Antioxidants in Infection

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Summary       Endogenous oxidation reactions are essential for the normal biochemistry of life and are especially critical for leukocyte microbial killing mechanisms in host defense to infectious diseases. However, reactive oxidative intermediates can damage normal tissues unless kept under antioxidant control. Three selected examples of oxidant-antioxidant systems involved in infectious diseases are discussed, regulation of molecular iron catalyzed oxidations, superoxide scavengers and inhibitors of nitric oxide synthase in septic shock, and the use of glutathione replacement therapy in HIV infection and AIDS. The data suggest that antioxidants, and therapy based on increasing antioxidant potential, have a major impact on clinical infectious diseases.

Key Words    anti-oxidants infection, oxygen radicals, septic shock, iron, nitric oxide, glutathione.

INTRODUCTION

The physiological and pathophysiological roles of oxidant stress and the importance of antioxidants in clinical medicine have only just begun to be appreciated in the field of infectious diseases. For example, in a recent Symposium organized by the joint Malnutrition Panels of the U. S. Japan Cooperative Medical Sciences Program (1), a session entitled “Antioxidants in Health and Disease” made mention only of the interactions of vitamin E and malaria, a paradoxical situation in which increased vitamin E levels protect parasites within infected red cells from oxidant mediated elimination (2). Other references to oxidant-antioxidant systems in infection were lacking. The purpose of this review is to highlight the role of oxidants-antioxidants in infectious diseases by discussing three selected examples of increasing current interest, and by so doing to stimulate further research in this important subject.
There are two important principles to elaborate before discussing specific examples of oxidant-antioxidant systems in infection. First, antimicrobial host defenses are strongly dependent on induced oxidant stress to the microorganisms causing infection. It has long been known that a burst of leukocyte produced reactive oxygen intermediates (ROIs) and free radicals is essential for antimicrobial host defenses (3). The oxidative burst results in the generation of reactive oxygen species such as \( \text{H}_2\text{O}_2 \), the superoxide anion \( \text{O}_2^- \cdot \), singlet oxygen \( ^1\text{O}_2 \), and the hydroxyl radical \( \text{OH}^- \cdot \), as well as secondary reaction products such as hypochlorous acid generated from interactions of \( \text{H}_2\text{O}_2 \) and \( \text{Cl}^- \) ion within cells. Patients with genetic diseases in which the capability to activate oxidative mechanisms is impaired, such as chronic granulomatous disease, are known to be at increased risk of infection. In addition, recent data demonstrate that nitric oxide (NO) and \( \text{O}_2^- \cdot \) react to produce peroxynitrite \( \text{OONO}^- \) and the free radical of nitric dioxide \( \text{NO}_2^- \) which are essential for macrophage tumoricidal and bactericidal activities \textit{in vitro} (4, 5). While the importance of the latter mechanisms for host defense \textit{in vivo} remains to be conclusively shown, they appear likely to be of clinical relevance.

Secondly, it is well established that uncontrolled oxidative reactions are harmful to normal tissues, and result in chemical damage to proteins, lipids and nucleic acids (6). Therefore it is absolutely necessary to control the reactions which release ROI's and free radicals by biological antioxidants and scavengers in biological systems. However, this control must be balanced, for, as Olson has noted, “the complete quenching of free radicals in cells also quenches life” (7). It is therefore necessary to think of oxidant stress and antioxidants as a system, in which failure to balance the two may result in disease. In this context, the broader definition of antioxidants proposed by Krinsky (8) is very appropriate: biological antioxidants are “compounds that protect biological systems against the potentially harmful effects of processes or reactions that can cause excessive oxidations.”

IRON AND IRON CONTAINING COMPOUNDS AND IRON BINDING PROTEINS

Iron, a highly reactive metal, is present in the body in ample quantities. In the normal reduction of molecular oxygen to water, oxygen radicals are formed which, in turn, will react with free iron by the Haber-Weiss reaction to form the highly reactive hydroxyl radical. This is a predictably tissue damaging reaction and uncontrolled, except for the fact that the binding of nearly all free iron into iron centered structures (eg heme) and iron-metalloproteins severely limits the amount of free iron available for these reactions. Iron-metalloproteins comprise various enzymes necessary for life, including those functioning to carry oxygen, to detoxify reactive oxygen species, to replicate DNA, and various proteins designed simply to transport iron in a non-reactive state. Because the
simultaneous need for iron to serve in its many functional roles (including generation of ROI's for control of infection) and to segregate the metal so that it cannot catalyze excessive tissue damaging oxidations is essential for health, iron metabolism is a good example to illustrate the principle of balanced oxidant-antioxidant systems.

