Nitrosative protein oxidation is modulated during early endotoxemia

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\textbf{A R T I C L E I N F O}

\textbf{Article history:}
Available online 2 December 2010

\textbf{Keywords:}
Sepsis
Lipopolysaccharide
Nitrosative
S-nitrosylation
Nitration
Blood pressure

\textbf{A B S T R A C T}

Formation of nitric oxide and its derivative reactive nitrogen species during endotoxemia has been implicated in the pathogenesis of the associated cardiovascular dysfunction. This stress can promote nitrosative post-translational modifications of proteins that may alter their activity and contribute to dysregulation. We utilized the ascorbate-dependent biotin-switch method to assay protein S-nitrosylation and immunoblotted for tyrosine nitration to monitor changes in nitrosative protein oxidation during endotoxemia. Hearts from lipopolysaccharide (LPS)-treated rats showed no apparent variation in global protein S-nitrosylation, but this may be due to the poor sensitivity of the biotin-switch method. To sensitize our monitoring of protein S-nitrosylation we exposed isolated hearts to the efficient trans-nitrosylating agent nitrosocysteine (which generated a robust biotin-switch signal) and then identified a number of target proteins using mass spectrometry. We were then able to probe for these target proteins in affinity-capture preparations of S-nitrosylated proteins prepared from vehicle- or LPS-treated animals. Unexpectedly this showed a time-dependent loss in S-nitrosylation during sepsis, which we hypothesized, may be due to concomitant superoxide formation that may lower nitric oxide but simultaneously generate the tyrosine-nitrating agent peroxynitrite. Indeed, this was confirmed by immunoblotting for global tyrosine nitration, which increased time-dependently and temporally correlated with a decrease in mean arterial pressure. We assessed if tyrosine nitration was causative in lowering blood pressure using the putative peroxynitrite scavenger FeTPPS. However, FeTPPS was ineffective in reducing global protein nitration and actually exacerbated LPS-induced hypotension.

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\textbf{Introduction}

Sepsis and endotoxemia is associated with systemic inflammation that injures and compromises the function of tissues, including those of the cardiovascular system [1]. Given the prevalence, high risk of mortality and financial healthcare burden associated with sepsis [2], it is important to define its pathogenesis at the molecular level. Cardiovascular dysfunction during sepsis includes hypotension, decreased contractility and decreased myocardial compliance. Alterations in circulating factors including prostanoids, cytokines and nitric oxide (NO) are thought to contribute to this etiology [3,4]. NO and its related derivatives may have causative roles in compromising cardiovascular performance during sepsis [5].

NO is a major regulator of cardiovascular function; controlling blood pressure [6], muscle tone as well as platelet aggregation [7,8]. However, during inflammation NO production is drastically enhanced by NO synthase (NOS) stimulation, as well as by increased inducible NOS (iNOS) expression and activity triggered by circulating endotoxins and cytokines [9]. Enhanced nitric oxide production can deplete substrates needed for its biosynthesis resulting in uncoupling and superoxide formation [10,11]. Inflammatory cells such as neutrophils and macrophages may also respond to endotoxins by generating reactive oxygen species (ROS) as well as reactive nitrogen species (RNS) [12]. Simultaneous formation of both superoxide and nitric oxide can lead to the generation of peroxynitrite (ONOO\(^{-}\)), a RNS that can oxidize protein thiols and nitrate tyrosine residues [13]. Increased tyrosine nitration, which may occur during times of nitroxidative stress such as sepsis [14], can be monitored using pan-specific antibodies to this modification [15]. A causative role for nitroxidative species, and so perhaps the nitroxidative protein modifications, in the pathogenesis of sepsis is supported by scavengers of OONO\(^{-}\) and other oxidants attenuating dysfunction [16–19].

