BMC Pharmacology



Research article Open Access

Peroxynitrite-mediated inactivation of heme oxygenases Robert Kinobe[†], Yanbin Ji[†] and Kanji Nakatsu*

Address: Department of Pharmacology and Toxicology, Faculty of Health Sciences, Queen's University, Kingston, Ontario, K7L 3N6, CANADA Email: Robert Kinobe - kinober@post.queensu.ca; Yanbin Ji - yyjj@post.queensu.ca; Kanji Nakatsu* - nakatsuk@post.queensu.ca

* Corresponding author †Equal contributors

Published: 21 October 2004

BMC Pharmacology 2004, 4:26 doi:10.1186/1471-2210-4-26

Received: 14 July 2004 Accepted: 21 October 2004

This article is available from: http://www.biomedcentral.com/1471-2210/4/26

© 2004 Kinobe et al; licensee BioMed Central Ltd.

This is an open-access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Abstract

Background: Endogenous nitric oxide (NO) and carbon monoxide (CO) are generated by nitric oxide synthase and heme oxygenase, respectively. Like NO, CO has been accepted as an important cellular signaling molecule in biological systems. An up-regulation in both gene and protein expression of heme oxygenase-I (HO-I) under oxidative/nitrosative stress has been well documented, and the protective role of HO-I and HO-2 against oxidative damage is proposed. However, data on the direct effect of reactive oxygen/nitrogen species (ROS/RNS) on HO function is incomplete. Using gas chromatography to quantify carbon monoxide (CO) formation from heme oxidation, we investigated the effects of peroxynitrite (ONOO-) on the *in vitro* catalytic activity of rat spleen (HO-I) and brain (HO-2) microsomal heme oxygenases.

Results: Exposure to ONOO-led to concentration-dependent but reversible decreases in the activity of microsomal rat spleen and brain HO activity. Spleen HO activity was 100-fold more sensitive to ONOO-dependent inactivation compared to that of the brain, with IC_{50} values of 0.015 \pm 0.005 mM and 1.25 \pm 0.25 mM respectively. Inhibition of both rat spleen and brain microsomal HO activity was also observed with tetra-nitromethane, a tyrosine nitrating agent, as well as two NO donors, S-nitrosoglutathione (GSNO) and diethylamine NONOate (DEANONOate). However, no additive effect was found following the application of NO donors and ONOO-together.

Conclusion: These results indicate that ONOO- may regulate HO-I and HO-2 activities by mechanisms that involve different interactions with these proteins. It is suggested that while nitration of tyrosine residues and oxidation of sulfhydryl groups may be involved, consideration should be given to other facets of ONOO- chemistry. This inhibition of HO activity offers a mechanism for cross talk between the nitric oxide synthase and HO systems.

Background

Heme oxygenases (HO, EC 1.14.99.3) are a highly conserved family of proteins that catalyse the oxidative cleavage of heme at the α -meso carbon to yield equimolar amounts of iron, carbon monoxide (CO) and biliverdin. Biliverdin is subsequently reduced to bilirubin by biliverdin reductase. Three distinct isozymes of HO (HO-1, HO-1)

2 and HO-3) have been identified. HO-1 (the inducible isoform) is predominantly expressed in the spleen, the primary site of heme catabolism but has been detected in many different tissues including the liver and the kidney. Several substances and conditions may induce the expression of HO-1. The involvement of HO and products of heme catabolism have been studied extensively with

respect to oxidative stress, ischemia, hypoxia and protection against transplant rejection [1-4]. HO-2 (the constitutive isoform) is predominantly expressed in the testes and the brain where HO-2 dependent CO production is thought to aid neuronal function [5-7]. HO-3 is also a constitutive isoform, which shares 90% homology with HO-2 but has very limited catalytic function [8].

Increasing amounts of evidence suggest that products of heme catabolism have cytoprotective roles such as antiinflammation, anti-apoptosis and anti-proliferation. Biliverdin and bilirubin have anti-oxidant and antiinflammatory properties [9-11], while iron is known to regulate transferrin, ferritin and nitric oxide synthase gene expression [12,13]. An up regulation in the synthesis of transferrin and ferritin enhances the binding, transport and storage of iron thus serving as an important control mechanism against the oxidative effects of iron. CO, like NO, has been accepted as an important cellular signalling molecule in biological systems. For example, both NO and CO are known to activate soluble guanylyl cyclase, resulting in elevated cGMP and the cGMP-mediated dilatation of blood vessels. CO also mediates vasodilation by directly activating the calcium-dependent potassium channels in vascular smooth muscle cells [14,15]. In addition, CO inhibits platelet aggregation and proliferation of vascular smooth muscle cells, inhibits apoptosis, and stimulates angiogenesis [16-19]. Because of the diversity in the effects of heme catabolism, several studies have suggested that induction of HO-1 expression by oxidising agents may serve as a defense mechanism against oxidative stress in vivo. For example, an induction in the expression of HO-1 prevents superoxide associated endothelial cell sloughing in diabetic rats [20]. Similarly, it has been demonstrated that ONOO-, a potent oxidizing agent generated by the interaction of NO and superoxide radical [21], causes a concentration-dependent increase in HO-1 protein expression and enzyme activity in rat aortic endothelial cells [22-24], as well as human colorectal adenocarcinoma cells [25]. These studies indicate that ONOO regulates the expression of HO-1 and that the heme oxygenase pathway contributes to protection against the cytotoxic effects of ONOO-which are due to its reactivity with cellular macromolecules such as the covalent modification of tyrosine, cysteine, methionine or tryptophan residues, oxidation of nucleic bases or the scavenging of cellular antioxidants such ascorbate and urate [26]. It is proposed that the ONOO-mediated HO-1 induction might occur via an interactive signalling mechanism that modulates oxidative stress responses but direct effects of ONOO on HO activity have not been studied. In this work, we examined the effect of ONOOon HO catalytic activity in rat spleen and brain microsomes respectively.

