The dark side of dioxygen biochemistry
Joan Selverstone Valentine*, Diana L Wertz, Thomas J Lyons, Lee-Loung Liou, Joy J Goto and Edith Butler Gralla

The cellular biochemistry of dioxygen is Janus-faced. The good side includes numerous enzyme-catalyzed reactions of dioxygen that occur in respiration and normal metabolism, while the dark side encompasses deleterious reactions of species derived from dioxygen that lead to damage of cellular components. These reactive oxygen species have historically been perceived almost exclusively as agents of the dark side, but it has recently become clear that they play beneficial roles as well.

Addresses
Department of Chemistry and Biochemistry, UCLA, Los Angeles, CA 90095-1569, USA
*e-mail: jsv@chem.ucla.edu
Current Opinion in Chemical Biology 1998, 2:253–262
http://biomednet.com/elecref/1367593100200253
© Current Biology Ltd ISSN 1367-5931

Abbreviations
Co Q or Q coenzyme Q, ubiquinone
GPx glutathione peroxidase
GSH reduced glutathione
GSSG oxidized glutathione
HNE 4-hydroxy-2-nonenal
MDA malondialdehyde
NADH reduced nicotinamide adenine dinucleotide
NADPH NADPH phosphate
NO nitric oxide
PUFA polyunsaturated fatty acid
ROS reactive oxygen species
SOD superoxide dismutase

Introduction
The chemical biology of dioxygen encompasses a large variety of reactions, most of them highly beneficial to the organisms in which they occur, but some of them deleterious. Several examples of beneficial reactions are covered in other articles in this issue, for example, reactions of dioxygen catalyzed by enzymes such as cytochrome ϵ oxidase or monooxygenase or dioxygenase enzymes. By contrast, the major emphasis in this review will be on reactions from the ‘dark side’ of dioxygen biochemistry, that is, those that cause oxidative damage in vivo, and on the biological systems that have evolved to defend against such sources of oxidative stress (see Figure 1).

Dioxygen, O2, is a powerful oxidizing agent, and the energy that fuels most nonphotosynthetic biology is obtained by reducing it to two water molecules in enzyme-catalyzed reactions. It must be supplied continuously to respiring cells but does not diffuse fast enough on its own to supply each cell of multicellular organisms. Consequently, proteins such as hemoglobin or myoglobin that bind, transport, store, and release dioxygen have evolved to aid in its rapid delivery. Dioxygen is also used as a source of oxygen atoms in a large variety of enzyme-catalyzed biosynthetic reactions of organic substrate molecules. The same oxidizing power of dioxygen that is the basis of respiration, however, also makes dioxygen simultaneously an agent of toxic oxidative stress [1**].

The dioxygen molecule is unusual in having two unpaired electrons in its most stable form. Consequently, its direct reactions with other molecules are generally slow in the absence of catalysts or radical initiators and are therefore not the primary causes of oxidative stress. Instead, superoxide, hydrogen peroxide, organic peroxides, hydroxyl radical, peroxynitrite, and other energetic molecules derived from their further reactions appear to be the agents of oxidative damage. These molecules or ions are collectively termed ‘reactive oxygen species’, or ‘ROS’. Their ultimate source appears to be mitochondria, where side reactions of dioxygen with components of the respiratory chain reduce it to superoxide. Superoxide may itself cause damage (see below) or may react further to give other ROS.

Antioxidant systems exist in cells to protect against ROS [1**]. Antioxidants in aqueous compartments, for example the cytosol and the extracellular fluids, consist of low molecular weight antioxidants such as glutathione, ascorbate (vitamin C) [2*] and urate, reductase enzymes that catalyze the regeneration of the reduced forms of these antioxidant molecules [3] and antioxidant enzymes such as superoxide dismutases (SOD) [4**], catalases and peroxidases. One class of molecules that is particularly susceptible to oxidative damage is the polyunsaturated lipids that are present in membranes of higher organisms. Unprotected, these molecules are highly susceptible to free-radical autoxidation reactions (Figure 2) that are a significant threat to membrane integrity and function. But the presence of abundant membrane-soluble free radical chain-breaking antioxidants such as α-tocopherol (vitamin E) and reduced ubiquinone (coenzyme Q, Co Q or Q) [5**], along with coupled enzymatic systems that use reduced nicotinamide adenine dinucleotide phosphate (NADPH) to keep them reduced, provide excellent protection against such damage, which only occurs when these defenses are depleted or overwhelmed.

