Mechanisms of circulatory and intestinal barrier dysfunction during whole body hyperthermia

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Am J Physiol Heart Circ Physiol 280: H509–H521, 2001.—This work tested the hypotheses that splanchic oxidant generation is important in determining heat tolerance and that inappropriate NO production may be involved in circulatory dysfunction with heat stroke. We monitored colonic temperature (Tc), heart rate, mean arterial pressure, and splanchic blood flow (SBF) in anesthetized rats exposed to 40°C ambient temperature. Heating rate, heating time, and thermal load determined heat tolerance. Portal blood was regularly collected for determination of radical and endotoxin content. Elevating Tc from 37 to 41.5°C reduced SBF by 40% and stimulated production of the radicals ceruloplasmin, semiquinone, and penta-coordinate iron(II) nitrosyl-heme (heme-NO). Portal endotoxin concentration rose from 28 to 59 pg/ml (P < 0.05). Compared with heat stress alone, heat plus treatment with the nitric oxide synthase (NOS) antagonist Nω-nitro-L-arginine methyl ester (L-NAME) dose dependently depressed heme-NO production and increased ceruloplasmin and semiquinone levels. L-NAME also significantly reduced lowered SBF, increased portal endotoxin concentration, and reduced heat tolerance (P < 0.05). The NOS II and diamine oxidase antagonist aminoguanidine, the superoxide anion scavenger superoxide dismutase, and the xanthine oxidase antagonist allopurinol slowed the rates of heme-NO production, decreased ceruloplasmin and semiquinone levels, and preserved SBF. However, only aminoguanidine and allopurinol improved heat tolerance, and only allopurinol eliminated the rise in portal endotoxin content. We conclude that hyperthermia stimulates splanchic oxidant generation and that inappropriate nitric oxide (NO) production is involved in splanchic vascular dysfunction with heat stroke. We developed these hypotheses based on previous work from our laboratory (18) demonstrating that hyperthermia stimulates splanchic production of NO and biomarkers of cellular oxidative stress.

With the use of electron paramagnetic resonance (EPR) spectroscopy to directly detect radicals in vivo (18), we observed that hyperthermia progressively increased portal venous content of the following: semiquinone radical, a biomarker of mitochondrial reductive stress (52); ceruloplasmin, an acute phase antioxidant protein that acts to reduce metal-catalyzed oxidant production (29); and penta-coordinate iron(II) nitrosyl-heme (heme-NO) (27). The respective characteristics of these radicals suggest that heat stroke stimulates NO and reactive oxygen species (ROS) production within splanchic viscera leading to transition metal activation and cellular oxidative stress. Indeed, our group (19) and others (36, 44) have proposed that metal-catalyzed oxidative stress is involved in hyperthermia-related pathlogy.

The purpose of the present study was to investigate mechanisms of •NO and ROS production in vivo during environmental heat stroke. Figure 1 details a working model that we propose for the etiology of heat illness. We (17) have previously shown that hyperthermia produces cellular hypoxia and metabolic stress within the

HEAT STROKE IS THE MOST CATASTROPHIC FORM of debilitating illness resulting from environmental heat stress.

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liver and intestine. We propose that subsequent cellular biochemical events secondary to hypoxia (e.g., increased cytosolic Ca\(^{2+}\)) promote mitochondrial ROS production and stimulate cellular oxidase and constitutive nitric oxide (NO) synthase (NOS) enzymatic activities. Stimulating mitochondrial ROS production and oxidase activity increases cellular production of O\(_2\)\(^{-}\) and H\(_2\)O\(_2\). O\(_2\)\(^{-}\) can activate transition metals, thereby increasing metal-catalyzed cellular oxidative stress. The increased flux of \(\text{NO}\) produced by Ca\(^{2+}\)-dependent NOS is protective through its cell regulatory and antioxidant actions.

C: as heat stress continues, hypoxic cells export oxidases into the extracellular space, where metal-catalyzed oxidative stress can produce multifocal cellular injury and inflammation. Damage to the intestine increases intestinal permeability to endotoxins, contributing to local inflammation and inducible NOS (iNOS) activation. The large increase in \(\text{NO}\) flux from iNOS promotes reactive nitrogen species (RNS) generation. \(\text{NO}\) inappropriate production of ROS and RNS leads to "nonprogrammed" splanchnic dilation, systemic hypotension, and circulatory shock.

