Glutamine is highly effective in preventing in vivo cobalt-induced oxidative stress in rat liver

Soledad Gonzales, Ariel H. Polizio, María A. Erario, María L. Tomaro

INTRODUCTION
Oxidative stress is the result of excessive production of oxidant species and/or depletion of intracellular antioxidant defenses, leading to an imbalance in the redox status of the cell.

Glutamine (Gln) is a multifaceted amino acid used as an energy substrate for most cells,[1] it is also a precursor for nucleotides[2], and it is the most abundant free α-amino acid found in plasma and in the free amino acid pool of the body.[3] One of the most important characteristics of glutamine is that it plays a critical role in glutathione biosynthesis. Glutamine provides glutamate to the glutathione system, which is one of the main sources of the antioxidant defense system in the cell.[4,5]. Therefore, Gln is a natural product that plays a leading role in the protection against oxidative stress injury[6,7].

It is accepted that CoCl₂ produces oxygen-derived free radicals, which leads to a greater oxidative stress damage[8]. Moreover, it has been demonstrated that cobalt salts activate the expression of several stress-responsive proteins, such as heme oxygenase[9,10].

Heme oxygenase (HO) is the rate-limiting microsomal enzyme that catalyzes heme degradation, which leads to the formation of carbon monoxide, iron and biliverdin, the latter being converted into bilirubin by the cytosolic enzyme biliverdin reductase[11,12]. All these products are biologically active because iron is an important gene regulator[13] and a pro-oxidant[14], bilirubin is a potent antioxidant[15] and CO has properties similar than nitric oxide[16,17]. Three isoforms of HO have been described in mammals: HO-1, the inducible enzyme[18], HO-2, the constitutive isoform[19] and the more recently identified HO-3[20]. HO-1 can be induced by a number of stressful stimuli including its own substrate heme, various heme proteins, heavy metals, glutathione depletion, UVA radiation, hypoxia, hyperoxia, ischemia reperfusion and many others[20,24].

There is compelling evidence that the biological damage attributed to reactive oxygen species (ROS) is dependent on the presence of iron such as heme-derived intracellular iron.[28]. Within most cells ferritin constitutes the major storage
site for non-metabolized intracellular iron and therefore plays a critical role in regulating the availability of iron to catalyze such harmful reactions as the peroxidation of lipids and the Fenton reaction generating the highly reactive hydroxyl radical (HO·).

Reactive oxygen species occur in tissues and may damage DNA, proteins, carbohydrates, and lipids. These potentially deleterious reactions are controlled by a system of antioxidant defenses which eliminate pro-oxidants and scavenge free radicals. Protection against oxidation is provided by various intracellular compounds such as glutathione, and antioxidant enzymes including catalase (CAT), superoxide dismutase (SOD), and glutathione peroxidase (GSH-Px).[8]

Previous studies carried out in our laboratory have demonstrated that CoCl₂ developed oxidative stress in rat liver and consequently a significant induction of HO-1 enzyme occurred.[8] The present study was performed in order to evaluate the antioxidant properties of glutamine, its capacity to modify glutathione (GSH) levels and its relationship to HO-1.

MATERIALS AND METHODS

Materials
NADPH, reduced glutathione (GSH), oxidized glutathione (GSSG), 5,5'-dithio-bis-(2-nitrobenzoic acid), thiobarbituric acid, glutathione reductase, ferritin from rat liver, rabbit anti-horse spleen ferritin, 4-chloro-1-naphthol, hydroquinone, 1,10-phenanthroline and glutamine were from Sigma Chemical Company (Saint Louis, MO); peroxidase-conjugated goat anti-rabbit immunoglobulins was from DAKO Chemical Company (Saint Louis, MO); peroxidase-conjugated goat anti-horse spleen ferritin, 4-chloro-1-naphtol, hydroquinone, 1,10-phenanthroline and glutamine were from Sigma Chemical Company (Saint Louis, MO); peroxidase-conjugated goat anti-rabbit immunoglobulins was from DAKO (Denmark). All other chemicals were of analytical grade.