Much of the clinical literature on iron has focused on the problem of iron deficiency and its treatment with iron (9). There is an expressed concern that iron administration may increase the availability of the metal for use by invading pathogens, especially when circulating iron-binding proteins are reduced and unable to fully bind extracellular iron, such as occurs in patients with protein-energy malnutrition. In addition, the physiological shift of iron from extracellular to intracellular compartments under the influence of cytokines produced during inflammation and infection has been viewed as a host defense strategy to further restrict access to iron by the infectious agent. The potentially favorable effects of reduced iron availability, however, may be offset by the reports of diminished immune function in iron deficiency states, which would increase host vulnerability to infections (10). While there remains major controversy as to the relative importance of these two different impacts of iron, the balance of evidence that pathogens have efficient mechanisms of obtaining iron, even under the most restrictive of in vivo conditions, and that they use low iron as signal to regulate virulence genes, including those controlling iron access systems, suggests that states of iron deficiency are more harmful to the host than to the pathogen (10).

On the other hand, states of iron excess, such as chronic hemolysis associated with hemoglobinopathies, hemochromatosis, and transfusion siderosis are associated with excessive infection morbidity and mortality. Here too controversy exists about the importance of free iron levels. On the one hand, because the pathogens which emerge under these conditions are often those lacking an effective system of iron acquisition, the availability of extra free iron may be essential for their virulence (11, 12). On the other hand, evidence of impaired immune function in iron overload states (13, 14) suggests that this condition increases susceptibility to infection, presumably by catalyzing membrane damaging lipid peroxidations which impairs cell function. In effect, a ying-ying situation exists in which either iron deficiency or excess results in impaired immune function and increased infections, with the latter being due to excessive oxidative reactions. Hence, normal iron metabolism is a natural balance of oxidative reactions catalyzed by iron and anti-oxidative iron trapping mechanisms which promotes optimal host defense to infection without damaging the host in the process.

**ROIs AND NITROGEN OXIDE RADICALS IN SEPTIC SHOCK AND SHOCK LUNG**

The pathogenesis of septic shock is an example of what may occur when complex regulatory mechanisms are uncontrolled; and represent a setting in

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which "if a little is good, a lot may be worse." A multiplicity of interactive
responses are turned on by infection, and attempts to visually present these
interactions lead to some of most complicated diagrams ever devised in biology.
The reader is referred to a recent review for a much more detailed presentation
of these complexities (15). For the purpose of the present discussion, we need
only consider a more simplified schema:

INITIATION → PROPAGATION → INTERACTION → DAMAGE

Septic shock is usually initiated by infection with gram negative bacterial
pathogens which release large quantities of endotoxin (lipopolysaccharide, LPS),
but also occurs during infection by gram positive pathogens which do not contain
LPS. One common feature is that LPS and peptidoglycan from the gram
positive organisms both result in the release of a similar group of cytokines from
leukocytes. Both gram negative and positive pathogens also activate the
complement system, a cascade of interacting proteins required to generate key
intermediates in the inflammatory response, eg C5a, the activated fragment of
the 5th component of complement. These mediators activate leukocyte
metabolism and endothelial cell adhesion, and upregulate the production of
ROI’s and free radicals within phagolysosomes, from increased arachidonate
metabolism, and by activation of nitrogen oxide synthase (NOS), increasing the
production of inducible NO. There are also important interactions among these
mediators, for example the ability of some cytokines to regulate the synthesis of
one another, the effects of released platelet activating factor (PAF) and other
mediators to increase ROI production by neutrophils and platelets, and the
ability of ROI’s to react with one another, leading to even more tissue damaging
products such as OH•, OONO−, and hypochlorous acid.

Traditional therapy for the complex pathophysiology of septic shock has
depended on antimicrobial therapy to eradicate the infection and pressor agents
to maintain tissue perfusion. Unfortunately, the release of biologically active
cell wall products from antibiotic damaged or dying organisms, including LPS
and peptidoglycan, can make the clinical situation worse initially. Induction of
the mediators of inflammation by these microbial constituents can increase the
volume of fluids which leak across damaged endothelium, leading to shock lung.
In addition, the use of pressors to combat hypotension may actually reduce blood
flow to tissues and either directly cause damage or set the stage for a classical
ischemia-reperfusion ROI induced damage (16). It is not surprising that
traditional therapy has limited effectiveness. Recent attempts to use monoclonal
antibodies against LPS to interrupt the cycle by inactivating the toxic effects of
LPS have improved responses in gram negative septic shock, but only to a limited
extent (17). Therefore, further experimental studies have been undertaken to
more directly control oxidative damage and reduce the NO mediated
vasodilation, which may be critical to the pathogenesis of septic shock (15).

