Review article:

CLASSIFICATION OF OXIDATIVE STRESS BASED ON ITS INTENSITY

Volodymyr I. Lushchak*

Department of Biochemistry and Biotechnology, Vassyl Stefanyk Precarpathian National University, 57 Shevchenko Str., Ivano-Frankivsk, 76025, Ukraine

* Corresponding author: Department of Biochemistry and Biotechnology, Vassyl Stefanyk Precarpathian National University, 57 Shevchenko Str., Ivano-Frankivsk 76025, Ukraine. Tel/fax.: +38 0342 59 61 71. E-mail: lushchak@pu.if.ua

ABSTRACT

In living organisms production of reactive oxygen species (ROS) is counterbalanced by their elimination and/or prevention of formation which in concert can typically maintain a steady-state (stationary) ROS level. However, this balance may be disturbed and lead to elevated ROS levels called oxidative stress. To our best knowledge, there is no broadly acceptable system of classification of oxidative stress based on its intensity due to which proposed here system may be helpful for interpretation of experimental data. Oxidative stress field is the hot topic in biology and, to date, many details related to ROS-induced damage to cellular components, ROS-based signaling, cellular responses and adaptation have been disclosed. However, it is common situation when researchers experience substantial difficulties in the correct interpretation of oxidative stress development especially when there is a need to characterize its intensity. Careful selection of specific biomarkers (ROS-modified targets) and some system may be helpful here. A classification of oxidative stress based on its intensity is proposed here. According to this classification there are four zones of function in the relationship between “Dose/concentration of inducer” and the measured “Endpoint”: I – basal oxidative stress (BOS); II – low intensity oxidative stress (LOS); III – intermediate intensity oxidative stress (IOS); IV – high intensity oxidative stress (HOS). The proposed classification will be helpful to describe experimental data where oxidative stress is induced and systematize it based on its intensity, but further studies will be in need to clearly discriminate between stress of different intensity.

Keywords: free radicals, reactive oxygen species, reactive nitrogen species, system response

Abbreviations: BOS – basal oxidative stress; HOS – high intensity oxidative stress; 8-OHG – 8-hydroxyguanine; G6PDH – glucose-6-phosphate dehydrogenase; GPx – glutathione-dependent peroxidase; GR – glutathione reductase; GSH – glutathione reduced; GSSG – glutathione oxidized; IDH – NADP+-isocitrate dehydrogenase; IOS – intermediate intensity oxidative stress; LOOH – lipid peroxides; LOS – low intensity oxidative stress; MDA – malonic dialdehyde; NADP-ME – NADP-malic enzyme; NOE – no observable effect point; 6PGDH – 6-phosphogluconate dehydrogenase; 8-oxodG – 8-oxo-7,8-dihydro-2′-deoxyguanosine; 8-oxoGua – 8-oxo-7,8-dihydroguanine; PPP – pentose phosphate pathway; RNS – reactive nitrogen species; ROS – reactive oxygen species; RS – reactive species; ROSISP – ROS-induced ROS-sensitive parameter; O₂⁻ – superoxide anion radical; SOD – superoxide dismutase; TBA – thiobarbituric acid; TBARS – thiobarbituric acid reactive substances; TRR – thioredoxin glutathione reductase; ZEP – zero equivalent point.
INTRODUCTION

Free radicals were discovered by Moses Gomberg (born in 1866, Yelizavetgrad, Russian Empire, now Kirovohrad, Ukraine) more than a century ago (Gomberg, 1900). For a long time it was believed that they did not exist in biological systems due to their short life time resulting from high chemical activity. In the late 1930s, however, German researcher Leonor Michaelis proposed that all oxidation reactions involving organic molecules would be mediated by free radicals (Michaelis, 1939). This actually incorrect prediction stimulated interest in the role of free radicals in oxidative biological processes. In the early 1950s, free radicals were detected in biological systems (Commoner et al., 1954) and virtually immediately were applied to diverse phenomena including human pathologies (Gerschman et al., 1954), and aging (Harman, 1956). Discovery of the presence of free radicals in biological systems was the first critically important finding in the field of free radical research in living organisms. Since that time, our knowledge of the involvement of free radicals in living processes has increased enormously. In the 1970s, Sies and Chance used noninvasive spectrophotometric methods and not only evaluated the operation of catalase in vivo, but also provided information on steady-state hydrogen peroxide levels in perfused rat liver (Sies and Chance, 1970). This work was virtually the first attempt to characterize ROS homeostasis in animal tissues. In the 1980s, it became clear that the generation and elimination of free radicals in living organisms are normally well-balanced and imbalances between these two processes underly many pathologies.

At the beginning of free radical research in living organisms, serious debates took place because it was supposed that if free radicals really did exist in biological systems, the latter should possess systems controlling the levels of reactive species (RS), particularly reactive oxygen species (ROS), i.e. some mechanisms for their elimination should exist. Therefore, the second principal discovery in free radical research in biological systems was extremely important. In 1969, McCord and Fridovich described a new function for an already well-known protein – erythrocuprein (hemocuprein); this enzyme catalyzed the dismutation of the superoxide anion radical and subsequently was renamed superoxide dismutase (McCord and Fridovich, 1969). The third critically important discovery in the field showed that free radicals were not always deleterious but actually had positive biological functions as well. Studies disclosed their involvement in combating infection as part of the cellular immune response, where ROS and reactive nitrogen species (RNS) operate in concert with reactive halogen species to fight invading microorganisms (Babior et al., 1973; 1975; Britigan et al., 1987; Ferrari et al., 2011; Rossi et al., 1985; Sies, 2014). Finally, identification of the signaling functions of ROS and RNS was the fourth principle discovery about free radical biology (Jacob et al., 2006; Lushchak, 2011a, b; Palmer et al., 1987; Scandalios, 2005; Sies, 2014; Stone and Yang, 2006; Veal et al., 2007; Winterbourn and Hampton, 2008). These four discoveries, along with the deciphered mechanisms of finely regulated RS production and their involvement in diverse homeostatic processes, were used to propose and develop Denham Harman’s Free Radical Theory of Aging (Harman, 1956, 2009). Today, it seems that of all theories of aging, Harman’s Free Radical Theory of Aging is the most consistent and, moreover, the most experimentally supported aging concept. However, it is also challenged by certain experimental data and therefore needs further investigation.

Generally, the main problems in the investigation of free radical processes in living organisms are related to: (i) the high reactivity and low stability of free radicals; (ii) their low concentrations; (iii) the absence of technical tools for reliable evaluation of absolute and sometimes even relative levels of free radicals in vivo; (iv) their low chemical specificity; (v) the huge diversity of reactions that
radicals can take part in; (vi) complicated spatiotemporal distribution in the cell; (vii) for multicellular organisms, the heterogeneity of cells in organs and tissues; (viii) changes in free radical processes depending on an organism’s physiological state.