ROS cause oxidative damage of proteins [6**,7*,8**], lipids and lipoproteins [9**], nucleic acids [10**,11**], carbohydrates and other cellular components [6**,10**] under conditions of oxidative stress, but the exact chemical identity of the particular damaging agent and
A schematic overview of some of the pathways leading to oxidative stress and of antioxidants that defend against them in a typical eukaryotic cell (center; mito, mitochondrion; ER, endoplasmic reticulum). There are four classes of oxidative damage: (a) Site-specific oxidative damage involving metal-catalyzed generation of hydroxyl radical from hydrogen peroxide which results in strand breaks and base damage in DNA. Similar events could occur wherever metal ions bind adventitiously. (b) Lipid peroxidation which damages membranes as well as producing toxic products such as MDA (malondialdehyde) and HNE (4-hydroxy-2-nonenal) which react with other cell components. (c) Damage to proteins resulting from direct oxidations by reactive oxygen species (ROS) or reactions with the products of lipid metabolism (for example, HNE, MDA). (d) Direct reactions of superoxide itself with certain iron–sulfur cluster prosthetic groups in exposed positions which result in full or partial disassembly of the cluster, inactivation of the enzyme, and release of iron. (Iron released in this manner may go on to catalyze more hydroxyl radical generation at specific locations.) (e) A schematic representing the major source of superoxide and hydrogen peroxide in the cell – leakage of electrons from the electron transport chain. I, III, and IV represent complexes I (NADH dehydrogenase), III (coenzyme Q: cytochrome c oxidoreductase) and IV (cytochrome oxidase) of the electron transport chain. Q, coenzyme Q; C, cytochrome c. (f) Defensive molecules are listed according to whether they are present in aqueous or lipid compartments. SOD, superoxide dismutase; GSH, reduced glutathione.

Until recently, the best characterized of the reactions involved in biological oxidative damage were those that appear to be due to attack either of hydroxyl radical, HO·, or of a metal-bound species with a similarly high degree of hydrogen atom abstraction reactivity. These ROS...
are generated in metal-catalyzed reactions of hydrogen peroxide [12] and are known to be capable of initiating free radical autoxidation of lipids and damaging protein and DNA when they are generated in close proximity to such sites [13••]. Superoxide anion (O$_2^-$), the one electron reduction product of molecular oxygen, is believed to be a key player in hydroxyl radical generation in vivo because its dismutation is the primary source of cellular H$_2$O$_2$ and possibly also because it can play the role of reducing agent for catalytic metal ions in the Fenton reaction. Superoxide on a chemical level is a rather sluggish reactant, however, and until recently there has been little hard evidence for toxicity due to direct reactions of O$_2^-$ itself. In the past few years, however, targets damaged specifically by superoxide have been identified. Certain iron–sulfur cluster-containing enzymes are known to be directly inactivated by superoxide in vivo and in vitro. These include the TCA cycle enzyme aconitate and dihydroxyacid dehydratase, an enzyme involved in branched chain amino acid synthesis in Escherichia coli [14••,15].

As might be expected, increases in oxidative stress cause cells to synthesize higher levels of antioxidants, antioxidant enzymes, repair enzymes, and other molecules that mitigate the effects of such stress. During the past few years, considerable progress has been made in characterizing the biochemical mechanisms involved in signal transduction and regulation of cellular responses to changes in levels of oxidative stress [16••,17,18]. Of particular interest in this area are new findings concerning the link between oxidative stress, mitochondrial function, and the signaling pathways of apoptosis (a form of programmed cell death) [19••]. Finally, the evidence linking oxidative damage to a large number of human diseases is beginning to accumulate.

