

White Paper

Modern Methods in Oxidative Stress Research

Redox biology and oxidative stress

Redox reactions are powerful chemical processes that involve the reduction and oxidation of proteins and metabolites found in living things. The mechanisms that regulate them are key to maintaining homeostasis and the balance between good health and disease pathology.

Oxidative stress is the state where the delicate balance of redox biology is upset, and the pathology of oxidative stress are the cellular consequences to such an imbalance.

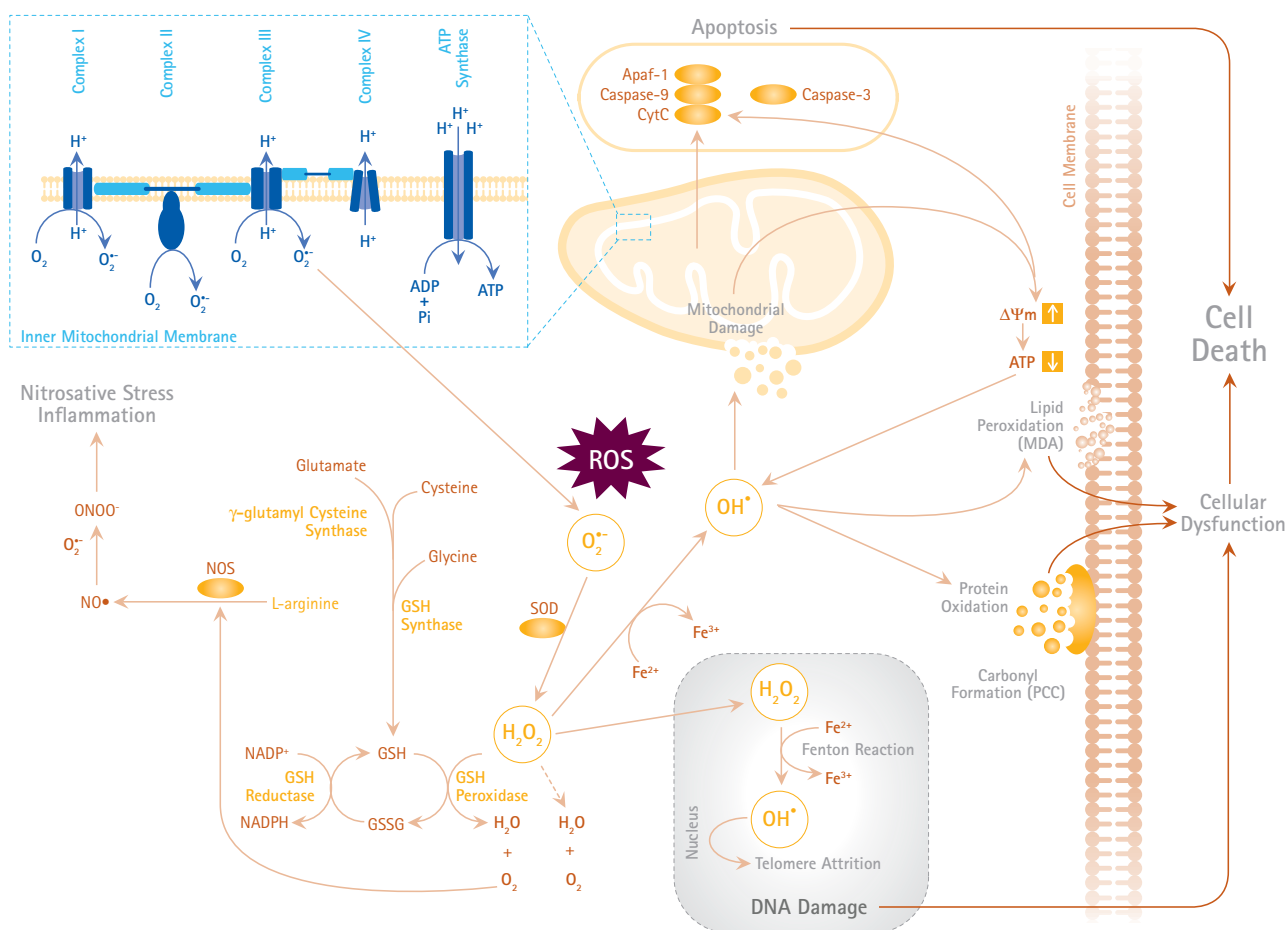


Figure 1: Oxidative stress can be linked to inflammation, DNA damage, mitochondrial damage, and cell death.

Reactive oxygen species (ROS) and reactive nitrogen species (RNS) are naturally occurring, short-lived, chemically reactive molecules produced by all living things. Structurally, they are chemical radicals and non-radical molecules derived from molecular oxygen species such as superoxide (O_2^-), singlet oxygen, and peroxide, or from reactive nitrogen species like nitric oxide (·NO) and peroxynitrite (ONOO·), for ROS and RNS, respectively^{1,4}. In mammalian cells, major endogenous sources of ROS are mitochondria and the oxidative phosphorylation reactions and respiratory processes contained within them, but other organelles such as the endoplasmic reticulum and even nuclear membranes can generate ROS/RNS as well.

ROS and RNS species also arise from a host of enzymatic reactions catalyzed by various enzymes including the cytochrome P450 enzymes, various oxidases, lipoxidases, and nitric oxide synthases. Besides endogenous sources, exogenous sources of ROS/RNS abound, both from our natural environment, like the UV radiation found in sunlight, to various food, pollutants, and toxins. Other sources include gamma irradiation, X-rays, xenobiotics, drugs, and poisons, all of which can lead to the production of ROS/RNS compounds in living systems.

In addition, our immune system has adapted the power of ROS/RNS for self-defense. ROS have a bactericidal effect on invading bacteria, and thus form a major component of our innate immune systems, though they can also contribute to and augment inflammation and immune disorders if left unchecked. Moreover, ROS/RNS are essential for various other biological functions, including cell survival, cell growth, proliferation, and differentiation^{24, 28}.

Common sources of ROS/RNS species*

Exogenous Sources

- γ irradiation
- UV irradiation
- Ultrasound
- Food
- Drugs (e.g. narcotic drugs and anesthetizing gases)
- Pollutants
- Xenobiotics
- Toxins

Endogenous Sources

- Cells (e.g. WBC, basophils, monocytes)
- Immune response
- ROS enzymes (e.g. NO synthase)
- Xanthine oxidase
- Metabolism and oxidative respiration (e.g. mitochondria)
- Injury (e.g. ischemic processes, burns)
- Metabolic disorders (intestinal disorders, aging)
- Diseases (chronic inflammation, cancer, hemochromatosis, thalassemia)

*Adapted from Kohen, R. et al. (2002)

Oxidative stress research approaches

Traditionally, oxidative stress research has focused on understanding how living cells handle the impact that ROS and RNS have on macromolecular homeostasis, and the traditional belief is that the pathology associated with oxidative stress is in direct response to ROS/RNS interactions. This connection is logical because ROS/RNS have the ability to react with a variety of cellular targets, including enzyme active sites, nucleic acids, and lipids. Diverse pathophysiologies are associated with ROS/RNS activity including diseases ranging from Alzheimer's disease, cancer, and cardiovascular disease, to diabetes and sepsis^{16, 25, 27}.

More recently, oxidative stress research has focused on the role of ROS/RNS in redox cell signaling, and how the disruption of normal redox signaling leads to a signaling dyshomeostasis and disease. ROS/RNS are known to be critical conveyors of signals (such as in NO signal transduction in the eye or in direct AP-1 and NF κ B activation) and activators of important kinase signaling cascades including MAP kinases or Jak/Stat pathways. Thus, they directly regulate cell growth as well as cell death and autophagic cellular responses²⁴. The challenge for scientists studying redox signaling is that it constitutes an essential part of normal homeostasis, so that any experimental and or therapeutic design must be carefully constructed to distinguish pathological redox signaling from normal physiological redox activity.

Key oxidative stress biomarkers, enzymes, and associated diseases

Proteins, nucleic acid, lipids, carbohydrates, and other players that are essential to fundamental cellular function and living processes are also participants in redox biology. The potential number of targets with which ROS/RNS could interact or apply influence upon is vast. The following tables provide only a small sampling of markers, proteins, and diseases associated with redox biology and oxidative stress. The lists illustrate the great depth and breadth of research targets and potential pathologies in this field of study.