Due to the reasons listed above and many other reasons, investigations of the processes involving RS and interpretation of experimental results are very complicated. For example, in many cases the same compounds at the same concentrations may increase or not affect the observable level of RS-modified molecules or increase/decrease activities of antioxidant enzymes, and yet all of these different states have been declared to represent the state of oxidative stress after introduction of this definition in 1980. In the present paper, using data from my lab as well as the literature, I propose explanations for the frequent contradictions in results found when analyzing RS-induced stresses. This paper will focus only on primary oxidative stress induced by ROS because it seems to be the simplest situation for description and analysis and the best-studied stress induced by ROS, the most commonly studied types of radicals. The state of secondary oxidative stress induced indirectly such as by heat shock, energy exhaustion, starvation, overfeeding and others, will not be covered here in order to simplify presentation of the key ideas.

WHAT ARE FREE RADICALS AND REACTIVE OXYGEN SPECIES?

From the chemical point of view, a free radical is any atom or molecule or its part (particle) possessing unpaired excited electron(s) in external molecular or atomic orbitals. The negative electrical charge of electron(s) may be counterbalanced by the positive nuclear charge of positrons resulting in a neutral particle, or if not counterbalanced results in anion or cation radicals. However, in biology there is another popular understanding of free radicals, less accurate, but widely used and, since we work in this field we will also use this broadly accepted understanding of free radicals. So, according to common biological understanding, a free radical is an unstable particle (atom or molecule or its part) possessing unpaired electron(s) in external atomic or molecular orbitals (Halliwell and Gutteridge, 1989).

From the biological point of view, the dioxygen molecule (O₂) is a biradical, because it contains two electrons with the same spin in an external antibonding molecular orbital. Due to Hund’s restriction rules, these should be located in different orbitals and, therefore, are not paired. They can be identified by the technique of electron paramagnetic resonance because they interact with an electromagnetic field (Malanga and Puntarulo, 2011). Molecular oxygen can be reduced via a four-electron mechanism with acceptance of four protons yielding two water molecules (Figure 1).

![Figure 1: Reduction of molecular oxygen via four- and one-electron schemes](image)

In this case, the free biradical is simply converted to a nonradical species due to acceptance of the four electrons and four protons. However, there is another way to reduce molecular oxygen – this is one-electron successive reduction (Figure 1). Receiving one electron, O₂ is converted to the superoxide anion radical (O₂⁻), containing one unpaired electron in an external antibonding orbital. Accepting a second electron and two protons, converts the superoxide anion radical into hydrogen peroxide (H₂O₂); H₂O₂ has a non-radical nature and is chemically more active than molecular oxygen, but less active than O₂⁻. Formation of the most reactive of oxygen species, the hydroxyl radical (HO’), results from the further reduction of H₂O₂
leading to its dismutation. Finally, acceptance of a fourth (final) electron and one more proton HO\(^+\) forms a water molecule. Usually, the chance to directly and separately bind an electron and proton is negligible, and this reaction generally occurs via the abstraction of a hydrogen atom from any substrate. Since \(O_2^{\cdot-}\), \(H_2O_2\), and HO\(^+\) are chemically more reactive than molecular oxygen, they are collectively called ROS but only \(O_2^{\cdot-}\) and HO\(^+\) are actually free radicals, whereas \(H_2O_2\) is not. Therefore, in biological research, the term “free radicals” is frequently replaced by “reactive oxygen species” (ROS), which is a more general term and includes both free radical and non-radical species. Singlet oxygen and various peroxides as well as many other oxygen-containing compounds are also included as ROS. It must be added that generally ROS are more chemically active due to cancelling of restriction of the ground state (triplet) oxygen. Finally, it should be noted that in many cases the terms “oxygen free radicals” and “reactive oxygen species” are used interchangeably; in many cases this is not correct and authors should pay attention to the correct use of these terms.

**GENERATION AND ELIMINATION OF REACTIVE OXYGEN SPECIES**

It is believed that in eukaryotic organisms more than 90% of ROS are produced by the mitochondrial electron-transport chain (Sies, 2014; Skulachev, 2012). Some amounts of ROS are also formed by electron transport chains in plasmatic (Lüthje et al., 2013), nuclear (Vartanian and Gurevich, 1989) and endoplasmic reticulum (Brignac-Huber et al., 2011) membranes. ROS generation takes place because some active electrons “escape” electron transport carriers and reduce molecular oxygen to yield \(O_2^{\cdot-}\). Superoxide is then spontaneously or enzymatically converted to \(H_2O_2\). The latter accepting one more electron is converted to HO\(^+\) and OH\(^-\) in reactions that are frequently catalyzed by transition metal ions (Fe\(^{2+}\) or Cu\(^{2+}\)). Finally, HO\(^+\) and OH\(^-\) receiving hydrogen atom or proton, respectively, are converted to water.

Many oxidase enzymes, such as oxidases of xanthine, carbohydrates, aldehydes, monoamines and amino acids also form ROS.

Figure 1 demonstrates the relationship between molecular oxygen, water and ROS. \(O_2^{\cdot-}\) can spontaneously interact with an electron donor and be converted to \(H_2O_2\). This reaction is substantially accelerated by superoxide dismutase (SOD; EC 1.15.1.1):

\[
O_2^{\cdot-} + O_2^{\cdot-} + 2H^+ \xrightarrow{SOD} O_2 + H_2O_2
\]

**Equation [1]**

Reduction of \(H_2O_2\) leads to the formation of HO\(^+\) and OH\(^-\). Hydroxyl radical is the most reactive of all the ROS mentioned above. There is no enzymatic system to defend living organisms against HO\(^+\), and, therefore, prevention of its formation is the most efficient way of protection against this highly reactive oxidant. There are several enzymatic systems dealing with \(H_2O_2\). Catalase (EC 1.11.1.6) dismutates \(H_2O_2\) to water and molecular oxygen:

\[
2H_2O_2 \xrightarrow{\text{catalase}} 2H_2O + O_2
\]

**Equation [2]**

There is also a large family of peroxidases that degrade other hydroperoxides as well as \(H_2O_2\). For example, glutathione-dependent peroxidases (GPx, EC 1.11.1.9) can reduce \(H_2O_2\) and lipid peroxides (LOOH) at the expense of reduced glutathione:

\[
H_2O_2 + 2GSH \xrightarrow{\text{GPx}} 2H_2O + GSSG
\]

**Equation [3]**

\[
LOOH + 2GSH \xrightarrow{\text{GPx}} LOH + GSSG + H_2O
\]

**Equation [4]**

The level of reduced glutathione is maintained/replenished by the reduction of glutathione disulfide by glutathione reductase (GR, EC 1.6.4.2):
In some organisms, such as *Drosophila*, thioredoxin glutathione reductase (Dm TrxR-1, or TRR, EC a1.8.1.B1) replaces GR for the replenishment of GSH [Eq. 5].