PEG-SOD, polyethylene glycol-conjugated superoxide dismutase; cNOS, constitutive NOS; L-NAME, N\(^{\text{\text{\text{"E}}}2}\)-nitro-L-arginine methyl ester.

To test these ideas, in vivo experiments were designed to examine the effects of 1) antagonizing NOS enzymatic activity, 2) increasing NOS substrate availability, 3) limiting superoxide anion bioavailability with polyethyleneglycol (PEG)-conjugated copper and...
zinc-containing superoxide dismutase (PEG-SOD), and 4) antagonizing the heat-sensitive ROS-producing enzyme xanthine oxidase with allopurinol.

MATERIALS AND METHODS

Animals

Male Sprague-Dawley rats (Harlan, Indianapolis, IN) weighing 300–350 g served as subjects. Rats were purchased in groups of 15 and housed in the University of Iowa Animal Care Facility at an ambient temperature of 22–24°C on a 12:12-h light-dark cycle. Animals were handled and familiarized with laboratory procedures for 3–5 days before use as experimental subjects. Food and water were provided ad libitum.

Rats were randomly assigned to the following groups: 1) sham-operated normothermic control (n = 4), 2) saline-treated normothermic control (n = 4), 3) noninjected hyperthermic control (n = 4), 4) saline-treated hyperthermic control (n = 10), 5) N’-nitro-l-arginine methyl ester (l-NNAME)-treated (25, 50, and 125 μM) heat exposed (n = 24), 6) aminoguanidine-treated (25 and 50 μM) heat exposed (n = 16), 7) l-arginine-treated (100, 200, and 300 μM) heat exposed (n = 24), 8) PEG-SOD-treated (10 and 30 mg/kg body wt) heat exposed (n = 16), 9) PEG-SOD-treated normothermic control (n = 4), and 10) allopurinol-treated (10, 20, and 30 mg/kg body wt) heat exposed (n = 24).

Experimental Design

Experiments were conducted between 800 and 1300 h. After surgical preparation, anesthetized rats were exposed to 40°C ambient temperature (T\textsubscript{a}) in a temperature-controlled environmental chamber while they received intravenous injections of the designated pharmacological agent. Thermocouples placed around the animal continuously monitored T\textsubscript{a}.

Preliminary experiments established that enzyme blockade was not maintained for the duration of an experiment if a single bolus injection of a reagent was used. Therefore, rats received injections (10 μl/100 g body wt iv) at time 0 (beginning of heat exposure) and at 15-min intervals until termination of an experiment (~90 min). Because of its long vascular half-life (41), PEG-SOD was given as two injections within the first 15 min of heating.

Colonic temperature (T\textsubscript{c}), heating time, mean arterial pressure (MAP), heart rate (HR), and superior mesenteric artery (SMA) blood flow were continually monitored. From these data, thermal load and splanchnic resistance were calculated. Portal venous and femoral artery blood samples were collected for analyses of radical and endotoxin content at four time points: 1) immediately before heat exposure (T\textsubscript{c} ~ 37°C), 2) at T\textsubscript{a} = 41.5°C, 3) immediately after SMA flow increased, and 4) after MAP fell below 100 mmHg. Heat exposure was terminated when MAP fell below 100 mmHg. Experiments were terminated when MAP fell below 60 mmHg.

Surgical Preparation

Animals were anesthetized with pentobarbital sodium (Nembutal; 50 mg/kg ip), injected with atropine (100 μl), and tracheotomized to ensure a patent airway. An incision was made in the inguinal region of the right hindlimb. The femoral artery was isolated for ~1 cm of its length and fitted with a catheter [polyethylene (PE)-50; Clay Adams, Parsippany, NJ] filled with heparinized saline (100 U/ml) to monitor HR and MAP and to sample blood. The leg incision was closed, and a midline laparotomy was performed. The portal vein was isolated ~2 mm from the liver, and a second catheter for sampling blood and delivering pharmacological agents was placed in the portal vein through a tributary vessel. Tributary vessels were chosen as cannulation sites to avoid impeding portal blood flow. Catheter placement was confirmed at the conclusion of each experiment.

