Stressor-Induced Increase in Microbicidal Activity of Splenic Macrophages Is Dependent upon Peroxynitrite Production

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Exposing mice to a social stressor called social disruption (SDR) that involves repeated social defeat during intermale aggression results in increased circulating cytokines, such as interleukin-1α (IL-1α) and IL-1β, and increased reactivity of splenic CD11b+ macrophages to inflammatory stimuli. For example, upon lipopolysaccharide stimulation, macrophages from stressor-exposed mice produce higher levels of cytokines than do cells from nonstressed controls. Moreover, the SDR stressor enhances the ability of these macrophages to kill Escherichia coli both in vitro and in vivo, through a Toll-like receptor 4-dependent mechanism. The present study tested the hypothesis that stressor-enhanced bacterial killing is due to increases in the production of peroxynitrite. Male mice were exposed to the SDR stressor or were left undisturbed. Upon stimulation with E. coli, splenic macrophages from SDR-exposed mice expressed significantly increased levels of inducible nitric oxide synthase mRNA and produced higher levels of peroxynitrite. Blocking the production of peroxynitrite abrogated the SDR-induced increase in microbicidal activity. Studies in IL-1 receptor type 1 knockout mice indicated that the increased microbicidal activity and peroxynitrite production was dependent upon IL-1 signaling. These data confirm and extend the importance of IL-1 signaling for stressor-induced immunopotentiation; the finding that inhibiting superoxide or nitric oxide production inhibits both peroxynitrite production and killing of E. coli demonstrates that peroxynitrite mediates the stressor-induced increase in bacterial killing.

Macrophages play a fundamental role in innate immunity and are highly efficient phagocytes important in the first wave of immune defense. Multiple studies now indicate that macrophage activity can be strongly modified by environmental factors, such as the physiological response to social and psychological stressors (10, 21, 52). The mammalian response to stressors involves activation of many physiological systems, including the sympathetic nervous system (SNS) and the hypothalamic-pituitary-adrenal (HPA) axis (18). These stressor-induced physiological responses have been shown to impact immune system functioning and the outcome of pathogenic challenge. Most studies have focused on the ability of stressors to suppress immunity, and it is now known that factors such as HPA axis-derived glucocorticoid (GC) hormones can suppress important transcription factors, such as NF-κB (61). As a result, prolonged elevations of glucocorticoids have been shown to suppress macrophage activity (2, 10, 21, 53).

Exposing mice to a widely used and well characterized social stressor involving intermale aggression, called social disruption (SDR), reduces splenic macrophage sensitivity to the suppressive effects of stressor-induced GC hormones. This reduced sensitivity is associated with the development of splenomegaly due to an accumulation of myeloid-derived cells (29). These splenic cells are also more reactive to microbial antigens; they produce higher levels of cytokines upon lipopolysaccharide (LPS) challenge (3, 8, 56) and have enhanced microbicidal activity, in part through Toll-like receptor (TLR) signaling (7). While the physiological parameters by which stressor exposure enhances immune activity are becoming more well defined and involve stressor-induced SNS activity (18, 23, 28, 36) and alterations to the intestinal microbiota (1, 6), the cellular events responsible for the increased ability of splenic macrophages to kill bacteria have not been widely studied. Therefore, the purpose of the present study was to determine key cellular events necessary for stressor exposure to enhance the ability of splenic macrophages to kill microbes.

Macrophages use a variety of mechanisms and effector molecules to kill microbes (54). Their primary method, however, involves the rapid production of reactive molecules associated with an “oxidative burst.” These reactive molecules are then able to damage proteins and DNA of microbial cells (13, 40). A major enzyme responsible for reactive molecule production is inducible nitric oxide synthase (iNOS). Shortly after phagocytosis, iNOS is expressed de novo in the cytosol of macrophages where it converts the amino acid arginine into citrulline and the reactive molecule, nitric oxide (49). Nitric oxide then diffuses across the phagosome membrane, where it can damage the engulfed microbe. In addition, the NADPH oxidase complex located on the phagosome membrane produces superoxide anion, which rapidly combines with nitric oxide to produce a highly destructive intermediate, peroxynitrite. Peroxynitrite is a powerful oxidant that can damage and inactivate many crucial bacterial proteins (47). Since these molecules are vital to macrophage killing, and because previous
studies have suggested that SDR exposure enhances iNOS gene expression and the assembly of the NADPH oxidase complex (7), we hypothesized that SDR-induced increases in bacterial killing involve heightened expression of iNOS and subsequent peroxynitrite production.