Covalent adduction of NO to protein cysteinyi thiols (termed S-nitrosylation) has emerged as an important regulator of cell signaling [20,21]. Protein S-nitrosylation can be induced by small
nitrosothiol containing compounds such as nitrosoglutathione (GSNO), nitrosocysteine (CysNO) [22,23], and specific reactive nitrogen species such as the nitrosionium ion (NO+) and dinitrogen trioxide (N₂O₃) [24]. S-nitrosylation regulates the activity of many proteins including caspases, kinases, Hsp90 and the ryanodine receptor [25,26]. Whilst the role of S-nitrosylation in sepsis is poorly understood, mice null for S-nitrosogluthathione reductase have enhanced global S-nitrosylation and an increased susceptibility to endotoxin mediated mortality [27]. Furthermore plasma nitrosothiol concentration increase during sepsis [28].

Here we assessed changes in myocardial nitrosative (S-nitrosylation and tyrosine nitration) protein modifications in a lipopolysaccharide (LPS) murine model of sepsis. We found no evidence for increased global protein S-nitrosylation, which may have been anticipated as outlined above. In fact, a more sensitive approach to measuring S-nitrosylation of specific proteins showed sepsis actually decreased this modification below controls. Concomitantly, however there was an increase in global protein tyrosine nitration which correlated temporally with systemic hypotension in LPS-treated mice.

Materials and methods

All procedures were performed in accordance with the Home Office Guidance on the Operation of the Animals (Scientific Procedures) Act 1986 in UK.

Analysis of S-nitrosylation in isolated HEK cells and perfused rat hearts using the ascorbate-dependent biotin-switch method

Confluent human embryonic kidney 293 (HEK293) cell were treated with phosphate buffered saline (PBS) alone or PBS containing either 300 μM CysNO, 100 μM hydrogen peroxide or 500 μM diamide for 10 min. Langendorff perfused isolated rat hearts were stabilized for 30 min before being perfused with Krebs buffer alone or Krebs buffer containing either 10 μM CysNO, 50 μM S-nitroso- N-acetyl-DL-penicillamine (SNAP) or 500 μM SNAP for 20 min at constant flow. After perfusion hearts were immediately snap frozen in liquid nitrogen.

Hearts and treated HEK293 cells were processed using a modified version of the biotin-switch method to determine the level of protein S-nitrosylation [29]. In brief, hearts were homogenized in Tris–HCl, pH 7.4 containing 100 mM maleimide, 0.2 mM neocuproine, 1 mM diethylenetriaminepentaacetic acid (DTPA) and protease inhibitors to generate a 10% weight/volume homogenate, which was then supplemented with sodium dodecyl sulfate (SDS) to give a concentration of 2.5%. In one experiment a Krebs perfused rat heart was lysed into Tris–HCl, pH 7.4 containing protease inhibitors. This sample was split in two groups, to one Tris–HCl, pH 7.4 alone was added and to the other Tris–HCl, pH 7.4 containing 200 μM SNAP. After incubation at room temperature for 20 min these samples were supplemented with 100 mM maleimide, 0.2 mM neocuproine, 1 mM DTPA and 2.5% SDS. Treated HEK293 cells were lysed directly into Tris–HCl, pH 7.4 containing 100 mM maleimide, 0.2 mM neocuproine, 1 mM DTPA and 2.5% SDS. All lysates were then incubated at 50 °C for 25 min to block free thiols and then desalted to remove any unincorporated maleimide. Samples were then split into two groups, to one 30 mM ascorbate was added and to the other both 30 mM ascorbate and 0.1 mM N-(6-(biotinamido)hexyl)-3-(2’-pyridylidithio)-propionamide (biotin-HPDP). After 1 h of incubation in the dark at room temperature samples were desalted to remove unincorporated biotin-HPDP, followed by addition of equal volume of 2× Laemml buffer (4% SDS, 20% glycerol, 0.004% bromphenol blue, 0.125 M Tris–HCl, pH 6.8). Samples were analyzed by resolving on SDS–PAGE gels followed by Western blotting then immunostaining using streptavidin-HRP.

