The effect of reactive oxygen and nitrogen species on the structure of cytoglobin: A potential tumor suppressor

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Abstract

Many current anti-cancer therapies rely on increasing the intracellular reactive oxygen and nitrogen species (RONS) contents with the aim to induce irreparable damage, which subsequently results in tumor cell death. A novel tool in cancer therapy is the use of cold atmospheric plasma (CAP), which has been found to be very effective in the treatment of many different cancer cell types in vitro as well as in vivo, mainly through the vast generation of RONS. One of the key determinants of the cell's fate will be the interaction of RONS, generated by CAP, with important proteins, i.e. redox-regulatory proteins. One such protein is cytoglobin (CYGB), a recently discovered globin proposed to be involved in the protection of the cell against oxidative stress. In this study, the effect of plasma-produced RONS on CYGB was investigated through the treatment of CYGB with CAP for different treatment times. Spectroscopic analysis of CYGB showed that although chemical modifications occur, its secondary structure remains intact. Mass spectrometry experiments identified these modifications as oxidations of mainly sulfur-containing and aromatic amino acids. With longer treatment time, the treatment was also found to induce nitration of the heme. Furthermore, the two surface-exposed cysteine residues of CYGB were oxidized upon treatment, leading to the formation of intermolecular disulfide bridges, and potentially also intramolecular disulfide bridges. In addition, molecular dynamics and docking simulations confirmed, and further show, that the formation of an intramolecular disulfide bond, due to oxidative conditions, affects the CYGB 3D structure, thereby opening the access to the heme group, through gate functioning of His117. Altogether, the results obtained in this study (1) show that plasma-produced RONS can extensively oxidize proteins and (2) that the oxidation status of two redox-active cysteines lead to different conformations of CYGB.

1. Introduction

The use of cold atmospheric plasma (CAP) has attracted a lot of attention over the past few years and has already found many biomedical applications, including wound healing [1], sterilization [2], biofilm removal [3], blood coagulation [4], and more recently cancer...
2.1. Expression and purification of recombinant CYGB

The expression of recombinant wild-type CYGB was performed as previously described [31]. Briefly, human cDNA of CYGB was cloned in the pET3a expression vector. Inclusion bodies were solubilized in 6 M guanidinium hydrochloride and insoluble material was eliminated. Afterwards, CYGB was reconstructed by adding free hemin and dialyzed overnight. The samples were then purified with an AKTA purifier system on a Hitrap DEAE-Sepharose column (both GE Healthcare, Life Sciences). The concentrated material was loaded onto a Sephacryl S-200 High resolution column (GE Healthcare). The final purity of the pooled CYGB was checked by absorbance spectra and SDS-PAGE.

2.2. CAP treatment of recombinant CYGB

A recombinant CYGB stock solution (19.3 mg/mL; 901 µM) was diluted either in deionized water or 50 mM tris-buffer (pH 8) to a final concentration of 20 µM. Plasma treatment was performed on 500 µL aliquots in a 24-well plate (on ice) for different time points (30 s, 1 min, 3 min, 5 min, 10 min) using the kINPen IND plasma jet (neoplas GmbH). The applied gas flow rate and gap distance were set at 31 L/min, and 10 mm, respectively, for all treatments. Argon was used as feed gas. After treatment, the evaporated volume of the aliquots was calculated and replaced by either water or tris-buffer. The treated samples were immediately put on ice before being handled for further analysis.

The pH of both used solvents was measured before and after treatment with CAP at different lengths of time to assess the effect of CAP treatment on the pH of the samples. The pH was measured with the pHenomel pH 1100H pH-meter (VWR).

2.3. Mass spectrometry

2.3.1. Native and denatured MS

Before native MS analysis, 50 µL of each sample was buffer exchanged using a Micro Bio-Spin P-6 Gel column (Bio-Rad) equilibrated with 100 mM aqueous ammonium acetate (pH 6.8), according to the manufacturer's protocol. Afterwards, the samples were further diluted to a concentration of 10 µM.

3 µL of sample solution were loaded into an in-house produced, gold coated nano-electrospray ionization (ESI) glass capillaries and mounted in the needle holder of the instrument. MS spectra were acquired on a Synapt G2 HDMS instrument (Waters, Wilmslow, UK) using the following instrument parameters: 1.6 kV spray capillary voltage; 100 V sampling cone; 3 V extractor cone; 6.3 mbar backing pressure; 15 V and 5 V collision energy in the trap (2.4 × 10⁻² mbar) and transfer (2.5 × 10⁻² mbar) cell, respectively; 45 V trap DC bias; 180 mL/min and 90 mL/min He and IMS gas flow, respectively. The ions were separated in the ion mobility (IM) cell, using nitrogen gas and a wave velocity of 750 m/s and a wave height of 40 V.