Previous studies have shown that interleukin-1 (IL-1) is increased in the circulation and secondary lymphoid organs of stressor-exposed mice (30). This increased IL-1 is thought to be responsible for the insensitivity of splenic macrophages to the suppressive effects of glucocorticoid hormones; IL-1R1 knockout mice did not develop glucocorticoid insensitivity after exposure to the SDR stressor. Whether IL-1R1 signaling also impacts the microbial activity of splenic macrophages, however, has not been tested. This is a logical hypothesis, because it has already been shown that Toll-like receptor signaling is necessary for the enhanced microbial activity to occur (7), and IL-1R1 signaling shares significant downstream pathways with Toll-like receptors (11, 45, 50). Moreover, IL-1R1 signaling is known to induce iNOS gene expression in part through signaling through the mitogen-activated protein kinases (MAPKs) (19, 35, 39, 59). Thus, we tested the hypothesis that SDR-induced increases in microbial activity and peroxynitrite production were IL-1R1 dependent.

MATERIALS AND METHODS

Animal handling. Male C57BL/6 and CD-1 mice between the ages of 4 and 6 weeks were purchased from Charles River Laboratories (Wilmington, MA), IL-1R1-/- mice were bred at Ohio State University and were descendants of a breeding pair originally obtained from Jackson Laboratory (Bar Harbor, ME). They were shown to exhibit a normal phenotype here. After the 2-h incubation, the nonadherent cells were washed from the adherent splenocytes after 20 and 90 min, respectively. The cells were incubated for 2 h to allow the macrophages to adhere.

Bacteria. Escherichia coli strain K-12 (ATCC 10798), originally obtained from the American Type Culture Collection (Manassas, VA), was grown from frozen stocks in Trypycase soy broth overnight at 37°C.

Microbial assay. A total of 5 × 10⁴ CD11b+ monocytes/macrophages from individual animals were plated in duplicate in a volume of 1 ml per well on 24-well tissue culture plates to determine phagocytosis and bacterial killing by the adherent splenocytes after 20 and 90 min, respectively. The cells were incubated for 2 h to allow the macrophages to adhere. After the 2-h incubation, the nonadherent cells were washed from the plates by three rinses with 1 ml of RPMI 1640. After the final wash, 1 ml of a mixture containing 5 × 10⁴ CFU of E. coli in RPMI 1640 was added to each well, and the plates pulse-centrifuged to help pull the E. coli onto the adherent cells. The E. coli were opsonized by adding 5% fresh serum (pooled from nonstressed HCC mice and stressed SDR mice) to the bacterial suspension. After the bacteria were incubated for 20 min, the extracellular bacteria were washed away by pipetting 1 ml of RPMI 1640 into the well for three washes. After the final wash, 1 ml of 1% Triton X-100 was added to one of the wells to lyse the splenocytes. The lysate was collected into a sterile tube for pour plate enumeration of phagocytized bacteria. The duplicate well was washed three times with RPMI 1640 supplemented with 10% FBS to remove extracellular bacteria. After the final wash, 1 ml of RPMI 1640 supplemented with 10% FBS was added to the duplicate wells. After the plates were incubated for an additional 70 min. After 70 min, the cells were again washed and then lysed with 1% Triton X-100 to quantify bacteria remaining alive within the macrophages via standard pour plate analysis.

Pharmacological inhibitors of macrophage function. To inhibit the production of nitric oxide by iNOS during the microbicidal assay, 500 μM N⁶-β-methyl-L-arginine acetate salt (L-NMMA; Sigma-Aldrich, St. Louis, MO) was added to the RPMI 1640 supplemented with 10% FBS during both the 2-h adherence and during coincubation with E. coli. To scavenge peroxynitrite, 25 μM 5,10,15,20-tetrasulfonato(naphthalenedisulfonato-III) chloride (FeTPPS; Sigma-Aldrich) was added during the 2-h adherence and during coincubation with E. coli. To inhibit NADPH oxidase production of superoxide, 300 μM apocynin was added during the 2-h adherence and during coincubation with E. coli.