The superior mesenteric artery was next isolated for ~1 cm of its length, and a miniaturized Transonic Doppler flow probe was positioned around the vessel to monitor SMA blood flow. The abdominal incision was closed, and a colonic temperature (T\textsubscript{c}) probe (36-gauge copper-constantan wire in PE-100 tubing) was inserted 7–8 cm past the anal sphincter to monitor T\textsubscript{c}.

Rats were allowed to stabilize for 30 min after surgery. Baseline HR, MAP, and SMA blood flow data were collected for an additional 30 min. During this 60-min period, T\textsubscript{a} was maintained at 37.0 ± 0.2°C with a heating pad. At the end of the control period, baseline portal venous and femoral artery blood samples were collected, and the T\textsubscript{c} of the environmental chamber was elevated to 40°C with a relative humidity of 30%.

Sample Collection and Handling

Cardiovascular data. MAP was determined by connecting the femoral artery catheter to a Gould P23 ID pressure transducer that electronically averages the pulsatile signal. HR was determined from the number of electronic pulsations sensed by a cardiotachometer (Beckman Instruments). SMA blood flow was determined using a pulsed Transonics Doppler flowmeter (University of Iowa Bioengineering Resource Facility). MAP, HR, and SMA blood flow were continuously monitored, and the mean value over each 60-s period was recorded.

Radicals and serum endotoxin. Portal venous and femoral artery blood samples (350 μl each) were collected in sterile, 1-ml Monoject syringes (2). Whole blood (200 μl) was immediately delivered into a quartz EPR tube (inner diameter 3 mm) and frozen at 77 K pending EPR analyses as previously described (18). The remaining 150 μl of blood was processed for quantitation of gram-negative bacterial endotoxin concentration per manufacturer’s instructions [limulus amebocyte lysate (LAL), quantitative chromogenic LAL, BioWhittaker, Walkersville, MD]. Briefly, cell and serum fractions were isolated by centrifugation in a refrigerated clinical microcentrifuge. The serum was extracted and delivered to pyrogen-free storage tubes (BioWhittaker), diluted 1:10 with pyrogen-free water, and heated for 10 min at 70°C to remove nonspecific LAL inhibitors present in blood products (8). Processed serum was stored at ~80°C pending endotoxin assay. Serum from all samples was tested in triplicate. All materials coming into contact with blood or test materials not purchased as pyrogen-free from BioWhittaker were rendered pyrogen-free by heating at 200°C for 4 h.

EPR Conditions

EPR spectra were recorded with a Bruker ESP 300 EPR spectrometer (Bruker Instruments, Karlsruhe, Germany) equipped with an ER035M gaussmeter, an ER4111VT variable-temperature unit, and an EIP-625A microwave-frequency counter. Signal averaging (multiple scans of the same sample) was used to improve the signal-to-noise ratio. Sample volume and geometry were kept constant to allow for comparisons of radical concentration between samples. All spectra were collected at 100 K, with data reported as the normalized average of 20 scans. EPR conditions were the
following: receiver gain $5.00 \times 10^5$, modulation frequency 100 kHz, modulation amplitude 4.0 Gauss (G), microwave frequency 9.43 GHz, microwave power 10 mW, and scan rate 6.0 G/s.

**Calculations**

Thermal stress was quantified by determining thermal load, a product of $T_c$ and time spent above 40.4°C (22). $T_c$ was recorded at 1-min intervals, and thermal load ($°C \times \text{min}$) was calculated as the following: the sum of the time interval (in min.) $\times (T_c \text{above} 40.4°C - 40.4°C)$. Heating rate (in °C/min) was calculated as the following: (maximum $T_c$ attained during heat exposure – baseline $T_c$/total heating time (in min).

Splanchnic vascular resistance is equal to pressure (in mmHg) divided by SMA flow (kHz shift). The percent change in splanchnic vascular resistance was calculated using the following formula: $[(R_c - R_c/\text{baseline}) \times 100$, where $R_c$ is baseline MAP divided by the mean SMA flow value in the control period and $R_c$ equals MAP divided by the SMA flow value in the test period (20).