Methods

Animals and treatments Female albino Wistar rats (160-180 g) were housed under standardized conditions with controlled temperature (22±3℃) and humidity (60%) and exposure to a 12-h light/12-h dark cycle. They were fed regular pelleted rat chow and given tap drinking water ad libitum. Rats were anesthetized by inhibition of adrenochrome formation rate at 480 nm for 10 min. The supernatant, a suspension of preserved organelles, was used as homogenate. Catalase activity was determined by measuring the decrease in absorbance at 240 nm,[26] and glutathione peroxidase activity following NADPH oxidation at 340 nm,[27] and superoxide dismutase activity by inhibition of adenochrome formation rate at 480 nm.[28]

One unit in the SOD assay is defined as the amount of enzyme that inhibits 50% of epinephrine auto-oxidation.

Lipid peroxidation Lipid peroxidation in liver was determined by measuring the rate of production of thiobarbituric acid reactive substances (TBARS), expressed as malondialdehyde equivalents.[30] One volume of homogenate was mixed with 0.5 volume trichloroacetic acid 150 g/L and centrifuged at 2 000 r/min for 10 min. The supernatant (1 mL) was mixed with 0.5 mL thiobarbituric acid (0.7 g/L) and boiled for 10 min. After cooling, sample absorbance was read spectrophotometrically at 535 nm. Malondialdehyde concentration was calculated using a ε value of 1.56×10⁵ mol/L cm.

Endogenous hepatic GSH content Total glutathione (GSH plus GSSG) was determined in liver homogenates after precipitation with 20 mL/L perchloric, and using yeast-glutathione reductase, 5,5'-dithio-bis-(2-nitrobenzoic acid) and NADPH and reading at 340 nm. Oxidized glutathione (GSSG) was determined by the same method in the presence of 2-vinylpyridine. GSH was calculated from the difference between total glutathione and GSSG.[30]

Ferritin content determination For ferritin assay the homogenate was prepared using 1 g of tissue in 10 V of 1 mmol/L phenylmethylsulfonyl fluoride, 0.2 mmol/L EDTA and 50 mmol/L potassium phosphate buffer (pH 7.4). Homogenates were centrifuged at 20 000 g for 20 min and supernatant fractions centrifuged at 150 000 g for 90 min. The microsomal pellet obtained was washed and resuspended in 20 mmol/L potassium phosphate buffer (pH 7.4), containing 135 mmol/L KC1, 1 mmol/L phenylmethylsulfonyl fluoride and 0.2 mmol/L EDTA to a protein concentration of 10 mg/mL. Microsomal HO-1 was obtained from similar procedures as described elsewhere[30]. The 150 000 g supernatants obtained from the microsomal preparation were fractionated by addition of ammonium sulfate (AS), and the 40-60% AS fraction dissolved in 10 mmol/L potassium phosphate buffer (pH 7.4) and dialyzed against the same buffer using this preparation as biliverdin reductase. Heme oxygenase activity was determined as described elsewhere[30]. The standard incubation mixture in a final volume of 200 µL contained 10 µmol/L potassium phosphate buffer (pH 7.4), 60 nmol. NADPH, 50 µL HO-1 (0.5 mg protein), 50 µL biliverdin reductase (0.42 mg protein), and 200 mmol/L hemin. Incubations were carried out at 37℃ during 30 min. Activity was determined by measuring bilirubin formation, which was calculated as the difference in absorbance measured at 455 and 520 nm, employing an ε value of 50 mmol/L cm (vis₃₈₅, 455 nm)[30]. Superoxide dismutase (SOD), catalase (CAT) and glutathione peroxidase (GSH-Px) activities were determined spectrophotometrically in liver homogenates prepared in a medium consisting of 140 mmol/L KC1 and 25 mmol/L potassium phosphate buffer (pH 7.4), and centrifuged at 600 r/min for 10 min. The supernatant, a suspension of preserved organelles, was used as homogenate. Catalase activity was determined by measuring the decrease in absorbance at 240 nm,[27], glutathione peroxidase activity following NADPH oxidation at 340 nm,[28], and superoxide dismutase activity by inhibition of adenochrome formation rate at 480 nm.[28]