RNA isolation and cDNA synthesis. Total splenocytes from individual SDR and HCC control mice were cultured in duplicate sets at 5 × 10⁴ adherent macrophages per ml in 24-well tissue culture plates at 37°C and 5% CO₂ for 2 h. After the 2-h incubation, the nonadherent cells were washed off by three rinses with RPMI 1640. After the final wash, 1 ml of RPMI 1640 supplemented with 10% FBS was added to half of the wells, and the other half of the cells were stimulated with 1 μg of LPS/ml in 1 ml of RPMI 1640 supplemented with 10% FBS. After incubation for 90 min,
the cells were washed with RPMI 1640, and 1 ml of TRIzol reagent (Gibco, Rockville, MD) was added to each well to isolate RNA. RNA was isolated according to the TRIzol protocol provided by the manufacturer (Gibco). Total RNA was reverse transcribed using a commercially available kit (Promega, Madison, WI) according to the manufacturer’s instructions. Briefly, 1 µg of total RNA was combined with 5 mM MgCl2, a 1 mM concentration of each deoxyxynucleoside triphosphate, 1 × reverse transcriptase buffer, 1 U of RNasin/µl, and 15 U of avian myeloblastosis virus reverse transcriptase/µg and primed with 1 µg of random hexamers and diethyl pyrocarbonate-H2O to a volume of 20 µl. The reaction was first incubated for 10 min at room temperature and then at 42°C for 1 h. This was followed by 5 min of incubation in boiling water and then a cooling period of 5 min on ice. The volume was adjusted to 50 µl by adding 30 µl of nuclelease-free water.

**RT-PCR.** Primers and probes were synthesized by Applied Biosystems (Foster City, CA). The 3′-3′ sequences were as follows: iNOS (forward, CAG CTT GGC TGT ACG AACT CTT; reverse, TGA ATG TGA TGT TGG CTT CCG; probe: CGG GCA GCC TGT GAG ACC TTT GA) and 18S (forward, CGG GTA CCA CAT CCA AGG AA; reverse, GCT GGA ATT CAG CTG GGC TGT ACA AAC CTT; probe, TGA ATG TGA TGT TTG). The PCR mixture consisted of 2.5 µl of cDNA, 2.5 µl of primer mix (900 nM), 2.5 µl of probe, 5 µl of sterile distilled H2O, and 12.5 µl of TaqMan universal master mix (Applied Biosystems) for a final volume of 25 µl. After an initial 2 min cycle at 95°C, followed by 10 min at 95°C, the reaction ran for 40 total cycles consisting of a 15-s denaturing phase (90°C) and a 1-min annealing/extension phase (60°C). The change in fluorescence was measured using an Applied Biosystems 7000 Prism sequence detector (Applied Biosystems) and analyzed using Sequence Detector v1.0. The relative amount of transcript was determined using the comparative threshold cycle (Ct) method as described by the manufacturer. In these experiments, gene expression in the untreated cells from the spleens of the nonstressed HCC mice was used as the calibrator. Gene expression, therefore, is expressed as the fold increase in expression over nonstressed HCC control mice.

**Peroxynitrite and superoxide measurement.** After spleen cells from individual animals were processed to a single cell suspension, CD11b+ cells were magnetically separated from the total cells using anti-CD11b+ microbeads and MACS MS separation columns according to the manufacturer’s directions (Miltenyi Biotec, Auburn, CA). The separated CD11b+ cells were suspended in 1 ml of RPMI 1640 supplemented with 10% FBS and counted using a Coulter counter. After counting, the cells were treated with 1 × 106 cells/ml and were plated in two sets of duplicate wells on opaque 96-well tissue culture plates in a volume of 200 µl/well. The cells were incubated for 2 h at 37°C and 5% CO2, to allow the cells to adhere. After the 2-h incubation, the adherent cells were rinsed three times with HBSS without phenol red to remove nonadherent cells. To all wells, 100 µl of HBSS without phenol red containing 25 µM dihydroethidium (DHE; Sigma-Aldrich), a fluorophore (31,51). Upon stimulation, the vehicle-treated cells from mice exposed to SDR produced significantly higher amounts of peroxynitrite upon stimulation with PMA/LPS/IFN-γ than did cells from nonstressed HCC control mice [F(6,186) = 20.77, P < 0.001]. This was evident at every time point tested (P < 0.05). Importantly, when cells from mice exposed to the SDR stressor were cocultured with E. coli, they killed significantly more E. coli over the 90 min time course than did cells from nonstressed control mice (Fig. 1D). This effect was evident when the CFU/ml were determined [F(1,24) = 44.39, P < 0.001]. To determine whether increased nitric oxide production was responsible for the stressor-induced increase in peroxynitrite production and the enhanced macrophage bactericidal activity, cells from SDR-stressor-exposed mice were stimulated with LPS and the highest dose of IFN-γ producing more nitrite than all other groups (P < 0.05; Fig. 1B). Moreover, the increase in nitrite production was associated with a significant increase in the production of peroxynitrite (Fig. 1C). Cells from mice exposed to SDR produced significantly higher amounts of peroxynitrite upon stimulation with PMA/LPS/IFN-γ than did cells from nonstressed HCC control mice [F(6,186) = 20.77, P < 0.001]. This was evident at every time point tested (P < 0.05). Importantly, when cells from mice exposed to the SDR stressor were cocultured with E. coli, they killed significantly more E. coli over the 90 min time course than did cells from nonstressed control mice (Fig. 1D). This effect was evident when the CFU/ml were determined [F(1,24) = 44.39, P < 0.001].