**Statistical Analyses**

Thermal responses, endotoxin values, and cardiovascular data were analyzed by one-way analysis of variance. Groups were categorized according to Tukey’s multiple comparisons procedure. Significant differences were identified with Dunnett’s test. Differences were considered significant at the $P < 0.05$ level.

**RESULTS**

**Normal Heat Stress Responses**

Our initial set of experiments was designed to characterize the typical radical, hemodynamic, and splanchnic endotoxin responses of the heat-stressed rat. To examine splanchnic hemodynamics and splanchnic radical production in parallel, Fig. 2 presents SMA blood flow and portal EPR spectra versus $T_c$. There were no differences in the responses of noninjected hyperthermic rats and saline-treated hyperthermic animals; therefore, data from these two groups were pooled and are referred to as hyperthermic controls.

EPR spectra of portal blood collected before heat stress displayed a weak composite signal, suggesting the presence of multiple radicals. The narrow $g = 2.005$ feature (peak-to-peak line width $\sim 10$ G) represents semiquinone radical (Fig. 2A) (18). The broad deflection at $g = 2.06$, indicative of a transition metal, represents ceruloplasmin (Fig. 2B) (4).

Elevating $T_c$ from 37 to 41.5°C stimulated a 127 ± 7% increase in splanchnic resistance (Table 1), a 40 ± 2% reduction in splanchnic blood flow (SBF) (Fig. 2 and Table 1), and increased circulating ceruloplasmin and semiquinone radical levels (Fig. 2B). The decline in SBF was nonlinear in nature, with a sharp decrease occurring at $T_c > 40°C$ (Fig. 2). Heart rate and MAP (Fig. 3, control) progressively increased with rising $T_c$, peaking at 589 ± 16 beats/min and 170 ± 2 mmHg, respectively (Table 1).

At $T_c$ of 41.5°C, we observed the evolution of a triplet EPR signal centered at $g = 2.012$ (splitting constant $\sim 17.5$ G) (Fig. 2, B and C) that represents heme-NO (18, 27). EPR evidence of heme-NO was not observed in arterial blood (data not shown), suggesting that this was a local phenomenon within the splanchnic region. Splanchnic resistance fell and SBF rapidly increased at $T_c$ of 41.8 ± 0.2°C. EPR of portal blood collected immediately after the initial rise in SBF shows a strong heme-NO signal (Fig. 2C), suggesting that overproduction of NO may contribute to declining splanchnic resistance. Only the ceruloplasmin signal was observed in arterial blood samples (data not shown), again suggesting that the rise in heme-NO levels was a local event within the splanchnic circulation. HR and MAP (Fig. 3, control) declined at $T_c$ of 42.0 ± 0.2°C 5–7 min after rising SBF. Total heating time for hyperthermic controls was 77 ± 4 min, and thermal load was 34.4 ± 3.2°C × min (Table 1). We interpret...
oxidative stress and lower heat tolerance. L-NAME treatment also increased ceruloplasmin and semiquinone radical levels (Fig. 5), marginally elevating the Tc at high Tc (Fig. 5), suggesting that antagonizing NOS contributes to the observed splanchnic dilation. In addition, 50 and 125 μM L-NAME doses decreased peak HR (Fig. 3 and Table 1) and lowered heat tolerance (Table 1).

### L-NAME

These experiments tested the hypothesis that globally antagonizing NO synthase (NOS) with L-NAME (37) would lower heme-NO accumulation, delay splanchnic dilation, and improve heat tolerance. We chose concentrations of 25 and 50 μM aminoguanidine based on the work of Grisham et al. (16) and Corbett et al. (6), who demonstrated in vivo the relative specificity of aminoguanidine for NOS II over NOS I and III. Vascular radical profile was similar between 25 and 50 μM aminoguanidine treatments; therefore, EPR spectra from the 25 μM-treated group are presented.

These experiments tested the hypothesis that L-arginine supplementation would increase hyperthermia-induced NO production, blunt the rise in splanchnic resistance, and preserve SBF. Contrary to our hypothesis, L-arginine...
nine dose dependently increased splanchnic resistance and MAP and lowered SBF (Table 1); events that occurred independent of alterations in splanchnic heme-NO production (data not shown). We speculate that the increase in resistance and MAP may be due to arginine-stimulated splanchnic vasoconstrictor release (32).