Dioxygen biochemistry is a large and active field and we could not hope to cover all of the important publications of the past year. Instead we have chosen those that appear to us to represent major themes of current research on biological oxidative stress. The articles that we have identified as being of special interest are those dealing with areas of research in which we believe a detailed understanding of the chemistry underlying the biological phenomena is beginning to emerge.

Production of reactive oxygen species in vivo

Most of the hydrogen peroxide and other ROS generated during the normal metabolism of a typical eukaryotic cell is derived from superoxide that is formed from reduction of dioxygen by components of the mitochondrial electron transport chain, primarily ubisemiquinone (QH$^*$) in complex III and secondarily NADH dehydrogenase (complex I), in what are believed to be side reactions of electron transport (Figure 1) [20,21]. In addition, however, there also exist specialized systems whose primary purpose is to generate superoxide and ROS for use in defense systems that protect against pathogens. An example is the NADPH oxidase system found in leukocytes, which catalyzes the one-electron reduction of dioxygen by NADPH to form superoxide [22]. An oxidative burst mechanism utilizing a similar NADPH oxidase system is also observed in plants [23••].

Reactive oxygen species in regulation and signaling

Cells respond to changing levels of oxidative stress by inducing or suppressing the expression of various genes ranging from those encoding antioxidant systems to those encoding components of the respiratory chain. Several...
regulatory proteins of this type found in *E. coli* (for example, OxyR, SoxR and FNR [16,17]) have been particularly well characterized with respect to the chemical reactions involved. Formation of a disulfide is believed to occur in OxyR [18] while protein-bound iron–sulfur clusters undergo redox reactions in SoxR [24] and FNR [25].

ROS have also been shown to function as secondary messengers in signaling pathways in higher organisms. Thus a cell may respond to a stimulus that is not due to changes in oxidative stress by generating ROS that diffuse to a target, react, and thereby transduce the signal [26,27,28,29].

Numerous recent studies have implicated production of ROS in the signaling pathways of apoptosis, and it is intriguing to speculate that the link may occur at the mitochondria, which are the source of most of the ROS produced in eukaryotic cells [10,29]. Mitochondria have recently been found to play a central role in apoptosis [30], by releasing cytochrome *c* into the cytosol where it causes activation of the protease caspase-3 as a part of the apoptotic pathway. In addition, ROS may play a role in modulating the mitochondrial permeability transition pore [31], which may be involved in delivery of the apoptotic signal.

**Antioxidant enzymes**

Superoxide dismutases (SODs) are antioxidant enzymes that catalyze the disproportionation of superoxide to give dioxygen and hydrogen peroxide (Equation 1) [4,21].

\[
\text{SOD} \quad 2O_2^- + H^+ \rightarrow O_2 + H_2O_2
\]

Peroxidases catalyze the two-electron reduction of hydroperoxide or organic peroxides to water and alcohol in addition to oxidized cofactor. For instance, glutathione peroxidase (GPx), which uses glutathione (GSH) as the reducing agent (Equation 2), is believed to be a major cytosolic antioxidant in most eukaryotic cells.

\[
\text{GPx} \quad ROOH + 2GSH \rightarrow ROH + H_2O + GSSG
\]

Some of the most startling recent findings concerning antioxidant enzymes are the recent demonstrations that mice lacking the genes for either CuZnSOD (the cytoplasmic SOD containing Cu and Zn in its active site) [32] or cellular GPx [33] develop relatively normally; however, fibroblasts from the CuZnSOD-deficient mice are markedly more sensitive to redox cycling drugs than the wild type cells [34]. By contrast, yeast lacking the gene for CuZnSOD grow poorly in air and die rapidly in stationary phase [35], and both yeast and mice lacking the genes for the mitochondrial MnSOD have drastically reduced life spans [35–37]. Recent studies of transgenic plants with altered levels of small molecule antioxidants and antioxidant enzymes have also provided valuable information concerning antioxidant systems in those organisms [38,39–41].