Results

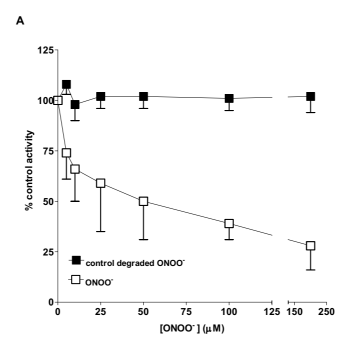
Effect of ONOO- and TNM on Microsomal HO activity

Exposure of either rat spleen or brain microsomes to ONOO resulted in decreases of HO activity in a concentration dependent manner (Figure 1A and Figure 1B). HO-1 (spleen microsomes) was more sensitive towards ONOO-mediated inactivation compared to HO-2 (brain microsomes) (Figure 1A and Figure 1B). The IC₅₀ values for inhibition of HO activity in rat spleen and brain microsomes were 0.015 ± 0.005 mM and 1.25 ± 0.25 mM, respectively. The same concentration of degraded ONOOdid not show any effects on HO activity in rat spleen or brain microsomal fractions. Maximal inhibition for microsomal HO-1 and HO-2 activity was approximately 70%, beyond which there was no further inhibitory effect with increasing concentrations of ONOO. To investigate whether effects of ONOO were due to nitration of tyrosine residues, spleen and brain microsomal fractions were treated with TNM, a known tyrosine-nitrating agent. It is shown that HO activity in rat spleen as well as brain microsomal fractions was significantly reduced by TNM albeit at higher concentrations compared to ONOO- (Figure 2).

Effects of sulfhydryl modifying reagents on HO activity

Since ONOO may affect protein targets by the oxidation of sulfhydryl groups as well as the nitration of tyrosine residues, we examined and compared the effect of other sulfhydryl modifying reagents on HO activity. Rat spleen or brain microsomal protein was treated with different sulfhydryl modifying reagents such as GSNO, DEA-NON-Oate, NEM and H2O2, and a combination of equimolar concentrations of ONOO- and GSNO or ONOO- and DEA-NONOate. We found that GSNO caused a concentration dependent inactivation of both HO-1 and HO-2. Consistent with the effect of ONOO-, GSNO was more active against HO-1 than HO-2 (Table 1). In contrast, DEA-NONOate and NEM caused a significant decrease in the catalytic activity of HO-1 at the two concentrations tested (0.2 mM and 2 mM), but did not inhibit HO-2 (Table 1). Pre-treatment with equimolar concentrations of ONOO- and DEA-NONOate or GSNO did not result in further inactivation of HO-1 or HO-2 than ONOO alone. H₂O₂ at concentrations as high as 2 mM had no significant effect on the catalytic activity of either isozymes.

To test whether effects of ONOO and the sulfhydryl modifying reagents on HO-1 and HO-2 are reversible, ONOO and DEA-NONOate were pre-incubated with rat spleen and brain microsomal protein for 20, 60 and 120 minutes prior to evaluation of HO activity. The inactivation of HO-1 by ONOO and DEA-NONOate and HO-2 by ONOO was reversible following prolonged pre-incubation time (60–120 minutes) Table 2. However, total HO activity in



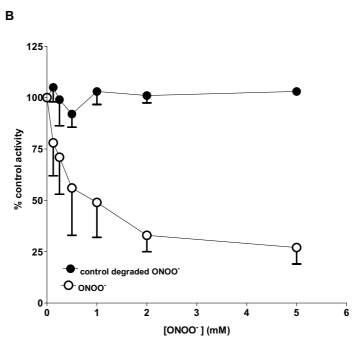


Figure I ONOO--mediated inactivation of HO-1 and HO-2. Rat spleen (Figure 1A) and brain microsomes (Figure 1B) (50–100 μg protein) were treated with indicated concentrations of ONOO or degraded ONOO in 100 mM potassium phosphate, pH 7.4 at room temperature for 10 seconds. The reaction was stopped by dilution of the reaction mixture to a protein concentration of (0.5 mg/mL) spleen microsomes and (1 mg/mL) brain microsomes in 100 mM phosphate buffer containing 1 mM NADPH and 50 μM methemalbumin. Incubations of the pretreated microsomal fractions were performed for 15 min and enzyme activity was determined by the quantitation of CO formed in the reaction mixture. Data are presented as the mean \pm SD of triplicate experiments. The rates of CO formation in the control reactions were 12 \pm 1 and 5 \pm 1 pmoles CO/min/mg protein for spleen and brain microsomes respectively. IC₅₀ values for HO-1 and HO-2 were 0.015 \pm 0.005 mM and 1.25 \pm 0.25 mM respectively.