**PEG-SOD**

SOD scavenges $O_2^-$, directly blocking $O_2^-$-mediated chemistry and indirectly increasing the bioavailability of $\textit{NO}$. PEG-SOD is a large molecule with a vascular half-life of 18–24 h (41). With the use of PEG-CuZn-SOD, these experiments tested the hypothesis that heat stress stimulates $O_2^-$ production, which is involved in regulating splanchnic hemodynamics and in cellular events leading to semiquinone radical production and ceruloplasmin release. Vascular radical profile was similar between 10 and 30 mg/kg PEG-SOD treatments; therefore, spectra from 10 mg/kg-treated animals are presented.

Treatment with PEG-SOD markedly reduced splanchnic constriction (Table 1) and maintained SBF at or near euthermic control levels throughout heat exposure (Fig. 7). PEG-SOD also blunted the rise in ceruloplasmin and semiquinone radical levels (Fig. 7, B vs. A) and significantly increased the magnitude of rise in $T_c$ required to elicit heme-$\textit{NO}$ accumulation and splanchnic dilation (42.5 vs. 41.5°C for hyperthermic controls, $P < 0.05$) (Fig. 7 vs. Fig. 2). Once resistance declined in this group, SBF rose to $44 \pm 8\%$ above euthermic baseline. Peak HR and MAP were signifi-
cantly increased (Fig. 8), and HR was preserved at or near peak levels until termination of an experiment \( T_c = 43.3 \pm 0.3 {}^\circ C \) (Fig. 8).

Both 10 and 30 mg/kg PEG-SOD significantly increased thermal load (Table 1). However, neither dose increased total heating time, and both the rate of rise in \( T_c \) and the maximum \( T_c \) attained before shock were greater than that of hyperthermic controls (Table 1), suggesting that lowering splanchnic constriction negatively impacted heat-dissipating capacity in this group.

**Allopurinol**

Allopurinol is a competitive antagonist of xanthine oxidase, an enzyme that directly produces \( O_2^- \) and hydrogen peroxide (H\(_2\)O\(_2\)). The intestine and liver are rich in xanthine oxidase protein, which is localized primarily to hepatic sinusoidal endothelial cells and intestinal epithelial cells (23). These experiments tested the hypothesis that heat stress can stimulate xanthine oxidase production of ROS that influence splanchnic hemodynamics and participate in cellular events leading to semiquinone radical production and ceruloplasmin release. Vascular radical profile was similar between 10 and 30 mg/kg allopurinol treatments; therefore, spectra from 10 mg/kg-treated animals are presented.

Treatment with allopurinol markedly decreased splanchnic semiquinone radical and ceruloplasmin levels (Fig. 9). EPR spectra of portal blood collected at 41.5°C (Fig. 9B) and 42.7°C (Fig. 9C) showed little change in ceruloplasmin or semiquinone radical concentrations from 37°C levels. As with PEG-SOD, allopurinol also

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**Fig. 4.** The influence of the NOS antagonist L-NAME (LN) on vascular responses during heat stress. Compared with hyperthermic controls, treatment with the NOS antagonist L-NAME (25 \( \mu \)M; \( n = 8 \)) significantly reduced SBF at \( T_c \leq 39.0{}^\circ C \) and rapidly increased semiquinone radical and ceruloplasmin formation (A vs. B, respectively), suggesting that blocking NOS activity increased metal activation and cellular oxidative stress. L-NAME treatment also blunted, but did not eliminate, the rise in heme-NO and delayed splanchnic dilation until \( T_c \sim 42{}^\circ C \). Note that a strong heme-NO signal was evident immediately before the rapid rise in SBF at 41.8°C, consistent with NO involvement in splanchnic dilation. *Significantly different from control, \( P < 0.05 \).

