Can Gas Replace Protein Function? CO Abrogates the Oxidative Toxicity of Myoglobin

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Abstract

Outside their cellular environments, hemoglobin (Hb) and myoglobin (Mb) are known to wreak oxidative damage. Using haptoglobin (Hp) and hemopexin (Hx) the body defends itself against cell-free Hb, yet mechanisms of protection against oxidative harm from Mb are unclear. Mb may be implicated in oxidative damage both within the myocyte and in circulation following rhabdomyolysis. Data from the literature correlate rhabdomyolysis with the induction of Heme Oxygenase-1 (HO-1), suggesting that either the enzyme or its reaction products are involved in oxidative protection. We hypothesized that carbon monoxide (CO), a product of Mb auto-oxidation, might attenuate Mb damage, especially since CO is a specific ligand for heme iron. Low density lipoprotein (LDL) was chosen as a substrate in circulation and myosin (My) as a myocyte component. Using CO as an antioxidant, LDL and My, the study compared the antioxidant potential of CO in Mb-mediated oxidation with the antioxidant potential of Hp in Hb-mediated oxidation. The main cause of LDL oxidation by Hb was found to be heme which readily transfers from Hb to LDL. Hb prevented heme transfer by sequestering heme within the Hb-LDL complex. Hemin barely transferred from Mb to LDL, and oxidation appeared to stem from heme iron redox in the intact Mb. Mb underwent oxidative crosslinking by Mb both in air and under N2. These reactions were fully arrested by CO. The data are interpreted to suit several circumstances, some physiological, such as high muscle activity, and some pathological, such as rhabdomyolysis, ischemia/reperfusion and skeletal muscle disuse atrophy. It appears that CO from HO-1 attenuates damage by temporarily binding to deoxy-Mb, until free oxygen exchanges with CO to restore the equilibrium.

Introduction

Hemoglobin (Hb) and myoglobin (Mb) are the two major respiratory hemoproteins (RH) burdened with the task of maintaining aerobic life: Hb shuttles oxygen from the lungs to tissues, while Mb maintains a store of oxygen in muscles, ready for use. Structural similarities in their globin (gl) backbones and a common ferrous heme iron at their active site indicate similar functionalities [1,2]. Both globins undergo conformational changes between oxy and deoxy states, and both possess a ferrous iron (FeII) at the center of a heme group to provide the conditions for oxygen binding. The oxygen binding task of RH necessitates a heme ferrous iron, even though the ferric iron valency is the more stable and indeed is used as ground state of other metabolic hemoproteins. To keep the iron in its ferrous state, a reducing environment is maintained both in myocyte and in the nucleus-free red blood cell (RBC). Should a cell rupture, the expelled oxyferrous RH is no longer insulated within a protective reducing environment and spontaneously undergoes auto-oxidation to the ferric (FeIII) form [3]. The auto-oxidation reaction generally yields a reactive oxygen species, like hydrogen peroxide, which goes on to wreak oxidative havoc in the cell: \( \text{H}_2\text{O} + \text{gl-Fe}^{\text{II}} + \text{O}_2 \rightarrow \text{gl-Fe}^{\text{III}} + \text{H}_2\text{O}_2 \). This process is well documented in the literature [4]. Moreover, in pathologies with increased hemolysis like malaria [5] or sickle cell anemia auto-oxidation is increased [1,3]. Both RH also possess redox activity based on FeIII to FeII redox [2–4].

Despite structural and functional similarities, major differences exist in the cellular habitats and turnover of the two proteins. It has been pointed out that deleterious Hb redox activity is dominated by its tendency to release heme once oxidized, while Mb redox activity tends to be based more on the intact hemoprotein [5,6]. Besides structural differences, their physiology and modus operandi differ: Hb is sequestered into a RBC lacking a nucleus and containing essential reducing enzymes. RBCs are prone to frictional damage as they journey through narrow capillaries, and, since the cell wall may get damaged, Hb may be released. Cell-free Hb is oxidatively destructive [1–7], necessitating the average RBC to remain in regular circulation for only ~12 weeks (and much shorter in pathological cases), after which it is replaced by a newly-formed one. In contrast, myocytes have a low turnover rate, and only if rhabdomyolysis occurs, extracellular Mb accumulates in the kidneys, causing renal failure [8,9].
(2014) CO Abrogates the Oxidative Toxicity of Myoglobin