The variety of known SODs has expanded in recent years as has information concerning their properties and function(s). Two particularly novel types of SODs were recently identified: nickel-containing SODs isolated from several strains of *Streptomyces* [42,43,44] and a monomeric CuZnSOD found in the periplasm of *E. coli* [45]. As for function, CuZnSOD was found to protect mammalian calcineurin from inactivation *in vitro* [46] and yeast calcineurin from inactivation *in vivo*, and it was discovered that a chaperone for copper was required for proper insertion of copper into CuZnSOD in yeast [47]. In the absence of this protein CuZnSOD is present in an inactive, copper-free form. Two bacterial genes unrelated to those known to encode SODs have been shown to rescue SOD-deficient *E. coli* strains [48,49].

The development of synthetic and engineered SODs has continued to advance. Two types of manganese-containing synthetic SODs, one with porphyrin ligands [50] and the other with a variety of macrocyclic ligands [51], were shown to have activity *in vivo*. In an elegant example of protein redesign, Pinto et al. [52] re-engineered the protein thioredoxin into an active iron-containing synthetic SOD enzyme.

A novel antioxidant function has been proposed for surface-exposed methionine residues that may be oxidized to methionine sulfioxide, with little effect on the properties of the protein, and then be re-reduced to the thioether by methionine sulfioxide reductase [53,54].

**Oxidative damage to biological molecules**

**Proteins**

Chemical studies of radical-mediated protein oxidation have demonstrated oxidative modification of protein sidechains, backbone cleavage, and protein–protein dimerization [6,7,8]. Sulfur-containing sidechains are particularly vulnerable to oxidation at sulfur, but most of the other oxidative pathways lead to carbonyl-containing products such as aldehydes and ketones, which are commonly measured using the 2,4-dinitrophenylhydrazine assay for protein carboxyls [7] (see lipids and carbohydrates section below for more discussion of the use of this reagent).

Recent calculations suggest that certain amino acids are more susceptible than others to irreparable damage, and that susceptibility depends upon the conformation of the domain in which they are found. For example, serine is postulated to be repairable when contained in an α helix but not when in a β sheet [55]. A recent report by Stadtman and co-workers [56] describes an observed increase in the surface hydrophobicity of proteins
that occurs with aging, an effect that is attributed to radical-mediated oxidative reactions.

Perhaps the most dramatic discovery in recent years concerning mechanisms of oxidative damage in biological systems is the facile reaction of superoxide with solvent-exposed iron–sulfur clusters in enzymes such as aconitase and other hydro-lyase enzymes containing 4Fe–4S clusters (Figure 3) [14**,15]. The reaction of superoxide with these centers has been demonstrated to inactivate such enzymes both in vitro and in vivo and to increase levels of intracellular free iron, which can catalyze oxidative damage to DNA [57**] (see below). This mechanism is the first clear cut example of a direct reaction of superoxide, rather than of a reactive oxygen species derived from it, leading to damage of a cellular component in vivo.

Possibly related is the recent observation of activation of iron regulatory protein-1 (IRP-1, which is identical to cytosolic aconitase) by hydrogen peroxide [58].

### Lipids and carbohydrates

Lipid peroxidation not only threatens the integrity and function of membranes and membranous proteins but also produces a variety of toxic aldehydes and ketones, one of the worst of which, trans-4-hydroxy-2-nonenal (HNE), is produced in high yield. HNE and malondialdehyde (MDA), another common toxic product formed upon peroxidation of lipids, are known to react via a Michael addition with nucleophilic sidechains of proteins and can result in protein cross-linking (Figure 4). Oxidative damage to carbohydrates can also produce products that are reactive with proteins and can result in damage [6**].