**Fig. 5.** Higher doses of L-NAME decreased heat tolerance. Compared with hyperthermic controls, both 50 and 125 \( \mu \)M L-NAME (\( n = 8 \) rats/group) significantly reduced SBF and increased semiquinone radical and ceruloplasmin formation (A vs. B, respectively). Note that EPR spectra collected at 38 and 41.3°C show only evidence of semiquinone radical and ceruloplasmin, consistent with increased metal activation and cellular oxidative stress with NOS blockade. In addition, both 50 and 125 \( \mu \)M L-NAME completely blocked the rise in heme-NO levels (C) and eliminated the hyperthermia-related splanchnic dilation. EPR spectra collected from animals treated with 50 and 125 \( \mu \)M L-NAME were similar; therefore, spectra from 50 \( \mu \)M-treated animals are presented. *Significantly different from control, \( P < 0.05 \).
increased the Tc at which heme-NO accumulation and splanchnic dilation were observed (Tc of 42.5 vs. 41.5°C for controls, P < 0.05). Indeed, the vascular radical profile from allopurinol-treated animals and the magnitude of thermal stress required for radical generation were similar to those of animals treated with PEG-SOD (compare Figs. 9 and 7). However, there is a notable absence of EPR-detectable ceruloplasmin with allopurinol treatment (Fig. 9, A–D).

Allopurinol did not alter the magnitude of change in resistance or SBF (Fig. 9 and Table 1) but shifted the SBF curve rightward, significantly slowing the rate of rise in resistance and the rate of decline in SBF (Fig. 9). On splanchnic dilation, SBF increased to +20 ± 4% above euthermic baseline. Allopurinol also increased peak HR, MAP (Fig. 8), and thermal load (Table 1). HR was maintained at or near peak levels until termination of an experiment (Fig. 8).

Portal Endotoxin

Portal venous blood collected before heat exposure tested positive for gram-negative bacterial endotoxins at a concentration of 28 ± 7 pg/ml (Fig. 10). Arterial blood tested negative for endotoxin (data not shown), suggesting that the liver clears endotoxins under these conditions. Heat stress alone and heat stress plus saline treatment significantly increased portal endotoxin.
concentration to 59 ± 7 and 62 ± 4 pg/ml, respectively (P < 0.05) (Fig. 10). Arterial blood tested negative for endotoxin, suggesting that the liver capacity to clear endotoxins remains intact at Tc of 41.5°C. Compared with hyperthermic controls, neither PEG-SOD (58 ± 5 pg/ml), aminoguanidine (60 ± 5 pg/ml), nor L-arginine (59 ± 7 pg/ml) treatments altered 41.5°C portal endotoxin concentration, whereas L-NAME (78 ± 4 pg/ml) significantly increased portal endotoxin levels. Only allopurinol treatment significantly reduced 41.5°C endotoxin levels (29 ± 8 pg/ml, P < 0.05) (Fig. 10).

DISCUSSION

The purpose of the present study was to test the hypotheses that splanchnic oxidant generation is an

![Fig. 8.](image)

Fig. 8. SOD appears to be involved in HR and MAP responses during heat stress. Compared with hyperthermic controls, PEG-SOD (A and B) and the xanthine oxidase inhibitor allopurinol (C and D) dose dependently increased the peak HR (A and C) achieved during heat stress and significantly increased heat tolerance. Allopurinol also dose dependently increased peak MAP (D) above controls, whereas only the higher concentration of PEG-SOD (30 mg/kg) elevated peak MAP (B) above control. Rats received injections (10 μl/100 g body wt iv) at time 0 (beginning of heat exposure) and at 15-min intervals until termination. *Significantly different from control, P < 0.05.

![Fig. 9.](image)

Fig. 9. Allopurinol (All) preserves splanchnic constriction and increases heat tolerance. Left: allopurinol (10 and 30 mg/kg) shifted the SBF curves rightward, indicating that it slowed the rate of decline in SBF. In addition, allopurinol treatment decreased semiquinone radical and ceruloplasmin formation, suggesting that metal activation and oxidative stress were lowered in this group. Allopurinol also significantly increased the Tc at which heme–NO accumulation and splanchnic dilation were observed (from 41.3 ± 0.2 to 42.7 ± 0.2°C), compare Fig. 9, B and C, with Fig. 2, B and C. On splanchnic dilation, blood flow rose to +20 ± 4% above euthermic control. EPR spectra collected from animals treated with 10 and 30 mg/kg allopurinol were similar; therefore, spectra from 10 mg/kg-treated animals are presented. *Significantly different from control, P < 0.05.
important determinant of heat tolerance and that \( \cdot \text{NO} \) is involved in splanchic vascular dysfunction with heat stroke. The major results of the present experiments support these proposals, suggesting that environmental heat stress can stimulate oxidase production of ROS that contribute to metal activation and circulatory and intestinal barrier dysfunction. In addition, the present data demonstrate that intact NOS activity is required for normal heat tolerance, whereas overproduction of \( \cdot \text{NO} \) may be responsible for the non-programmed loss of splanchic resistance that precedes heat stroke. We speculate that cellular hypoxia and derangements in intracellular \( \text{Ca}^{2+} \) control may be central to the observed increase in ROS and \( \cdot \text{NO} \) production (Fig. 1).