Table 1. Common biomarkers in oxidative stress research*

Biomarker	Indication	Common Sample Source
Protein Targets		
Protein Carbonylation	Protein Oxidation	Blood, Cells, Tissue extracts
Protein Carbamylation	Protein Modification	Blood, Cells, Tissue extracts
Protein -SH oxidation	Protein Oxidation	Blood, Cells, Tissue extracts
3-nitrotyrosine modification	Protein Nitration	Blood, Cells, Tissue extracts
3-chlorotyrosine modification	Protein Oxidation	Blood, Cells, Tissue extracts
Dityrosine formation	Protein Oxidation	Blood, Cells, Tissue extracts
Advanced Glycation End Products (AGE) accumulation	Protein Glycation	Blood, Cells, Tissue extracts
Advanced Oxidation Protein Products (AOPP)	Protein Oxidation	Blood, Cells, Tissue extracts
Lipid Targets		
8-isoprostaglandin oxidation	Lipid Peroxidation	Blood, Bodily fluids, Cells, Tissue extracts
Malondialdehyde (MDA)	Lipid Peroxidation	Blood, Bodily fluids, Cells, Tissue extracts
8-iso-Prostaglandin F ₂ α (8-isoprostane)	Lipid Peroxidation	Blood, Bodily fluids, Cells, Tissue extracts
4-Hydroxynonenal (4-HNE)	Lipid Peroxidation	Blood, Cells, Tissue extracts
Thiobarbituric acid reactive substances (TBARS) (MDA assay)	Lipid Peroxidation	Blood, Bodily fluids, Cells, Tissue extracts
Oxidized HDL	Lipid Oxidation	Blood, Bodily fluids
Oxidized LDL	Lipid Oxidation	Blood, Bodily fluids
Nucleic Acid Targets		
8-hydroxyguanosine (8-OHG)	DNA Oxidation	Blood, Bodily fluids, Cells, Tissue extracts
8-hydroxydeoxyguanosine (8-OHdG)	DNA Oxidation	Blood, Bodily fluids, Cells, Tissue extracts
Apurinic/aprimidinic (Abasic/AP) sites	DNA Oxidation	Cells, Tissue extracts
8-hydroxyadenine	DNA Oxidation	Blood, Bodily fluids, Cells, Tissue extracts
5-hydroxycytosine	DNA Oxidation	Blood, Bodily fluids, Cells, Tissue extracts
Benzopyrene (BPDE) DNA Adduct	DNA Damage	Cells, Tissue extracts
DNA double-strand breaks (DSBs)	DNA Oxidation/DNA Damage	Cells
Universal Targets		
Hydrogen Peroxide	General ROS	Blood, Bodily fluids, Cells, Tissue extracts
Nitric oxide	General RNS	Blood, Bodily fluids, Cells, Tissue extracts
Myeloperoxidase (MPO)	General ROS	Blood, Bodily fluids, Cells, Tissue extracts
S-glutathionylation	General ROS/RNS	Blood, Bodily fluids, Cells, Tissue extracts
Catalase	Anti-oxidant	Blood, Cells, Tissue extracts
Glutathione peroxidase GPX-1	Anti-oxidant	Blood, Bodily fluids, Cells, Tissue extracts
GSH	Anti-oxidant	Blood, Cells, Tissue extracts
Superoxide Dismutase	Anti-oxidant	Blood, Bodily fluids, Cells, Tissue extracts
Ferric Reducing Ascorbate (FRASC)	Anti-oxidant	Blood, Bodily fluids, Cells, Tissue extracts

*Adapted from Babusikova E. et al. (2013)

Table 2. Selected enzyme families studied in oxidative stress

Alcohol dehydrogenase (AdhE)	Glyoxalase Enzymes
Aldehyde Oxidase Enzyme family	Myeloperoxidase/MPO
Catalase	NAD(P)H dehydrogenase Enzymes (NQO1, NQO2)
COX (Cyclooxygenase) Enzymes	NADPH Oxidases
FeS Enzymes (i.e. succinate dehydrogenase, aconitase, dihydroxyacid dehydratase)	Nitric oxide synthase Enzyme family (NOS, eNOS, nNOS)
GAPDH (Glyceraldehyde 3-phosphate dehydrogenase)	Peroxiredoxin Enzyme family
Glutaredoxin Enzyme family	SOD (Superoxide Dismutase)Enzyme Family
Glutathione Peroxidase	Thioredoxin Reductase Enzyme Family

Table 3. Diseases associated with oxidative stress*

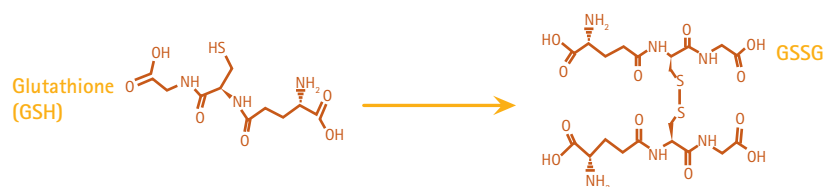
Disease	Systems Affected	Link to Oxidative Stress
Macular degeneration	Eyes, retina	ROS and intermediates
Diabetes	Multiple organs	ROS multiple enzyme dysfunction
Chronic fatigue	Multiple organs	Inflammation, C-reactive protein (CRP), ROS
Autoimmune disorders (Lupus, RA, MS)	Immune system, Multiple organs, Joints	Inflammatory ROS
Asthma	Lung	Inflammatory ROS
Neurodegenerative Diseases (Alzheimer's, Parkinson's, Amyotrophic Lateral Sclerosis Disease)	Brain, CNS, PNS	ROS/RNS, inflammation
Nephritis	Kidney	ROS, Glutathione dysfunction
Melanoma	Skin	DNA damage, lipid peroxidation, ROS
Myocardial infarction	Heart	ROS
Metabolic Syndrome	Multiple organ	ROS/RNS, inflammation

*Adapted from Rahal, A. et al. (2014)

Oxidative stress research methodology

As indicated in the preceding tables, there are numerous biomarkers, protein targets, enzymatic reactions and pathological consequences linked to biological redox reactions and oxidative stress. Fortunately, the approaches available for quantifying, imaging, and analyzing the cellular and physiological responses governing a cell's redox biology and oxidative state are equally broad. Oxidative stress is essentially driven by an imbalance or dyshomeostasis in vital redox chemistries. Most assays and reagents developed to study oxidative stress either use the direct chemical imbalance to detect the oxidative state, or they detect the downstream consequences of the oxidative condition on the proteins, lipids, DNA, or cellular processes.

For example, reduced glutathione (GSH) is considered to be one of the most important scavengers of reactive oxygen species, and its ratio with oxidized glutathione (GSSG, shown in Figure 2) is a well-accepted biomarker of oxidative stress³⁰. Normally, GSH acts as an important ROS scavenger and works to maintain the level of toxic compounds like peroxides at a low level, as well as to regenerate important cellular anti-oxidant molecules like ascorbic acid (vitamin C) which help to maintain redox homeostasis⁶.

**Figure 2: GSH to GSSG diagram***

*Adapted from Zitka, O. et al. (2012)

The GSH:GSSG ratio in normal, healthy cells is known to be greater than 100:1. However, under various models of oxidative stress, this healthy ratio can plummet to just 10:1 or lower^{7,30}. These levels can be measured either directly via chromatography, or biochemically by exploiting the existing chemistry with indicators or dyes, and quantifying the signal. For example, by including the chromogenic dye compound DTNB as an indicator (EMD Millipore Cat. No. 371757 or APT250), one can measure the level GSH:GSSG in cell extracts by ELISA (Figure 3).

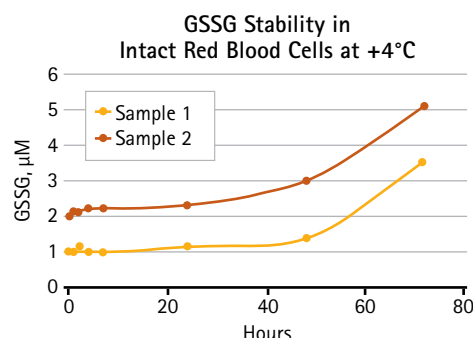


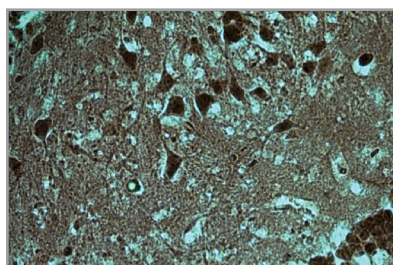
Figure 3: Measuring the level of GSSG using a chromogenic dye.*

*Representative data using EMD Millipore Cat. No. 371757

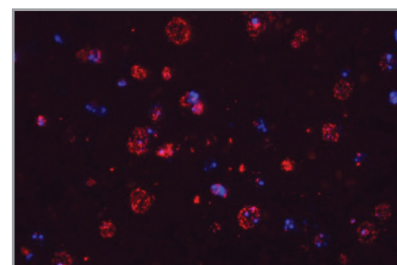
More recently, GSH:GSSG levels have been measured using live cell dyes that become fluorescent only when reacted under oxidized conditions, such as monochlorobimane (mBCl) or newer dyes like ThiolTracker™ Violet. These can be used to measure levels dynamically, thus enabling researchers to analyze oxidative stress in real time^{3,5}.

Detection of modified targets of ROS/RNS activity

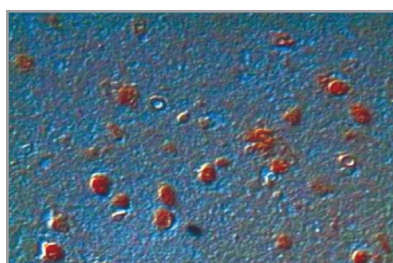
The consequence of ROS activity can be detected directly by measuring physical changes. For example, the modification of guanine in DNA to 8-hydroxydeoxyguanosine (8-OdHG) or the tyrosine in proteins to nitrotyrosine may be detected by specific antibodies or by altering the ROS modification through linking it to a chemical tag and then employing immunodetection or other means. Figure 4 shows the use of antibodies to detect ROS byproducts, and Table 4 lists some commonly used antibodies (see appendix for a wider selection).



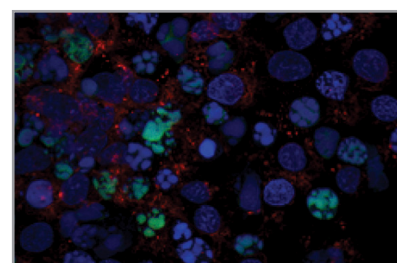
Protein oxidation detection using Nitrotyrosine antibody (Catalog No. AB5411)



Lipid peroxidation detection using 4-Hydroxynonenal antibody (Catalog No. AB5605)



DNA oxidation detection using 8-Hydroxydeoxyguanosine antibody (Catalog No. AB5830)



Detection of DNA damage by oxidative stress using anti-phospho-histone H2A.X (Ser139), clone JBW301, Alexa Fluor® 488 conjugated antibody (Catalog No. 05-636-AF488)

Figure 4: Modifications to DNA, lipids, and proteins caused by oxidative stress were analyzed using immunohistochemical (IHC) and immunofluorescence (IF) techniques.