Identification of the primary sites of oxidative damage in living organisms is a major challenge since antioxidant protections differ considerably depending on the nature of the ROS and the site of attack. It is therefore important to note that products of lipid and/or carbohydrate oxidation can often react with proteins and that the resulting adducts contain carbonyl groups and therefore are reactive with 2,4-dinitrophenylhydrazine in the assay for protein carbonyls [6**7**]. Thus detection of high levels of protein carbonyls does not necessarily indicate that the proteins themselves are being directly oxidized by ROS; the carbonyls may instead result from reactions of undamaged proteins with toxic products of lipid or carbohydrate oxidation. Methodology to identify the exact nature of protein carbonyls in biological samples is being developed to address this issue [59**].

An interesting and different type of biological reaction of HNE is its reaction as a mechanism-based inhibitor of cytochrome P450 [60].

### Nucleic acids

Elevated levels of oxidative stress have long been known to result in oxidation of DNA, and recent results suggest that free intracellular iron is involved in this oxidation [57**61]. A widely accepted theory is that ‘free’ iron may bind loosely to various sites in the DNA, where it can act as a catalyst for the generation, from hydrogen peroxide, of a very reactive species that reacts with DNA in the immediate vicinity. Among the species suggested to attack DNA are hydroxyl radical, an iron ferryl radical and an iron-bound hydroxyl radical [10**]. The reactions leading to the reactive species are referred to as ‘Fenton chemistry’ and are well characterized reactions in vitro,
with the amino groups of other proteins to produce cross-linked reductants such as superoxide, ascorbate and NADH may to cause iron deposition in the nucleus [57]. Increases in cellular iron have recently been reported in the cell. Clusters leads to an increased concentration of ‘free’ iron its ability to cause iron release from vulnerable 4Fe–4S reducing agent in the Fenton reaction. It is proposed that carbonyls. be reactive with 2,4-dinitrophenylhydrazine in assays for protein Michael acceptor for various nucleophilic protein sidechains. The is detrimental to proteins as a result of its ability to function as a toxic aldehydes and ketones such as HNE. (b) This compound may have an altered hydrophobicity of nitrosating species (Equation 3) [70].

\[
\begin{align*}
R & - S - H + NO + oxidant \rightarrow R - S - NO + ? \\
& (3)
\end{align*}
\]

Nitrosylation of thiols has been reported to result in modification of protein function [68] and to occur in signal transduction pathways [69]. The mechanism of of nitrosating species (Equation 3) [70].

\[
\begin{align*}
R & - S - H + NO + oxidant \rightarrow R - S - NO + ? \\
& (3)
\end{align*}
\]

Nitric oxide has also been reported to react with proteins containing iron–sulfur clusters [71–73].

Reactive oxygen species in disease

There is convincing evidence for the involvement of oxidative damage in several classes of diseases [8**], although the identity of the ROS formed, the nature of the chemical reactions involved in their formation and subsequent reactions, and the primary targets of oxidative damage are often not yet identified. In the case of diseases that lead to neurodegeneration, redox metal ion involvement has frequently been implicated [74*,75*,76,77*,78,79,80*,81]. Atherosclerosis [9**] and diabetes [82] are other diseases in which oxidative damage has been implicated.

Drug action involving reactive oxygen species

The pharmacological action of many drugs involves ROS. Interesting examples are isoniazid, a drug used to treat the pathogen Mycobacterium tuberculosis [83**], and the antitumor antibiotic agent leinamycin [84]. Photodynamic therapy is another example of drug action involving a ROS, in this case, singlet dioxygen, ¹O₂ [85].
Conclusions
The past several years have seen significant advances in our understanding of biochemical pathways responsible for the oxidative stress and biological damage associated with various disease states and with living in air. At the same time, it has become increasingly clear that ROS, in addition to their role as toxic agents, are also used in certain signal transduction pathways, drug metabolism, and in different types of biological defense mechanisms. It has also become clear that the biochemistry of NO and O2 is often strongly linked.