The samples were also measured under denaturing conditions by diluting the sample 1:1 with acetonitrile containing 1% formic acid. The MS parameters for measuring the denatured samples were: 1.4 kV spray capillary voltage; 25 V sampling cone; 1 V extractor cone; 2.7 mbar backing pressure; and 5 V collision energy in trap (2.00 × 10⁻² mbar) and transfer (2.2 × 10⁻² mbar) cell. Settings for IM separation were the same as under native conditions. Additional native and denaturing measurements on the tris-diluted CYGB samples were also performed in 10 mM dithiothreitol (DTT).

2.3.2. In solution protein digestion and MS

To detect oxidative modifications at amino acid level, a tryptic digest was performed on each treated sample following our own protocol. Briefly, 40 µg of plasma-treated sample was used. 1% RapiGest SF Surfactant (Waters) was added to a final concentration of 0.1% before incubation for 5 min at 100 °C. 200 mM TCEP (tris (2-carboxyethyl) phosphine) was added to reduce the sample. After 1 h incubation at 55 °C, 375 mM 2-iodocetamide was added and further incubated for 30 min in the dark to alkylate the sample. After alkylation, the sample was precipitated with pre-chilled acetone and incubated overnight at
− 20 °C. The acetone-precipitated pellet was resuspended in 100 mM triethylammonium bicarbonate to a final concentration of 1 μg/μL before MS grade trypsin protease (Thermo Scientific) was added to a final protease-protein ratio of 1:40 (w/w). Tryptic digest was performed overnight at 37 °C. The enzymatic reaction was stopped by freezing the samples at − 20 °C. The digested samples were measured using liquid chromatography − ion trap tandem mass spectrometry (LC-MS/MS).

The peptide mixture was separated by reversed phase C18 (RP-C18) liquid chromatography on a nanoAcquity UPLC system (Waters) using a Symmetry C18 trap column (5 μm particle size; 280 μm × 20 mm) connected to a BEH C18 analytical column (1.7 μm particle size; 100 μm × 100 mm) (Waters). Before loading, each sample was dissolved in 10 μL mobile phase A (0.1% formic acid in 100% water) and in total 1 μg of protein digest was loaded onto the column. A linear gradient of mobile phase B (0.1% formic acid in 100% acetonitrile) from 5% to 55% in 50 min followed by a steep increase to 100% mobile phase B in 3 min was used at a flow rate of 400 nL/min. The nano-UPLC system was coupled online with the mass spectrometer using a PicoTip Emitter (New objective) and a nanospray ion source (Thermo Scientific). The LTQ ion trap mass spectrometer (Thermo Scientific) was set up in MS mode, where a full MS1 scan (300–2000 m/z) was followed by collision-induced dissociation (CID) and a subsequent MS2 scan (100–2000 m/z). All scans were acquired in the linear ion trap with an automatic gain control (AGC) set at 3 × 10⁴ ions for MS1 and 1 × 10⁴ ions for MS2 scans. The normalized collision energy was 31% in CID, and a dynamic exclusion list of 30 s for data dependent acquisition was used. Between every sample a blank run was performed to exclude any possible carry-over effect.

Proteome Discoverer (2.1 SP1) software (Thermo Scientific) was used to perform database searching against the UniProt reference human database using both the Sequest and Mascot algorithms. Following settings were applied: precursor mass tolerance of 500 ppm, fragment mass tolerance of 0.50 Da. Trypsin was specified as digesting enzyme and 2 missed cleavages were allowed. Cysteine carbamidomethylation was defined as fixed modification and oxidations (on residues M/Y/D/F/H/W/N and P) were set as variable modifications. The results were filtered using the following settings: only medium and high confident peptides with a global FDR < 5% based on a target-decoy approach [16] and first ranked peptides were included in the results.

2.4. UV−vis spectroscopy

UV−vis circular dichroism (CD) and absorption spectra were collected using a Chirascan Plus CD (LAAPD solid-state detector) spectrometer (Applied Photophysics Ltd.) operating with a bandwidth of 1 nm, a step size of 1 nm, a scanning speed of 1 nm/s, a sample temperature of 20 °C and a cuvette with a path length of 0.5 mm or 2.0 mm. The spectral region sensitive to secondary structure 180−280 nm was measured using a 0.5 mm path length and the spectral region 225–600 nm comprising the Soret band and Q-bands was measured with a path length of 2.0 mm.

2.5. Resonance Raman spectroscopy

Resonance Raman (RR) spectra were acquired using a Dilor XY-800 spectrometer in low-dispersion mode using a liquid N₂-cooled CCD detector. The excitation source was a Spectra Physics (Mountain View) BeamLok 2060 Krypton laser operating at 413.1 nm. The spectra were recorded at room temperature and the protein solutions were magnetically stirred at 500 rpm in order to avoid local heating and photochemical decomposition. The slit width used during the experiments was 200 μm. In general, 10–12 spectra were acquired with an integration time of 240 s each. Spikes due to cosmic rays were removed by omitting the highest and lowest data points for each frequency and by averaging the remaining values. Typical sample concentrations were in the order of 20 μM.