Table 4. Selected antibodies used in oxidative stress research

Antibody Target	Catalog Number
Aldehyde dehydrogenase 3A (Aldh3A1)	ABS454
γ H2A.X	16-202A
4-Hydroxynonenal (4-HNE)	AB5605
8-Hydroxydeoxyguanosine (8-OHdG)	AB5830
8-Hydroxyguanosine	MAB3560
Lysyl Oxidase (LOX)	ABT112
Cu/Zn SOD	07-403
Dual Oxidase 2	MABN787
Myeloperoxidase	475915
NADPH Oxidase	ABC459
Neuronal Nitric Oxide Synthase (nNOS)	AB5380
Nitrotyrosine	AB5411
Peroxiredoxin 1	07-609
Superoxide dismutase 1 (SOD1)	AB5482
Thioredoxin 1 (TRX)	AB9328

Protein carbonylation and the formation of the protein carbonyl group is a common modification observed under oxidative stress. High levels of protein carbonylation have been associated with various diseases, including Alzheimer's disease, diabetes, autoimmune disease, arthritis, sepsis, and respiratory distress⁸. Protein carbonylation can be detected and measured by chemically linking the existing carbonyl groups to a detection molecule, most commonly dinitrophenol (DNP), and then detecting that entity with an antibody. Because they are commercially available, researchers have access to assays that include most common methods including western blot, IHC, ELISA, and flow cytometry. An example of detection via chemically altering the ROS modification is shown in Figure 5.

EMD Millipore applies this antibody-based technology to various platforms to enable ROS activity detection: Western blot (OxyBlot™ Protein Oxidation Detection Kit; Catalog No. S7150), ELISA (OxyELISA™ Oxidized Protein Quantitation Kit; Catalog No. S7250), ICC (OxyICC™ Oxidized Protein Detection Kit; Catalog No. S7350), IHC (OxyIHC™ Oxidative Stress Detection Kit; Catalog No. S7450), and flow cytometry (Nitrotyrosine Assay Kits; Catalog Nos. 17-10006 & 17-376). This is a useful technique, and applying it to immunohistochemistry, for instance, enables the localization of the affected tissue in relation to healthy tissues, which can be useful in measuring the effect of ischemic conditions such as stroke. With appropriate controls and data collection methods, staining can be semi-quantitative as well.

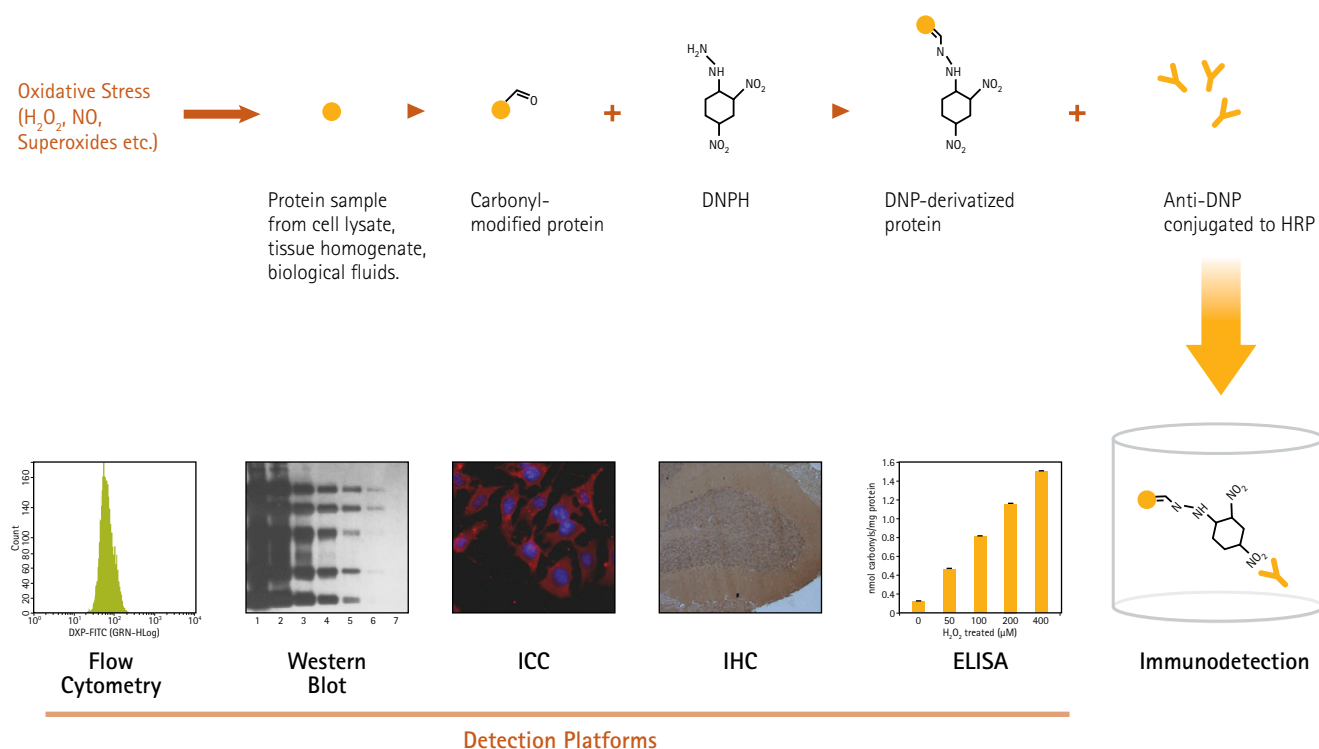


Figure 5: Diagram of assay used in various OxyBlot™, OxyELISA™, OxyICC™, OxyIHC™ and FlowCelect® protein oxidation detection assays*. Method detects the DNP epitope using an anti-DNP specific antibody (e.g. MAB2223) and has been adapted for ready-to-use kits for applications in flow cytometry, western blot, ELISA and immunocytochemistry, and immunohistochemistry.

There is a wide range of chemical dyes that exploit the existing oxidative chemistry or tagging its consequences to examine and quantify ROS/RNS species in living cells and high throughput, flow-based platforms. Table 5 below lists a selection of dyes used in oxidative stress research for the detection of the various ROS/RNS species.

Table 5. Selected dyes used in oxidative research

Reactive Oxygen and Nitrogen Species	Detection Reagents	Sample	Detection Method
Hydrogen peroxide H_2O_2	CM-H2DCFDA	Living cells	Fluorescent
	Dihydrocalcein AM/Calcein	Living cells	Fluorescent
	Dihydrorhodamine 123	Living cells	Fluorescent
	Lucigenin	Living cells	Fluorescent
	Amplex ultrared	Living cells; lysates	Fluorescent
	2,4-dinitrophenylhydrazine (DNPH)/ Dinitrophenol (DNP)	Lysates/ Fixed cells	Fluorescent/Colorimetric/ ECL/Antibody
Hydroxyl radical $HO\bullet$	3'-(p-Aminophenyl) fluorescein (APF)	Living cells	Fluorescent
	3'-(p-Hydroxyphenyl) fluorescein (HPF)	Living cells	Fluorescent
	CM-H2DCFDA	Living cells	Fluorescent
	Proxyl fluorescamine	Living cells	Fluorescent
	TEMPO-9-AC	Living cells, tissues	Fluorescent
Hypochlorous acid $HOCl$	Aminophenyl fluorescein (APF)	Living cells	Fluorescent
	Dihydrorhodamine 123	Living cells	Fluorescent
	Luminol	Lysates	Luminescent
Nitric oxide NO	DAF-FM	Living cells	Fluorescent
	1,2-diaminoanthraquinone (DAA)	Living cells	Fluorescent
	2,3-Diaminonaphthalene	Lysates	Fluorescent
Peroxyl radical $HOO\bullet$	BODIPY FL EDA	Lysates	Fluorescent/ECL/Antibody
	BODIPY 665/676	Living cells/Lysates	Fluorescent
	diphenyl-1-Pyrenylphosphine (DPPP)	Lysates/Living cells	Fluorescent
Peroxynitrite anion $ONOO^-$	3'-(p-Aminophenyl) fluorescein (APF)	Living cells	Fluorescent
	3'-(p-Hydroxyphenyl) fluorescein (HPF)	Living cells	Fluorescent
	Dihydrorhodamine 123	Living cells	Fluorescent
Superoxide anion $O_2^{\bullet-}$	MTT/NBT/XTT/WST	Living cells	Colorimetric
	Dihydroethidium (DHE)	Living cells	Fluorescent

Measuring pathways and complex cellular responses to oxidative stress

Aside from measuring ROS/RNS species and their consequences upon macromolecules themselves, significant efforts in research are focused upon the impact that oxidative stress has on the living cell's health and metabolism. The impact of ROS/RNS species on cell metabolism can be measured using bead-based multiplex assay technology, such as the Luminex® platform, where levels of individual analytes can be quantified by comparing signals with a standard curve. Multiplex assays have readouts that are similar to traditional single analyte ELISAs, but they are more complex because of the difficulties associated with antibody interactions and multi-target matrix effects. Analytically validated multiplex analyte panels and arrays address these complexities.

Traditionally a mainstay of cytokine and inflammation research, multiplex assays are also widely used in toxicology and new arenas like oxidative stress. For example, multiplex assays are used to measure the upregulation of anti-oxidant associated enzymes such as superoxide dismutases, catalases, glutathione peroxidases, and peroxiredoxins. These enzymes

are used by cells to defend themselves against ROS damage by clearing the ROS/RNS molecules. In normal conditions, superoxide dismutase and catalase convert the superoxide and hydrogen peroxide ROS radicals into oxygen and water to minimize the damage to the cells. However, when the ROS clearance pathway is impaired, accumulation of ROS can cause oxidative stress to the cells and induce DNA damage, lipid peroxidation, and enzyme inactivation. An example of the kind of data generated through multiplex assays for such anti-oxidant enzymes is shown in Figure 6.