Affinity capture of S-nitrosylated proteins using the biotin-switch method

Homogenate from three control hearts were combined together and the homogenate for three CysNO perfused hearts, to give one sample for control and one for CysNO perfused. Both sets of homogenate were processed using the ascorbate-dependent biotin-switch method described above. Biotin–HPDP modified proteins were purified using streptavidin–agarose. In brief, after biotin–switch analysis homogenates were combined with streptavidin–agarose, 1% triton and protease inhibitors followed by 3 h incubation at 4 °C on a rotary shaker. Streptavidin–agarose beads were pelleted and then resuspended in fresh buffer before being added to empty spin column. Each spin column was washed six times with Tris–HCl, pH 7.4 containing 1% triton. Modified proteins were eluted using buffer containing 100 mM 2-mercaptooethanol. The same protocol was used to compare control or LPS-treated rats, utilizing 100 mg of cardiac tissue as input for the biotin-switch assay.

Mass spectrometry

Gel slices were treated with trypsin using published methods [30,31], modified for use with an investigator ProGest robotic digestion system (Genomic Solutions). Digested peptides were separated by using nano-flow liquid chromatography (Ultimate 3000™, Dionex) on a reverse-phase column (PepMap 100, 75 μm I.D., 15 cm length, Dionex) at a flow rate of 300 nl/min. The column was attached to a linear ion trap mass spectrometer (LTQ-XL, ThermoFinnigan) set to full ion scan mode over the mass-charge (m/z) range 300–2000. Tandem mass spectrometry (MS/MS) was carried out on the top six ions in each MS scan using the data-dependent acquisition mode with dynamic exclusion enabled. Generated spectra were matched to database entries (UniProt Knowledgebase) using TurboSEQUEST software (Bioworks 3.4, ThermoFinnigan). Protein and peptide identifications were verified using Scaffold (version 1.0, Proteome Software Inc., Portland, OR). Peptides with above 95.0% probability of identification were accepted as specified by the Peptide Prophet algorithm [32]. Protein identifications were accepted if probability was greater than 99.0% and contained at least two identified peptides as assigned by the Protein Prophet algorithm [33].

Radiotelemetry

Mean arterial pressure (MAP) was assessed by remote radio- telemetry in conscious freely moving mice as described [34]. Briefly, randomly selected C57Bl/6 mice at 12 weeks of age were anesthetised with 2% isoflurane in 1 l of oxygen per minute with pre and post-operative analgesia (buprenorphine, 0.1 mg/kg). Radio-telemetry probe catheter (TA11PA-C10, O.D 0.4 mm, Data Science International Inc., St. Paul, MN) was implanted into the aortic arch via the left carotid artery. Following 1 week recovery, mice housed in individual cages were placed above the telemetric receivers with output to a computer. Cardiovascular variables and activity were routinely monitored and recorded by scheduled sampling for 10 s every 5 min (Dataquest LabPRO Acquisition System version 3.01, Data Sciences International, St. Pauls, MN).

LPS treatment and heart preparation

Six week old Wistar rats were given an intraperitoneal (i.p.) injection of either saline or 10 mg/kg Escherichia coli bacterial lipopolysaccharide (LPS, E. coli 0111:B4, Sigma-Aldrich, UK). Rats
injected with LPS were anesthetized at 2, 4 or 8 h followed by removal and then storage of isolated hearts in liquid nitrogen. Rats injected with saline had their hearts removed at 8 h after the initial injection. Telemetered C57Bl/6 mice were injected i.p. with 9 mg/kg E. coli bacterial lipopolysaccharide (E. coli 0111:B4) or an equivalent volume of sterile saline as described [35]. Some mice were given an i.p. injection of 100 mg/kg 5,10,15,20-tetrakis(4-sulfonatophenyl)porphyrinato iron (III) (FeTPPS) 1 h before LPS treatment. During blood pressure measurement studies mice were culled 18 h after LPS or saline treatment. For analysis of protein nitration, mice were anesthetized 6 h after injection followed by removal and then storage of isolated hearts in liquid nitrogen.