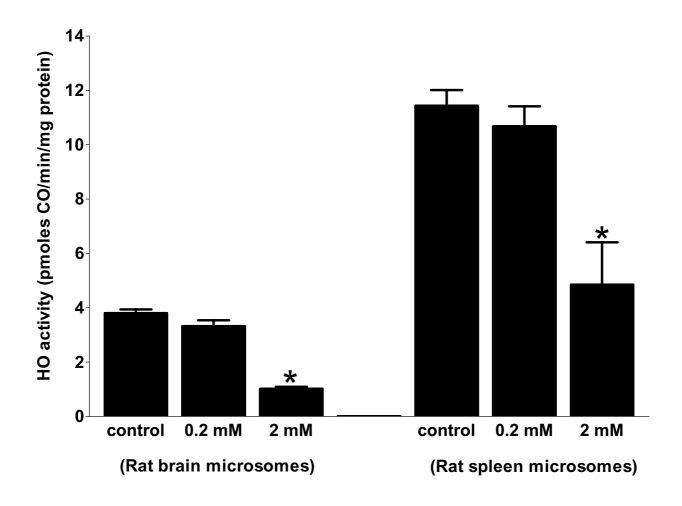


Figure 2 Effect of TNM-mediated nitration on the catalytic activity of HO. HO-I (rat spleen microsomes and HO-2 (rat brain microsomes). Microsomal protein (50–100 μg) was incubated with 0.2 mM or 2 mM (final concentration) of TNM in 100 mM potassium phosphate buffer, pH 7.4, at 37°C for 20 minutes. The reaction was stopped by dilution of the reaction mixture to a protein concentration of (0.5 mg/mL) in 100 mM phosphate buffer containing I mM NADPH and 50 μM methemalbumin. Incubations of the pretreated microsomal protein were done for 15 min and enzyme activity was determined by the quantitation of CO formed in the reaction mixture. Data are presented as the mean \pm SD of triplicate experiments. The rate of CO formation in control reactions was 11.4 \pm 0.6 and 4.0 \pm 0.14 pmoles CO/min/mg protein for HO-1 and HO-2 respectively. The asterisk denotes significant inhibition of the respective HO activity using a one-way ANOVA, P \leq 0.05.

brain and spleen microsomal protein was not affected by the prolonged pre-incubation for 60–120 minutes in the absence of ONOO and DEA-NONOate.

Discussion

Production of ONOO in vivo is a consequence of oxidative and nitrosative stress, and ONOO-mediated tissue injury is thought to be involved in the pathogenesis of

many conditions including atherosclerosis, ischaemia/reperfusion, shock, Alzheimer's disease, diabetes and multiple sclerosis [27-30]. Continuous challenge and exposure to oxidative/nitrosative stressors has led to the evolution of numerous defense mechanisms in biological systems and one such mechanism that has been elucidated by many researchers is the HO/CO system [4,9-11]. ONOO- causes a concentration-dependent increase in

Table I: Effect of NO donors and sulfhydryl modifying reagents on the catalytic activity of HO-I (spleen microsomes) and HO-2 (brain microsomes)

Heme oxygenase activity (pmoles CO/min/mg protein)						
Experimental conditions		HO-I (Spleen microsomes)	HO-2 (Brain microsomes)			
Control		11.9 ± 0.6	3.1 ± 0.5			
ONOO-						
(0.2 mM	5.4 ± 0.9*	2.7 ± 0.6			
	2 mM	3.8 ± 0.9*	1.4 ± 0.3*			
GSNO						
	0.2 mM	6.5 ± 0.3*	2.9 ± 0.5			
	2 mM	4.2 ± 0.2*	1.4 ± 0.1*			
DEA-NONOate						
	0.2 mM	4.7 ± 0.4*	2.9 ± 0.7			
	2 mM	3.6 ± 0.4*	2.4 ± 0.5			
NEM						
	0.2 mM	8.7 ± 0.8	2.9 ± 0.3			
	2 mM	3.4 ± 0.5*	2.6 ± 0.4			
H ₂ O ₂						
	0.2 mM	11.3 ± 0.2	3.0 ± 0.4			
	2 mM	11.8 ± 1.5	3.2 ± 0.6			
ONOO- and GSNO						
	0.2 mM	4.3 ± 0.6*	2.8 ± 0.4			
	2 mM	4.0 ± 0.2*	1.3 ± 0.2*			
(ONOO- and DEA-NONOate)						
,	0.2 mM	4.7 ± 0.1*	2.6 ± 0.2			
	2 mM	3.9 ± 0.6*	1.4 ± 0.3 *			