Furthermore, an advantage of multiplex assay platforms is that analyte targets from multiple pathways can be measured together, thereby allowing the quantification of pathway interactions. For example, drugs and various compounds can act as sources of oxidative stress, which is a major cause of cell cytotoxicity-mediated cell death in the liver. Figure 7 shows multiplexed quantitation of biomarkers of drug-induced liver injury (DILI), a measure of liver cytotoxicity, in rat liver following treatment with various small molecule drug compounds such as acetaminophen (APAP) and thioacetamide (TAA).

Figure 6: Multiplexed analysis of oxidative stress associated enzymes in human cell lines and heart tissue. Lysates from different human cell lines (HUVEC, HEK 293, HeLa, and HepG2) and heart tissue were analyzed with the MILLIPLEX® MAP Human Oxidative Stress Magnetic Bead Panel (Catalog No. H0XSTMAG-18K) according to the assay protocol. The Median Fluorescence Intensity (MFI) was measured with the Luminex® system.

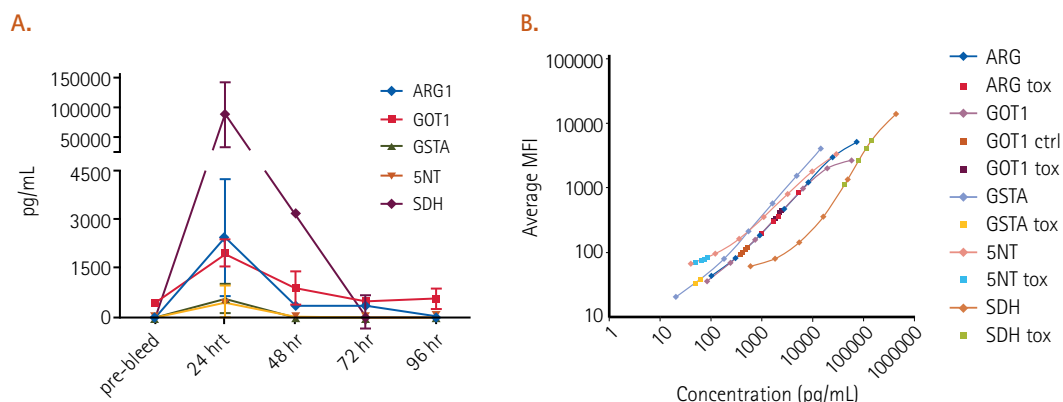
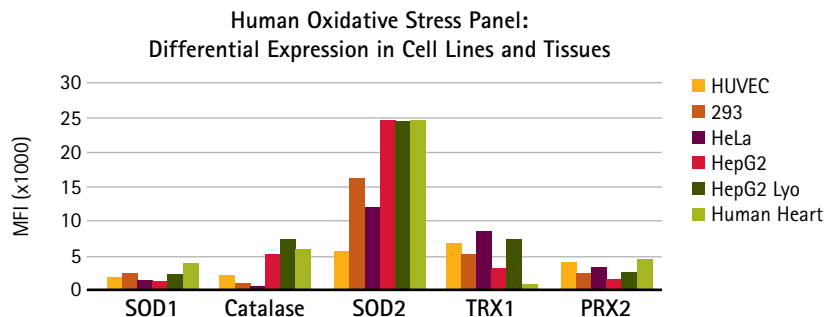


Figure 7: Multiplexed quantitation of biomarkers of drug-induced liver injury in rat following treatment with acetaminophen or thioacetamide. (A) Time-course measurements of blood protein concentrations of hepatotoxicity biomarkers ARG1, GOT1, GST-α, 5'-NT and SDH in rat acetaminophen injury model. (B) Blood protein concentrations of ARG1, GOT1, GST-α, 5'-NT and SDH are elevated in rat thioacetamide liver injury model. The MILLIPLEX® MAP Rat Liver Injury Magnetic Bead Panel (Catalog No. RLI1MAG-92K) was used to quantify blood protein concentration.

The results demonstrate that such multi-analyte panels can be used for quantitative immunoassays that simultaneously and precisely measure circulating hepatotoxicity biomarkers and prominent cytokines, demonstrating the value of this very powerful approach for investigators to monitor oxidative stress and its impact on cellular pathways.

As demonstrated in the previous example, drugs and other small molecules are often sources of oxidative stress. In biological research, small molecule inhibitors are often used to modulate an experimental system to assist researchers in sorting out the myriad complex protein and chemical interactions that are occurring at any one time.

In oxidative stress research, a wide range of small molecules are available that modulate a variety of important cellular pathways impacted by oxidative stress. Small molecule reagents include inducers of oxidative stress such as peroxynitrite, modulators like the free radical scavenger TEMPOL, and specific inhibitors of oxidative stress enzymes (e.g. nitric oxide synthase NOS or Glutathione S-Transferase (GST) and others). By combining specific inhibitors of relevant pathways, such as kinase inhibitors for signaling pathways, or mitochondrial inhibitors for bioenergetic studies, researchers can examine how the various cellular pathways interact and intersect during oxidative stress.

Table 6. Selected small molecules used in oxidative stress research

Compound	Oxidative Stress-Related Target
AAPH (2,2'-Azobis(2-amidinopropane) dihydrochloride)	Oxidative Stress Inducer
DPPH Free Radical	Oxidative Stress Inducer
Peroxynitrite	Oxidative Stress Inducer
Apocynin	NADPH Oxidase (NOX) Inhibitor
7-Nitroindazole	Nitric Oxide Synthase (NOS) Inhibitor
Diphenyleneiodonium Chloride	Nitric Oxide Synthase (NOS) Inhibitor
HX1	Myeloperoxidase Inhibitor
Ferrostatin-1	Ferroptosis Inhibitor (non-apoptotic cell death inhibitor)
TEMPOL	Free-radical scavenger
Luteolin	Anti-oxidant
BEC, Hydrochloride	Arginase Inhibitor
LY 83583	Guanylate Cyclase Inhibitor
ML171	NADPH Oxidase (NOX) Inhibitor
VSC2	Nrf2 Cytoprotective Pathway Activator
SB203580	p38 MAPK Inhibitor

Measuring gene expression in oxidative stress research

With redox biology and ROS/RNS species playing such vital roles in living systems, their impact on gene expression has been well studied. Direct signaling by ROS/RNS can impact gene expression, in turn affecting cell health and cell proliferation; for example, NO-mediated signaling can regulate gene expression^{12,22,23}. Traditional methods for examining gene expression follow what have become standard approaches, from quantitative PCR to RNA sequencing.

The most frequently studied targets are those related to ROS/RNS creation, containment, or processing (for example, antioxidants such as glutathione peroxidases and other enzymes listed in Table 1), as well as those involved in the oxidative stress gene response (apolipoprotein E, catalase, cytoglobulin). Table 7 provides a small listing of the popular ROS/RNS associated genes examined.

Table 7. Selected Genes involved in ROS Metabolism & Gene response

ALOX12	ANGPTL7	AOX1	APOE	ATOX1	BNIP3	CAT	CCL5
CCS	CSDE1	CYBA	CYGB	DGKK	DUOX1	DUOX2	DUSP1
EPHX2	EPX	FOXO1	GLRX2	GPR156	GPX1	GPX2	GPX3
GPX5	GPX6	GSS	GTF2I	IPCEF1	KRT1	LPO	MBL2
MPO	MPV17	MSRA	MT3	MTL5	NCF2	NME5	NOS2
NUDT1	OXR1	OXSRI	PDLIM1	PNKP	PRDX2	PRDX5	PREX1
PRG3	PRNP	RNF7	SCARA3	SEPP1	SFTPD	SGK2	SIRT2
SOD1	SOD2	SOD3	SRXN1	STK25	TPO	TTN	TXNRD2

With our growing understanding of gene regulation and RNA metabolism, miRNAs are increasingly being researched for their role in oxidative stress and ROS/RNS regulation. Table 8 lists some of the miRNAs that are known to touch oxidative stress and redox biology.

Table 8. New miRNA targets associated with oxidative stress*

MicroRNAs	Association with oxidative stress
miR-150	downregulated
miR-142-5p	downregulated
miR-122	downregulated
miR-34c	upregulated
miR-34-5p	upregulated
miR-29b	upregulated
miR-743a	involved
miR-335	involved
miR-34a	involved
miR-200c	involved
miR-145	involved
miR-205	involved
miR-320	involved
Let-7	involved
miR-23	involved
miR144	involved
miR-451	involved

Adapted from Ganguly, N.K. et al. (2014)

Lastly, new detection technologies, such as SmartFlare™ RNA detection probes, enable measurement of oxidative stress response in single, live cells, using flow cytometry, imaging flow cytometry, or other cell analysis methods. Such technologies offer researchers the ability to track these miRNA and also mRNA corresponding to ROS/RNS metabolic and response genes in heterogeneous cell populations.

Measuring mitochondrial health in oxidative stress

Mitochondrial function is critical to cellular health. Ironically, mitochondria are responsible for up to 90% of the endogenous ROS present in the normal healthy cell¹¹, so it is not surprising that oxidative stress research focuses much attention on tracking mitochondrial ROS and measuring mitochondrial function and health. Because mitochondria are dynamic organelles that move and fuse inside living cells, the advancements made in fluorescent and potentiometric dyes have made it possible to study not only the number and size of mitochondria in cells but also their health, membrane potential, and the impact of oxidative stress and cell death processes.

Mitochondrial membrane potential ($\Delta\psi_m$) is a critical measurement as it relates to the cell's capacity to generate ATP by oxidative phosphorylation. The maintenance of the mitochondrial membrane potential is accomplished by a delicate balance of electrical charges, pH, and ion gradients. Oxidative stress and the associated ROS impact these electrical and chemical gradients, and when the ionic fluxes surpass the ability of the mitochondria to buffer them, the mitochondrial membrane potential can collapse and lead to bioenergetic stress and cell death²⁰.