2.6. Computer simulations (molecular dynamics and docking)

Molecular dynamics (MD) and docking simulations were performed in order to elucidate the functionality of monomeric CYGB with and without an intramolecular disulfide bridge, at the molecular level. The simulations were carried out using the GROMACS [32] program package (version 5.1), applying the GROMOS 45a3 force field [33]. The coordinate file (with the initial configuration) of monomeric CYGB was obtained from the Protein Data Bank (PDB ID: 1VSH) [34]. The structure of CYGB without an initial disulfide bond, i.e. CYGBSH−SH, was placed in a dodecagonal box, with size chosen in such a way that there initially was 1.0 nm buffer distance between the protein and the boundary of the simulation box. This was done in order to avoid long distance interactions (i.e. Coulomb and van der Waals interactions) between CYGBSH−SH and its periodic images. The box was then filled with water molecules surrounding the CYGBSH−SH molecule, employing the SPC water model [35], and sodium counter ions were added to neutralize the system. Periodic boundary conditions were applied in all directions.

In order to study the effect of the above mentioned disulfide bond on the conformation of CYGB, this disulfide linkage was manually created using Avogadro software [36]. Subsequently, energy minimization was applied to optimize the modified structure, using the steepest descent algorithm. The measured length of the disulfide bond after the energy minimization was 2.05 Å, which is exactly the same as the disulfide bond length given in literature [37,38]. The abovementioned MD procedure was then repeated (i.e., the creation of the box and addition of water molecules, as well as counter ions) to create the simulation system with CYGB containing a disulfide bridge, i.e. CYGBS−S.

Thus, two model systems were used (i.e., CYGBSH−SH and CYGBS−S) covered with water as a starting structure for the calculations. The systems were minimized in three steps, by applying three optimization algorithms, i.e., steepest descent, conjugate gradient and low-memory Broyden-Fletcher-Goldfarb-Shanno (L-BFGS) [39]. Subsequently, a 500 ps equilibration run was performed employing the NVT ensemble (i.e., a system with constant number of particles N, volume V and temperature T). Next, a 150 ns production run was performed using NPT dynamics (i.e., a system with constant number of particles N, pressure P and temperature T). The NPT simulations were carried out for 300 K and 1 bar pressure, employing the velocity-rescaling thermostat with a coupling constant of 0.1 ps and the isotropic Parrinello-Rahman barostat [40,41], with a compressibility and coupling constant of 4.5 × 10⁻⁵ bar⁻¹ and 2 ps, respectively. A 1.4 nm cut-off distance was applied for the van der Waals interactions. The long range electrostatic interactions are described by the particle mesh Ewald (PME) method [42], again using 1.4 nm cut-off for the real-space interactions in combination with a 0.15 nm spaced-grid for the reciprocal-space interactions.

In the docking simulations, binding hot spots of equilibrated CYGBSH−SH and CYGBS−S were predicted using the FTMMap family of the web server [43]. FTMMap samples the positions of 16 small probing organic molecules and identifies potential ligand binding regions of the protein based on the lowest binding free energy. These individual probing molecules may bind at various locations of the protein, and thus their clusters specify binding hot spots (so-called binding pockets) of the protein. The probing molecules used in the FTMMap server are: ethane, ethanol, isopropanol, isobutanol, acetonitrile, methanamine, N,N-dimethylformamide dimethyl ether, benzaldehyde, benzene, cyclohexane, phenol, acetamide, acetone, acetaldehyde and urea. They vary in shape, size and polarity, and can be considered as drug-like molecules [43]. The hot spot of the protein was identified by means of a mapping algorithm that consisted of the following four steps: (1) each probe molecule samples on a grid around the fixed protein. The energy function considers van der Waals energy with attractive and repulsive contributions and an electrostatic as well as solvation term characterized by the Poisson−Boltzmann continuum model. The algorithm chooses
the 2000 best positions of each probing molecule and proceeds to the next stage. (2) These best positions of the probe molecules with the protein complex are energy minimized using the CHARMM potential [44] with an analytical continuum electrostatics model [45]. During the energy minimization run, the protein atoms are fixed while the probing molecules are free to move. (3) The minimized structures are clustered within a 4 Å radius, considering the lowest energy structure. For each probe molecule, six clusters are retained with the lowest energy, which are ranked by their Boltzmann averaged energies. (4) The clusters of the 16 different probe molecules are then clustered into consensus clusters. This clustering stage defines the center of mass for each cluster probe. The highest ranked consensus cluster is selected, using the distance between the centers of mass of the cluster centers as measure for the distance and 4 Å as the clustering radius. VMD [46], and Pymol [47] visualizing tools were used for preparation of the images.