Microsomal protein (50–100 μ g) was incubated with 0.2 mM or 2 mM (final concentration) of ONOO-, GSNO, DEA-NONOate, NEM, H_2O_2 , equimolar concentrations of ONOO- and GSNO or ONOO- and DEA-NONOate in 100 mM potassium phosphate buffer, pH 7.4, at 37°C for 20 minutes. The reaction was stopped by dilution of the reaction mixture to a protein concentration of (0.5 mg/mL) in 100 mM phosphate buffer containing 1 mM NADPH and 50 μ M methemalbumin. Incubations of the pretreated microsomal fractions were done for 15 min and enzyme activity was determined by the quantitation of CO formed in the reaction mixture. Data are presented as the mean \pm SD of triplicate experiments. Asterisks denote significant inhibition of the respective HO activity using a one-way ANOVA, $P \le 0.05$.

HO-1 protein expression suggesting that the HO pathway contributes to protection against the cytotoxic effects of ONOO-[22-24]. Most studies in this field, however, have focussed on the induction of HO-1 mRNA and/or protein expression under different conditions of oxidative/nitrosative stress rather than enzyme activity. Considering the cellular toxicity of ONOO- and the inhibitory effect of ONOO- on numerous enzyme systems, we sought to investigate the direct effect of ONOO- on the catalytic activity of two microsomal HO isozymes.

Rat spleen and brain microsomal fractions were used because of the predominant expression of HO-1 in the spleen and HO-2 in the brain. We have shown that ONOO-inhibits the activity of both HO-1 and HO-2 in a concentration-dependent manner (Figure 1). HO-1 was found to be more sensitive to ONOO- treatment compared to HO-2. Lower concentrations of ONOO- (15 μ M, final concentration) decreased HO-1 activity by 50% while a much higher concentration (1.25 mM) was required to decrease HO-2 activity by the same magni-

tude. Generally, HO catalytic function is dependent on an accessory enzyme NADPH-cytochrome P450 reductase (CPR), which serves as a redox partner during the oxidative break down of heme and the conversion of NADPH to NADP+. It is possible therefore, that ONOO- may have altered HO activity indirectly by inactivating CPR. The ONOO- dependent inactivation of recombinant CPR in bacterial membranes has been noted [32], but our attempt to supplement rat microsomal HO-1 and HO-2 with recombinant rat CPR in the presence of a detergent did not attenuate the effect of ONOO- on microsomal HO-1 and HO-2 (data not shown). This suggests that the effects of ONOO on HO activity may be mediated by mechanisms that are independent of the effect of ONOOon microsomal CPR activity. The differential effect of ONOO on the catalytic activity of HO-1 and HO-2 may suggest a selective mechanism on the catalytic function of the inducible enzyme. This may have direct implications in biological systems where ONOO is generated at high concentrations. The ONOO-mediated increase in HO-1 gene and/or protein expression may be a cellular response at the acute phase. Subsequently, total HO activity may then be maintained in certain ranges in order to retain intracellular oxidative responses and the balance of redox states for normal cellular function. This idea is consistent with other studies, which have shown that induction of HO-1 mRNA and protein expression is not always followed by a proportionate increase in catalytic function. For example, by using quantitative RT-PCR it was found that a 3.8-fold increase in sarcoma-induced HO-1 mRNA expression yields only 2.1-fold increase in total HO activity [31]. Overall, the ONOO-mediated regulation of the catalytic function of HO-1 and differences in the sensitivity of HO-1 and HO-2 may be attributed to disparity in the amino acid compositions and the structure of the two proteins.

Mechanistically, the effects of ONOO on protein function are mainly due to its reactivity and covalent modification of tyrosine and/or cysteine residues [26]. This often leads to impaired protein function with very few exceptions such as the recently observed nitration and stimulation of the enzymatic activity of microsomal glutathione S-transferase (MGST) [33]. To probe mechanisms of ONOOmediated inhibition of HO activity observed here, we examined the effects of TNM, another tyrosine-nitrating chemical compound on rat spleen and brain microsomal HO activity. Significant inhibition of HO catalytic activity by TNM was observed for spleen as well as brain microsomal fractions albeit at higher concentrations compared to ONOO (Figure 2). This suggests that both TNM and ONOO- may inactivate HO activity by nitration of tyrosine residue(s) that is/are important for conservation of HO-1 and HO-2 function. This is consistent with data from the analysis of the complete amino acid sequences of rat and human HO-1 [34,35] and HO-2 [36,37]. However, there was no difference in the effects of TNM on HO-1 and HO-2 (Figure 2), suggesting that the differential effects observed with ONOO could be due to modification of amino acid residues other than tyrosine. Oxidation of cysteine residues as a putative mechanism underlying ONOO-mediated inactivation of HO was investigated. Effects of GSNO, DEA-NONOate, NEM and H₂O₂, or a combination of equimolar concentrations of ONOO and GSNO or DEA-NONOate on HO activity were tested. GSNO showed similar inhibitory effects to ONOO-but a combination of equimolar concentrations of ONOO and DEA-NONOate or GSNO did not have any additive effect on the ONOO-mediated inactivation of HO-1 or HO-2. In addition, the inactivation of HO-1 by ONOO and DEA-NONOate and HO-2 by ONOO were reversible following prolonged pre-incubation time (60–120 minutes) Table 2. This differential effect between HO-1 and HO-2 following exposure to ONOO-, GSNO, DEA-NONOate or NEM is suggestive of qualitative as well as quantitative differences in the distribution of target amino acids such as