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ice-cold 10 mmol/L HEPES pH 7.9 solution containing 10 mmol/L KCl and 0.5 mmol/L dithiothreitol. Homogenates were centrifuged at 10 000 g for 5 min. Standard horse ferritin diluted in the range of 0.25-15 ng/50 μL. Tris-Na (TS) and homogenates (diluted to approximately 50 μg protein/50 μL TS) were applied in triplicate onto nitrocellulose membranes (MSI, Westboro, MA) presoaked in TS using a vacuum dot blot as described by Roskams and Connors[32]. Briefly, membranes were blocked for 1 h at 25 °C with 3 g/L Molico instant non-fat dry milk in TS, rinsed for 5 min thrice with TS and incubated overnight at 4 °C with primary antibody: rabbit anti-horse ferritin. Membranes were then rinsed and incubated with secondary antibody (peroxidase conjugated, goat anti-rabbit immunoglobulins) for 1 h at 25 °C, rinsed again for 5 min three times with TS and developed with a solution containing α-chloronaphthol in methanol and hydrogen peroxide. Blots were quantified by computerized densitometry. Blots were quantified by Gel-Pro® analyzer 3.1 version, Media Cybernetics.

**Ferritin iron determination** Homogenates were prepared as described above (ferritin content). Ferritin iron levels were determined as following: supernatants were heated at 70 °C for 10 min, centrifuged at 15 000 g for 15 min, and resulting supernatants stored at -20 °C. The liver ferritin extract prepared was subjected to acid hydrolysis with 2.8 mol/L HCl at 90 °C for 1 h after which precipitated proteins were removed by centrifugation at 15 000 g for 15 min. One millilitre of the resulting supernatant was incubated with 20 μL of 20 g/L hydroquinone and 20 μL of 10 g/L o-phenanthroline and the optical density at 505 nm was determined. A standard curve was generated based on the absorbance of standard solution of ferrous sulfate at pH 3. As with the liver ferritin extract, the standard iron solution was carried through the acid hydrolysis procedure as well. All glassware for the ferritin iron assay was acid washed and all chemicals and reagents were ultra pure.

**Western blot analysis for HO-1 expression** Samples of homogenate obtained for HO-1 activity assays were also analyzed by Western immunoblot technique as previously described[33]. An amount of protein (50 μg) from homogenates of control and treated rats was run in sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis using a 12% acrylamide resolving gel (Mini Protein II System, BioRad, Hertz, UK). Separated proteins were transferred to nitrocellulose membranes and non-specific binding of antibodies was blocked with 3% non-fat dried milk in PBS, pH 7.4 for 1 h at room temperature. Membranes were then probed with polyclonal goat anti HO-1 antibody (Santa Cruz, Bio Tech., CA), (1:300 dilution in Tris-buffered saline, pH 7.4) overnight at 4 °C. Immune complexes were detected using donkey anti-goat secondary antibody (1:1 500), (Santa Cruz, Bio Tech., California), and were visualized using ECL reagent (Amersham, Pharmacia). Intensity of bands was analyzed with Gel-Pro® analyzer 3.1 version, Media Cybernetics.

**Protein determination** Protein concentration was evaluated by the method of Lowry et al[34], using bovine serum albumin as standard.

**Statistical analysis** Figures in the text and tables indicate mean±SD. Differences between control and treated were analyzed using the Student’s t-test, taking P<0.05 as significant.