To determine whether increased nitric oxide production was responsible for the stressor-induced increase in peroxynitrite production and the enhanced macrophage bactericidal activity, cells from SDR stressor-exposed and nonstressed HCC control mice were treated with L-NMMA, which blocks nitric oxide production (31,51). Upon stimulation, the vehicle-treated cells from mice exposed to SDR produced significantly higher levels of peroxynitrite than did vehicle-treated cells from HCC control mice [F(6,66) = 21.90, P < 0.001] (Fig. 2A). However, after L-NMMA was added to the culture, there was no difference in peroxynitrite production in stimulated cells from stressor-exposed and nonstressed control mice [F(6,66) = 0.14, not significant; Fig. 2A]. This finding is consistent with data obtained in the microbici-
treated splenic macrophages from mice exposed to the SDR stressor—FeTPPS, which is a peroxynitrite scavenger (38, 43, 44). Vehicle-peroxynitrite (12, 27), we next confirmed whether the enhanced oxide, can lead to the production of reactive molecules other than stressor-exposed or the nonstressed HCC control mice were In this case, neither the splenic macrophages from the SDR

"...E. coli..."}

reactive oxygen species are partly responsible for the enhanced bactericidal activity in CD11b+ cells from SDR-exposed mice. Since peroxynitrite is the reaction product of nitric oxide and superoxide anion, we assessed the role of superoxide anion production in contributing to the increase in peroxynitrite that is needed for the stressor-induced increase in bacterial killing. Superoxide anion production was induced by in vitro stimulation of CD11b+ splenocytes from stressed and nonstressed animals with...
PMA/LPS/IFN-γ. Vehicle-treated cells from SDR stressor-exposed animals produced significantly more superoxide anion than did cells from nonstressed HCC control mice or from cells treated with apocynin to block NADPH oxidase activity \[F(6,60) = 57.10, P < 0.05; \text{Fig. 4B}\]. Importantly, stressor-exposure increased killing of E. coli in vehicle-treated mice \[F(1,27) = 27.20, P < 0.001; \text{Fig. 4C}\], whereas inhibition of NADPH oxidase activity eliminated the stressor-enhanced bacterial killing during \textit{in vitro} culturing \[F(1,28) = 3.21, \text{not significant}; \text{Fig. 4D}\].

**IL-1 signaling contributes to enhanced bactericidal activity in CD11b+ cells from SDR-exposed mice.** Previous studies have shown that IL-1α and IL-1β mRNA expression and protein production are increased in secondary lymphoid organs following SDR (30). Also, elimination of IL-1R1 signaling through the use of an IL-1R1 knockout (IL-1R1−/−) mouse strain demonstrated that some of the effects of SDR on the immune system, namely, trafficking of CD11b+ cells to the spleen and the development of glucocorticoid insensitivity in this cell population, fail to appear (30). Therefore, we further examined whether IL-1R1 signaling was necessary for the SDR stressor-enhanced iNOS gene expression, superoxide anion and peroxynitrite production, and microbicidal activity. Splenic macrophages from wild-type C57BL/6 mice had a significant increase in iNOS mRNA after \textit{E. coli} stimulation in comparison to iNOS mRNA in splenic macrophages from nonstressed control mice \[t(15) = 2.19, P < 0.05; \text{Fig. 5A}\].