critical tyrosine and/or cysteine residues in the tertiary structures of these proteins. The amino acid sequences of human and rat HO-2 reveals a conserved core of cysteine residues (Cys 264-Pro 265, Cys 281-Pro 282) [38,39], but there are no cysteine residues in the HO-1 amino acid sequence. Despite this fact, results from our experiments show that HO-1 is more sensitive to inactivation by the NO donors GSNO and DEA-NONOate, and NEM (Table 1). We considered the possibility that ONOO might exert its effects by interaction with the substrate as has been reported for the interaction of NO with heme [40]. This possibility seems unlikely as such a mechanism would be expected to affect the catalytic rate of HO-1 and HO-2 equally. Another consideration is that the chemistry of ONOO- is richer than is usually discussed; in their review Alvarez and Radi [41] describe the interactions of ONOOwith a number of functional groups that are not often mentioned in the consideration of the interactions of ONOO- with various proteins. Thus, ONOO- inhibition of HO activity could be a result of an interaction with amino acids other than cysteine or tyrosine. This possibility of ONOO interactions with other amino acids may also explain the reversibility of the inhibition of HO activity by ONOO-. Furthermore, it is possible that ONOO-inhibition of HO-1 and HO-2 occurs courtesy of different facets of ONOO chemistry. Differences in the relative potency of DEA-NONOate, NEM, GSNO and ONOO may also be attributed to differences in the chemical properties and the electrophilic potentials of these reagents. For instance, the nitric oxide radical (NO•) generated from DEA-NON-Oate is less reactive and unstable compared to NO+ produced from GSNO.

Collectively, our data indicate that ONOO may be important in the cross-talk between HO and NO systems. While tyrosine residues for HO-1 and/or sulfhydryl groups for HO-2 should be considered as potential targets for ONOO interactions, other amino acids should be studied in the elucidation of mechanism(s) of ONOO action on HO isozymes.

Conclusion

This study documented for the first time the ONOO-mediated inactivation of HO-1 and HO-2. The IC_{50} for HO-1 was approximately 80-fold less than that for HO-2. While conserved tyrosine residues (HO-1) and/or sulfhydryl group(s) (HO-2) may play a critical role in maintaining the functional capacity of heme oxygenases, the consideration of other amino acids is suggested. In addition, the higher sensitivity of HO-1 to ONOO-mediated inactivation may indicate dual regulatory mechanisms on the catalytic function of the inducible enzyme under the conditions of oxidative/nitrosative stress.

Table 2: Time dependent reversibility of the effect of ONOO and DEA-NONOate on the catalytic activity of HO-I (spleen microsomes) and HO-2 (brain microsomes)

		Heme oxygenase activity (pmoles CO/min/mg protein) at different princubation times			
Experimental conditions		20 min	60 min	I20 min	
Spleen microsomes (HO-I)	ONOO-				
	Control	10.4 ± 1.0	10.8 ± 0.8	8.7 ± 0.9	
	0.2 mM	3.6 ± 1.4*	10.5 ± 0.8	8.3 ± 1.1	
	DEA-NONOate				
	Control	10.6 ± 2.0	11.5 ± 1.3	9.4 ± 0.2	
	0.2 mM	6.6 ± 0.7*	11.1 ± 1.4	8.9 ± 0.3	
Brain microsomes (HO-2)	ONOO-				
	Control	3.6 ± 0.5	4.2 ± 0.4	4.4 ± 0.3	
	2 mM	1.9 ± 0.1*	4.4 ± 0.3	3.8 ± 0.3	
	DEA-NONOate				
	Control	3.6 ± 0.3	4.1 ± 0.3	4.0 ± 0.3	
	2 mM	3.0 ± 0.3	3.9 ± 0.1	3.6 ± 0.3	

Microsomal protein (50–100 μ g) was incubated with 0.2 mM or 2 mM (final concentration) of ONOO and DEA-NONOate in 100 mM potassium phosphate buffer, pH 7.4, at 37°C for 20, 60 and 120 minutes. The reaction was stopped by dilution of the reaction mixture to a protein concentration of (0.5 mg/mL) in 100 mM phosphate buffer containing 1 mM NADPH and 50 μ M methemalbumin. Incubations of the pretreated microsomal fractions were done as described in materials and methods. Data are presented as the mean \pm SD of triplicate experiments. Asterisks denote significant inhibition of the respective HO activity using a one-way ANOVA, P \leq 0.05.