Finally, the oxidized coenzyme NADP⁺ is reconverted to NADPH by several enzymes. This mainly involves pentose phosphate pathway (PPP) enzymes, glucose-6-phosphate dehydrogenase (G6PDH, EC 1.1.1.49) and 6-phosphogluconate dehydrogenase (6PGDH, EC 1.1.1.43):

\[
\text{NADP}^+ + \text{glucose-6-phosphate} \rightarrow \text{NADPH} + 6\text{-phosphogluconolactone}
\]

Equation [6]

\[
\text{NADP}^+ + 6\text{-phosphogluconate} \rightarrow \text{NADPH} + \text{ribuloso-5-phosphate} + \text{CO}_2
\]

Equation [7]

In some tissues, particularly the brain, malate dehydrogenase (oxaloacetate-decarboxylating) utilizing NADP⁺ called also as NADP-malic enzyme (NADP-ME, EC 1.1.1.40) catalyzing reaction [Eq. 8] may also be important producer of NADPH:

\[
\text{NADP-ME}\hspace{1cm} (S)-\text{malate} + \text{NADP}^+ \rightleftharpoons \text{pyruvate} + \text{CO}_2 + \text{NADPH}
\]

Equation [8]

NADP⁺-isocitrate dehydrogenase (IDH, threo-DS-isocitrate: NADP⁺ oxidoreductase (decarboxylating); EC 1.1.1.42) also provides substantial NADPH amounts in some cases:

\[
\text{Isocitrate} + \text{NADP}^+ \rightarrow \text{2-oxoglutarate} + \text{CO}_2 + \text{NADPH}
\]

Equation [9]

The above enzymatic systems are responsible for elimination of \(O_2^-\) and \(H_2O_2\), and, therefore, prevention of \(HO^-\) formation. Usually, these enzymes are grouped in two sets – the first set contains primary antioxidant enzymes that directly deal with ROS (SOD, catalase, and other peroxidases), whereas the second set includes associated or auxiliary antioxidant enzymes, assisting the first group. For example, these provide the reducing equivalents needed for ROS elimination (e.g. GR, TRR, G6PDH, 6PGDH, NADP-ME, IDH, etc.). The antioxidant enzymes and other proteins involved in antioxidant defense collectively form a group called high molecular weight (mass) antioxidants. Other antioxidants belong to a group of low molecular weight (mass) antioxidants. This includes compounds of molecular mass usually less than 1000 carbon units, (overall molecular mass < 1000) such as vitamins C and E, carotenoids, anthocyanins, glutathione (GSH), uric acid and many other natural or synthetic compounds. It should be noted that low molecular mass antioxidants may protect organisms against \(HO^-\). In concert, low and high molecular mass antioxidants form a unique and very efficient system to maintain ROS levels within in a certain range (Sies, 1993).

Under homeostatic conditions in organisms, the operation of two systems, the generation and elimination of ROS, is well balanced due to which the steady state ROS, at least \(H_2O_2\), level is kept below 10 nM (Sies, 2014). However, even if the elimination systems work ideally, some ROS escape them resulting in basic level of modification of cellular components. Due to that reality, we always find some amount of ROS-modified biomolecules in unstressed organisms. This is the so-called basic level of ROS-induced modification of cellular components.

**COMMONLY USED MARKERS OF ROS-INDUCED MODIFICATION OF CELLULAR COMPONENTS**

It seems that despite their high chemical reactivity most generated ROS do not lead to serious negative physiological consequences for organisms. That is mainly due to the action of highly efficient systems of ROS neutralization operating in concert with reparation and elimination of ROS-modified mole-
cules. Thus, a certain level of ROS-modified molecules always exists, that may be called the basal steady-state (stationary) level (Lushchak, 2010, 2011a, b, 2012; Sies, 2014). Reactive oxygen species can modify most types of biomolecules including proteins, lipids, carbohydrates, nucleic acids, metabolic intermediates, etc. It is widely accepted that the use of only one type of modification to assess oxidative damage during oxidative stress is not sufficient. That is due to the different sensitivity, dynamics, and nature of ROS-promoted modifications. Instead, in order to evaluate the intensity of ROS-involving processes, several approaches for the evaluation of particular oxidatively modified molecules have been selected. They reflect the level of products of interaction between ROS and cellular components of different natures. “Classically”, several essential markers are used. They are: (i) for lipids – the formation of malonic dialdehyde, isopsoralens, and lipid peroxides; (ii) for proteins – protein carbonyl groups; and (iii) for DNA – 8-oxoguanine. Malonic dialdehyde is commonly measured via its reaction with thiobarbituric acid (TBA). However, this reaction is not specific and many other compounds react with TBA under the assay conditions. The array of products formed is collectively called thiobarbituric acid reactive substances (TBARS) to reflect this low specificity. Certain amino acids, carbohydrates, aldehydes and other compounds interfere with the reaction measurement and, therefore, this method should be used with precaution and discussed taking into account the highlighted issues (Lushchak et al., 2011). In the last decade, an HPLC technique was applied to evaluate MDA levels and this method, along with immunochemical identification (Claeson et al., 2001) can now be recommended as more reliable than the TBARS assay. There are also many other approaches to evaluate the intensity of ROS-induced lipid peroxidation and the measurement of lipid peroxides (Claeson et al., 2001), 4-hydroxynonenal (Zimniak, 2011) or exhaled carbohydrogens (Mayne, 2003) are just some of them. Selection of methods depends on many things, particularly tools available (Abele et al., 2011; Halliwell and Gutteridge, 1989).

Probably the most popular method for detection of ROS-modified proteins is the one based on the formation of additional carbonyl groups with their visualization due to their interaction with 2,4-dinitrophenylhydrazine (Lenz et al., 1989; Lushchak, 2007; Lushchak et al., 2011). The hydrazones formed are measured spectrophotometrically. Specific antibodies that interact with carbonyl groups on proteins (Lenz et al., 1989; Wehr and Levine, 2012) have also been developed. In some cases, there is also the possibility to evaluate the amount of di-tyrosines and other products of free radical-induced oxidation of proteins (Babusíková et al., 2008; Catalgo et al., 2011).

Oxidation of nucleic acids also forms an array of products, but in this case there are some favorites that are relatively easy to quantify. These are mainly oxidatively modified guanine derivatives, of which 8-hydroxyguanine (8-OHG) is the most commonly used marker (Lovell and Markesbery, 2008; Lovell et al., 2011), but 8-oxo-7,8-dihydro-2′-deoxyguanosine (8-oxodG) and 8-oxo-7,8-dihydroguanine (8-oxoGua) (Olinski et al., 2007) can also be measured.