3. Results

3.1. Spectroscopic changes upon plasma treatment of CYGB

As an initial study of the structural effects of plasma treatment on CYGB, UV–vis absorption and CD spectra were recorded and are presented in Fig. 1. The UV–vis absorption spectrum of the untreated sample contains a Soret (γ) band at 417 nm, and β and α bands in the visible region of the spectrum at 535 and 565 nm, respectively, indicating that the heme iron is in a ferric low-spin HisF8-Fe(III)-HisE7 hexa-coordination state [48]. The treatment of CYGB with plasma induces significant spectroscopic changes. Most notably is the time-dependent disappearance of the Soret band, but also the steady decrease of the intensity of the β and α bands with longer treatment times. When comparing the spectra, acquired in the two solvents that were employed, both similarities and differences can be observed. For both, a time-dependent decrease in intensity of the Soret band can be observed. However, the Soret band of the samples diluted in tris-buffer is still clearly visible after 5 min exposure to plasma. Even after 10 min, there is still positive intensity discernible at 417 nm in tris-buffer, while this is not the case in water. Even more so, in water, a new peak appears at 387 nm after 5 min of treatment, indicating that free heme is present in the solution [49]. In tris-buffer, the same peak (383 nm) only starts to appear after 10 min treatment, again indicating the presence of free heme.

The CD spectra of CYGB before and after treatment are shown in Fig. 1C and D. All spectra display the typical double minima at 208 and 222 nm, characteristic for protein structures that are predominantly α-helical [50]. Plasma treatment, besides altering the intensity of the double minima, does not seem to influence the spectra significantly, suggesting that the secondary structure remains largely unaltered by the plasma treatment. As the \( \Theta_{222}/\Theta_{208} \) ratio is sensitive to structural changes, the ratio was plotted over treatment time (Fig. 1E) [50]. The plot shows that, with both solvents, the ratio drops depending on the treatment time. The change in the ratio, although very minor, indicates a slight unfolding of the protein.

pH measurements indicated that the pH of the samples dissolved in water significantly drops after plasma treatment, whereas with tris-buffer, no change in pH could be observed (Fig. S3). As the change in pH of water could have a profound effect on CYGB, and thus on the
broadening and a shift to higher m/z values, but also to a small increase in between the monomers (Fig. 3B), highlighting the presence of dimeric species that still remain after denaturation. This indicates that the dimerization occurs via covalent bonds. In Fig. 3(A), the m/z region between 600 and 700 is shown with the peak corresponding to the free heme group (616 m/z). A plasma treatment time of up to 3 min does not show much effect; however after 5 min a new peak appears at 661 m/z, indicating the binding of a molecule to the heme group (while also replacing a hydrogen atom) which we interpret as \( NO_2 \) (+45 Da).

We note that dimers are already present in the native spectra (Fig. 2) without plasma treatment, but almost entirely absent in the denatured spectrum (Fig. 3A). In order to investigate whether pre-existing noncovalent CYGB dimers are converted to covalent, disulfide-bound dimers under the influence of plasma oxidation, measurements in tris-buffer, under native conditions, were repeated after adding 10 mM of the reducing DTT to the samples. Fig. 4 shows the full spectra and zoomed in section of the monomeric + 9 peak (inset, highlighted in blue). As can be observed, addition of DTT leads to a loss of the dimeric species, indicating that dimerization is stabilized by disulfide bonds. Furthermore, the addition of DTT also reduces the plasma-induced oxidations. This is clearly visible when comparing the monomeric + 9 peaks in Fig. 4 with the ones depicted in Fig. 2, under otherwise identical conditions. The peak shifts after 5 and 10 min of plasma treatment again indicate binding of an additional ligand to the heme group.

### 3.2. Mass spectrometry experiments

Following the UV–vis and CD measurements, mass spectrometry (under native and denaturing conditions) was used to assess changes in the tertiary structure of CYGB samples as well as chemical modifications after treatment with plasma.

#### 3.2.1. Native MS

Fig. 2 shows the native mass spectra of untreated and treated CYGB. The spectrum of the untreated sample ("control") clearly exhibits a charge state distribution for monomeric and dimeric species, with the monomeric species accounting for the majority of the signal (see also Fig. S1). As can be seen, plasma treatment not only leads to peak broadening and a shift to higher m/z values, but also to a small increase of dimeric species. The peak shift and broadening is highlighted (in blue) for the monomeric + 9 species in the m/z region of 2400–2600 (Fig. 2, inset). Here, it can be seen that plasma treatment of only 30 s already causes a mass shift to higher m/z values and peak broadening, which indicates an unresolved distribution of oxidized species. Not much change is observed up to 3 min treatment of CYGB. However, a major peak shift is visible after 5 and 10 min of treatment, which is probably caused by binding of an additional ligand to the heme (see further, Fig. 3). In addition, a small fraction of apo-CYGB is also present, in agreement with UV–vis data that showed heme loss after 10 min as well.