**EPR Results**

EPR data from the present study demonstrate that environmental heat stress can increase circulating ceruloplasmin levels and stimulate splanchic semiquinone radical production. Ceruloplasmin is an acute phase protein synthesized by the liver that exhibits robust ferrooxidase activity, oxidizing loosely bound transition metals and decreasing their participation in cellular redox chemistry (24, 29). The involvement of catalytic metals in oxidant generation and cellular injury is a well-documented phenomenon (33). Accordingly, ceruloplasmin is viewed as a potent antioxidant protein. Hyperthermia-induced elevations in circulating ceruloplasmin and semiquinone radical levels suggest that metal-catalyzed oxidative stress may contribute to heat stress pathology. We speculate that hepatic tissues (46) and vascular endothelial cells (9) may be sources of catalytic metals under these conditions.

Mechanisms by which hyperthermia causes metal activation are unclear, but both \( \text{O}_{2}^{\cdot} \) (9) and \( \cdot \text{NO} \) (38) can reduce protein-bound metals, promoting their participation in redox chemistry. While the current results suggest that production of both \( \text{O}_{2}^{\cdot} \) and \( \cdot \text{NO} \) are stimulated by heat stress, the fact that PEG-SOD and allopurinol treatments lowered ceruloplasmin concentrations, whereas L-NAME increased ceruloplasmin, suggests that \( \text{O}_{2}^{\cdot} \) may be the effector molecule that activates transition metals during hyperthermia. It is unlikely, given the large size of the PEG-SOD complex and the relatively short duration of these experiments, that PEG-SOD could be impacting mitochondrial \( \text{O}_{2}^{\cdot} \) production. Rather, these results imply that extramitochondrial sources of \( \text{O}_{2}^{\cdot} \), such as cellular oxidases, may be involved. Data from experiments using allopurinol as an intervention support this conclusion. Collectively, we interpret these results as evidence that heat stress stimulates oxidase production of \( \text{O}_{2}^{\cdot} \) and \( \text{H}_{2}\text{O}_{2} \), leading to transition metal activation within splanchic tissues. Adaptive responses include hepatic release of ceruloplasmin and local \( \cdot \text{NO} \) production.

The present study strongly suggests that constitutive synthesis of \( \cdot \text{NO} \) is essential for successful adaptation to acute heat stress, whereas NOS II activation may underlie hyperthermia-induced splanchic dilation. Antagonizing NOS with L-NAME dose dependently lowered heme-\( \cdot \text{NO} \) accumulation and SBF, increased semiquinone radical levels, and lowered heat tolerance. In contrast, aminoguanidine treatment slowed but did not eliminate heme-\( \cdot \text{NO} \) accumulation, significantly delayed splanchic dilation, and improved heat tolerance more than any other intervention. On the basis of previously cited in vivo work (6, 16), we speculate that aminoguanidine antagonized inducible but not constitutive \( \cdot \text{NO} \) synthesis in these experiments. We interpret these results as evidence that NOS II activation contributes to hyperthermia-related splanchic dilation. Alternatively, because aminoguanidine also inhibits diamine oxidase, these results may indicate that antagonizing amine metabolism during heat stress lowers NOS activation. However, while the precise mechanism of action for aminoguanidine remains unclear, these results suggest that constitutive \( \cdot \text{NO} \) production protects normal heat tolerance, whereas NOS II activation may compromise thermoregulatory processes by lowering peripheral resistance.