Most of the mitochondria-specific dyes used in oxidative research depend upon a healthy mitochondrial membrane potential for entry and accumulation. The mitochondrial dyes are positively charged lipophilic compounds that equilibrate across membranes in a polarized fashion. In other words, the amount of dye that accumulates is proportional to the polarization of the membrane potential. As the dyes enter and become concentrated, many undergo a fluorescence shift that can be used to monitor the status of the mitochondrial membrane potential and, by extension, mitochondrial function.

Table 9. Common fluorescent dyes to study mitochondria

Fluorescent Dye	Ex Max	Em Max	Comment/ Function
DASPEI	461	589	Excellent for staining live mitochondria in living cells
DiOC6	488	501	Indicator of mitochondrial membrane potential, widely used but must use very low concentrations
JC-1	498	527/599	Indicator of mitochondrial membrane potential; spectral shift depends upon dye concentration; commonly used in apoptosis studies
MitoSense Red (DiIC1(5))	638	659	Indicator of mitochondrial health, useful when combined with other cytoplasmic dyes like CF488A and 7-AAD for cell health studies
MitoSOX Red	488	580	A live-cell permeant fluorogenic dye which targets the mitochondria and reacts with superoxide radicals and fluoresces yellow/red
NAO	495	522	Uptake is not dependent upon mitochondrial membrane potential, good for measuring mass
RHOD 123	507	529	Indicator of mitochondrial membrane potential and ATP production, fast but can be self-quenching
TMRE TMRM	533	576	Indicator of mitochondrial membrane potential but more permeable than RHOD 123, low mitochondrial binding, effective at low concentrations (1-30 nM)

*Adapted from Marin-Garcia, J. et al. (2013)

MitoSOX red is a live-cell-permeant fluorogenic dye that targets the mitochondria because it binds mtDNA, but also reacts with superoxide radicals present in excess during oxidative stress and fluoresces yellow/red. CF647 is a bright, far red fluorescent dye. An example of how mitochondrial health is examined using dyes such as these is shown in Figure 8. Here, the FlowCelect® MitoStress Kit (Catalog No. FCCH100109) was used to collect information on both oxidative stress and apoptotic state in a population of cells using flow cytometry.

The FlowCelect® MitoStress Kit includes MitoSOX Red as well as Annexin V conjugated to CF647 (which binds to phosphatidylserine (PS) on the surface of apoptotic cells). As shown in Figure 8, healthy control cells do not

demonstrate Annexin V signal, while apoptotic cells will exhibit positive 665 nm fluorescence as Annexin V binds exposed PS. The FlowCelect® MitoStress kit can thus distinguish the following populations: 1) Live, healthy, cells with little or no superoxide; 2) Stressed cells with oxidized MitoSOX Red (indicator of accumulated superoxide) but no Annexin V staining; 3) Stressed and Early Apoptotic cells with oxidized MitoSOX Red and Annexin V binding; and 4) Apoptotic Cells with Annexin V binding only. Because the assay is flow cytometry-based, it is rapid and requires minimal sample. Researchers can exploit this biochemistry to their advantage to assess levels of oxidative stress (measured via the MitoSOX red fluorescence) and the state of apoptosis (measured by CF647 fluorescence).

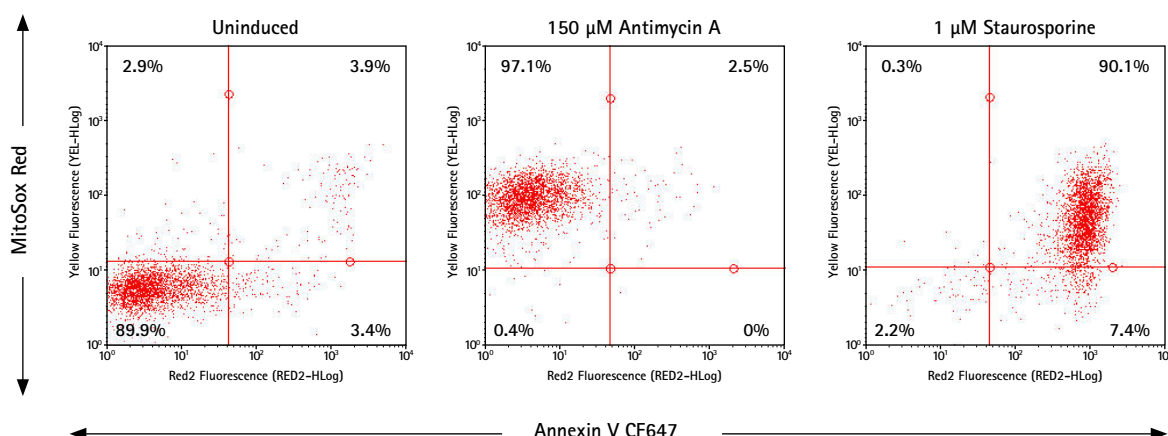


Figure 8: Dot plot analysis of Jurkat cells treated with different inducers and then stained using FlowCelect® MitoStress Kit. Untreated Jurkat cells (left plot), or those treated with 150 μ M Antimycin A (inhibitor of Mitochondrial complex III which increases mitochondrial ROS production, middle plot), or with 1 μ M Staurosporine (right plot) were stained using FlowCelect® MitoStress Kit and analyzed by flow cytometry. Samples which undergo a change in mitochondrial superoxide production show an increase in yellow/red fluorescence and samples which demonstrate increased apoptosis show an increase in red2 fluorescence.

In the FlowCollect® MitoDamage Kit (Figure 9), MitoSense Red is combined with Annexin V conjugated to the dye CF488A, and combined with the cell impermeant DNA intercalator 7-Aminoactinomycin-D (7-AAD), a dye that distinguishes dead from live cells. The simultaneous use of these reagents enables the simultaneous determination of viability along with early, mid and late apoptosis in one simple assay. Healthy cells with intact mitochondrial membrane potentials demonstrate robust fluorescence at 650 nm from MitoSense red, while cells which have impaired mitochondrial membrane potential demonstrate lower 650 nm fluorescence. The early-apoptosis indicator Annexin V is labeled with the green fluorophore CF488A, so healthy cells exhibit no green fluorescence, while apoptotic cells fluoresce green. Finally, by including the 7-AAD dye, permeability changes typically observed later in apoptosis as well as in necrotic

cell death can also be determined. 7-AAD is excluded from live, healthy cells as well as early apoptotic cells and these cells have low red fluorescence, but 7-AAD can pass into late stage and necrotic cells thus these cells exhibit strong 7-AAD fluorescence. Therefore in such an analysis four different cell populations can be distinguished: 1) Live cells with normal mitochondrial membrane potential; 2) Cells with dissipated membrane potential but no Annexin V or 7AAD staining; 3) Early apoptotic cells with dissipated membrane potential and Annexin V binding; and 4) Late apoptotic cells or dead cells with dissipated membrane potentials. The capacity for this assay to place each cell into one of these populations provides a complete picture of mitochondrial damage to cell health and enables correlation of mitochondrial damage to cell health.