### 3.2.2. Denatured MS

Plasma-treated samples were also measured under denaturing conditions, in order to determine if the observed dimers were noncovalent complexes, or covalently linked, e.g. via disulfide bridges (Fig. 3). As can be seen in Fig. 3B (bottom panel, “control”) and Fig. S2, unfolding leads to a much broader charge state distribution for the monomer that no longer contains the heme group. The spectra also show minor peaks in between the monomers (Fig. 3B), highlighting the presence of dimeric species that still remain after denaturation. This indicates that the dimerization occurs via covalent bonds. In Fig. 3(A), the m/z region between 600 and 700 is shown with the peak corresponding to the free heme group (616 m/z). A plasma treatment time of up to 3 min does not show much effect; however after 5 min a new peak appears at 661 m/z, indicating the binding of a molecule to the heme group (while also replacing a hydrogen atom) which we interpret as \( NO_2 \) (+45 Da).

Next, we aimed to map plasma-induced oxidative modification sites using LC-orbitrap MS/MS of trypsin-digested samples. Table 1 gives an overview of different types of amino acids that were found to be oxidized after plasma treatment of a specific duration. CAP was able to oxidize all different amino acids that were selected for the analysis (Table 1), with some seeming to be more readily oxidized than others. Of the investigated amino acids, methionine, tryptophan, and tyrosine were the three found to be preferably oxidized, as reported previously [51]. Surprisingly, no clear treatment-time-dependent increase in the number of oxidized amino acids could be observed, indicating that oxidation may occur rapidly. In the suppl. Table S2, the complete amino acid sequence of CYGB is given together with the specific location of each identified modification, for each exposure time. Table S2 shows that some of the amino acids, at certain locations, were never oxidized. For example, two of the six methionine and two of the five tyrosine (Met66, Met72, Tyr59, and Tyr123) were not found to be oxidized, under any condition.

Table 2 gives the total number of identified peptide fragments (so-called ‘Peptide Spectrum Matches’, PSM), together with the number of oxidized PSMs. Here, a clear treatment-time-dependent increase in oxidized peptides can be observed, illustrating the increasing oxidation potential with increasing treatment time. It is also noticeable (Tables 1, 2 & S2) that in the control samples some amino acids (M, Y and F) are also oxidized. This oxidation is likely due to the exposure of the sample to the surrounding atmosphere, or may occur during electrospray.

### 3.4. MD and docking simulations

CYGB possesses two cysteine residues (Cys38 and Cys83), which are known to be able to form disulfide bridges, either inter- or intramolecularly [25,26]. As CYGB is proposed to be a monomer in vivo [52], and the formation of an intramolecular disulfide bridge can
influence the activity of CYGB, we chose to investigate the effect of the formation of an intramolecular disulfide bridge in between the cysteine amino acids of monomeric CYGB (i.e., CYGBS-S) with MD simulations. We have shown experimentally that an intermolecular disulfide bridge is present in the dimeric fraction. However, we can also speculate that the monomeric fraction may contain an intramolecular disulfide bridge.

**Fig. 3.** Denaturing MS of CYGB. ESI spectra of 10 μM CYGB (diluted 1:1 in acetonitrile containing 1% formic acid) after different duration of plasma treatment (20 μM CYGB in 50 mM tris-buffer). (A.): Zoomed in m/z region from 600 to 700. The peak at 616.3440 m/z represents the free heme group. 5–10 min of treatment induces binding of a molecule, i.e., NO₂ to the heme group (661.3398 m/z). (B.): Full spectrum ranging from 800 to 4000 m/z showing the apo-form of the protein. The appearance of smaller peaks (asterisks) in between the monomers highlights the presence of covalent dimers in the sample. The theoretical mass of CYGB in the apo-form (without heme) is 21 404.62 Da compared to the experimental mass of 21 407.29 ± 4.16 Da in the control.

**Fig. 4.** Native MS after reduction with DTT. Native MS spectra of 10 μM CYGB in 100 mM ammonium acetate after adding 10 mM DTT. Monomeric +9 is highlighted. Inset: the zoomed in section of the monomeric +9 peak. Addition of DTT leads to a loss of the dimers via reduction of disulfide bridges. DTT also seems to reduce the number of oxidations (narrowing of the peaks). See also Fig. 2 for comparison.

**Table 1**

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The monomeric fraction may contain an intramolecular disulfide bridge.

**Fig. 5.** Native MS spectra of 10 μM CYGB in 100 mM ammonium acetate after adding 10 mM DTT. Monomeric +9 is highlighted. Inset: the zoomed in section of the monomeric +9 peak. Addition of DTT leads to a loss of the dimers via reduction of disulfide bridges. DTT also seems to reduce the number of oxidations (narrowing of the peaks). See also Fig. 2 for comparison.
Table 2

Oxidative modifications of CYGB. A list of the number of oxidized peptide fragments before and after plasma treatment. The number of Peptide Spectrum Matches (PSM) is the total number of identified peptide spectra matched for the protein. Samples were measured with Ion trap LC-MS/MS and spectra were analyzed with Mascot and Sequest algorithms.