Most of the ROS formed in normal eukaryotic cells are derived from superoxide formed at the mitochondria. The central role the mitochondria play in cell life and cell death is undoubtedly intimately linked to dioxygen biochemistry, but remains incompletely understood to date. The details (sources, targets, exact chemical identities, and concentrations of reactive oxygen species) are subjects of active inquiry. Elucidation of the chemical mechanisms and pathways associated with oxidative damage is particularly important given the pivotal role dioxygen and ROS play in cellular biochemistry.

References and recommended reading
Papers of particular interest, published within the annual period of review, have been highlighted as:
• of special interest
•• of outstanding interest

   This book contains an excellent collection of review articles by several of the leaders in the field of biological oxidative stress research. It provides broad coverage of relevant topics and a welcome emphasis on plant systems.

   Ascorbate is a biological antioxidant of major importance. Nevertheless, it can act as a reductant of metal ions (such as iron and copper) that promote Fenton reactions of hydrogen peroxide to produce dangerous oxidants, it has frequently been claimed that ascorbate can, under certain conditions, act as a pro-oxidant in vivo. This paper reports evidence to the contrary indicating that ascorbate is an antioxidant and not a pro-oxidant in human plasma in vivo, even under conditions of iron overload.

   Stoichiometric antioxidants such as ascorbate and glutathione (GSH) are re-reduced and thus recycled many times in vivo. The reduction of oxidized GSH (GSSG) is catalyzed by GSH reductase. Less is known about mechanisms of reduction of dehydroascorbate to ascorbate in vivo, but it is widely believed that the biological reductant is GSH. This paper reports an alternative mechanism in which the selenoenzyme thioredoxin reductase catalyzes reduction of dehydroascorbate by reduced nicotinamide adenine dinucleotide phosphate (NADPH).

   This is a timely minireview of superoxide and superoxide dismutases.


Yeast lacking coenzyme Q were hypersensitive to polynsaturated fatty acids and lipid hydroperoxides were elevated. The hypersensitivity was rescued by addition of antioxidants. The paper provides dramatic evidence for the antioxidant action of reduced ubiquinone in vivo and of the toxicity of the autoxidation products of unsaturated fatty acids.

   A comprehensive review of biochemical mechanisms of oxidative damage caused by reactive oxygen species, with emphasis on proteins as targets for such damage.

   A minireview on the same subject as [6*].

   A review of radical-mediated protein oxidation with emphasis on the physiology and pathology of the resulting damage to cellular proteins.

   A timely minireview of LDL oxidation.

    A timely minireview of mechanisms of DNA oxidation catalyzed by iron.

    A timely minireview of biochemical DNA oxidation.


   This paper reports the presence of a single histidine residue that is at or near the surface of the protein ceruloplasmin and that coordinates copper ions. The presence of the surface bound copper confers pro-oxidant properties on that protein. Mutagenesis of the surface histidine residue to an arginine results in loss of the property. This study provides a precedent for surface binding of a redox-active metal by a protein that can enhance its ability to act as a catalyst of oxidative damage because of its accessibility to other components of the cell.

   A comprehensive review of the properties of such proteins, including a detailed discussion of the sensitivity of some of them to oxidation by superoxide or other oxidants.


   SoxR contains 2Fe-2S clusters and acts as a redox-sensitive transcriptional activator.


   This interesting review points out that the best method to guard against reactive oxygen species formed in vivo is not antioxidants but reduction in the amount of superoxide formed at the mitochondria that subsequently reacts to give the reactive oxygen species. Topics discussed include mechanisms of superoxide formation in the cell, respiration as a tool to lower dioxygen concentrations, uncoupling as a mechanism to prevent superoxide formation, mitochondrial pores, and the link between the state of the mitochondrion and apoptosis.


This paper shows that the 2Fe-2S clusters of SoxS are reversibly oxidized in vivo in response to superoxide generation. This oxidation correlates with the transcriptional activity of soxS, the target of SoxS. Thus SoxS is the first transcription factor containing iron-sulfur clusters whose activity has been demonstrated to be a direct function of the redox state of the cluster.