Methods

Materials

Hemin, ethanolamine, bovine serum albumin (BSA), β -NADPH, N-ethylmaleimide (NEM), tetra-nitromethane (TNM) and reduced glutathione (GSH) were purchased from Sigma Chemical Co. (St. Louis Mo.). Diethylamine NONOate (DEA-NONOate) was obtained from Calbiochem Inc. (Darmstadt, Germany). All other chemicals were reagent grade from a variety of commercial sources.

Preparation of rat spleen and brain microsomes

Adult male Sprague-Dawley rats (250–300 g) were obtained from Charles River Canada Inc. (St-Constant, Que). The animals were cared for in accordance with the principles and guidelines of the Canadian Council on Animal Care. Microsomal fractions were prepared according to previously described procedures [42,43]. Spleen microsomes were used as a source of HO-1 while brain microsomes were used as a source of HO-2.

Preparation of ONOO- and treatment of microsomal protein

ONOO- was synthesised from acidified nitrite and $\rm H_2O_2$ according to the method of Beckman [44] and microsomal HO-1 (spleen microsomes) and HO-2 (brain microsomes) were treated with ONOO-following the procedures described by Ji and Bennett [33]. Briefly, microsomes (50 –100 µg protein in 100 mM potassium phosphate, pH 7.4) were exposed to ONOO-at room temperature at indicated concentrations for 10 seconds by

adding a small volume during vigorous mixing. HO activity was initiated by diluting the reaction mixture to a protein concentration of (0.5–1 mg/ml) in 100 mM phosphate buffer containing substrates (50 μM methemalbumin, and 1 mM NADPH). To control for the potential effect of nitrite and nitrate that may be formed during the incubation of ONOO-, ONOO- was allowed to decompose in phosphate buffer prior to incubation with microsomal protein and determination of enzyme activity for some of the reactions.

Treatment of microsomes with NO donors, TNM and sulfhydryl modifying reagents

GSNO was prepared by reacting equimolar amounts of sodium nitrite and GSH according to the method described by Ji *et al.*, [43]. Microsomal protein (50 – 100 μ g) was incubated with 0.2–2 mM GSNO, DEA-NON-Oate, NEM or H₂O₂ in 100 mM potassium phosphate buffer, pH 7.4, at 37 °C for 20 minutes. The reaction was stopped by dilution of the reaction mixture to a protein concentration of (0.5–1 mg/mL) in 100 mM phosphate buffer containing (50 μ M methemalbumin, and 1 mM NADPH).

In vitro Assay for HO activity

HO activity in rat spleen and brain microsomal fractions was determined by the quantitation of CO formed from the oxygen, CPR and NADPH-dependent degradation of methemalbumin as previously described [45,46]. The reaction mixture contained, (0.5–1 mg/mL microsomal

protein, 50 μ M methemalbumin, and 1 mM NADPH) in 100 mM phosphate buffer. All incubations for the assay of HO activity were performed under the conditions for which the rate of CO formation (pmol CO. mg⁻¹ protein. min⁻¹) was linear with respect time and microsomal protein concentration. CO formation was monitored by gas chromatography according to the method described by Vreman and Stevenson [46].

Statistical analysis

Data are presented as the mean \pm SD from triplicate experiments and statistical analyses were performed by one-way ANOVA. P values ≤ 0.05 was considered to be significant.

Abbreviations

HO, heme oxygenases; CPR, NADPH-cytochrome P450 reductase; CO, carbon monoxide; ONOO-, peroxynitrite; GSNO, S-nitrosoglutathione; DEA-NONOate, diethylamine NONOate; NEM, N-ethylmaleimide; TNM, tetranitromethane; ROS/RNS, reactive oxygen/nitrogen species; RT-PCR; reverse transcriptase polymerase chain reaction; H₂O₂, hydrogen peroxide.

Authors' contributions

RK, YJ and KN were involved in the experimental design and data collection. All authors read and approved the final manuscript.

Acknowledgements

This work was supported by grant MOP 64305 from the Canadian Institutes of Health Research.

References

- Prabhakar NR: Endogenous carbon monoxide in control of respiration. Respir Physiol 1998, 114:57-64.
- Maines MD: The heme oxygenase system: a regulator of second messenger gases. Annu Rev Pharmacol Toxicol 1997, 37:517-554.
- Hancock WW, Buelow R, Sayegh MH, Turka LA: Antibodyinduced transplant arteriosclerosis is prevented by graft expression of anti-oxidant and anti-apoptotic genes. Nat Med 1998, 4:1392-1396.
- Soares MP, Lin Y, Anrather J, Csizmadia E, Takigami K, Sato K, Grey ST, Colvin RB, Choi AM, Poss KD, Bach FH: Expression of heme oxygenase-1 can determine cardiac xenograft survival. Nat Med 1998, 4:1073-1077.
- Zakhary R, Poss KD, Jaffrey SR, Ferris CD, Tonegawa S, Snyder SH: Targeted gene deletion of heme oxygenase 2 reveals neural role for carbon monoxide. Proc Natl Acad Sci U S A 1997, 94:14848-14853.
- Ewing JF, Weber CM, Maines MD: Biliverdin reductase is heat resistant and coexpressed with constitutive and heat shock forms of heme oxygenase in brain. J Neurochem 1993, 61:1015-1023.
- Trakshel GM, Kutty RK, Maines MD: Resolution of the rat brain heme oxygenase activity: absence of a detectable amount of the inducible form (HO-1). Arch Biochem Biophys 1988, 260-732-739
- McCoubrey WK Jr, Huang TJ, Maines MD: Isolation and characterization of a cDNA from the rat brain that encodes hemoprotein heme oxygenase-3. Eur J Biochem 1997, 247:725-732.
- 9. Nakagami T, Toyomura K, Kinoshita T, Morisawa S: A beneficial role of bile pigments as an endogenous tissue protector:

