPG-rich environment as occurs within the red cell; therefore, an intrinsically low oxygen affinity is a valuable characteristic for efficient delivery of oxygen to the tissues for a red cell replacement. This holds especially true for doubly cross-linked hemoglobin as the modified protein demonstrates nearly complete insensitivity to allosteric regulation by inositol hexaphosphate (IHP) (Table 1).

Thermal stability data for the modified proteins clearly show that not all cross-links impart equal stability: α99-fumaryl-α99, β2 and α2, β82-trimesyl-β82 responded differently to the conditions. Yang et al. showed that α99-fumaryl-α99, β2 metHb and α2, β82-fumaryl-β82 metHb are equally stable. Comparison with those results suggests that stability depends on the structure of the link as well as the sites that it spans. Furthermore, additional cross-links do not necessarily impart cumulative stability, as observed by Ueda et al. Despite its dual cross-links, our doubly cross-linked Hb fully denatures at the same temperature as does α2, β82-trimesyl-β82. In principle, with different melting temperatures for fumaryl (singly) cross-linked Hb and doubly cross-linked Hb, separation for the purpose of purification might be possible. However, we were unable to achieve separation of cross-linked species by heat treatment, although this does remove unmodified proteins. We suspect that heterogeneity changes the melting points of the individual components, as observed by Yang and Olsen and justified by Despa and coworkers.

The relative thermal stabilities of the four COHb species are different under aerobic conditions, with α99-fumaryl-α99, β2 and α99-fumaryl-α99, β82-trimesyl-β82 demonstrating equal stability (Fig. 9). Rapid auto-oxidation is a characteristic of fumaryl cross-linked oxyHb species. However, in light of the stability that modified COHb possesses under aerobic conditions, we recognize its potential for carbonmonoxHb-based applications.

Conclusions

We have shown that functional hemoglobin bis-tetramers are accessible by an efficient three-step procedure: protect, functionalize and connect by a “click” reaction. Double cross-linking, which is a non-commutative process, proceeds with maximum efficiency by directed sequential cross-linking, producing regioselective modification of the β-subunits with an azido moiety poised for solubility-directed coupling by reaction with bis-alkynes. This production of a pure product is a step forward in bis-tetramer design and large scalability, which has been a particular challenge for the alternative of producing protein expression from bacterial sources. Finally, the oxygenation properties of the cross-linked hemoglobin bis-tetramer include cooperativity and oxygen affinity that would be suitable for circulation as an alternative to red cells in transfusions, while the extended structure can be expected to prevent extravasation that would produce vasoactivity.

Acknowledgements

We thank the Canadian Blood Services for support through an operating grant.
References