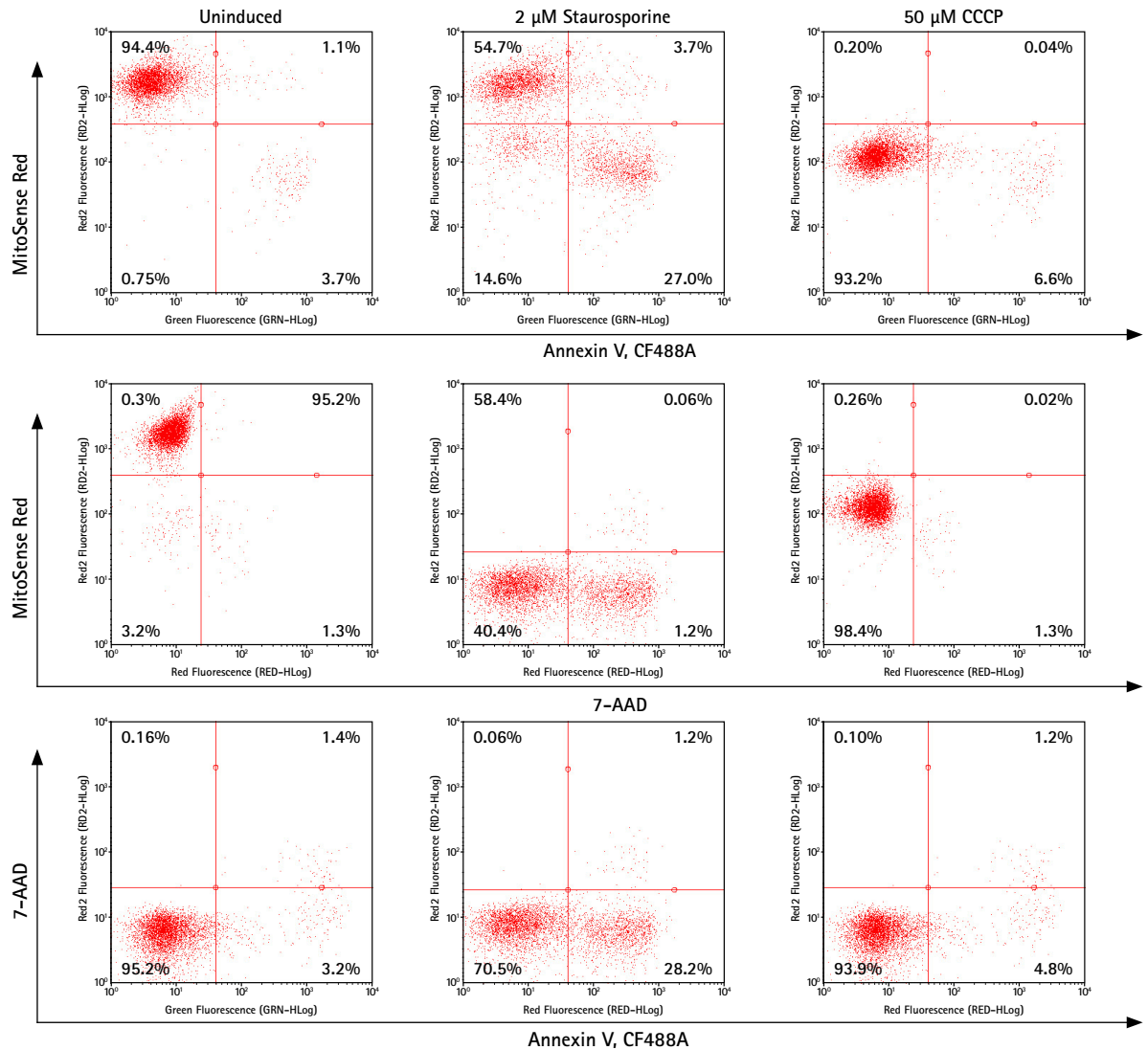


Figure 9: Dot plots depicting Jurkat cells treated with multiple inducers and stained using the FlowCollect® MitoDamage Kit. Jurkat cells were treated with zero (left), 2 μM Staurosporine (middle) and 50 μM CCCP (right) and then stained using the MitoDamage Kit. Plots show the percentage of positive cells for 1) Apoptosis (Annexin V binding) and mitochondrial membrane potential change (Red2), 2) Cell death and mitochondrial membrane potential change, and 3) apoptosis and cell death.

Platforms for dynamic oxidative stress and cell death research

The preceding sections have shown that the reagents and platforms used in oxidative stress research are quite varied and span the spectrum of methods and materials available today. Simple western blots and immunohistochemical approaches, to single and multiplexed target ELISAs, flow cytometry, and molecular DNA/RNA approaches have all been successfully employed in oxidative stress research.

As research uncovers more detail regarding how cells maintain redox homeostasis and how the various elements of exogenous and endogenous sources of ROS/RNS move cells into redox dyshomeostasis, it is becoming increasingly clear that these complex interactions will require sophisticated technological approaches. Within the last decade, two very powerful platforms, the CellASIC® ONIX Microfluidic System and the ImageStream® Mark II Imaging Flow Cytometer, have emerged as robust and innovative technologies for studying oxidative stress.

CellASIC® ONIX Microfluidic Platform: A dynamic live cell assay platform to elucidate the mechanisms underlying autophagy elicited by oxidative stress

Autophagy (self-eating) is a complex cellular process essential for cell survival under stressed conditions. Recent evidence has also suggested that tumor cells use the autophagic pathway to promote survival under oxidative stress conditions such as nutrient deprivation or hypoxia. Traditionally, the use of static end-point assays has limited our understanding of the dynamics of autophagy during cell stress and recovery phases. To gain a better understanding of the autophagic process, the CellASIC® ONIX Microfluidic System can be used to monitor both the rate of autophagosome formation and changes in lysosomal degradative processes during autophagy in live cells. By combining live cell analysis with cell lines stably expressing fluorescently-tagged markers such as LC3-GFP that are specific for autophagosomes, the cellular dynamics of autophagy can be visualized at the single cell level in real time.

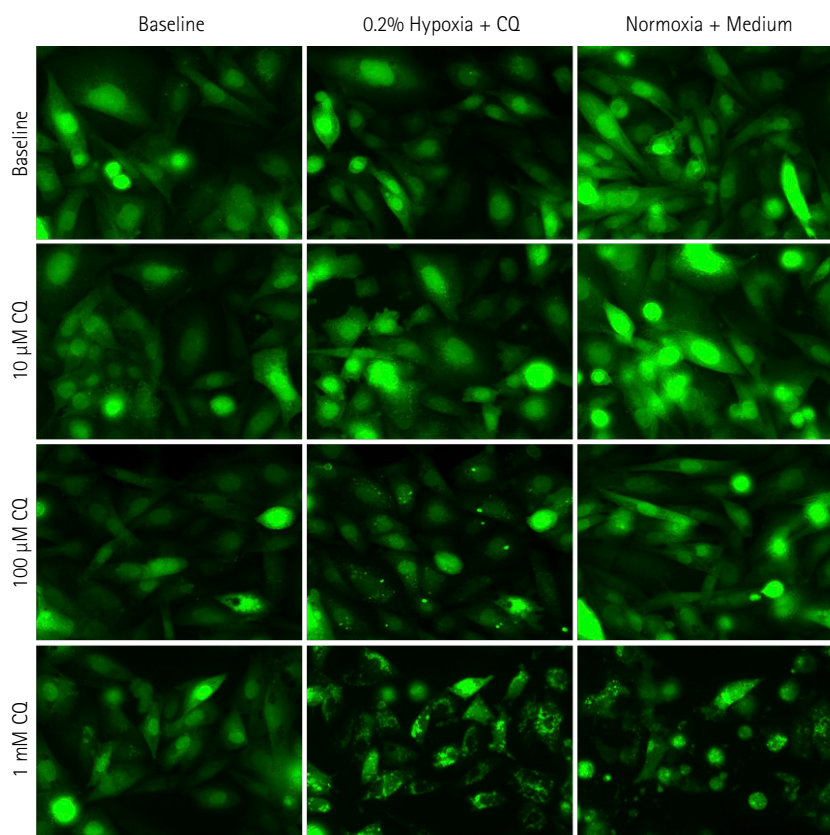


Figure 10: Images of the LC3-GFP CHO reporter cells cultured on the CellASIC® ONIX system taken during each phase of the hypoxia-induced autophagy assay. Cultures were first perfused with standard growth medium under normoxic conditions for 60 minutes, followed by continuous CQ (10 μ M, 100 μ M, or 1 mM) perfusion in the presence of severe hypoxia (0.2% O₂) treatment for 3 hours, followed by removal of the stressors and reestablishment of normoxia in standard growth medium for another 16 hours. An Olympus IX-71 inverted microscope was used for the entire process; all images were taken under the 40x objective.

Moreover, because the CellASIC® ONIX Microfluidic System can temporally regulate media and gas exchange within a culture chamber, this affords unprecedented control of the cellular growth microenvironment. An example of results obtained from this system is shown in Figure 10, where LC3-GFP CHO reporter cells were exposed to starvation or hypoxic conditions—both of which foster oxidative stress—in the presence of various stressor compounds and monitored for autophagy.

Because the CellASIC® ONIX Microfluidic System maintains live cell conditions, changes in LC3 levels (as measured by autophagosome counts) were monitored throughout the culture duration using a standard fluorescence microscope. The real-time analysis capabilities of the CellASIC® ONIX platform permits the manipulation and assessment of not only oxidative stress events but also of the autophagic process that follows.

Amnis brand ImageStream® Mark II Imaging Flow Cytometer: high resolution images of thousands of cells undergoing oxidative stress

Oxidative stress research techniques and methods rely heavily on identifying markers and utilizing detection chemistries to help scientists quantify, record, and compare the cellular and organismal responses to oxidative stress. With the advances made in fluorescent chemistries and the increasing use of live cell analysis as an experimental stage, combined with the need for statistically significant data from populations, the advantages of a system that collects images in flow are clear.

The ImageStream® Mark II Imaging Flow Cytometer platform combines the power of digital fluorescence microscopy with the speed and sensitivity of flow cytometry to enable researchers to approach scientific questions with experimental approaches that are not possible with either technique alone. For example, ionizing radiation is an exogenous source of oxidative stress that results in DNA damage. One of the most common methods of detecting DNA damage as measured by the DNA double strand breaks (DSBs) that are the hallmark of ionizing radiation is to detect the phosphorylation of histone H2A.X using a phosphospecific antibody that has been labeled with a fluorescent tag¹⁵. Phosphorylated H2A.X (γ -H2A.X) facilitates recognition and repair of DNA double strand

breaks that may occur from exposure to ionizing radiation. Staining irradiated cells for γ -H2A.X reveals nuclear foci that are readily observed microscopically in a dose-dependent manner (Figure 11).

Irradiated cells were analyzed for the number of spots in the nuclear region using advanced masking techniques that identify punctate staining. Morphological measurements employed in this analysis including object shape, size, and punctate fluorescence spot counting emphasize the advantages of quantitative multiparametric image analysis on large numbers of cells enabled by the ImageStream® system.

As discussed, oxidative stress can have an impact on cell health and even activate apoptotic or autophagic pathways. While traditional flow cytometry solutions are helpful for resolving populations based on degree of fluorescent signal from stress markers, discrimination of apoptotic cells from healthy cells is enhanced on the ImageStream® system, which uses the image of the apoptotic cell nuclei as the differentiating data. For example, Figure 12 shows the dramatic changes in nuclear morphology that are hallmarks of apoptosis. When cells begin to die by apoptosis, fragmentation and condensation of the DNA occurs. This makes possible the automated identification of apoptotic cells by measurement of the area and the intensities of the brightest portions of the nuclear image. The bright, punctate nuclear satellites found in apoptotic cells can be quantitatively distinguished from the appearance of a normal, healthy nucleus.