<table>
<thead>
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<th>Sample</th>
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<th>#Oxidation PSM</th>
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<td>138</td>
<td>54</td>
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Relative to CYGBSH (cf. the yellow and green dashed lines). In order to calculate the distances between the helices, residues Ala82 and Ala88 in the F-helix and Ala107 and Ala112 in the E-helix were chosen. These amino acids are positioned at the end of the black dashed lines but they are not shown in the figure for the sake of clarity. The calculated distances between these amino acids located in the E- and F-helices are also given in Table S1, indicating the minor change in the modified CYGBSH. It is also obvious from Fig. 5 that there is a negligible shift of the heme group in the modified CYGBSH molecule compared to native CYGBSH. Hence, the formation of the disulfide bridge between the cysteine residues does not alter the local orientation of the heme group and imidazole planes of the histidines (i.e., His81, His113) bound to the heme: the imidazole planes do not rotate versus each other. This was also reported in literature by experimental investigations [53].

Comparing the consecutive MD frames of both monomeric CYGBSH and CYGBS indicates that the His117 residue located nearby the heme group (see Fig. 6) substantially changes its orientation in CYGBS. Indeed, the calculated distance between the heme iron and one of the nitrogen atoms of His117 (located in the imidazole ring) almost doubled in CYGBS (see Table S1), which is due to the conformational change caused by the intramolecular disulfide bond. Besides the change in orientation of His117, it is noticeable that the propionate group of the heme closest to His117 flips its position upon disulfide bond formation (facing downward in CYGBS, see Fig. 6(b) and (d)). To get more insight into the structural changes caused by the intramolecular disulfide bond, the binding hot spots of both CYGBSH and CYGBS were investigated as described in Section 2.

10 CYGB structures with 1 ns time interval were extracted using the last 10 ns trajectory of the equilibrated (so-called production) run. These structures were employed to examine the structural dependencies of the binding hot spots of CYGB. Here, only one of the results out of ten are presented, as the other nine results were very similar. Three different pockets (with lowest binding energies) were obtained in both CYGBSH and CYGBS, which are depicted as meshed regions in Fig. 6. The other two meshed regions are not shown as we are focusing on the catalytic site of CYGB (i.e., the heme pocket). As mentioned above, the formation of a disulfide bond between the cysteine residues leads to a change of the CYGB conformation, which results in a doubling of the distance between the highly localized His117 residue and the heme group in CYGBS (see Table S1 and cf. Fig. 6(a) and (c) or Fig. 6(b) and (d)). Accordingly, this results in the heme group to become more accessible to the probing molecules (see overlap between heme group and meshed regions in Fig. 6(c) and (d)). In addition, the flipping of the position of the heme propionate upon disulfide formation further contributes to the accessibility of the heme pocket (Fig. 6(b) and (d)). Our binding hot spot analysis indeed shows that in the case of CYGBS the binding pocket is located around the heme group, whereas in CYGBSH it is found with limited access to the heme group, i.e., around the helices (cf. Fig. 6(a) and (c), or Fig. 6(b) and (d)). This again confirms that in CYGBSH the access to the heme group is more limited than in CYGBS.

4. Discussion
In this study, it was shown that the treatment of recombinant CYGB with CAP resulted in chemical modifications of the protein. The spectral characteristics described for the untreated recombinant CYGB are identical to the ones described previously, showing that the protein is in the hexacoordinated ferric form [48]. Upon plasma treatment, spectral changes occur, which appear to be dependent on the duration of the treatment. These changes were more prominent when water was used as a solvent than in tris-buffer. This suggests that in water, the effect of plasma, which is mainly through reactive species production, is greater than in tris-buffer. However, there is one main difference between the two solvents that were employed: deionized water has no buffering capacity, whereas tris-buffer does. As plasma treatment induces many different reactive species (i.e. \( \text{OH}^-, \text{H}_2\text{O}_2^-, \text{NO}^-, \text{O}_2^-, \text{ONOO}^-, \text{NO}_2^-\)), these species can contribute to the acidification of liquids. Indeed, pH measurements highlighted that plasma treatment induces a significant drop in the pH of deionized water, whereas no such effect could be observed with tris-buffer (Fig. S2). Thus, with water, there is a secondary effect of plasma treatment, through the acidification of the sample. Low pH can, depending on the protein, lead to protein denaturation and aggregation. In Fig. 1(A), after 5 min treatment, the Soret peak at 415 nm has almost completely disappeared, and a new peak has emerged at 387 nm, indicating free heme in the solution [49]. Additionally, the ultracentrifugation of the 5 min treated sample showed a clear brown precipitate, indicating the aggregation and denaturation of the sample upon plasma treatment. In tris-buffer, on the other hand, no clear precipitate after ultra-centrifugation of the 5 min treated sample could be observed, yet still, clear spectral changes were observed. Altogether, this confirms that in water, plasma treatment acidifies the solution, leading to structural changes in the protein, and the eventual (partial) denaturation of the protein sample. This has some implications for the further interpretation of the results from this study. As the main aim of this study was to investigate the effects of plasma-generated RONS on CYGB, the results obtained from treating CYGB with plasma in tris-buffer are of higher relevance and will be discussed further in more detail (see Suppl. Figs. S6 & S7 for MS spectra in water).