The favorable effects of ROI scavengers and antioxidants in experimental
endotoxin shock or E. coli sepsis has provided the most compelling evidence for
the key role of oxy-radicals in the pathogenesis of septic shock. For example,
Yoshikawa has shown that either superoxide dismutase (SOD) or catalase reduce the rapid hypotensive effects of LPS injections (18), and Broner et al. have reported that SOD, but not N-acetyl cysteine (NAC), reduces endotoxin induced mortality. However, both confirmatory (19, 20) and contradictory (21, 22) data have been published. Some of the discrepancies may be due to the use of different models (23) and the different pharmacology and pharmacokinetics of the inhibitory agents employed, and there remains much more work to be done. In particular, little direct evidence has been obtained thus far in humans and further study is both required and warranted.

A parallel set of studies has begun to evaluate the role of nitric oxide (NO) in shock. This is a stable, freely diffusible, free radical gas derived from the oxidation of arginine by the enzyme nitric oxide synthase (or NOS), which appears to be a major regulator of normal vascular tone (24). NO is now known to be identical to endothelial-derived relaxing factor, and it is the active metabolite of nitroglycerin and other organic nitrate coronary artery dilators. It binds to the iron heme center of guanyl cyclase and activates production of cGMP which, in turn, regulates a cGMP dependent protein kinase leading to vascular smooth muscle relaxation. Neutrophils and macrophages contain a distinct, inducible NOS, which is activated by cytokines to produce NO. Because cytokines also activate oxidative reactions leading to superoxide production, and NO is attacked by $O_2^- \cdot$ to form peroxynitrite, which decomposes to highly reactive radicals such as $OH^-$ and $NO_2^-$, a highly microbicidal milieu develops. Overproduction of NO, especially at an endothelial site, could theoretically lead to inappropriate vasorelaxation and a drop in blood pressure.

Can this happen in septic shock? This possibility has been suggested (25), in part because C5a activation of neutrophils leads to the expression of surface receptors mediating neutrophil attachment to vascular endothelium. Several experimental studies have shown that NOS inhibitors greatly affect shock caused by LPS infusion. Thus, use of $N^G$-methyl-L-arginine (L-NMA) in a dog model of endotoxemia (which resulted in a 33% drop in peripheral vascular resistance and a 54% decrease in systolic blood pressure) led to restoration of normal values within 1.5 min of administration of the drug (26). In a rodent model, a dose dependent effect of L-NMA was reported, in which 3 mg/kg L-NMA was ineffective, 30 mg/kg reversed endotoxin induced shock, and 300 mg/kg accelerated the fall in blood pressure initiated by endotoxin infusion (27).

Similar findings have been reported when the shock syndrome was induced by a cytokine, tumor necrosis factor (TNF) (28). This may be the pathway by which glucocorticoids exert an effect in septic shock, since the transcriptional activation of both TNF and inducible NOS is prevented by steroids (29). Two patients with refractory septic shock have recently been reported who received treatment with NOS inhibitors, either L-NMA or $N^G$-nitro-L-arginine methyl ester (L-NAME) (30). In both patients a rapid dose-dependent rise in blood pressure was observed in association with a rise in systemic vascular resistance.
Continuous infusion of L-NAME in one patient allowed blood pressure maintenance with reducing doses of noradrenaline, and by 48 h both drugs could be stopped. However, this patient died 2 days later of “recurrent intra-abdominal sepsis, adult respiratory distress syndrome, and disseminated intravascular coagulation” (DIC). This has raised the possibility that NOS inhibitors are two edged swords, including the improvement of blood pressure in septic shock, but also perhaps resulting in harmful vasoconstriction in the visceral circulation leading to bacterial translocation across a damaged bowel, coronary artery constriction leading to compromised cardiac function and lowered cardiac output, platelet aggregation and promotion of DIC, and reduction in NO dependent microbial activity (31).