**Immunoblotting**

Samples were resolved using poly-acrylamide gel electrophoresis then transferred onto PVDF membranes using Western blotting. Samples generated using the biotin-switch method were probed using streptavidin-HRP. LPS treated samples were immunostained using a 3-nitrotyrosine antibody (Millipore, clone 1A6). Affinity purified biotinylated proteins after the biotin-switch method from LPS-treated rat heart homogenate was immunostained using a myosin heavy chain antibody (Abcam ab15-100).

**Results**

To establish if the ascorbate-dependent biotin-switch assay could specifically detect S-nitrosothiols the method was evaluated in HEK cells treated with CysNO, hydrogen peroxide (H2O2) or diamide as shown in Fig. 1A. Treatment with CysNO, a known mediator of protein S-nitrosylation [36], generated a large increase in ascorbate-dependent protein labeling as anticipated. H2O2 or diamide, which preferentially induce protein sulfenic acid or disulfide bond formation, respectively, generated no detectable protein labeling. This helps verify the specificity of this method for S-nitrosothiols, as well as confirm this methodology for detecting protein S-nitrosylation actually worked in our hands. We then used this method to monitor protein S-nitrosylation in hearts from saline- or LPS-treated rats; but did not find any variation in global protein S-nitrosylation as shown in Fig. 1B. However, as the biotin-switch method has poor sensitivity [37], subtle changes in S-nitrosylation may have been missed.
We reasoned that the sensitivity of detecting protein S-nitrosylation may be improved by first identifying proteins in the heart that were targets of this modification. We might then use sensitive antibodies to these S-nitrosylation targets and assess whether they are present in streptavidin-agarose affinity-captured proteins in hearts from saline- or LPS-treated animals that had undergone the biotin-switch labeling protocol.

The ability of NO donors SNAP or CysNO to induce detectable S-nitrosylation in the Langendorff perfused rat heart were compared. SNAP only generated protein S-nitrosylation when directly incubated with heart homogenate in vitro and not when perfused through the coronary vessels of isolated rat hearts (Fig. 1C). In contrast perfusion with CysNO induced significant S-nitrosylation as visualized by the biotin-switch method (Fig. 1D). These S-nitrosylated target proteins were affinity captured with streptavidin-agarose and resolved by SDS-PAGE and visualized with colloidal Coomassie blue (Fig. 1E), which confirmed multiple unique proteins in the samples from CysNO-treated hearts compared to controls. The unique bands were excised from the SDS–PAGE gel and identified using tandem mass spectrometry. This led to the identification of several potential targets for S-nitrosylation as listed in Table 1.

We then used a sensitive antibody to one of these newly identified S-nitrosylation targets, namely myosin heavy chain, to screen for its abundance in streptavidin-agarose affinity captures prepared from hearts from mice treated with LPS. This showed that myosin heavy chain becomes de-nitrosylated in a time-dependent manner following LPS-treatment (31 ± 12% decrease over 8 h) as seen in Fig. 2A. This loss of protein S-nitrosylation was unexpected as we had anticipated increased S-nitrosylation due to LPS-induced nitrosative stress as discussed above. Consequently, we hypothesized that the loss in S-nitrosylation may be explained by a loss in S-nitrosylating derivatives due to the biochemical conversion of NO to OONO·. Accordingly, we analyzed 3-nitrotyrosine (a biomarker of OONO·) content of hearts from control or LPS-treated animals using Western immunoblotting. This analysis (Fig. 2B) showed LPS treatment enhanced tyrosine nitration at 2 h and more prominently (~4-fold increase above control) at 4 h, before returning to basal control level by 8 h. Protein tyrosine nitration following LPS-treatment was particularly apparent in a 13 kDa protein, and was also prominent in other anti-nitrosylation immunoblots analyzing related samples from LPS-treated animals (Fig. 3B).