Certainly, there are many more different markers of ROS-induced modification of cellular constituents, but those listed here are the most widely used and applied approaches.

OXIDATIVE STRESS:
DESCRIPTION AND DEFINITIONS

As described above, under normal conditions, living organisms maintain a basal steady-state (stationary) ROS level within a certain range. Homeostasis is provided due to the fact that systems of ROS generation are counterbalanced by prevention and elimination systems along with any other components interacting with ROS (Figure 2).
Figure 2: The dynamics of levels of reactive oxygen species in biological systems. The basic steady-state (stationary) level of reactive oxygen species fluctuates over a certain range under normal conditions. However, under stress ROS levels may increase beyond the normal range resulting in acute or chronic oxidative stress. Under some conditions, ROS levels may not return to their initial range and stabilize at a new quasi-stationary level.

However, under certain circumstances this balance may be shifted resulting in an enhanced ROS steady-state level even up to 100 nM (Sies, 2014). Certainly, this has consequences due to enhanced oxidative modification of diverse macromolecular components of an organism. The state, when ROS levels exceed the basal values leading to functional disturbances has been called oxidative stress. It was first defined in 1985 by Prof. Helmut Sies: Oxidative stress “came to denote a disturbance in the prooxidant-antioxidant balance in favor of the former” (Sies, 1985). The next year he published a definitive review summarizing the accumulated knowledge at the time about ROS effects on nucleic acids, proteins, lipids, and carbohydrates, as well as relationships between ROS and inflammation, carcinogenesis, ageing, radiation damage and photobiological effects (Sies, 1986). Later H. Sies modified the mentioned above definition to “An imbalance between oxidants and antioxidants in favor of the oxidants, potentially leading to damage, is termed “oxidative stress” in order to emphasize the damage to certain cellular components (Sies, 1997). Finally, the definition was modified to also underline ROS-based signaling “a disruption of redox signaling and control” (Sies and Jones, 2007). More recently one more definition was proposed: “Oxidative stress is a situation where the steady-state ROS concentration is transiently or chronically enhanced, disturbing cellular metabolism and its regulation and damaging cellular constituents” (Lushchak, 2011b). This reflects the steady-state level of products of ROS-promoted processes along with ROS effects on the functioning of living organisms. Figure 2 shows two different scenarios which result from perturbations of ROS-related processes. If the capacity of antioxidant system is not overwhelmed, the stress-enhanced ROS level can return to its initial state. In many cases, an induction of ROS-regulated genes may be needed to cope with the enhanced ROS levels. Generally, if ROS steady-state levels return to the initial value within minutes/hours after stress induction, if organisms are capable and have enough resources for the corresponding response, the stress is called “acute oxidative stress” (Lushchak, 2011b). However, sometimes ROS levels do not return to the initial range but stabilize at a somewhat higher level or just extend the steady-state ROS range existing under normal conditions and in this case the stress does not last for prolonged time period. This scenario is called “chronic oxidative stress” and frequently occurs under pathological conditions. Finally, after some perturbations, particularly as a consequence of substantial physiological or pathological shifts or chronic intoxication, the steady-state ROS level does not return to its initial range but stabilizes at a higher level, called a quasi-stationary or quasi-steady-state one. It seems that this scheme characterizes most known scenarios of perturbations in ROS homeostasis.

One of the main questions remaining is: how to define oxidative stress over the basal state of ROS levels and operation of organisms?
PROBLEMS IN INTERPRETATION OF EXPERIMENTAL DATA WITH INDUCED OXIDATIVE STRESS

Oxidative stress induced by external factors, particularly primary oxidative stress can be caused by the direct effects of ROS on living organisms. In the simplest case, unicellular organisms or cells in cultures are subjected to certain RS. Curve 1 of Figure 3 shows a schematic of a typical response by a cellular endpoint to different concentrations of oxidants producing hydrogen peroxide or direct $H_2O_2$ application. Endpoint parameters of interest such as cell survival or activity of antioxidant enzymes are frequently used for evaluation of ROS effects on living organisms. At very low concentrations, ROS do not affect these parameters (zone I). However, they can be altered by ROS addition in a concentration-dependent manner. An increase in concentration of the inducer, although at first glance may seem paradoxical, enhances cell survival and activity of antioxidant enzymes (zone II) (Bayliak et al., 2006; Semchyshyn and Lozinska, 2012). These effects are primarily related to the activation of many cellular processes, particularly directed to increase cell resistance to oxidative or general stresses. Up-regulation of antioxidant enzymes is a perfect example of this. So, at these levels, the oxidant assists to develop the adaptive response in order to improve biological functions. This sort of relationship between toxicant/oxidant and the measured end parameter (endpoint) has been called “hormesis” (Calabrese, 2008; Le Bourg, 2009; Rattan, 2008; Ristow and Schmeisser, 2011; Ristow and Zarse, 2010). The cellular response to ROS is measurable up to a maximum level at a certain ROS concentration followed by decrease in the endpoint parameter back to control (basal) level. Increases in oxidant concentration may reduce the measured parameter back to approximately zero or to some other horizontal asymptote. To underline the behavior of curve 1, zone II may be divided for zone IIa where the endpoint parameter increases and zone IIb where the parameter decreases to “no observable effect” (NOE) point. A further increase in inducer dose results in curve 1 passing through the NOE and decreased levels of the endpoint parameter in zones II and IV.

Figure 3: Relationships between the dose of an inducer of oxidative stress and commonly used endpoint (end parameter) parameters that may be measured. Zone I – no observable effects are registered due to very low intensity oxidative stress (basal intensity oxidative stress – BOS); zone II – low intensity (mild) oxidative stress (LOS) with a slightly enhanced level of oxidatively modified molecules and enhanced activity of antioxidant enzymes in response to oxidative stress; zone III – intermediate intensity oxidative stress (IOS); and zone IV – high intensity (strong) oxidative stress (HOS). Curve 1 – ROS-induced ROS-sensitive function (ROSISP), curve 2 – level of oxidatively modified components. Abbreviations: NOE – no observable effect point; ZEP – zero equivalent point – the level of components of interest corresponds to the initial (basic) level in the absence of inducers of oxidative stress.

It is critically important to note that the whole dose dependence of curve 1 in Figure 3 is connected with the interaction between ROS and certain cellular components. This interaction leads to oxidative modification of cellular components which is reflected by curve 2 in Figure 3. These characteristics of cellular response to different concentrations of oxidants are frequently found in experiments. Interestingly, of presence of these complicated relationships can frequently be misleading and can result in discrepancies in the interpretation of experimental data, especially if only a single dose of oxidant is evaluated (as compared with analysis of mul-
tiple points on a dose-response curve). The, complicated behavior of the system is explained by the many components involved and different sensitivity of cellular components to ROS-induced modification, their localization, and target accessibility to ROS, subject to repair, reduction and degradation pathways.