### RESULTS

**Assessment of oxidative stress parameters** Oxygen reactive species are regarded as initiators of peroxidative cell damage. TBARS measurement was used as an assay for lipid peroxidation in vitro. A significant increase 50±3% in lipid peroxidation was observed 1 h after CoCl2 treatment, reaching a peak 100±5% 3 h later and decreasing thereafter. Control levels were regained 15 h after the injection (Figure 1).

Reduced glutathione is a leading substrate for enzymatic antioxidant functions and is capable of non-enzymatic radical scavenging. It could therefore be expected that if CoCl2 induces the formation of oxidant species, it will also affect GSH-liver levels. Data in Figure 1 showed that GSH concentration in the liver of treated animals decreased by roughly 32±2% in respect to controls, 1 h after CoCl2 treatment. Liver glutathione levels reached a minimum (40±2% of control value) 3 h after injection, increasing thereafter to approach control levels 9 h later. Control animals failed to show any significant changes in the evaluated parameters up to the end of the 36 h observation period (data not shown).

**Effect of CoCl2 treatment on HO-1 activity** Increased TBARS content as well as GSH decrease appeared as closely related events, taking place several hours before HO-1 induction. Figure 2 shows that HO-1 activity was only evidenced 6 h after CoCl2 administration, with a peak value at 12 h (14-fold over control values), to decrease up to 36 h after treatment, when enzymatic activity was similar to control values.

**Effect of CoCl2 treatment on ferritin and ferritin iron contents** A positive relation between ferritin and liver ferritin iron levels was observed. As shown in Figure 3, the concentration of liver ferritin iron was 40±2% greater than controls 18 h after CoCl2 injection, while a 50±3% increase was found 6 h later, and its concentration remained high up to 28 h.

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**Figure 1** Time course of cobalt chloride effect on lipid peroxidation and on liver GSH content. Data are mean±SD, n = 6. ⁎P<0.01 as assessed by Student’s t-test. Values measured in control (vehicle-injected) animals were the same as those at 0 time.

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after cobalt treatment. Likewise, 18 h after CoCl<sub>2</sub> injection, and 12 h after HO induction, ferritin levels were 20±1% higher than those of control animals (Figure 3). An increase by about 67±5% was obtained 24 h after treatment, and this increment was sustained for at least 28 h after CoCl<sub>2</sub> administration (Figure 3).

**Effect of glutamine pretreatment on hepatic TBARS levels, glutathione content, antioxidant enzyme activities and HO-1 activity and expression**

Administration of glutamine 24 h before cobalt treatment entirely prevented TBARS increases as well as GSH decreases, which showed similar levels than control animals (Figures 4A and B, and partially prevented the increase in HO-1 activity (Figure 4C). The activity of the antioxidant enzymes CAT, GSH-Px and SOD diminished 12 h after treatment by about 30±5%, compared to control animals, but no effect on antioxidant enzyme activities was observed after Gln pretreatment (Table 1). On the other hand, administration of Gln alone had no effect on HO-1 activity and oxidative stress parameters (Figure 4, Table 1).

The behavior of HO-1 expression was similar to that observed with HO-1 activity. Therefore, a marked increase in its expression was obtained 12 h after CoCl<sub>2</sub> injection, which was partially prevented by Gln administration (Figure 5).

| Table 1 Effect of glutamine on antioxidant enzyme activities in cobalt-treated rat liver (mean±SD, n = 6) |
|---------------------------------|
| Treatment | CAT (pmol/mg protein) | GSH-Px (U/mg protein)<sup>a</sup> | Total SOD (U/mg protein) |
|-----------|------------------------|-------------------------------|-------------------------|
| Control   | 2.5±0.2                | 0.16±0.01                     | 7.6±0.4                 |
| CoCl<sub>2</sub> | 1.7±0.1<sup>b</sup> | 0.11±0.01<sup>b</sup> | 5.7±0.4<sup>b</sup>   |
| Gln       | 2.4±0.3                | 0.15±0.01                     | 7.2±0.7                 |
| Gln + CoCl<sub>2</sub> | 1.8±0.1<sup>a</sup> | 0.11±0.01<sup>a</sup> | 5.1±0.4<sup>a</sup>   |

Rats were killed 12 h after the beginning of the experiment. <sup>a</sup>One unit of the enzyme represents the decrease of 1 mmol of NADPH/min under assay conditions. <sup>b</sup>P<0.05 vs control group.