Measurement of mean arterial pressure in telemetered mice treated with LPS showed a maximal decrease around 6 h followed by slow time-dependent recovery towards basal (Fig. 3A). The development of this hypotension after LPS treatment temporally correlated with the increase in protein tyrosine nitration shown in Fig. 2B. In an attempt to prevent LPS-induced tyrosine nitration an intraperitoneal injection of the OONO· scavenger FeTPPS was administered 1 h before LPS treatment. However, this potentially protective intervention actually had the reverse effect and enhanced protein nitration ~2-fold above control and also potentiated the LPS-induced hypotension. Treatment of mice with just the scavenger FeTPPS alone also significantly enhanced protein nitration ~2-fold above control.

### Discussion

Sepsis induces a nitrooxidative stress [14], which may be causal in the accompanying cardiovascular dysfunction. A logical potential consequence of nitrooxidative stress is the post-translational modification of redox active proteins. Nitrosative species may target redox active protein thiols or tyrosine residues to induce S-nitrosylation or 3-nitrotyrosine [38]. Accordingly, we monitored these nitrooxidative protein modifications in a rodent model of sepsis.

Evidence continues to accumulate supporting protein S-nitrosylation as a regulatory mechanism, with aberrant changes in this modification being causatively implicated in disease progression [21]. We monitored changes in protein S-nitrosylation using the biotin-switch method but did not find an increase in this modification, although we were confident the method was working as cells or isolated hearts exposed to CysNO generated a large signal using this labeling protocol. When the biotin-switch assay has been compared to the tri-iodide chemiluminescence measurement of protein S-nitrosylation, it was found to be relatively insensitive. High levels of S-nitrosylation (in the nanomole per milligram protein range) were required to generate an S-nitrosylation signal using the biotin-switch method, and physiological activators of NO synthesis did not increase the signal [37]. Thus the lack of a change in global nitrosylation can be explained by the poor sensitivity of the ascorbate-dependent biotin-switch method. A further confounding factor is that the biotin-switch assay like most proteomic approaches is hampered by the large variation in the relative abundance of proteins [38]. Whilst the ability of proteomic methods to assess the full complement of proteins in the experimental system has great benefits, a major problem can be that stoichiometric changes in low abundance proteins are often masked against a background signal resulting from non-stoichiometric modifications of high abundance proteins. Thus low abundance proteins may change their S-nitrosylation status during sepsis but the change in amplitude of this signal is smaller than the basal signals generated from abundant proteins.

<table>
<thead>
<tr>
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<th>Protein name</th>
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This problem caused by high abundance proteins generating a background greater than signals induced by stoichiometric S-nitrosylation of a low abundance protein can be overcome by identifying the proteins in the system susceptible to modification and then individually assessing them. Indeed, this is what we did, identifying cardiac proteins susceptible to S-nitrosylation using CysNO treatment to induce a robust, detectable signal well above background. These proteins were then affinity captured and identified using LC–MS. Armed with this knowledge we then assessed the abundance of these proteins in affinity preparations of cardiac S-nitrosylated proteins from control or LPS-treated animals. In this way we found a loss in myosin heavy chain S-nitrosylation during sepsis; not the increase we had anticipated based on the nitroxidative stress associated with sepsis [5]. Loss of myosin heavy chain S-nitrosylation during the onset of endotoxemia may itself have a functional effect, potentially contributing to contractile dysfunction during sepsis. This possibility is supported by recent work showing S-nitrosylation of skeletal and cardiac myosin caused a dose-dependent and oxygen-dependent reduction in actin filaments velocity over myosin, which double contractile force [39]. Future studies identifying and then mutating the site of S-nitrosylation to mimic or block its formation may help clarify the importance of this precise modification. Other targets of S-nitrosylation would also have to be examined on a case-by-case basis, again requiring further work. However, it is worthwhile considering that the decrease in myosin heavy chain S-nitrosylation detected in cardiac tissue of LPS treated animals could be due to increased protein ubiquitination and subsequent degradation, or by enhanced S-nitroso-glutathione reductase (GSNOR) expression. The enzyme GSNOR is known to reduce GSNO thereby decreasing its bioavailability to transnitrosylate target proteins.