Analysis of many hundreds of reliable publications with the term “oxidative stress” in them lets us categorize given interpretations. Induction of oxidative stress is usually evidenced by: (i) enhanced levels of oxidized cellular constituents; (ii) increased levels or activities of antioxidant and associated enzymes; (iii) decreased levels or activities of antioxidant and associated enzymes, and, finally, by a combination of the above mentioned responses. In certain cases, the levels of ROS-modified molecules may also be decreased due to their elimination by specific systems (this case is a relatively rare event and complicates the description; therefore it will not be covered here). Why do such different responses, sometimes opposite, all lead to a conclusion of the induction of oxidative stress? The first case from the above list (enhanced levels of oxidized cellular constituents) usually does not raise serious questions if evaluated correctly and if several markers are measured simultaneously. The most common practice includes evaluation of oxidized lipids (e.g. lipid hydroperoxides) and oxidized proteins (e.g. protein carbonyls). Other parameters like glutathione disulfide levels or the ratio of oxidized to total or reduced glutathione are also measured as well as oxidized nucleic acids or various complexes formed from pairings of carbohydrates, proteins or nucleic acids. Much more complicated is the situation with respect to the levels or the activities of antioxidant enzymes. As mentioned above, under oxidative insults, enzymes may demonstrate some or all of the potential responses: decreased, increased, or not changed. Decreased activities are usually discussed from the point of view of enzyme inactivation by ROS. Indeed, many antioxidant and associated enzymes have been shown to be inactivated by ROS (Belluzzi et al., 2012; Hermes-Lima and Storey, 1993; Lee et al., 2012; Lushchak and Gospodaryov, 2005; Semchyshyn and Lozinska, 2012; Semchyshyn and Lushchak, 2004; Yang and Ming, 2012). Specific mechanisms may differ substantially, but a decrease in activity is a common event. Increases in the activities of antioxidant and associated enzymes under oxidative stress are usually connected with their de novo synthesis (Lushchak, 2010, 2011a) or activation of preexisting inactive molecules (Bayliak et al., 2006, 2007; Semchyshyn, 2009). Although activation of inactive enzyme molecules is still debatable issue, up-regulation of their biosynthesis is well-established. The process of up-regulation may involve enhanced gene transcription, protein translation and posttranslational modification or maturation (Lushchak, 2011a; Sies, 2014; Stone and Yang, 2006). Several regulatory systems responsible for up-regulation of antioxidant and associated enzymes have been described in different organisms. These systems are regulated by transcription factors, the best-known ones being SoxR and OxyR in bacteria (Demple, 1991; Lushchak, 2001), Yap1 in budding yeasts (Lushchak, 2010), Rap2.4a and Npr1 in plants (Lushchak, 2011a; Srinivasan et al., 2009), and NrR2 and NF-kB in animals (Lushchak, 2011a; Wang et al., 2012). The molecular mechanisms involved in redox signaling by the listed above and other transcription factors are based on reversible oxidation of cysteine residues of sensor proteins (Lushchak, 2011a; Sies, 2014; Stone and Yang, 2006). These have been shown to be responsible for realization of adaptive responses to the introduction of inducers of oxidative stress at low or intermediate concentrations.

In summary, we can say that oxidative stress clearly presents when: (i) a steady-state level of ROS-modified cellular components is enhanced; (ii) ROS-regulated transcription factors are activated and antioxidant and associated enzymes are up-regulated; and finally, (iii) real evidence of
ROS-induced inactivation of antioxidants or their consumption is demonstrated.

Now the question is: how can all the accumulated information available in the literature be categorized? In the following section I am going to propose a system which may provide interpretation for all experimental results. The key idea used to systematically categorize the effects of oxidative stress is based on its different intensity due to the application of different doses/concentrations of inducers in different studies.

CLASSIFICATION OF INTENSITY OF OXIDATIVE STRESS: MILD, MODERATE OR STRONG?

Investigation of different modes of oxidative stress induction in all groups of organisms (e.g. bacteria, fungi, plants and animals) has always been complicated (Lushchak, 2011a). For example, a “classic” inducer of oxidative stress, hydrogen peroxide (H₂O₂), affects the levels of oxidized lipids and proteins in bacteria (Semchyshyn et al., 2005) and yeasts (Bayliak et al., 2006, 2007; Semchyshyn, 2009) often increasing the levels in one case and decreasing in another (due to the operation of specific defense and detoxification systems), but mainly showing enhanced levels. Activities of antioxidant and associated enzymes were similarly increased, decreased or not changed in different cases. In most of these cases, we were talking about induction of oxidative stress with the need to explain the obvious differences. This experience and discussion with many colleagues made it clear that there was a desperate need to sort the accumulated wealth of experimental data and determine why responses were so variable between different studies.

Curve 1 in Figure 3 shows the relationship between the dose of an oxidant effector and the endpoint parameter measured. The latter parameter may be different in different studies, e.g. cell survival, activation of ROS-sensitive regulatory proteins, activity of antioxidant enzymes, etc. For analysis we will use those which are ROS-sensitive and at the same time are induced/enhanced by ROS exposure at low concentrations. Curve 2 in Figure 3 shows the relationship between concentration of the oxidant and the level of oxidatively modified components. Those may be different products of oxidation of proteins, lipids, nucleic acids, carbohydrates, etc. These provide an integral marker of ROS-induced modification of cellular constituents. Next, we will analyze the behavior of curves 1 and 2 at different concentrations of inducers of oxidative stress.

At very low concentrations (zone I) no observable effects are seen – oxidant effects are near negligible and significant responses cannot be discerned. In living systems, ROS are always present and the introduction of additional small amounts of oxidant (e.g. levels similar to basal amounts in vivo or even slightly higher) does not disturb the cellular processes to an extent that may be detected using conventional assay approaches. However, a further increase in the concentration of the inducer (zone II) enhances the observable level of oxidatively modified components and, at the same time, increases the endpoint parameter measured – i.e. the ROS-induced ROS-sensitive parameter (ROSISP). The mechanisms responsible for this induction were discussed briefly above. In this zone, elevation of the inducer concentration results in the development of either a full response (zone IIA) or a reduction in the ROSISP level despite a concomitant increase in the levels of ROS-modified components (zone IIB). In other words, in zone II, we can see that the expression of the ROSISP rises to a maximum but then decreases again to the point when no observable effect (NOE) is seen. The levels of oxidatively modified components at the NOE point are substantially increased, but after that point, the ROSISP further decreases (zone III). Finally, in zone IV both measured functions converge to some plateau – i.e. virtually all available potential substrates are oxidized in this situation which results in the development of a near maximum response. Figure 3 represents an “idealized” relationship be-
tween the concentrations of the inducer, the levels of oxidatively modified components, and ROSISP but it can be seen these relationships can account for many different dose dependency relationships that have been reported in the literature.