Two plasma proteins Haptoglobin (Hp) and Hemopexin (Hx) scavenge extracellular Hb and heme respectively. Hp has an extremely high affinity for Hb and, once bound, escorts it to the liver where the Hp-Hb complex binds to the scavenger receptor, CD163 [10–12]. Hx has a high and specific affinity for heme (heme-FEIII) and, after binding, clears it out via the CD91 receptor in the liver. The two plasma proteins work in tandem [1]. At the liver, the enzyme Heme Oxygenase (HO) converts heme to biliverdin, iron and carbon monoxide (CO) [13–15]. Intense research from recent decades has shown that genes for the induced enzyme, HO-1, are expressed in most cells, and that enzyme levels are elevated under conditions which weaken the hemo-globin bond, like stress, oxidative stress, sepsis, malaria, hypertension. In cases with increased hemolysis, like sickle cell anemia which was intensively studied, it was established that CO produced from HO-1 induced by the loosely bound (free) hemin, is in fact salutary [16–18]. Another hemolytic situation studied this concern is malaria, where added or endogenously formed CO has been demonstrated therapeutic [19,20]. While much research has concentrated on the body’s defense mechanisms against Hb and its breakaway heme, practically nothing is known about the ways in which the body deals with oxidatively toxic Mb. Mb escapes into the acellular space as a consequence of muscle trauma and can also wreak harm in the cell. Oxidative damage, once thought to be the result of free iron released from Mb, was later studied by observing the Mb-induced LDL oxidation model and found to be the consequence of intact Mb redox cycling activity [21]. A study, based on a rat-model of rhabdomyolysis, discovered that Mb-mediated oxidative damage in the kidney is followed by the rapid induction of HO-1 [22]. The same group reported that, by induction of HO-1 prior to rhabdomyolysis via administration of either Hb or hemin, full protection was provided against the damage [23]. These data imply that the enzyme HO-1 and/or its products, bilirubin and CO, provide protection. Moreover, one of our previous studies indicated that a mutual presence of CO and peroxide, hydrophilic CO, provide protection. Moreover, one of our previous studies indicated that a mutual presence of CO and peroxide, hydrophilic as OH₂O₂ or hydrophobic as in oxidized LDL, blocks oxidative damage by Hb and Mb [24].

Based on the information available, we reasoned that CO might act as a specific inhibitor, arresting damage caused by Mb. Hence, the current study was devoted to a comparison of Hp shielding Hb and the protection afforded by CO against Hb and Mb. Importantly, care was taken to maintain conditions as close to in vivo as possible. These include: mild oxidative conditions comprising a few micromolar H₂O₂, a low concentration (a few μM) of hemoprotein and no free oxygen [25].

Materials and Methods

All experimental protocols were approved by the Institutional Animal Care and Use Committee (IACUC) at the Hebrew University of Jerusalem, which adheres to the Israeli guidelines which follows the NIH/USA animal care and use protocols.

Blood donors for the preparation of LDL and Hb (see below), were fully informed concerning this study and signed a written consent form; ethical approval for this study was obtained through the research ethics board of Tel-Aviv University.

Frozen muscle acetone powder isolated from rabbit leg muscle was kindly provided by Prof. A. Muhlrad of the Hebrew University of Jerusalem.

Bovine catalase, 5-dimethylaminonapthaleine-1-sulfonil (dansyl) chloride, phenyl methyl sulfonil fluoride (PMSF), EDTA, KBr, agarose, AAPH, 2,2'-Azobis(2-amidino propane) hydrochloride, horse heart myoglobin and human haptoglobin (1-1 and 2-2) were purchased from Sigma Chemical Co., St. Louis, MO, USA.

Hydrogen peroxide (Merck, Darmstadt, Germany); DE-52 cellulose (Whatman International, Maidstone, England); PD-10 desalting columns (Amersham Pharmacia Biotech, Buckinghamshire, England); chemicals for SDS-PAGE (Bio-Rad Laboratories, Richmond, CA); gases: carbon monoxide, at least 99.5%, was supplied by Gordon Gas and Chemicals Ltd., Israel, nitrogen, at least 99.9%, was supplied by the Israel Oxygen Center.