Comparing protein S-nitrosylation induced by SNAP and CysNO highlights the classic form of NO itself as a poor S-nitrosylating agent, and is consistent with reports of other groups [40]. The poor membrane permeability of SNAP prevents direct transnitrosylation NO donation to intracellular target protein thiols. Instead SNAP, a relatively unstable donor, rapidly breaks down extracellularly and releases NO which then diffuses across the membrane. This form of free authentic NO is a poor S-nitrosylating agent and instead is scavenged by heme containing proteins such as soluble guanylate cyclase [41,42]. In contrast, CysNO is more stable and readily crosses the membrane through the L-amino acid transporter [40], serving as a powerful intracellular nitrosylating agent by direct transfer of NO to a target protein thiol. Clearly a change in cellular protein S-nitrosylation status may not directly reflect a change in free classical NO, but rather a change in low molecular weight nitrosothiols (CysNO, GSNO) or reactive nitrosylating species such as dinitrogen trioxide (\(\text{N}_2\text{O}_3\)) or nitrosonium ion (\(\text{NO}^+\)).

We speculated that the depletion in S-nitrosylation following LPS-treatment (identified using the myosin heavy chain screening approach) may be due to a loss in S-nitrosylating species of NO due to biochemical conversion to OONO\(^-\) [43]. This could be potentiated by superoxide generation under septic conditions, which can occur when NOS enzymes uncouple due to substrate depletion [10,11]. Endotoxins may also enhance gp91phox containing NADPH oxidase activity to further elevate superoxide. We compared tyrosine nitration, a biomarker of OONO\(^-\) [13], in hearts from control or LPS-animals. LPS-treatment of rats increased global protein tyrosine nitration which was maximal after 4 h and returned to basal at 8 h. In mice maximal hypotension occurred 6 h after LPS treatment and thus coincided with maximal protein tyrosine nitration.

LPS-induced tyrosine nitration could be causatively linked with the associated hypotension. Consistent with this possibility, others found direct injection of OONO\(^-\) significantly lowered blood pressure in rats [44,45]. Thus there may be specific target protein(s)
that alter their function following nitration and contribute to cardiovascular dysfunction (hypotension, impaired cardiac output) during sepsis. This possibility was also supported by reports [17,18,46], showing the OONO\textsuperscript{-} scavenger FeTPPS attenuates dysfunction during sepsis. To examine the potential causal relationship between tyrosine nitration and blood pressure lowering we also utilized FeTPPS, hypothesizing that it would scavenge LPS-induced OONO\textsuperscript{-} formation and so limit protein tyrosine nitration and consequently attenuate the hypotension. However, we observed precisely the opposite effect; FeTPPS dramatically enhanced LPS-induced tyrosine nitration and exacerbated the hypotension. There are several possible reasons why our model of FeTPPS treatment enhanced cardiac protein nitration in injected mice. Firstly if the product is contaminated with free iron then this may enhance protein oxidation through Fenton-chemistry dependent oxidant formation. Secondly our method of administering the FeTPPS by intraperitoneal injection would give a large bolus dose of this drug to specific region of tissue, which would then diffuse outwards into the blood and other organs. This contrasts other studies that utilized intravenous treatments. Such a concentrated localized form of FeTPPS may be cytotoxic whereas intravenous injection would rapidly dilute this compound into the circulatory system potentially abrogating the risk of cytotoxic overload. Further complexity results from OONO\textsuperscript{-} potentially targeting redox active protein thiols, the oxidation of which can activate vasodilatory pathways [47,48].

To conclude, this study demonstrates dynamic change in the myocardial nitrosative environment during early endotoxia. Further work is required to clarify the potential role of protein S-nitrosylation or tyrosine nitration in the pathogenesis of this deleterious, life-threatening pathology.

Acknowledgments

We thank the Wellcome Trust for providing a Sir Henry Wellcome Fellowship (J.R.B.) and the Medical Research Council for project grant support (P.E.).

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