To our best knowledge, there have been no serious attempts to date to categorize oxidative stress depending on its intensity. Therefore, based on the information provided above, the following attempts to provide such an exercise using Figure 3. Zone I where no observable effects of added ROS are seen can be called "no stress at all" or "no stress". Zones II, III and IV where the stress can be observed are labeled "mild", "moderate" and "severe (strong)" oxidative stress, respectively. Under mild oxidative stress (zone II), an elevated level of ROS-modified molecules is observed, and the ROSISP situates above zero equivalent point (ZEP), which means that ROSISP is increased. For convenience, zone II may be subdivided for zone IIA where ROSISP is increasing from ZEP to its maximum level, and zone IIB where ROSISP decreases from the maximum to ZEP and crosses at the NOE point. Under moderate oxidative stress (zone III), the level of ROS-modified molecules is higher that under mild oxidative stress and the ROSISP situates below the ZEP, which means that it is decreased. Finally, under strong oxidative stress conditions (zone IV), the level of ROS-modified molecules reaches the maximum, and the ROSISP also situates below the ZEP and reaches minimum values. The entire concept is mainly related to simplified in vivo systems. Reactive oxygen species affect targets more or less nonspecifically, but induce defense systems specifically. The specificity of the pair is provided by properties of the affected target and the ROS that interacted with it.

It is also possible to propose more convenient classification from a semantic point of view. Using Figure 3, the four zones of for "Endpoint" vs "Dose of inducer" may be called: I – basal oxidative stress zone (BOS); II – low intensity oxidative stress (LOS); III – intermediate intensity oxidative stress (IOS); and IV – high intensity oxidative stress (HOS). The proposed classification may be helpful to describe experimental data where oxidative stress is induced and systematize it based on its intensity. I invite interested readers to discuss the issue in order to choose the most adequate and convenient classification system.

EXPERIMENTAL COMPLICATIONS

The previous section represents the "idealized" cellular response to exposure to an inducer of oxidative stress – a two-dimensional system with variable levels of inducer and cellular response. In reality, this ideal system is complicated by at least four factors. These are (i) the time course of the response, (ii) tissue/cell specificity, (iii) accessibility of targets to the inducer especially when dealing with multicellular organisms, and (iv) the physiological state of the organism.

It is clear that in order to develop a response to any oxidizing effector, some time is required. Moreover, time courses of various processes are usually different. Therefore, in addition to concentration dependency, the investigator has to study the development of the response over time.

Some additional points should be highlighted here. (1) If we measure several parameters to characterize oxidative stress as is the usual practice in most studies cases, the results from some of these parameters may classify the stress as strong, whereas others may indicate intermediate or even mild stress. In these cases, researchers should choose which intensity of the stress they deal with. Perhaps, some clues for selection can be provided by weighing all parameters evaluated and choosing the zone in which most of them are located; (2) How can we differentiate zones III and IV in Figure 3, i.e. the zones of intermediate (moderate) intensity oxidative stress (IOS) and high intensity oxidative stress (HOS)? Here, I propose to use an approach from biochemical kinetics. The relationship between oxidant dose and
the level of ROS-oxidized products usually follows a sigmoid or S-shaped curve converging to a horizontal asymptote (although the asymptote is virtually never reached experimentally). In biochemical kinetics, similar function is usually described by the Hill equation and the mathematical apparatus used to calculate the maximum parameter (saturation of the binding centers or maximum rate of the enzyme) may be applied. Since the maximum parameter is complicated to measure, the calculated one is used for our purposes. Actually, I recommend using the point where 90 % of the calculated maximum level is reached as a border between zones III and IV (IOS and HOS); (3) The intensity of oxidative stress changes with time after application of the stressor. For example, at the beginning the stress may be classified as HOS or IOS, but over time it might change to IOS or even LOS reflecting the organismal response or adaptation. For this reason, the researcher should be very accurate when defining the time course of the stress effects; (4) In some cases, acute oxidative stress may be a mild stress, whereas chronic oxidative stress would correspond with intermediate or high intensity oxidative stress. *not sure I understand this last point? Please check and revise if needed?

Probably, there is a need here to summarize once more biologically most relevant biomarkers to characterize oxidative stress. They are: (i) presence of ROS-modified molecules and products of ROS-promoted reactions (for lipids – malonic dialdehyde, isopсорalen, and lipid peroxides, for proteins – carbonyl proteins, and for nucleic acids – 8-oxoguanine); (ii) induction of defense systems (SoxR and OxyR in bacteria; Yap1 in yeasts; Npr1 and Rap2.4a in plants; and Nrf2 in animals). These events if not counterbalanced may lead to cell death via apoptosis or necrosis.

Studies of multicellular organisms add complications for oxidative stress researchers. The delivery of inducers and tissue/cell specificity in the response to inducers are the main problems here. The routes for inducer delivery can vary substantially and include uptake through routes including the alimentary system, skin, gills, and lungs, etc. The chemical properties of the inducers, the specificity of the absorption system, as well as inducer metabolism and excretion from the body all combine to define the dose of inducer that is experienced by each tissue type and, therefore, the tissue-specific (as well as whole organism) response(s) that occur. It should also be noted, that it is not always the original oxidant compounds that may affect the target organism or its tissues, but also the products of their chemical modification or biological catabolism that may actively determine the overall response. Another important aspect of the induction of oxidative stress by exogenous ROS is the issue of tissue specificity or even cell specificity in those tissues with multiple cell types. Each cell/tissue type may respond differently to oxidative stress inducers including experiencing different local doses, showing different thresholds for damage, possibly undergoing different types of damage, and being differentially important in determining the overall whole organism response to the stress.

The list above and many other experimental complications clearly demonstrate that the proposed classification system relies on many parameters, depends on specific conditions, the physiological state of the organisms, the parameters measured, etc. Obviously, the model will not always work out and this leads to the conclusion that it should be used in a prognostic manner. I suggest that the proposed system should be used not as “ideal” classification, but rather as working model to develop a reliable system of classification of oxidative stress with predictive strength and which can be used for quantitative evaluation.

CONCLUSIONS AND PERSPECTIVES

Oxidative stress has been extensively studied for about four decades. Substantial progress has been achieved to date – from descriptive characterization of this process to
delineation of molecular mechanisms underlining adaptive responses and targeted manipulations of expected responses. Up to the present, descriptive works still prevail, but more and more frequently studies assessing the molecular mechanisms involved are appearing (Hermes-Lima et al., 1998; Lushchak, 2011a, b; Ma, 2013; Nibali and Donos, 2013; Sies, 2014; Stone and Yang, 2006; Storey, 1996; Yang and Ming, 2012). In the light of this article, it is still important to characterize internal processes induced by ROS. Which specific targets are important for survival and for adequate responses to oxidative insults? Again, this depends on many circumstances. For example, the loss of transmembrane ion gradients as a result of high levels of lipid peroxides may be responsible in some cases, whereas in other situations, irreversible changes can be triggered by oxidative damage to mitochondrial or nuclear DNA. In many instances, ROS-triggered damage to cellular components may direct the cell to apoptosis or necrosis.