For spectrophotometric measurements a UV/vis 920 GBC, Dandenong, Australia, was used. Fluorescence measurements were carried out using Jasco FP-6200 spectrophotofluorimeter which allows four simultaneous kinetic measurements.

Protein preparation

Myosin was prepared from frozen muscle acetone powder according to the available procedure literature [26]. Its concentration was determined using ε = 2.83×10⁻⁸ M⁻¹ at 280 nm [27].

Hemoglobin was isolated from red blood cell lysates by ion-exchange chromatography using CM-52 cellulose [2] followed by desalting on PD-10 columns. This Hb was verified as oxyHb by UV-Visible absorption spectrum. Lack of catalase was certified by ferryl Hb formation in the presence of an equimolar amount of H₂O₂.

MetHb was prepared from the oxyHb [2] and before use it was purified by admixing with DE-52 cellulose for 5 min followed by centrifugation to remove globin-free hemin contaminants [28].

COHb was prepared by administration of carbon monoxide at atmospheric pressure for one minute and verified by its specific absorption peaks [2].

Myoglobin, which is supplied in the oxidized form, was prepared by applying commercial ferric Mb to a Sephadex G-25 column with a few grains of solid sodium dithionite spread on top to maintain Mb in its ferrous state. The immediate separation of the protein from dithionite prevented protein oxidation. The concentration of oxyMb was determined according to the literature [2].

Concentrations of all hemoproteins were expressed in heme equivalents throughout this study, except experiments which include Hp, where ζβ globin (2 hemes) was considered a unit interacting with Hp.

Haptoglobin (Hp): concentrations of commercial Hp were determined by repeated measurements using the Lowry method, ε₂₈₀ nm = 53.90×10⁻⁶ M⁻¹ for Hp1-1 and 58.65×10⁻⁶ M⁻¹ for Hp2-2 (per ζβ unit) [29].

LDL: native form (nLDL) was isolated from freshly drawn venous blood of healthy volunteers using established procedure [30]. To clear LDL from the KBr and EDTA added during the preparation, it was eluted through two PD-10 columns against argon-saturated phosphate buffered saline (PBS), pH 7.4. Protein concentration of isolated LDL was determined by the method of Lowry et al. [31]. nLDL was kept under argon at 0°C (an ice/water mixture) and was used up within 7 days. Vitamin E depleted LDL (dLDL) was prepared from nLDL according to the literature [32]. dLDL was kept under argon at 0°C to be used up within 24 hours.

Analytical procedures

Heme transfer from hemoprotein to LDL was measured using FRET (fluorescence resonance energy transfer), a method which exploits fluorescence quenching due to energy transfer from protein to dyesluted LDL [33]. The reaction mixtures contain dansylated LDL, H₂O₂ and alternately Hb and Mb. The anaerobic atmosphere was either nitrogen (control) or CO. Heme transfer kinetics was followed by observing fluorescence at 400 nm, as shown in an earlier study [6].
**Results**

Scientific literature is filled with experimental data related to Hb damage and protection. Yet most in vitro studies use high concentrations of peroxide, a low pH, room temperature and aerobic conditions which don’t exist in vivo. To relate in vitro to in vivo conditions, the current study used a low peroxide concentration (µM range) as well as a lack of free oxygen (hypoxia) [25].

**Hp attenuates heme loss while allowing oxidation of ferrous Hb**

The first experiment compares damage to the heme core by peroxide in the absence and presence of Hp (Type 1-1). Three reaction mixtures were examined: 1) oxyHb in buffer alone; 2) as the first plus equimolar quantities of hydrogen peroxide; 3) as the second plus Hp equimolar to oxyHb. Reaction components were added in the following order: Hb, H₂O₂ and Hp. All three mixtures were incubated at 37 °C for 30 minutes and absorption spectra in the Soret region were recorded.