The present work was not designed to evaluate mechanisms underlying the protective effects of \( \cdot \text{NO} \), but low flux \( \cdot \text{NO} \) production has been shown to enhance cellular antioxidant capacity, acting as a primary antioxidant (51) and as a chain-breaking antioxidant (21, 25, 50, 51). During heat stress, \( \cdot \text{NO} \) may act to buffer the rise in intracellular ROS levels and subsequent generation of lipid peroxides. In addition, \( \cdot \text{NO} \) opposes splanchic vasoconstrictive activity, thereby preserving SBF and potentially reducing tissue hypoxia and acidosis (17). Local hypoxia elicits vasoconstriction followed by relaxation in mesenteric arterioles (31), and acidosis is a key predictor of negative clinical outcomes from heat illness (35). Moreover, \( \cdot \text{NO} \) inhibits platelet aggregation (34), neutrophil adherence to activated
vascular tone; therefore, we speculate that both ROS and reactive nitrogen species may contribute to local circulatory dysfunction during heat stress.

**Endotoxin Results**

Results from the present study establish that severe heat stress can produce splanchnic endotoxemia. However, hepatic reticuloendothelial cell function remains intact up to 41.5°C because endotoxins were not detected in the systemic circulation. These effects were eliminated with allopurinol treatment and exacerbated with l-NAMe. This suggests that ROS produced by xanthine oxidase may be responsible for the observed rise in splanchnic endotoxin levels, whereas -NO may be a critical molecule that protects intestinal barrier function and/or contributes to reticuloendothelial cell endotoxin removal.

The heat sensitivity and physiological significance of xanthine oxidase was first demonstrated by Skibba et al. (44, 45), who reported that stimulation of hepatic xanthine oxidase activity produced metal-catalyzed oxidative injury in the hyperthermic liver. Subsequent works by Tan et al. (47) and Terada et al. (48) have shown that an ischemic insult to the intestine and liver can stimulate cellular export of xanthine oxidase. Once in the vascular compartment, xanthine oxidase can be bound by endothelial cell glycosaminoglycans (47), localizing the enzyme next to endothelial cell surfaces.

In the current work, heat stress reduced SBF by ~40% as Tc rose from 37 to 41.5°C. We (17) have previously shown that heat stress of this magnitude produces significant cellular hypoxia in the liver and intestine. Hypoxia can stimulate conversion of xanthine dehydrogenase to xanthine oxidase and increase its cellular export (23). Although the current project did not directly measure xanthine oxidase activity, our data suggest that the combined effects of heat and cellular hypoxia may have stimulated cellular xanthine oxidase activity, leading to intestinal injury and splanchnic endotoxemia.

The role of endotoxins in heat illness remains to be established. The current view of the involvement of splanchnic organs in the pathogenesis of heat stroke is that ischemic and/or thermal stresses can damage the intestinal wall, allowing endotoxins to escape the intestinal lumen and enter the portal circulation (1, 13). Thermal injury to the liver would then allow endotoxins to spill over into the systemic circulation, stimulating a cascade of events similar to systemic inflammatory response syndrome. However, it generally requires much higher Tc to produce spillover; therefore, the role of endotoxins in the current work remains debatable. However, it is noteworthy that pretreating experimental animals with steroids, antiendotoxin antibodies, or with antibiotics to reduce gut flora reduces heat stress lethality (3, 10–12). On the basis of the present work, we suggest that the above-listed interventions may have acted by reducing NOS II activation.

In summary, interventions that increased systemic antioxidant capacity (aminoguanidine, PEG-SOD, or allopurinol) decreased production of radical biomarkers of metal-catalyzed oxidative stress. Agents that reduced NOS II or cellular oxidase activity (aminoguanidine and allopurinol) also significantly improved cardiovascular performance and heat tolerance. The mechanisms responsible for these results could involve decreased cellular ROS production and transition metal activation in combination with reduced flux of -NO.

Additionally, these data establish that intact NOS activity is required for normal heat tolerance, whereas overproduction of -NO and H2O2 may be responsible for local circulatory dysfunction with heat stroke. Xanthine oxidase may be a critical heat-sensitive cellular oxidase that contributes to intestinal barrier dysfunction and splanchnic dilation. On the basis of these data, we conclude that splanchnic oxidant generation is an important determinant of heat tolerance and that -NO is involved in vascular dysfunction with heat stroke.

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