That there is a complex and perhaps dual role of NO in septic shock is also indicated by experiments in which infusion of NOS inhibitors or administration of NO or arginine have improved outcomes in various shock models (32–34). There is also the theoretical possibility, supported by limited clinical information, that inhalation of NO can result in vasodilation of pulmonary vessels in ventilated segments of the lung during adult respiratory distress syndrome (ARDS) associated with sepsis (35). In this manner, oxygen exchange across ventilated regions of the lung might be preferentially improved, with significant clinical improvement. Since NO does not react readily with O₂ to form toxic products, there seems little chance of local damage in this setting. However, the findings suggest that inhibition of NOS could worsen established ARDS in septic patients. These data make it clear that human studies need to be cautiously and carefully done, and that a balance in NO and ROI metabolism may be essential to optimize outcomes in septic shock.

On the other hand, some data suggest that ARDS may be the consequence of local production of ROI’s and free radicals. A necessary and critical feature of ARDS, a syndrome characterized by increased permeability of small pulmonary vessels leading to pulmonary edema, refractory hypoxemia, and respiratory failure, appears to be the recruitment of neutrophils to the lung and the production of ROI’s during activation of the cells (36). In one study, administration of superoxide dismutase conjugated to polyethylene glycol (which lengthens the half life of the enzyme and appears to increase its uptake by vascular endothelium) to guinea pigs with lung injury secondary to experimental E. coli sepsis significantly attenuated the damage (37). Another study has reported depressed levels of alveolar fluid glutathione levels in patients with ARDS (38), which suggests that antioxidant deficiency may contribute to lung damage in sepsis. This is consistent with prior experimental studies showing that administration of oxidant generating systems into the rabbit lungs in vitro or in vivo will result in an experimental form of ARDS resembling that seen in human patients (39, 40), and with recent reports that expiratory gas condensates of patients with ARDS contains measurable levels of H₂O₂ (41, 42). The potential interaction of reactive oxygen species and NO to damage the lung, and the ability of systemic or inhaled antioxidants (such as N-acetyl cysteine) to
prevent the oxidant mediated damage in ARDS, especially if NO inhalation therapy is likely to be employed, should be thoroughly studied.

GLUTATHIONE DEFICIENCY IN THE ACQUIRED IMMUNODEFICIENCY SYNDROME

Selenium nutrition was first investigated in patients with AIDS idiopathic cardiomyopathy in 1986 because of the reported relationship of selenium deficiency and cardiomyopathy described in Keshan Disease in China (43). Clinical improvement with the oral administration of selenium salts was described the following year (44). Two years later, reduced systemic thiol or glutathione (GSH) levels were reported in AIDS patients (45, 46), and subsequently, reduced intracellular GSH levels were detected in T lymphocytes, both CD4+ and CD8+ (47). While the extent of the median GSH reduction was modest, in the range of 60% of normal, this was actually due to the loss of a subset of CD4 cells with a high GSH level and not just a reduction in the GSH content of all T cells (48).

The potential significance of these findings has not been ignored, and increasing evidence for an inverse relationship between intracellular CD4 antioxidant capacity and replication of HIV has been reported. In vitro studies have shown that certain cytokines stimulate HIV replication (49), and that GSH, GSH-esters, or N-acetyl cysteine inhibit this event (50). Cytokine-stimulated upregulation of HIV proceeds via the activation of a nuclear transcription factor, NFKB (51), responding to ROI’s produced during cytokine stimulation (52). It was therefore rational to attempt to inhibit this process by exogenous antioxidant thiols, such as N-acetylcysteine, which in fact prevents the activation of NFKB by either phorbol esters or TNF (53). While the function of high GSH CD4+ cells is not clear, or even if they represent a distinctive subset of CD4+ cells, GSH deficiency has been shown to reversibly impair several immune responses in which these cells participate, including T cell proliferation and differentiation, MLR, CTL and NK activity (50). Herzenberg and colleagues have shown that GSH and CD4 cell number vary independently, however in as much as lowered GSH levels are observed even in asymptomatic HIV+ individuals (48), an influence of low levels may be exerted early in the course of the disease. Since the progression of asymptomatic HIV infection to AIDS is associated with a burst of virus proliferation and is accompanied by recurrent infections, with both ordinary as well as opportunistic pathogens, it is certainly reasonable to propose that oxidant stimuli generated by cytokine activation during these infections can directly contribute to the upregulation of the virus, especially if unopposed by adequate antioxidant capacity. If this is so, then restoration of antioxidant capacity might have a salutory effect, reduce virus replication, and delay clinical advance of HIV infection towards AIDS (48). Studies are currently under way to evaluate this possibility. If the hypothesis is proven correct, it will propel antioxidant therapy to the fore in HIV, and, no doubt, will encourage similar investigations in other infectious diseases.
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