As seen from the spectra (Fig. 1), H₂O₂ oxidized oxyHb to ferric-Hb, as indicated by a shift in the Soret peak from 414 nm to 405 nm [2]. Hp thus did not prevent ferrous to ferric iron oxidation. These findings are in correlation with previous studies carried out in air and high peroxide conditions [39]. In the two cases where the heme iron was oxidized (spectra 2 and 3), since the extinction coefficient of ferric-Hb is 1.43 times higher than that of oxyHb [2], if all the oxyHb were to undergo oxidation to ferric state, we would expect the peak to be higher. However, as seen from the spectra, some of the expected absorbance was lost, indicating that a portion of heme disintegrated. Nevertheless, the Soret peak formed in the presence of Hp (spectrum 3) is higher than the one formed in its absence (spectrum 2). This implies that the heme moiety was partially shielded from oxidative disintegration by the bound Hp, in agreement with previous data [12,33].

**Lipid oxidation** was monitored by lipid and/or protein parameters.

**Lipid fractionation by charge was carried out using fast anion exchange liquid chromatography (Mono QR HR 5/5 type) according to previous literature [34].**

Lipid oxidation in LDL was determined as TBARs (ThioBarbbituric Acid Reactive substances) according to the literature [35,36]. To minimize the effects of light scattering, a longer wavelength of 532 nm was measured. Additionally lipid oxidation was monitored by the formation conjugated dienes (CD) at 268 nm rather than 234 nm to minimize the effects of light scattering [6,37].

Fluorimetric bityrosines screening: this technique allows continuous monitoring of bityrosines formation resulting from the quenching of tyrosine radicals to form bityrosines [38]. Fluorescence excitation is at 327 nm and emission at 400 nm.

SDS-PAGE analysis of cross-linked protein: oxidation products of ApoB were followed by SDS-PAGE with β-mercaptoethanol using 6–12% acrylamide bilayer slabs. Gels were stained with Coomassie Brilliant Blue R-250 [6].

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**Figure 1.** Hp attenuates heme loss while allowing oxidation of ferrous Hb. Spectrum of OxyHb (5 µM) was recorded. H₂O₂ (to yield 6 µM) was added and the mixture was incubated in 37°C and the spectrum recorded again. Experiment was repeated in presence of Hp1-1 (6 µM): 1: time zero (with or without Hp); 2: following 30 min in absence of Hp; 3: following 30 min in presence of Hp. Representative spectra of 3 independent experiments are shown.

doi:10.1371/journal.pone.0104075.g001

**Hp inhibits Hb-induced LDL oxidation**

As seen in the previous experiment, a portion of heme disintegrated in solution. This portion may oxidatively attack neighboring components. To check if this is the case, we measured the oxidation of both protein and lipids in LDL under the same conditions as before. LDL was chosen as an in vivo substrate in circulation that is sensitive to oxidation by cell-free Hb. A mixture of Hb and H₂O₂ was incubated with LDL for three hours at 37°C, under nitrogen. Fig. 2A shows lipid oxidation quantified by the formation of TBARS. Comparing lanes 1 and 2 in Fig. 2A indicates that the association of Hb with Hpl-1 inhibited the formation of lipid peroxides. Protein oxidation was observed kinetically by examining the fluorescence of bityrosines formed in LDL. An example for differences in oxidation when ferrous Hb is employed can be seen in a previous study from our laboratory [40]. Fig. 2B shows that Hp almost fully inhibits the reaction. Bityrosines that form may be either intramolecular or intermolecular. As only one protein molecule exists per LDL particle, intermolecular cross-linking results in aggregation.

An SDS-PAGE was run in order to analyze the aggregation state of the protein. Lanes 1 and 2 in Figure 2C indicate that within 3.5 hours, little protein remained as the original 500 kDa monomer (apoB). Most of the protein had become cross-linked in a covalent intermolecular form (HMA – High Molecular-weight Aggregates). In contrast, in the reaction mixtures containing Hp, most of the protein remained as an apoB monomer, indicating that the presence of haptoglobin was able to prevent the formation of covalent aggregates of LDL.