From the absorption spectra of the samples treated in tris-buffer, it can be concluded that plasma induces a decrease in intensity of the Soret band at 417 nm. This could indicate two things: the first option would be that the protein is continuously degrading more with longer treatment times, with the eventual, almost complete denaturation of the
protein at 10 min treatment time. Although the appearance of a peak at 383 nm (after 10 min treatment) indicates the leaking of the heme out of the protein (and thus denaturing of the protein), CD and MS data (Figs. 1E and 2) however shows that the largest fraction of the protein is still folded and contains the heme group. The slight unfolding we see in the CD-spectra may be explained by the MD data where it is shown that the formation of an intramolecular disulfide bridge in CYGBs-S induces a minor shift of the E- and F-helices, relative to CYGBSH (see Table S1 & Fig. 5). Furthermore, measurement of the drift time of the monomeric (+9 peak) species showed that there was no significant alteration of the collision cross section (CCS) of the treated protein (Fig. S4), indicating that the three-dimensional shape, i.e. the tertiary structure, is not significantly affected upon plasma treatment, and thus excluding this option.

The second option would be that plasma treatment induces specific chemical modifications in the protein that lead to the observed spectral changes. The Soret band is emblematic for heme-containing proteins such as globins and various cytochromes, and arises primarily due to π-π* transitions in the heme group [55]. Binding of molecules to the heme structure or close environment could affect the (electronic) structure and lead to a decrease and/or shift of the Soret peak. From the MS data, it is clear that there is indeed covalent modification of CYGB upon plasma treatment. In Fig. 2, it can be seen that there is a peak shift and broadening, indicating that plasma treatment induces numerous oxidations, either at specific amino acids, or at the heme group. Moreover, the MS spectra of the denatured sample (Fig. 3) clearly show that, with 5 or 10 min treatment, there is a binding of a molecule, most likely NO2, to the protein bound heme group. Binding of NO2 has already been shown to occur in vitro in other globins, i.e. myoglobin [56], hemoglobin [57], and more recently leghemoglobin [58], by exposing the proteins to excess NaN02. In these studies, it was shown that the heme nitration originated from the substitution of a hydrogen atom by NO2 from one of the vinyl groups. The subsequent formation of a nitrovinyl group results in a modified visible spectrum (a Soret band with lower intensity) and a heme group with m/z 661, similar to the results that were obtained in this study. Addition of a 1000-fold excess of NaNO2 to CYGB (data not shown) led to the same modified spectra observed after plasma treatment of 5 and 10 min, confirming the nitration of the heme. Furthermore, resonance Raman (RR) experiments further confirmed the binding of NO2 to the heme (Fig. S5). The high-frequency region of the RR spectra revealed the appearance of a strong band at ~1325 cm⁻¹, which is specific for a nitroaromatic group [59–61], again indicating the binding of NO2 to the vinyl groups of the heme. Plasma treatment of liquids does indeed yield the formation of NO2 [62–64]. Thus, upon plasma treatment for 5 min and longer, NO2, and most probably other related nitrogen species, are sufficiently produced in order to induce the binding of NO2 to the heme. Furthermore, the fact that NO2 can bind to the CYGB heme could implicate that it also performs a similar biological role, i.e. the involvement of CYGB in NO homeostasis. Previous reports already proposed a role for CYGB in NO regulation through its NO dioxygenase activity [26]. However, under conditions of oxygen deprivation, CYGB could act as a source of NO by switching to a nitrite reductase function. The switching between these two functions is regulated through the oxidation of the two surface-exposed cysteine residues [65].

Besides nitration of the heme, plasma induces the oxidation of specific amino acid residues of CYGB. Here, the three most oxidized amino acids were methionine, tryptophan, and tyrosine. In Takai et al. [66] reported that plasma was able to chemically modify amino acids through the oxidation of their side chains, with sulfur-containing (i.e. Met and Cys) and aromatic amino acids (i.e. Trp, Tyr, and Phe) preferentially modified. Furthermore, they found that the relative reactivity towards plasma treatment was Met > Cys > Trp > Phe > Tyr. This is not entirely in agreement with our findings (Table 1). However, in this study, a complete protein was treated with plasma, whereas in the study of Takai et al., a mixture of amino acids was treated with plasma. Therefore, it is highly likely that in CYGB, due to its 3D-structure, some amino acids, e.g. Phe, are less solvent accessible compared to when individual amino acids are studied, leading to different results. This is further highlighted in Table S2, where it can be seen that certain locations of preferably oxidized amino acids, such as Met and Trp were never oxidized, indicating that the access of certain amino acids is hindered due to the tertiary structure of CYGB. This observed reactivity is in agreement with an increasing body of evidence from other experimental approaches which utilize ROS to cause oxidation of the solvent-accessible surface area of proteins, such as hydroxyl radical footprinting by synchrotron radiation [51] as well as fast photochemical oxidation of proteins [67] and other oxidative labelling techniques [68].