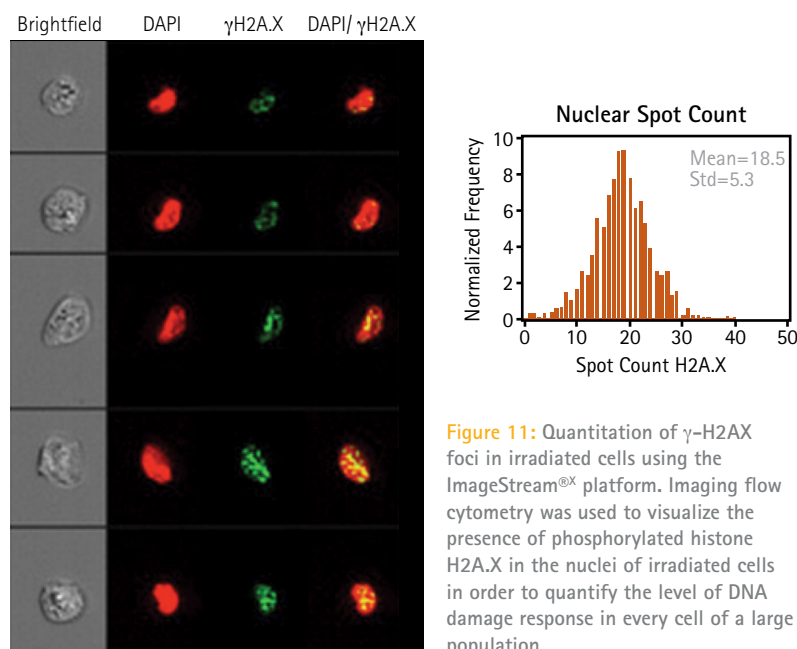


Figure 11: Quantitation of γ -H2A.X foci in irradiated cells using the ImageStream® platform. Imaging flow cytometry was used to visualize the presence of phosphorylated histone H2A.X in the nuclei of irradiated cells in order to quantify the level of DNA damage response in every cell of a large population.

A review of the published literature making use of the ImageStream[®] technology in oxidative stress demonstrates its utility in a variety of living systems from bacteria to cancers, addressing studies from cigarette smoke toxicity and its influence on DNA damage to transcription factor localization in *miR-144/451*^{-/-} mice^{26,29}. Furthermore, one the most useful aspects that an imaging platform like the ImageStream[®] instrument brings to researchers is the ability to precisely determine within a cell where a particular drug or compound, or protein is localizing. For instance, researchers examining a new chemical variant of an anti-tumor compound, which normally localized to the nucleus was found to be localized to the mitochondria and not the nucleus, thus providing an alternative explanation for its cytotoxicity than

previously thought¹³. In another example, researchers examining a new redox marker designed to study the dynamics of pathogenic bacteria during infection in both permissive and non-permissive human macrophages were able to discover and quantify redox marker differences in subpopulations of the human cells, while still being able to process individual images of individual bacteria. Because of the large sampling size and high image resolution available on ImageStream[®] imaging flow cytometer, researchers were able to observe redox heterogeneity between cells, and even between bacteria, providing a level of detail that had not been available before²⁶. Collectively, these studies highlight the many advantages of the ImageStream[®] platform, and the advances in research it is enabling.

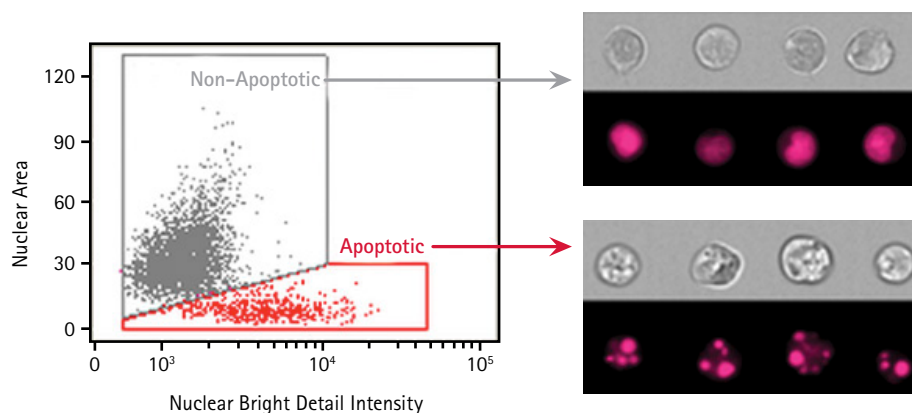


Figure 12: Apoptotic Index Using the ImageStream[®] Platform. Imaging flow cytometry used to quantify the degree of nuclear fragmentation in apoptotic cells.

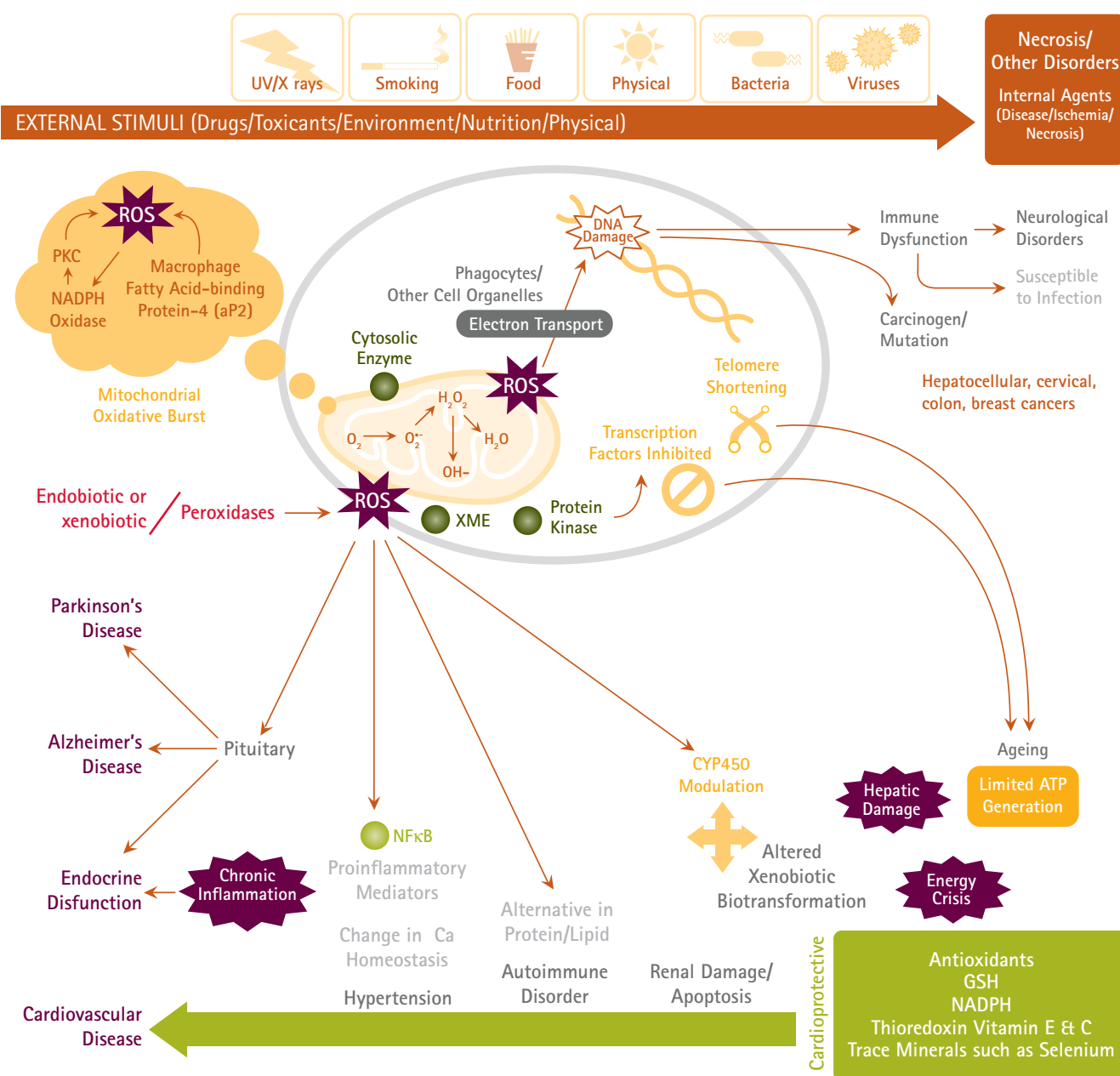
Concluding remarks

The importance and complexity of research into redox biology and that of oxidative stress cannot be underestimated. The potential damage inflicted upon biological systems by ROS/RNS and oxidative stress has been implicated in contributing to a number of important diseases including cancer, Alzheimer's disease, heart disease, and diabetes. Therefore, there is a pressing need to learn as much as we can about the factors that manifest, contribute, respond to, and control oxidative stress in living systems. As we learn more and more about the relationship that redox biology and ROS/RNS share in the chemistry of life, we have come to

understand their importance not only in the fundamental enzymatic and energy metabolism of living things, but also in health, developmental biology, and aging.

This brief review has attempted to outline some of the reagents, kits, assays, and platform options available to scientists studying redox biology and oxidative stress research. With them, scientists will be able to continue research and discovery into the cellular and environmental mechanisms that govern the production, maintenance, response and deposition of ROS/RNS species.