FNR is a 4Fe-4S-cluster-containing transcription factor that regulates a network of genes that facilitate adaptation to low dioxygen concentrations in E. coli. In this study, the transformation of the [4Fe-4S]$^+$ cluster into a [2Fe-2S]$^+$ cluster upon exposure to air in vitro was observed using electronic, EPR (electron paramagnetic resonance) and Mossbauer spectroscopies. Little 3Fe-4S cluster was observed. Thus the 4Fe-4S cluster in FNR appears to be more susceptible to direct reaction with dioxygen than the 4Fe-3S cluster in aconitase, which is oxidized rapidly by superoxide but not by dioxygen. These results suggest that the degree of reactivity of 4Fe-4S clusters toward oxidants can be modulated by their individual protein environments.


Two unique isozymes of SOD from S. griseus are described. SOD I was composed of four identical subunits of molecular weight 130,000 kDa. It was found to contain 0.9 nickel ions per subunit. Its absorption and electron paramagnetic resonance (EPR) spectra were characteristic of Ni(III). SOD II was composed of four identical subunits of molecular weight 22,000 kDa. It was found to contain both iron and zinc. Nickel SOD was found to be widely distributed in Streptomyces. For many years it was thought that all SODs contained either Fe, Mn, or Cu and Zn. One wonders what other novel SODs will be found in the future.


In this paper the authors showed that a specific protein carrier of copper, Lys7, was required for the proper metallation of CuZnSOD in Saccharomyces cerevisiae. A lys7 knockout strain showed an oxygen sensitive phenotype characteristic of CuZnSOD deficiency. This strain contained copper-deficient CuZnSOD protein but showed normal processing of other copper proteins. The oxygen sensitivity of the lys7 knockout strain is particularly striking since both the CuZnSOD protein and the copper ions are present in sufficient amounts but the copper is not inserted into the protein.


A whole cell EPR method was developed to determine relative amounts of ‘free’ intracellular iron in Escherichia coli. The results of this study indicate that superoxide is able to leach iron from proteins containing 4Fe−4S clusters and that superoxide stress increases the amount of ‘free’ iron in cells. The increase in iron was shown to be due to leakage from 4Fe−4S clusters rather than from iron storage proteins. This paper presents data strongly supporting a mechan-ism for superoxide toxicity in which superoxide damage of proteins similar to basement membranes 4Fe−4S clusters causes release of ‘free’ iron which then catalyzes oxidative damage of cellular components by hydrogen peroxide in the presence of cellular reducing agents. The relatively large amount of ‘free’ iron that can be detected by this method in E. coli is striking. It will be important to determine what form this iron takes in the cell (that is, oxidation state, state of aggregation, identity of any ligands) because Fe^{3+} itself is not soluble under these conditions.


4-hydroxy-2-nonenal (4-HNE) and malondialdehyde (MDA) are products of lipid peroxidation which are known to modify proteins through covalent alkylation of lysine, histidine, and cysteine amino acid residues. Antibodies were raised that recognize the different types of protein adducts that are formed upon reaction with these unsaturated aldehydes. Isolated hepatocytes were exposed to either carbon tetrachloride (CCL4) or iron/ascorbate. Levels of MDA and protein carbonyls increased dramatically in both cases, whereas 4-HNE levels were little changed. Immunoprecipitation experiments indicated that increases in MDA corresponded with increases in intensity and number of MDA-adducted proteins. Neither CCL4 nor iron/ascorbate elicited changes in 4-HNE or induced 4-HNE-modified proteins. Experiments of the type described in this paper will be very useful in establishing the details of oxidative damage pathways occurring in vivo.


DNA damage was induced by NAD(P)H in the presence of Cu(II). The damage was inhibited by catalase as well as by chelation of Cu(II) but not by hydroxyl radical scavengers. The results suggest that DNA damage occurs in the presence of Cu(II) by a mechanism that is not dependent on the hydroxyl radical formation. NAD(P)H can act as an endogenous reductant of DNA-bound Cu(II) to Cu(II) which then reacts with hydrogen peroxide to give oxidation of DNA. The oxidation is proposed to occur via a pathway involving copper(II) complex or by generation of hydroxyl radicals in a site-specific manner.