- anti-complement effects of biliverdin and conjugated bilirubin. Biochim Biophys Acta 1993, 1158:189-193.
- Willis D, Moore AR, Frederick R, Willoughby DA: Heme oxygenase: a novel target for the modulation of the inflammatory response. Nat Med 1996, 2:87-90.
- Wunder C, Potter RF: The heme oxygenase system: its role in liver inflammation. Curr Drug Targets Cardiovasc Haematol Disord 2003, 3:199-208.
- Pantopoulos K, Hentze MW: Nitric oxide signaling to iron-regulatory protein: direct control of ferritin mRNA translation and transferrin receptor mRNA stability in transfected fibroblasts. Proc Natl Acad Sci U S A 1995, 92:1267-1271.
- Chen L, Wang Y, Kairaitis LK, Wang Y, Zhang BH, Harris DC: Molecular mechanisms by which iron induces nitric oxide synthesis in cultured proximal tubule cells. Exp Nephrol 2001, 9:198-204.
- Furchgott RF, Jothianandan D: Endothelium-dependent and independent vasodilation involving cyclic GMP: relaxation induced by nitric oxide, carbon monoxide and light. Blood Vessels 1991, 28:52-61.
- Wang R, Wu L, Wang Z: The direct effect of carbon monoxide on KCa channels in vascular smooth muscle cells. Pflugers Arch 1997, 434:285-291.
- 16. Durante W, Schafer Al: Carbon monoxide and vascular cell function (Review). Int J Mol Med 1998, 2:255-262.
 17. Liu XM, Chapman GB, Peyton KJ, Schafer Al, Durante W: Carbon
- Liu XM, Chapman GB, Peyton KJ, Schafer AI, Durante W: Carbon monoxide inhibits apoptosis in vascular smooth muscle cells. Cardiovasc Res 2002, 55:396-405.
- Brouard S, Otterbein LE, Anrather J, Tobiasch E, Bach FH, Choi AM, Soares MP: Carbon monoxide generated by heme oxygenase I suppresses endothelial cell apoptosis. J Exp Med 2000, 192:1015-1026.
- Sato K, Balla J, Otterbein L, Smith RN, Brouard S, Lin Y, Csizmadia E, Sevigny J, Robson SC, Vercellotti G, Choi AM, Bach FH, Soares MP: Carbon monoxide generated by heme oxygenase-I suppresses the rejection of mouse-to-rat cardiac transplants. J Immunol 2001, 166:4185-4194. 2001 Mar 15
- Quan S, Kaminski PM, Yang L, Morita T, Inaba M, Ikehara S, Goodman AI, Wolin MS, Abraham NG: Heme oxygenase-I prevents superoxide anion-associated endothelial cell sloughing in diabetic rats. Biochem Biophys Res Commun 2004, 315:509-516.
- Pryor WA, Squadrito GL: The chemistry of peroxynitrite: a product from the reaction of nitric oxide with superoxide. Am J Physiol 1995, 268:2699-2722.
- Foresti R, Sarathchandra P, Clark JE, Green CJ, Motterlini R: Peroxynitrite induces haem oxygenase-1 in vascular endothelial cells: a link to apoptosis. Biochem J 1999, 339:729-736.
- 23. Foresti R, Clark JE, Green CJ, Motterlini R: Thiol compounds interact with nitric oxide in regulating heme oxygenase-I induction in endothelial cells. Involvement of superoxide and peroxynitrite anions. J Biol Chem 1997, 272:18411-18417.
- 24. Motterlini R, Foresti R, Bassi R, Calabrese V, Clark JE, Green CJ: Endothelial heme oxygenase-I induction by hypoxia. Modulation by inducible nitric-oxide synthase and S-nitrosothiols. J Biol Chem 2000, 275:13613-20.
- Hara E, Takahashi K, Takeda K, Nakayama M, Yoshizawa M, Fujita H, Shirato K, Shibahara S: Induction of heme oxygenase-I as a response in sensing the signals evoked by distinct nitric oxide donors. Biochem Pharmacol 1999, 58:227-236.
- Ducrocq C, Blanchard B, Pignatelli B, Ohshima H: Peroxynitrite: an endogenous oxidizing and nitrating agent. Cell Mol Life Sci 1999, 55:1068-1077.
- 27. Keynes RG, Garthwaite J: Nitric oxide and its role in ischaemic brain injury. Curr Mol Med 2004, 4:179-191.
- Li J, Li W, Su J, Liu W, Altura BT, Altura BM: Peroxynitrite induces apoptosis in rat aortic smooth muscle cells: possible relation to vascular diseases. Exp Biol Med (Maywood) 2004, 229:264-269.
- Good PF, Werner P, Hsu A, Olanow CW, Perl DP: Evidence of neuronal oxidative damage in Alzheimer's disease. Am J Pathol 1996, 149:21-28.
- Szabo C: DNA strand breakage and activation of poly-ADP ribosyltransferase: a cytotoxic pathway triggered by peroxynitrite. Free Radic Biol Med 1996, 21:855-869.
- 31. McAllister SC, Hansen SG, Ruhl RA, Raggo CM, DeFilippis VR, Greenspan D, Fruh K, Moses AV: Kaposi sarcoma-associated herpesvirus (KSHV) induces heme oxygenase-I expression and