As for cysteine, during the digestion of the protein sample prior to measuring with LC-MS/MS, the cysteines are reduced by TCEP and carbamidomethylated. This modification is introduced on purpose to prevent disulfide bridges from forming again, and was therefore not selected for the analysis of oxidative modifications [69]. Yet, from the
MS data (Figs. 2 and 4) it is clear that cysteine do become oxidized (i.e., formation of disulfide bridge). CYGB possesses two cysteine residues (Cys38 and Cys83), which are known to be able to form disulfide bridges, either inter- or intramolecularily [25,26]. Such a formation of CYGB dimers upon plasma treatment can indeed be observed in native mass spectra (Fig. 2), and even more clearly under denaturing conditions in Fig. 3. Moreover, the addition of the reducing agent DTT to both the untreated and treated samples led to the disappearance of the dimeric peaks in the spectrum (Fig. 4). However, there is still a significant portion of CYGB that stays in the monomeric form (Fig. 2). It is highly likely that the monomers that are still present after treatment have formed intramolecular disulfide bridges. Although the dimeric form of CYGB has been suggested to be less active than the monomeric form (see paragraph below), this observed dimerization after treatment could be an artefact of the experimental setup. In vivo, the concentration of CYGB lies somewhere between 0.1 and 5 μM, at which CYGB is proposed to be a monomer [70]. Higher concentrations, like the one that was used here, could bring CYGB molecules closer together and subsequently stimulate the formation of dimers. Nonetheless, from the abovementioned, we can conclude that plasma treatment leads to the formation of intermolecular disulfide bridges, thereby stabilizing the CYGB dimers, but presumably also intramolecular disulfide bridges in the remaining monomers. This would confirm the two cysteine residues as redox-active cysteines.

The state of the two surface-exposed cysteine residues, either free, intra- or intermolecularily linked, has been proposed to be critical in determining the biochemical and redox properties of CYGB [71]. A recent study demonstrated a clear difference in the ability of CYGB to bind lipids between the monomeric and dimeric form of the protein, with the monomer, containing an intramolecular disulfide bridge, more easily able to bind lipids [25]. This difference is suggested to occur because the presence of an intramolecular disulfide bond leads to the formation of a more open heme pocket structure, thereby facilitating access for external ligands, such as lipids, whereas an intermolecular disulfide bond does not alter the environment of the heme pocket [25]. This is perfectly in line with the MD and docking simulations performed in this study on monomeric CYGB. Here, it is seen that in CYGBSH-SN, the access to the heme group is more limited than in CYGBSH-S. We further found that the His117 residue plays a vital role in either limiting or increasing the access to the heme group, thereby acting like a gate function for CYGBSH-SN and CYGBSH-S. Thus, our investigations confirm, and further show, that the formation of an intramolecular disulfide bond affects the CYGB structure, thereby opening the access to the heme group, through gate functioning of His117, which in turn could lead to a higher binding affinity. In addition the redox state of the Cys58 and Cys83 residues appears to act as a ‘molecular switch’, activating the protein only under the correct conditions, i.e. oxidative stress. This would support CYGB’s proposed cytoprotective role in oxidative stress conditions, which is probably linked to the lipid peroxidase activity of the protein [25,49].

5. Conclusion

In conclusion, the treatment of CYGB with CAP leads to chemical modifications of the protein, yet the overall structure of the protein remains largely intact.

Short treatment times were already sufficient to induce the oxidative modification of mainly sulfur-containing and aromatic amino acids. Longer treatment times also lead to the nitration of the heme, through the binding of NO₂ to the vinyl groups, which proposes a role for CYGB in NO homeostasis. Upon plasma treatment the two cysteine residues of CYGB were oxidized, which stabilizes the dimeric CYGB, through intermolecular disulfide bridges, but most likely also monomeric CYGB containing an intramolecular disulfide bond. The simulations performed in this study also confirm the importance of the oxidation of Cys58 and Cys83 residues in regulating the ligand-binding properties, and thus also the activity, of CYGB. Furthermore, His127 was found to play a vital role in regulating the access to the heme group.

As mentioned in the introduction, the aim of this study was to elucidate the effects of plasma-generated (and in general) RONS on the structure and, to some extent, the function(s) of CYGB. Although some protein denaturing occurred, our results nonetheless confirm that (plasma-generated) RONS can indeed induce chemical modifications at specific sites, which lead to different conformations of CYGB. These different conformations are mediated by the oxidation status of two redox-active cysteines. As globins (more particularly hexacoordinated globins) are increasingly hypothesized to play a role in redox biology, findings here seem to further support the idea of CYGB functioning as a redox-active protein.

This study is of great interest because it provides further support and new insights that CYGB, on a molecular basis, has the potential to play a role in the cellular response against oxidative stress. However, further research on a cellular level is needed to validate these findings and to unravel the cellular pathways in which CYGB is involved. Such research should focus on elucidating the effects of (plasma-generated) RONS on cells and the role of CYGB in the response to these RONS.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.redox.2018.07.019.

References