**FIG 3** Use of a peroxynitrite scavenger prevented the stressor-induced increase in microbicidal activity. (A) The number of \textit{E. coli} remaining alive in the cultures at the 90-min time point was significantly lower in the presence of splenic macrophages from SDR-exposed mice compared to nonstressed controls. (B) When the peroxynitrite scavenger FeTPPS was added to the cultures, there was no difference in \textit{E. coli} levels in cultures containing cells from SDR exposed or from HCC control mice (*, P < 0.05; n = 7 HCC and n = 10 SDR from two different experiments).

**FIG 4** Stressor exposure increases both superoxide anion and nitric oxide production; these intermediates react to form peroxynitrite. (A) Stressor exposure increases the production of superoxide anion, and \textit{in vitro} inhibition of NADPH oxidase activity eliminates this increase \((n = 6 \text{ HCC and } n = 6 \text{ SDR from two different experiments); *, P < 0.05}\). (B) Inhibiting the activity of NADPH oxidase significantly reduces stressor-induced peroxynitrite production (*, P < 0.05 at 45, 60, 75, and 90 min; n = 6 HCC and n = 6 SDR from two different experiments). (C) Vehicle-treated stressor-exposed mice killed more \textit{E. coli} \textit{in vitro} than nonstressed controls \((n = 15 \text{ HCC and } n = 15 \text{ SDR from four different experiments); P < 0.05 versus HCC control}. (D) Reduction of peroxynitrite production through blocking the formation of superoxide anion significantly reduces the stress-induced increase in bacterial killing (*, P < 0.05 versus HCC; n = 15 HCC and n = 15 SDR from four different experiments).
This stressor-induced increase in iNOS mRNA was not evident in the IL-1R1−/− mice (t(4) = 1.16, not significant) (Fig. 5A). Likewise, cells from stressor-exposed wild-type C57BL/6 mice produced higher levels of both superoxide anion [F(6,120) = 9.64; P < 0.001; Fig. 5B] and peroxynitrite [F(6,95) = 11.01; P < 0.001; Fig. 5C]. Importantly, both superoxide and peroxynitrite production was significantly lower in the SDR-exposed IL-1R1−/− mice compared to wild-type mice exposed to SDR (P < 0.05). Stressor-exposed WT mice killed more E. coli than nonstressed controls [F(1,14) = 36.40; P < 0.001; Fig. 5D]. However, stressor-exposure failed to increase microbicidal activity in the splenic macrophages from IL-1R1−/− mice (Fig. 5E). The number of E. coli within the cells from stressor-exposed mice was similar to the number found within the nonstressed controls after 90 min in culture [F(1,127) = 0.21, not significant] (Fig. 5E), and neither the stressor-exposed nor the nonstressed IL-1R1−/− mice were able to kill E. coli (Fig. 5E).

**DISCUSSION**

This study confirms and extends previous reports showing that stressor exposure can enhance the bactericidal ability of macrophages by identifying key cellular events that are responsible for the enhanced killing. The SDR-enhanced killing was observed early after phagocytosis (<90 min), which is consistent with the amount of time needed to develop reactive intermediates (22, 24). Thus, we hypothesized that a possible target for stress-induced modification is the production of reactive molecules which begins immediately following phagocytosis. In support of this hypothesis, macrophages from stressed animals stimulated with whole E. coli expressed an ∼10-fold increase in iNOS gene expression over gene expression in E. coli-stimulated cells from nonstressed control mice. Moreover, when splenic macrophages from mice exposed to SDR were incubated with the iNOS inhibitor L-NMMA (a synthetic form of arginine that cannot be converted to produce nitric oxide), this increase in stress-induced killing was abolished. This finding is consistent with reports from others that have shown that stressor exposure can enhance macrophage phagocytosis (46) and microbicidal activity through the enhanced production of nitric oxide (23, 25, 32).