Overview of Oxidative Stress and Disease Development



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Appendix

Products for Oxidative Stress Research

Description	Cat. No
Antibodies	
Anti-4-Hydroxynonenal	AB5605
Anti-8-Hydroxydeoxyguanosine	AB5830
Anti-8-Oxoguanine, clone 483.15	MAB3560
Anti-AGE (Advanced Glycated End-products)	AB9890
Anti-Aldh3A1	ABS454
Anti-Cu/Zn-SOD	07-403
Anti-Cytochrome C, clone C-7	05-479
Anti-Dual oxidase 2, clone Duox S-12	MABN787
Anti-eNOS/NOS III, CT	07-520
Anti-Glutathione	AB5520
Anti-Glutathione, clone D8	MAB5310
Anti-Glutathione, detects both GSH & GSSG	AB5010
Anti-Glutathione: N-ethylmaleimide adduct, clone 8.1GSH	MAB3194
Anti-Glutathione-S-Transferase (GST), <i>S. japonicum</i> form	ABN116
Anti-Glutathione-S-Transferase, <i>S. japonicum</i> form	AB1372
Anti-Glutathione-S-Transferase, <i>S. japonicum</i> form	AB3282
Anti-Glutathione-S-Transferase, <i>S. japonicum</i> form, clone 8F6.2	MAB3117
Anti-Glutathione-S-Transferase-pi, mammalian form	AB8902
Anti-GPx1	ABN63
Anti-iNOS/NOS II, clone 13F5.1	MABN527
Anti-iNOS/NOS II, NT	06-573
Anti-iNOS/NOS II, NT	ABN26
Anti-KEAP1	ABS97
Anti-KEAP1, clone 7G4B10	MABC713
Anti-Lactotransferrin	07-685
Anti-Lysyl Oxidase (LOX)	ABT112
Anti-Mitochondrial dicarboxylate carrier, clone 1F5.1	MABN457
Anti-Mn-SOD	06-984
Anti-Myeloperoxidase Rabbit pAb	475915
Anti-Myeloperoxidase, human white cell	AB1224
Anti-NDUFB7	ABC391
Anti-Nitric Oxide Synthase I	AB1529
Anti-Nitric Oxide Synthase I	AB1632
Anti-Nitric Oxide Synthase I	AB5380
Anti-Nitric Oxide Synthase II	AB16311
Anti-Nitric Oxide Synthase II	AB5382
Anti-Nitric Oxide Synthase II	AB5384
Anti-Nitric Oxide Synthase III	AB16301
Anti-Nitric Oxide-Dopamine	AB5902
Anti-Nitric Oxide-Glutathione	AB5540
Anti-Nitrotyrosine	06-284
Anti-Nitrotyrosine	AB5411

Appendix

Products for Oxidative Stress Research

Description	Cat. No
Antibodies (continued)	
Anti-Nitrotyrosine	AB5532
Anti-Nitrotyrosine Magnetic Bead Conjugate	16-310
Anti-Nitrotyrosine, clone 1A6	05-233
Anti-Nitrotyrosine, clone 1A6, agarose conjugate	16-163
Anti-Nitrotyrosine, clone 1A6, Alexa Fluor® 488 conjugate	16-226
Anti-Nitrotyrosine, clone 1A6, Alexa Fluor® 555 conjugate	16-227
Anti-Nitrotyrosine, clone 1A6, HRP conjugate	16-207
Anti-Nitrotyrosine, clone 2A8.2	MAB5404
Anti-nNOS/NOS I	07-571
Anti-nNOS/NOS I	07-571-I
Anti-nNOS/NOS I, clone 2G1.1	MABN533
Anti-NOX4 Antibody	ABC459
Anti-Peroxiredoxin 1	07-609
Anti-Peroxiredoxin 2	07-610
Anti-Peroxiredoxin 3	07-611
Anti-Peroxiredoxin 4	07-612
Anti-Peroxiredoxin-2	ABN1011
Anti-Peroxiredoxin-3, clone EPR8115, Rabbit Monoclonal	MABN1162
Anti-Peroxiredoxin-4, clone CPTC-PRDX4-3	MABS37
Anti-Peroxiredoxin-5 (PRDX5), Human, clone 5 286 6F7	MABN300
Anti-Peroxiredoxin-5, (PRDX5), clone 5 288 2F4	MABN301
Anti-phospho eNOS (Ser1177)	07-428-I
Anti-phospho-eNOS (Ser1177)	07-428
Anti-phospho-eNOS/NOS III (Ser116)	07-357
Anti-phospho-eNOS/NOS III (Ser617)	07-561
Anti-phospho-eNOS/NOS III (Ser635)	07-562
Anti-phospho-eNOS/NOS III (Thr495), Rabbit Monoclonal	04-811
Anti-phospho-Histone H2A.X (Ser139), clone JBW301, FITC conjugate	16-202A
Anti-RAGE	AB5484
Anti-RAGE	AB9714
Anti-RAGE, azide free	AB5484Z
Anti-RAGE, clone DD/A11 or A11	MAB5328
Anti-RAGE, CT	AB5601
Anti-REDD1	ABC245
Anti-SOD1 (ALS mutant), clone MS785	MABN834
Anti-SOD1, clone 6F5	MABC684
Anti-SOD2	AB10346
Anti-SODD, NT	AB16518
Anti-SSBP1	ABN403
Anti-Superoxide Dismutase 1	AB5482
Anti-Superoxide Dismutase 1, aa 25-37	AB5480
Anti-Thioredoxin 1	AB9328

Appendix

Products for Oxidative Stress Research

Description		Cat. No
Small Molecules		
Oxidative stress Inducers	AAPH	100100
	DPPH Free Radical	300267
	Peroxynitrite	516620
Anti-oxidants and Free Radical Scavengers	(±)-Taxifolin Hydrate	580553
	α-Lipoic Acid	437692
	Carboxy-PTIO, Sodium Salt	217386
	EUK-8	341209
	Ferrostatin-1	341494
	FeTMPyP	341501
	FeTPPS	341492
	Glutathione Monoethyl Ester	353905
	Glutathione, Reduced, Free Acid	3541
	Luteolin	440025
	MCI-186	443300
	Mn-cpx 3	475867
	MnTBAP	475870
	MnTMPyP	475872
	Myeloperoxidase Inhibitor III, HX1	532280
	Myeloperoxidase Inhibitor-II	504908
	N- <i>tert</i> -Butyl-α-phenylnitron	203995
	PTIO	523350
	TEMPOL	581500
Arginase Inhibitors	BEC, Hydrochloride	197900
	DL-α-Difluoromethylornithine, Hydrochloride	288500
	NG-Hydroxy-L-arginine, Monoacetate Salt	399250
	Nω-Hydroxy-nor-L-arginine, Diacetate Salt	399275
Guanylate Cyclase Inhibitors	LY 83583	440205
	Methylene Blue	467250
	NS 2028	492030
	ODQ	495320
NADPH Oxidase (NOX) Inhibitors	Apocynin	178385
	NOX1 Inhibitor, ML171	492002
	NOX1 Inhibitor, NoxA1ds Set	532759
	NOX Inhibitor III, VAS2870	492200
	NOX Inhibitor VII, Thr101	500526
	NOX Inhibitor VIII, VAS3947	532336

Appendix

Products for Oxidative Stress Research

	Description	Cat. No
Small Molecules (continued)		
Nitric Oxide Synthase Inhibitors	1400W	100050
	1-Pyrrolidinecarbodithioic Acid, Ammonium Salt	548000
	7-Nitroindazole	483400
	Advanced Glycation Endproduct-BSA	121800
	AG126	658452
	Diphenyleneiodonium Chloride	300260
	L-N ⁵ -(1-Iminoethyl)ornithine, Dihydrochloride	400600
	L-NIL, Dihydrochloride	482100
	N ⁶ ,N ⁶ -Dimethyl-L-arginine, Dihydrochloride	311203
	N ⁶ ,N ⁶ -Dimethyl-L-arginine, Dihydrochloride	311204
	N ⁶ -Monomethyl-L-arginine, Monoacetate Salt	475886
	N ⁶ -Nitro-L-arginine Methyl Ester, Hydrochloride	483125
	Nitric Oxide Synthase, Neuronal Inhibitor I	490070
	nNOS - PSD-95 Interaction Inhibitor, ZL006	482740
	p-Nitroblue Tetrazolium Chloride	484235
	SKF-525A, Hydrochloride	567300
	S-Methyl-L-thiocitrulline, Dihydrochloride	472804
Nrf2 Related Products	Keap1-Nrf2 Interaction Probe, ML334	505987
	Nrf2 Activator	492040
	Nrf2 Activator II, AI-1	492051
	Nrf2 Activator III, TAT-14 Peptide	492042
	Nrf2 Activator IV, VSC2	530351

	Description	Cat. No
Kits and Assays		
	OxyBlot™ Protein Oxidation Detection Kit	S7150
	OxyELISA™ Oxidized Protein Quantitation Kit	S7250
	OxyICC™ Oxidized Protein Detection Kit	S7350
	OxyIHC™ Oxidized Protein Detection Kit	S7450
	Human Oxidative Phosphorylation (OXPHOS) Magnetic Bead Panel	HOXPSMAG-16K
	Rat/Mouse Oxidative Phosphorylation (OXPHOS) Magnetic Bead Panel	RMoxPSMAG-17K
	Cytochrome C ELISA kit	EAMT001-1KIT
	Human Fatty Acid Oxidation Magnetic Bead Panel 1	HFA01MAG-11K
	Human Fatty Acid Oxidation Magnetic Bead Panel 2	HFA02MAG-11K
	Human Oxidative Stress Magnetic Bead Panel	HOXSTMAG-18K
	Mitochondrial Complex I Activity Assay Kit	AAMT001-1KIT
	Mitochondrial Complex II Activity Assay Kit	AAMT002-1KIT
	Mitochondrial Complex IV (Human) Activity Assay Kit	AAMT004-1KIT
	Mitochondrial Complex IV (Mouse) Activity Assay Kit	AAMT006-1KIT
	Mitochondrial Complex V (ATP synthase) Activity Assay Kit	AAMT005-1KIT
	NovaQUANT® Human Mitochondrial to Nuclear DNA Ratio Assay	72620
	NovaQUANT® Human Oxidative Stress qPCR Kit	72627
	NovaQUANT® Mouse Mitochondrial to Nuclear DNARatio Assay	72621
	NovaQUANT® Mouse Oxidative Stress qPCR Kit	72628

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Lit No. RF1101EN00 BS-GEN-15-11491 7/2015

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