**A Hp with reduced activity prevents oxidation of dLDL**

There are certain in vivo conditions in which LDL is more vulnerable to oxidation. One condition is a lack of vitamin E within the particle, a vitamin E-depleted form (dLDL), rendering it more sensitive to oxidation. Another condition is a type of
haptoglobin which has a weaker protective effect against hemin release, Hp 2-2 [41]. To evaluate the degree of protection from cell-free Hb provided by Hp to the vasculature, we tested whether Hp 2-2 could protect dLDL. Ferric-Hb was incubated with dLDL, with and without Hp 2-2. As a double control, dLDL and Hp were each incubated alone. As dLDL is highly negatively charged, an agarose column was used to separate the components of the reaction mixtures according to charge. The column was developed using a salt gradient of 0–1 M NaCl, buffered with tris-HCl at pH 7.4, and protein fractions were eluted.

As seen from Fig. 3, the elution peaks of Hp alone (peak 4) and LDL alone (peak 1) appear at ~50 minutes, while the elution peak of oxidized LDL (peak 2) was at ~60 minutes. This may be due to an increase in negative charge on LDL, caused by its oxidation. In contrast, the presence of Hp in the reaction mixture caused the elution peak to remain at LDL's original position (peak 3).

CO attenuates Hb-induced LDL oxidation

In a previous study, we showed that the mutual presence of CO and peroxide arrests ferric-Hb in the carboxy, redox-inactive, ferrous form [24]. In the first part of the current study, we stated that oxidation of LDL takes place even under an anaerobic atmosphere. The following experiment determined whether ferric-Hb can oxidize LDL in a CO atmosphere.

LDL oxidation kinetics was traced in reaction mixtures containing ferric-Hb, LDL, and hydrogen peroxide. Air was replaced with either N₂ or CO. Protein oxidation was observed by the formation of conjugated dienes at 268 nm. Fig. 4 shows that the rates of Hb-induced oxidation in the presence of CO were reduced dramatically. Oxidation under CO proceeded more slowly. Suppression of oxidation by CO also occurred when the freshly prepared LDL was replaced by dLDL, although to a lesser extent.

Effect of CO on heme transfer from ferric-RH to LDL

As shown above, CO can diminish the oxidation of LDL by Hb. We wished to find out whether Hb-induced LDL oxidation is the direct outcome of intact hemoglobin or globin-free heme. To evaluate the heme transfer from globin to LDL, ApoB was dansylated, according to a previous study from our lab [6]. Heme that binds to LDL results in quenching of fluorescence of the dansyl probe. The results in Fig. 5 show that carbon monoxide appreciably diminishes the rate of heme transfer to LDL. In comparison, Mb exhibited only negligible quenching, with a...
further reduction of transfer under CO. This finding corresponds with results carried out previously [6].

CO locks HRP iron in a ferrous state

Previously, it was reported that HRP was completely unable to peroxidize LDL in the presence of CO [42], but no mechanism was suggested. We showed in the current study that Mb does not release heme easily, operating as an intact enzyme (Fig. 5). In Fig. 6, we show absorption spectra of HRP in the Soret region of reaction mixtures containing CO and peroxide at time zero and after one hour’s incubation. Fig. 6A depicts practically no change in the Soret peak of the heme following incubation in nitrogen. In contrast, Fig. 6B demonstrates that under CO, ferric heme was reduced to carboxy-ferrous heme, locking the enzyme in a redox inactive state. This mechanism provides an explanation for the previous observation that LDL oxidation is arrested in the presence of CO [42].

Mb peroxidase-like redox activity is specifically restrained by CO

As reported previously, of the two RH proteins Mb is closer in its enzymatic function to HRP than Hb [6]. Therefore, it was of interest to compare the effect of CO on Mb (main figure, Fig. 7) and Hb (insert) oxidative activity. The kinetics of LDL oxidation by Hb or Mb was traced under anaerobic conditions provided by N₂ (solid line) or CO (dot-dash line). The two upper lines – Hb and the two lower lines – Mb. (Ex/Em = 327/400 nm). A representative result of 4 independent experiments is shown.

doi:10.1371/journal.pone.0104075.g005

sharp contrast to the effect of CO on Hb, where oxidation still took place, albeit at a slower rate.

As stated in the introduction, an important difference between Hb and Mb is the physiological environment in which each RH resides and operates. Mb resides in the cell, dominated by other proteins. Therefore, in the next experiment, we traced the effect of Mb on myosin under oxidative conditions.