- activity in KSHV-infected endothelial cells. *Blood* 2004, 103:3465-73.
- Sergeeva SV, Slepneva IA, Khramtsov VV: Effect of selenolipoic acid on peroxynitrite-dependent inactivation of NADPHcytochrome P450 reductase. Free Radic Res 2001, 35:491-497.
- Ji Y, Bennett BM: Activation of microsomal glutathione s-transferase by peroxynitrite. Mol Pharmacol 2003, 63:136-146.
- 34. Shibahara S, Muller R, Taguchi H, Yoshida T: Cloning and expression of cDNA for rat heme oxygenase. Proc Natl Acad Sci U S A 1985, 82:7865-7869.
- 35. Shibahara S, Sato M, Muller RM, Yoshida T: **Structural organization of the human heme oxygenase gene and the function of its promoter.** *Eur J Biochem* 1989, **179:**557-563.
- Cruse I, Maines MD: Evidence suggesting that the two forms of heme oxygenase are products of different genes. J Biol Chem 1988, 263:3348-3353.
- McCoubrey WK Jr, Ewing JF, Maines MD: Human heme oxygenase-2: characterization and expression of a full-length cDNA and evidence suggesting that the two HO-2 transcripts may differ by choice of polyadenylation signal. Arch Biochem Biophys 1992, 295:13-20.
- 38. Ding Y, McCoubrey WK Jr, Maines MD: Interaction of heme oxygenase-2 with nitric oxide donors. Is the oxygenase an intracellular 'sink' for NO? Eur J Biochem 1999, 264:854-861.
- 39. Huang TJ, McCoubrey WK Jr, Maines MD: Heme oxygenase-2 interaction with metalloporphyrins: function of heme regulatory motifs. Antioxid Redox Signal 2001, 3:685-696.
- Juckett M, Zheng Y, Yuan H, Pastor T, Antholine W, Weber M, Vercellotti G: Heme and the endothelium. Effects of nitric oxide on catalytic iron and heme degradation by heme oxygenase. J Biol Chem 1998, 273:23388-23397.
- Alvarez B, Radi R: Peroxynitrite reactivity with amino acids and proteins. Amino Acids 2003, 25:295-311.
- Appleton SD, Chretien ML, McLaughlin BE, Vreman HJ, Stevenson DK, Brien JF, Nakatsu K, Maurice DH, Marks GS: Selective inhibition of heme oxygenase, without inhibition of nitric oxide synthase or soluble guanylyl cyclase, by metalloporphyrins at low concentrations. Drug Metab Dispos 1999, 27:1214-1219.
- 43. Ji Y, Akerboom TP, Sies H: Microsomal formation of S-nitrosoglutathione from organic nitrites: possible role of membrane-bound glutathione transferase. Biochem J 1996, 313:377-380.
- Beckman JS, Chen J, Ischiropoulos H, Crow JP: Oxidative chemistry of peroxynitrite. Methods Enzymol 1994, 233:229-240.
- Cook MN, Nakatsu K, Marks GS, McLaughlin BE, Vreman HJ, Stevenson DK, Brien JF: Heme oxygenase activity in the adult rat aorta and liver as measured by carbon monoxide formation. Can J Physiol Pharmacol 1995, 73:515-518.
- Vreman HJ, Stevenson DK: Heme oxygenase activity as measured by carbon monoxide production. Anal Biochem 1988, 168:31-38.

Publish with **Bio Med Central** and every scientist can read your work free of charge

"BioMed Central will be the most significant development for disseminating the results of biomedical research in our lifetime."

Sir Paul Nurse, Cancer Research UK

Your research papers will be:

- available free of charge to the entire biomedical community
- peer reviewed and published immediately upon acceptance
- cited in PubMed and archived on PubMed Central
- ullet yours you keep the copyright

Submit your manuscript here: http://www.biomedcentral.com/info/publishing_adv.asp