It is not likely that nitric oxide was directly responsible for the enhanced microbicidal activity since nitric oxide has only moderate microbicidal activity in comparison to other reactive species (13, 16, 47). Moreover, in the presence of superoxide anion, nitric oxide rapidly reacts with superoxide to produce peroxynitrite,
which is a powerful oxidant that is able to severely damage bacteria through oxidation and nitration of proteins (47, 50, 54). Studies in vitro have shown that ligating neurotransmitter receptors, namely, the α-adrenergic receptor, on macrophages increases peroxynitrite production (65), which has been suggested to be important for E. coli killing (14, 34, 63, 67). Because previous studies have indicated that NADPH oxidase, and thus superoxide, is enhanced in the spleens of mice exposed to the SDR stressor prior to intravenous E. coli challenge (7), and because of the observed stressor-induced increase in iNOS gene expression, we assessed whether splenic macrophages from mice exposed to the SDR stressor would produce higher levels of peroxynitrite. As predicted, peroxynitrite production was significantly increased. Importantly, blocking the effects of the iNOS enzyme in turn blocked the increase in peroxynitrite as well as bacterial killing. In addition, blocking superoxide anion production also blocked peroxynitrite production and bacterial killing. These data, along with the finding that scavenging peroxynitrite abolished the stressor-induced increase in the microbicidal activity of splenic macrophages, support the hypothesis that stressor-induced increases in microbicidal activity is ultimately due to the increased production of peroxynitrite from reactive nitrogen and oxygen intermediates.

The use of IL-1R1−/− mice indicates that IL-1 signaling is necessary for the stressor-induced increase in microbicidal activity to occur. This finding is important because studies in humans demonstrate that both prolonged natural stressors and acute laboratory stressors often result in increased circulating cytokines, such as IL-6, TNF-α, and IL-1 (57). This is consistent with findings from laboratory animals that have shown that a variety of different stressors, such as tail shock, acute restraint, and social interactions, also elevate circulating cytokines (25, 26, 48, 66). Exposure to the SDR stressor, in particular, has been shown to elevate circulating levels of IL-6 and TNF-α (3, 56), as well circulating and lymphoid tissue levels of IL-1α and IL-1β (30). The mechanisms by which stressor exposure enhances these cytokines are becoming more well defined and are thought to involve stressor-induced activation of the sympathetic nervous system (35), as well as stressor-induced alterations of the microbiota (1, 6). The biological importance of stressor-induced increases in cytokine levels has not been systematically studied, but our results suggest that increased macrophage activity is one biological outcome of stressor-induced increases in cytokine levels.

Signaling through the IL-1R1 is not the only way in which stressor exposure can enhance the activity of splenic macrophages. The expression of TLR2 and TLR4 is enhanced on splenic macrophages from mice exposed to SDR (7), and our previous results demonstrated that exposing mice to the SDR stressor prior to intravenous challenge with E. coli significantly increases the rate at which bacteria can be cleared from the blood and from the spleen (7). Importantly, the SDR-induced increase in clearance was not evident in C3H/HeJ mice that lack functional TLR4, suggesting that signaling through TLR4 is also necessary for the stressor-enhanced killing to manifest. This may not be surprising, since the IL-1R1 and TLR4 utilize many overlapping intermediates in their signaling cascades, and the proteins needed for producing reactive nitrogen and oxygen intermediates have close signaling links to TLR4 and IL-1R1 (17, 19, 39, 59).

Recently, a microarray study quantifying the expression of over 45,000 assayed transcripts found overexpression of key transcription factors, such as NF-κB, PU.1, and NRF2 and evidence of MAPK signaling (N. D. Powell et al., unpublished data). A key component of TLR4 and IL-1R1-induced iNOS and NADPH oxidase activation are the MAPKs (17, 19, 35). Thus, it is possible that the physiological response to the SDR stressor primes cells for enhanced MAPK signaling through an upregulation of TLR4 and IL-1R1 (7, 30). Such priming would in turn increase peroxynitrite-dependent microbicidal activity. This hypothesis will be further tested in future studies.

The physiological stress response has long been considered to be largely immunosuppressive, but accumulating evidence indicates that in some circumstances the stress response instead enhances immunity. This enhanced immune activity can be adaptive, particularly in laboratory animals challenged with bacteria (7, 15, 23, 32) or during squamous cell carcinoma (26). It must be recognized, however, that the reactive molecules modified upon stressor exposure have important roles in numerous physiologic responses that can tip the balance between health and disease (9, 20, 42, 64). Although important for their bactericidal effects, high levels of nitric oxide and superoxide anion can be destructive to the host, causing tissue damage and exacerbating inflammatory conditions such as sepsis, asthma, arthritis, ulcerative colitis, and periodontitis (37, 41, 58, 60, 62). Thus, beneficial outcomes, e.g., increased bacterial clearance, and negative outcomes, e.g., tissue damage, must be balanced when the stress-response modifies innate immunity. Understanding how specific stressors and their associated physiological stress response impact infection and the inflammatory response could lead to more effective treatments, particularly in susceptible patient populations.

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