Future progress in the field needs identification of the most crucial cellular targets for ROS action as well as further discovery of the underlying mechanisms and consequences of the interaction between ROS and cellular components. The mechanisms responsible for combating ROS and their regulation would be the second hot topic for ongoing studies of ROS metabolism. In recent years, it was discovered that ROS and ROS-regulated pathways are actively involved in modification of diverse cellular processes starting from core metabolism and hormonal signaling through to complicated processes such as fertilization, development, etc. The latter along with some biotechnological avenues would also extend ROS-related studies in practical directions. Therefore, much remains to be learned about the effects of ROS on biological systems, the adaptive strategies that overcome ROS attack, and the natural use of ROS in the signaling and regulation of metabolism.

Acknowledgements: The author is grateful to Drs. H. Sies, H. Semchyshyn, D. Abele, D. Gospodaryov and M. Bayliak for critical analysis of the manuscript and a number of suggestions and ideas. Long-term personal communications with Drs. H. Sies, K. Storey, J. Storey, R. Levine, M. Nikinmaa, A. Boldyrev, V. Skulachev, and M. Hermes-Lima stimulated the author’s interest to the field of oxidative stress. I wish to thank especially J. Storey who contributed greatly to the paper editing English. The idea of this work has been formulated and discussed with many colleagues owing to financial support and great facilities provided by the Institute for Advanced Study, Delft, Germany, for the author.

REFERENCES
Abele D, Vazquez-Medina J, Zenteno-Savin T. (eds): Oxidative stress in aquatic ecosystems. New York: Wiley, 2011.
Babior BM, Kipnes RS, Curnutte JT. Biological defense mechanisms. The production by leukocytes of superoxide, a potential bactericidal agent. J Clin Invest 1973;52:741–4.
Babior BM, Curnutte JT, Kipnes RS. Biological defense mechanisms. Evidence for the participation of superoxide in bacterial killing by xanthine oxidase. J Lab Clin Med 1975;85:235–44.
Babusíková E, Jesenák M, Dobrota D, Tribulová N, Kaplán P. Age-dependent effect of oxidative stress on cardiac sarcoplasmic reticulum vesicles. Phycol Res 2008;57:49–54.
Bayliak M, Semchyshyn H, Lushchak V. Effect of hydrogen peroxide on antioxidant enzyme activities in Saccharomyces cerevisiae is strain-specific. Biochemistry (Moscow) 2006;71:1013–20.
Bayliak MM, Semchyshyn HM, Lushchak VI. Possible accumulation of non-active molecules of catalase and superoxide dismutase in S. cerevisiae cells under hydrogen peroxide induced stress. Cent Eur J Biol 2007;2:326–36.
Belluzzi E, Bisaglia M, Lazzarini E, Tabares LC, Beltramini M, Bubacco L. Human SOD2 modification by dopamine quinones affects enzymatic activity by promoting its aggregation: possible implications for Parkinson's disease. PLoS One 2012;7(6):e38026.
Brignac-Huber L, Reed JR, Backes WL. Organization of NADPH-cytochrome P450 reductase and CYP1A2 in the endoplasmic reticulum–microdomain localization affects monoxygenase function. Mol Pharmacol 2011;79:549–57.

Britigan BE, Cohen MS, Rosen GM. Detection of the production of oxygen-centered free radicals by human neutrophils using spin trapping techniques: a critical perspective. J Leukoc Biol 1987;41:349–62.

Calabrese EJ. Hormesis and medicine. Br J Clin Pharmacol 2008;66:594–617.

Catalgo LB, Grimm S, Grune T. Protein carbonyl measurement by enzyme linked immunosorbent assay. In: Abele D, Vazquez-Medina J, Zenteno-Savin T (eds.): Oxidative stress in aquatic ecosystems (pp 432-9). London: Wiley & Sons Ltd, 2011.

Claeson K, Thorsen G, Karlberg B. Methyl malondialdehyde as an internal standard for the determination of malondialdehyde. J Chromatogr 2001;751:315–23.

Commoner B, Townsend J, Pake GE. Free radicals in biological materials. Nature 1954;174:689–91.

Demple B. Regulation of bacterial oxidative stress genes. Annu Rev Genet 1991;25:315–37.

Ferrari CK, Souto PC, França EL, Honorio-França AC. Oxidative and nitrosative stress on phagocytes' function: from effective defense to immunity evasion mechanisms. Arch Immunol Ther Exp (Warsz) 2011;59:441–8.

Gerschman R, Gilbert DL, Nye SW, Dwyer P, Fenn WO. Oxygen poisoning and X-irradiation: a mechanism in common. Science 1954;119:62362–6.

Gomberg M. An instance of trivalent carbon: triphenylmethyl. J Am Chem Soc 1900;22:757–71.

Halliwell B, Gutteridge JMC. Free Radicals in Biology and Medicine. Oxford: Clarendon Press, 1989.

Harman D. Aging: a theory based on free radical and radiation chemistry. J Gerontol 1956;11:928–300.

Harman D. Origin and evolution of the free radical theory of aging: a brief personal history, 1954–2009. Biogerontology 2009;10:773–81.

Hermes-Lima M, Storey JB, Storey KB. Antioxidant defenses and metabolic depression. The hypothesis of preparation for oxidative stress in land snails. Comp Biochem Physiol B 1998;120:437–48.

Jacob C, Knight I, Winyard PG. Aspects of the biological redox chemistry of cysteine: from simple redox responses to sophisticated signalling pathways. Biol Chem 2006;387:1385–97.

Le Bourg E. Hormesis, aging and longevity. Biochim Biophys Acta 2009;1790:1030–9.

Lee YN, Shim YJ, Kang BH, Park JJ, Min BH. Overexpression of human clusterin increases stress resistance and extends lifespan in Drosophila melanogaster. Biochem Biophys Res Commun 2012;420:851–6.

Lenn AG, Costabel U, Shaltiel S, Levine RL. Determination of carbonyl groups in oxidatively modified proteins by reduction with tritiated sodium borohydride. Anal Biochem 1989;177:419–25.

Lovell MA, Markesbery WR. Oxidatively modified RNA in mild cognitive impairment. Neurobiol Dis 2008;29:169–75.

Lovell MA, Soman S, Bradley MA. Oxidatively modified nucleic acids in preclinical Alzheimer's disease (PCAD) brain. Mech Ageing Dev 2011;132:443–8.