**DISCUSSION**

Since the pioneer work of Stocker et al<sup>[15]</sup>, when they demonstrated the antioxidant properties of bilirubin, several authors have proposed that the specific induction of HO-1 by various forms of oxidative stress is part of the defensive mechanism mounted by cells against stress injury.

In agreement, our results clearly demonstrated that after CoCl<sub>2</sub> treatment, there is an enhancement in TBARS levels and a decrease in reduced glutathione (GSH) contents. Both events lead to an induction of HO activity (Figures 1 and 2). Depletion of GSH is in fact an index of oxidative stress and it is also linked with the activation of transcriptional factors and regulation of gene expression<sup>[18]</sup>. Previous works...
Antioxidants, bile pigments, such as bilirubin, capable of acting as pro-oxidants and increasing the concentrations of active defense, operating by decreasing the levels of potential oxidative stress represents an antioxidant defense system decreased by induction of HO-1 has been correlated with a decrease of endogenous GSH.

It is however, worth to note that heme catabolism generates both pro- and antioxidant compounds, consequently influencing cellular sensitivity to oxidants. This issue has been extensively discussed by Ryter and Tyrrell. Several reports have proposed that heme oxygenase induction by various forms of oxidative stress represents an antioxidant defense, operating by decreasing the levels of potential pro-oxidants and increasing the concentrations of active bile pigments, such as bilirubin, capable of acting as antioxidants. Accordingly, the higher ferritin levels found in this study, subsequently to HO induction (Figure 3), may be due to HO-dependent release of iron from endogenous heme sources. In this way, there is an enhancement of cellular iron sequestering capacity that may confer increased resistance to oxidative stress. These results are in agreement with recent studies implicating heme oxygenase-dependent increase in ferritin. Even though HO is highly induced and therefore bilirubin levels were enhanced, the oxidative stress parameters were still observed. This fact could be due to the pro-oxidant effect of iron exceeds the antioxidant properties of bilirubin. When ferritin increased and therefore iron was subsequently sequestered, the oxidative stress parameters were not detected (Table 1, Figures 1 and 3).

Interestingly, during ROS scavenging, GSH is oxidized and form glutathione-protein mixed disulfides. The cell’s ability to reduce or synthesize GSH (via glutamate) is a key mechanism by which the oxidative stress can be regulated. Besides, it has been demonstrated that Gln preserves gut glutathione levels during intestinal ischemia/reperfusion.

Glutathione plays a key role in the protection against oxidative stress injury. Our present findings showed that treatment with glutamine, a well known precursor in the GSH biosynthesis, totally reverted the decrease in GSH levels and the increase in lipid peroxidation, and partially inhibited heme oxygenase activity (Figures 4A and B).

Surprisingly, no effect on antioxidant enzyme activities was found (Table 1). On the other hand, treatment of rats with cobalt produced a significant increase in HO-1 expression, and this effect was also significantly reverted by Gln pretreatment (Figure 5).

Taken together, our data suggest that oxidative stress caused in rat liver the following events: the induction of heme oxygenase produced heme cleavage, which results in increased intracellular free iron, when ferritin content and ferritin-bound iron also beginning to increase, in an attempt to limit iron availability. Induction of ferritin and the concomitant iron sequestration may protect rat liver from oxidative injury by restricting iron-catalyzed free radical reactions. When Gln was administered all these events seemed not to be necessary, to judge by the inhibition in HO-1 activity and expression.

To sum up, our results strongly suggest that the protective effect exerted by Gln was due to the enhancement of GSH, the major soluble antioxidant compound in the liver.

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