CO blocks Mb-induced myosin oxidation

OxyMb was mixed with excess hydrogen peroxide in order to peroxidize myosin. The formation of bityrosines was traced by its fluorescence emission as detailed in a previous study [38]. As seen from Fig. 8A, under nitrogen the reaction reached completion after 25 minutes. In contrast, under CO, a negligible amount of bityrosines formed, and the reaction reached completion after 10 minutes. To analyze which components of the mixture took part in the formation of bityrosines, reaction mixtures were incubated for 40 minutes to reach completion and analyzed with SDS-PAGE. As seen in Fig. 8B, the reaction resulted in the reduction of the myosin bands in favor of high molecular weight bands at the gel interface (lane 3). These bands, as shown in previous literature [43], represent intermolecular covalent aggregates. Evidently, myosin monomers (heavy and light chains) faded while aggregates were seen to appear more strongly. Replacing air
with nitrogen resulted in practically the same bands, as shown in lane 4. However, replacing nitrogen with CO resulted in a complete inhibition of aggregation, as seen from the gel: bold myosin monomer bands in lane 5 are favored over weak bands of aggregates.

Discussion

The study shows that in the absence of free oxygen and at low peroxide levels, ferrous-Hb undergoes oxidation to ferric-Hb. Under such conditions, hemin disintegrates rapidly, judging by the partial loss in absorbance of the ferric Soret band. In corroboration with a previous study [12], while the association of Hb with Hp did not inhibit the oxidation of ferrous to ferric iron, hemin was protected from disintegration. The current study further demonstrates that when a solution of cell-free Hb contains circulatory components of plasma, like hydrophobic LDL, hemin readily transfers from globin to these components. As a result, both lipids and the apoB protein of the LDL particle undergo oxidation (Fig. 2). Hp acts by trapping the heme in the Hp-Hb complex, such that it can no longer oxidize LDL. The Inhibition of heme-induced oxidation even holds true in the case of the less effective form, Hp 2-2, and the vulnerable form of LDL, dLDL (Fig. 3). These results clearly show that the efficiency of Hp in inhibition of Hb oxidative activity stems from preventing sensitive targets from associating with loosely-bound hemin [44]. The specific protection afforded to Hb by Hp fits in well with our knowledge of the dissociation of Hb into \( \alpha \beta \) dimers at low concentrations occurring \textit{in vivo}. Hp prevents the escape of the loosely-bound hemin since it masks the surface of the \( \alpha \beta \) dimer when it binds, covering a large part of the Hb interface [45]. The current study indicates that CO provides some protection against oxidation of LDL by Hb, but this protection is much less efficient than that of Hp.

The prominent difference between the two mechanisms by which Hb and Mb act, relates to the fact that heme transfer from Mb is negligible in comparison to Hb (Fig. 5). Differences in mechanisms of the two proteins are especially prominent when observing the kinetics of LDL oxidation and its arrest by CO (Fig. 7, main). LDL is oxidized by ferric-Mb at a constant rate, appropriate for enzymatic function. This differs completely from the multistage rate of Hb-induced LDL oxidation (Fig. 7 insert). Despite a common physiological function (oxygen binding to a divalent heme iron), Hb and Mb differ in the mechanism by which they evoke oxidation. Hb oxidative activity results fundamentally
from a weakening of the trivalent heme-globin bond [44]. As discussed earlier, ferric-Hb redox activity is manifested by the presence of components that bind hemin strongly, such as LDL [6,33]. Thus, only a high-affinity globin-binding protein, such as Hp, can efficiently trap hemin. On the other hand, as suggested in the past and indicated in the current study, Mb’s oxidative power stems from a protein-bound ferric heme whose activity is peroxidase-like: namely, fully dependent on heme iron redox capability. The differences in mechanism of oxidation induced by the two RH proteins are demonstrated schematically in Fig. 9.

As mentioned in the introduction, besides differences in structure/function, the physiological compartment in which RH are found is completely disparate. Hb is short-lived and becomes harmful when expelled from a ruptured cell to the vasculature. Mb, on the other hand, resides in a long-lived, nucleated myocyte,
filled with functional proteins and thus may cause oxidative
damage both inside and outside the myocyte.