A comprehensive review of reactions of NO and species derived from it with metalloproteins.


Peroxynitrite, formed from the reaction of nitric oxide (NO) with superoxide, has been implicated as a major agent of NO-induced cytotoxicity. This comprehensive review describes the experiments in support of this hypothesis.


This commentary critically evaluates the hypothesis that peroxynitrite plays a major role in cytotoxicity in vivo and concludes that the evidence currently available is not sufficient to reach that conclusion.


This paper concludes that the evidence is not conclusive that nitrotyrosine is specific as a biomarker of peroxynitrite formation in vivo. The presence of NO−/NOH− in biological systems suggests that the nitrating agent be referred to as a reactive nitrogen species.


It has only recently been realized that peroxynitrite reacts rapidly with CO2 to form an adduct, ONOO CO2-. This reaction occurs at physiologically relevant
concentrations and may represent the major fate of peroxynitrite when it is formed in vivo. This paper reviews this topic and concludes that considerably more knowledge about the lifetime and reactivity of this adduct is required before its biological significance can be assessed.


Nitric oxide (NO) stimulates Ras by increasing its guanine nucleotide exchange activity by nitrosation. In this report the authors determine that this S-nitrosylation specifically occurs at Cys118 of the protein. Mutation of this residue to a serine prevents NO induction of guanine nucleotide exchange and two mitogen-activated protein (MAP) kinases downstream of Ras.

70. Gow A, Buerk DG, Ischiropoulos H: A novel reaction mechanism for the formation of S-nitrosothiol in vivo. J Biol Chem 1997, 272:2841-2845. A mechanism for formation of S-nitrosothiols (R-SH) from NO and thiols in vivo is proposed in which NO reacts with R-SH to form RS-NOH, which reacts further with O2 to give RS=NO plus superoxide. Chemical evidence in support of the feasibility of this mechanism is presented.


This paper reports mapping of the distribution of iron in the brain using a method for detection of Cu(I) formation (76). The authors show that isoniazid-resistant KatG mutants compensate in isoniazid resistance, but results in increased peroxide sensitivity. In this report, the authors show that the activities of aconitase and complexes I, II, and III of the electron transport chain, which are Fe-S cluster containing enzymes, are reduced in heart tissue from two Freidreich ataxia patients relative to the average control activities.


Friedreich ataxia is a degenerative disease linked to mutations in the frataxin gene. This gene is believed to be involved in mitochondrial iron transport, based on studies of a homologous protein in yeast. In this report, the authors show that the activities of aconitase and complexes I, II, and III of the electron transport chain, which are Fe-S cluster containing enzymes, are reduced iron in tissue from two Freidreich ataxia patients relative to the average control activities.


A comprehensive review of proposed mechanisms of neurotoxicity associated with amyloid deposition in Alzheimer’s disease.

77. Sayre LM, Zagorski MG, Surewicz WK, Kraft GA, Perry G: •••


Friedreich ataxia is a degenerative disease linked to mutations in the frataxin gene. This gene is believed to be involved in mitochondrial iron transport, based on studies of a homologous protein in yeast. In this report, the authors show that the activities of aconitase and complexes I, II, and III of the electron transport chain, which are Fe-S cluster containing enzymes, are reduced iron in tissue from two Freidreich ataxia patients relative to the average control activities.


The catalase-peroxidase KatG in Mycobacterium tuberculosis activates the drug isoniazid to a form toxic to this bacterium. Loss of KatG function results in isoniazid resistance, but results in increased peroxide sensitivity. In this report, the authors show that isoniazid-resistant KatG mutants compensate for the loss of KatG function by mutations resulting in the overexpression of Ahp C, a putative alkyl hydroperoxidase.