Lüthje S, Möller B, Perrineau FC, Wöltje K. Plasma membrane electron pathways and oxidative stress. Antioxid Redox Signaling 2013;18:2163-83.

Lushchak VI. Oxidative stress and mechanisms of protection against it in bacteria. Biochemistry (Moscow) 2001;66:476–89.

Lushchak VI. Oxidative stress in yeast. Biochemistry (Moscow) 2010;75:281–96.

Lushchak VI. Free radical oxidation of proteins and its relationship with functional state of organisms. Biochemistry (Moscow) 2007;72:809–27.

Lushchak VI. Oxidative stress in animal. Biochemistry (Moscow) 2011;75:281–96.

Lushchak VI. Adaptive response to oxidative stress: Bacteria, fungi, plants and animals. Comp Biochem Physiol C 2011a;153:175–90.

Lushchak VI. Environmentally induced oxidative stress in aquatic animals. Aquat Toxicol 2011b;101:13–30.

Lushchak VI. Glutathione homeostasis and functions: potential targets for medical interventions. J Amino Acids 2012; 2012:736837.
Lushchak VI, Gospodaryov DV. Catalases protect cellular proteins from oxidative modification in *Saccharomyces cerevisiae*. Cell Biol Int 2005;29:187–92.

Lushchak VI, Semchyshyn HM, Lushchak OV. “Classic” methods for measuring of oxidative damage: TBARS, xylene orange, and protein carbonyls. In: Abele D, Vazquez-Medina J, Zenteno-Savin T (eds.): Oxidative stress in aquatic ecosystems (pp 420-31). London: Wiley & Sons Ltd, 2011.

Ma Q. Role of nrf2 in oxidative stress and toxicity. Annu Rev Pharmacol Toxicol 2013;53:401–26.

Malanga G, Puntarulo S. The use of electron paramagnetic resonance in studies of oxidative damage to lipids in aquatic systems. In: Abele D, Vazquez-Medina J, Zenteno-Savin T (eds.): Oxidative stress in aquatic ecosystems (pp 448-57). London: Wiley & Sons Ltd, 2011.

Mayne ST. Antioxidant nutrients and chronic disease: use of biomarkers of exposure and oxidative stress status in epidemiologic research. J Nutr 2003;133:933–40.

McCord JM, Fridovich I. Superoxide dismutase. An enzymic function for erythrocuprein (hemocuprein). J Biol Chem 1969;244:6049–55.

Michaelis L. Free radicals as intermediate steps of oxidation-reduction. Cold Spring Harb Symp Quant Biol 1939;7:33–49.

Nibali L, Donos N. Periodontitis and redox status: a review. Curr Pharm Des 2013;19:2687–97.

Olinski R, Siomek A, Rozalski R, Gackowski D, Fokskinski M, Guz J et al. Oxidative damage to DNA and antioxidant status in aging and age-related diseases. Acta Biochim Pol 2007;54:11–26.

Palmer RM, Ferrige AG, Moncada S. Nitric oxide release accounts for the biological activity of endothelium-derived relaxing factor. Nature 1987;327:524–6.

Rattan SI. Principles and practice of hormetic treatment of aging and age-related diseases. Hum Exp Toxicol 2008;27:151–4.

Ristow M, Schmeisser S. Extending life span by increasing oxidative stress. Free Radical Biol Med 2011;51:327–36.

Ristow M, Zarse K. How increased oxidative stress promotes longevity and metabolic health: The concept of mitochondrial hormesis (mitohormesis). Exp Geront 2010;45:410–8.

Rossi F, Della Bianca V, de Togni P. Mechanisms and functions of the oxygen radicals producing respiration of phagocytes. Comp Immunol Microbiol Infect Dis 1985;8:187–204.

Scandalios JG. Oxidative stress: molecular perception and transduction of signals triggering antioxidant gene defenses. Braz J Med Biol Res 2005;38:995–1014.

Semchyshyn H. Hydrogen peroxide-induced response in *E. coli* and *S. cerevisiae*: different stages of the flow of the genetic information. Cent Eur J Biol 2009;4:142–53.

Semchyshyn HM, Lozinska LM. Fructose protects baker's yeast against peroxide stress: potential role of catalase and superoxide dismutase. FEMS Yeast Res 2012;12:761–73.

Semchyshyn HM, Lushchak VI. Effect of protonofore 2,4-dinitrophenol on catalase activity of intact *Escherichia coli* bacteria. Ukr Biohim Zh 2004;76:42–8.

Sies H. Biochemistry of oxidative stress. Angew Chem Int Ed Engl 1986;25:1058-71.

Sies H. Strategies of antioxidant defense. Eur J Biochem 1993;215:213–9.

Sies H. Oxidative stress: oxidants and antioxidants. Exp Physiol 1997;82:291–5.

Sies H. Role of metabolic H2O2 generation: redox signalling and oxidative stress. J Biol Chem 2014;289:8735–41.

Sies H, Chance B. The steady state level of catalase compound I in isolated hemoglobin-free perfused rat liver. FEBS Lett 1970;11:172–6.

Sies H, Jones DP. Oxidative stress. In: Fink G (ed.): Encyclopaedia of stress (pp 45-8). San Diego, CA: Elsevier, 2007.

Skulachev VP. Mitochondria-targeted antioxidants as promising drugs for treatment of age-related brain diseases. J Alzheimers Dis 2012;28:283–9.
Srinivasan T, Kumar KR, Meur G, Kirti PB. Heterologous expression of Arabidopsis NPR1 (AtNPR1) enhances oxidative stress tolerance in transgenic tobacco plants. Biotechnol Lett 2009;31:1343–51.

Stone JR, Yang S. Hydrogen peroxide: a signaling messenger. Antioxid Redox Signaling 2006;8:243–70.

Storey KB. Oxidative stress: animal adaptations in nature. Braz J Med Biol Res 1996;29:1715–33.

Vartanian LS, Gurevich SM. [NADH- and NADPH-dependent formation of superoxide radicals in liver nuclei]. Biokhimiiya 1989;54:1020–5.

Veal EA, Day AM, Morgan BA. Hydrogen peroxide sensing and signaling. Mol Cells 2007;26:1–14.

Wang X, Tao L, Hai CX. Redox-regulating role of insulin: the essence of insulin effect. Mol Cell Endocrinol 2012;349:111–27.

Wehr NB, Levine RL. Quantitation of protein carbonylation by dot blot. Anal Biochem 2012;423:241–5.

Winterbourn CC, Hampton MB. Thiol chemistry and specificity in redox signaling. Free Radic Biol Med 2008;45:549–61.

Yang Z, Ming XF. mTOR signalling: the molecular interface connecting metabolic stress, aging and cardiovascular diseases. Obes Rev 2012;13:58-68.

Zimniak P. Relationship of electrophilic stress to aging. Free Radical Biol Med 2011;51:1087–105.