Single-chained extra-cellular Mb deposits in the kidney, causing
major damage there. In addition to the large amount of Mb, some
Hb also finds its way into the kidney following rhabdomyolysis. It
was shown in the rat rhabdomyolysis model that the presence of
residual amounts of Hb aided in countering an overdose of Mb,
causing oxidative damage [22]. These conditions induce HO-1.
Recent literature in combination with the current study suggest the
following explanation: ferric-Hb readily liberates globin-free
hemin, an inducer of HO-1. The product of the enzymatic
reaction, CO, spreads to the immediate vicinity, binding strongly
to ferrous Mb, rendering it inactive. This is summarized below:

\[
\text{Hb}^{II} \rightarrow \text{Hemin} \rightarrow \text{HO-1} \rightarrow \left\{ \text{CO} + \text{Mb}^{II} + \text{H}_2\text{O}_2 \rightarrow \text{COMb}^{II} \right\} * \\
\]

{* - combined reducing effect of CO and peroxide exhibited in a
previous study [24].

The minute amount of hemin dispelled from Mb in the cell
induces production of HO-1 which produces more CO. In the
myocyte, Mb resides in close proximity with contractile proteins
that are prone to oxidation. Therefore, their immediate protection
from injurious oxidants is necessary, since Mb acts as an
antioxidant, protecting myoglobin from oxidation [50]. Based on the findings of the current study, it can be
anticipated that as a result of heme dismantling by HO-1, the
antioxidant product, CO, arrests the oxidative activity of Mb.

Ordinarily, following strenuous activity, the muscle cell loses a
considerable amount of its Mb-bound oxygen, leaving part of it in
a deoxy state. Deoxy-Mb is less stable and more active in inducing
oxidation [31,52] and thus harmful. As stated above, under
oxidative conditions myoglobin releases a fraction of its hemin,
inducing HO-1, which in turn releases CO during heme
catabolism. A shortfall in free oxygen leads to the immediate
saturation of deoxy-Mb with CO. Carboxy-Mb is a most desired
complex for the provisional preservation of Mb as ferrous-Mb. In
support of the above-suggested scenario, a previous study of
skeletal muscle type points out that induction of HO-1 follows a
fiber-type specific pattern such that upon addition of free hemin,
induction is more pronounced in red muscle with a high Mb
content [53].

In this context, it has been shown that inhaled carbon monoxide
protects muscles from structural injury and energy depletion
following I/R, as well as modulates inflammation in tissue after
therapy, without affecting the production of HO-1, providing a
form of metabolic rescue [34].

Taking together the current and our previous study [6], it
appears that hemin binds more strongly to Mb than to Hb. The
transfer of heme to LDL is consequently different in the two
molecules. In Hb, hemin is trapped by ApoB protein and
thereafter disintegrates in conjunction with LDL oxidation. Unlike
Hb, where the transferred heme triggers oxidation, Mb acts as an
intact hemoprotein peroxidase, resembling HRP. Mb’s heme iron
redox is based on Fe^3+/^2^+ activity. Thus, in order to block redox
activity of Hb, a chaperone protein, Hp, is required, while in order
to block injurious Mb redox activity, a small, yet efficient electron
trap will suffice. The specific and high affinity of CO for ferrous
iron becomns its call. As shown in a previous study [24], CO
defused the peroxide and arrested the heme iron in a ferrous state.

Author Contributions

Conceived and designed the experiments: NS MS. Performed
the experiments: EAS AYS. Analyzed the data: EAS AYS NS MS.
Contributed reagents/materials/analysis tools: NS MS. Contributed to
the writing of the manuscript: EAS AYS NS MS.

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to block injurious Mb redox activity, a small, yet efficient electron
trap will suffice. The specific and high affinity of CO for ferrous
iron becomns its call. As shown in a previous study [24], CO
defused the peroxide and arrested the heme iron in a ferrous state.

Author Contributions

Conceived and designed the experiments: NS MS. Performed
the experiments: EAS AYS. Analyzed the data: EAS AYS NS MS.
Contributed reagents/materials/analysis tools: NS MS. Contributed to
the writing of the manuscript: EAS AYS NS MS.

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