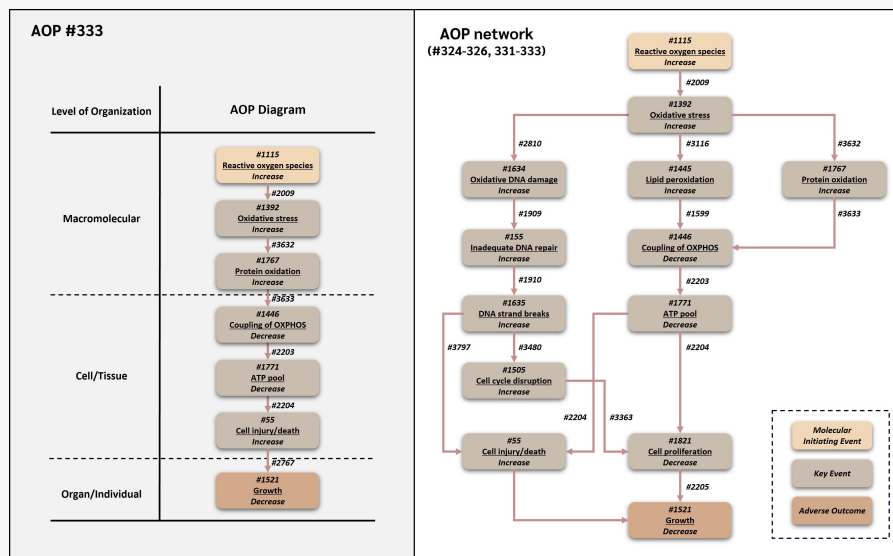


AOP ID and Title:

AOP 333: Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death

Short Title: ROS leading to growth inhibition via protein oxidation and cell death

Graphical Representation



Authors

You Song, Li Xie, Knut Erik Tollefsen

Norwegian Institute for Water Research (NIVA), Sognsveien 72, 0855, Oslo, Norway

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Coaches

Shihori Tanabe

Abstract

This adverse outcome pathway (AOP 333) describes a linear route by which increased reactive oxygen species (ROS) can lead to decreased organismal growth through oxidative stress-mediated protein oxidation, mitochondrial bioenergetic impairment, ATP depletion, and increased cell injury/death. Increased ROS is treated operationally as the molecular initiating event because it represents the earliest common measurable redox perturbation shared by many chemical and non-chemical stressors within the broader ROS-growth AOP network. Increased ROS leads to oxidative stress. When oxidant production exceeds antioxidant and repair capacity, proteins become oxidatively modified through processes such as carbonylation, thiol oxidation, glutathionylation, nitration, fragmentation, aggregation, or altered degradation. Oxidative modification of proteins involved in mitochondrial electron transport, ATP synthase activity, substrate transport, or maintenance of mitochondrial membrane potential can impair coupling of oxidative phosphorylation (OXPHOS). Decreased OXPHOS coupling reduces the cellular ATP pool. ATP depletion compromises ion homeostasis, membrane integrity, stress-response capacity, biosynthesis, and the execution of regulated cell death pathways, thereby increasing cell injury/death. Increased loss of viable cells can reduce tissue or organismal growth, particularly in developing, rapidly growing, or metabolically active systems.

AOP 333 reuses and connects established AOP-Wiki components from several AOP contexts. The upstream ROS and oxidative stress segment is associated with AOP 478, in which energy deposition generates free radicals and oxidative stress and includes oxidative damage to proteins as a downstream consequence (AOP-Wiki, 2026a). The relationship from decreased coupling of OXPHOS to decreased ATP pool is associated with AOP 263, an OECD-published AOP that links OXPHOS uncoupling to growth inhibition through ATP depletion and reduced cell proliferation (AOP-Wiki, 2026b; OECD, 2022; Song and Villeneuve, 2021). AOP 333 differs from AOP 332 by routing ATP depletion through increased cell injury/death rather than through decreased cell proliferation. The cellular injury/death module reuses the broadly shared AOP-Wiki KE 'Increase, Cell injury/death', which occurs in AOPs 12, 13, 17, 38, and 48 in neurotoxicity, oxidative stress, fibrosis, and excitotoxicity contexts (AOP-Wiki, 2026c-g). The AOP is relevant to environmental and human health contexts because ROS generation, oxidative stress responses, protein oxidation, mitochondrial ATP production, cell viability, and growth are broadly conserved among aerobic eukaryotes. The AOP can support mechanistic interpretation of oxidative stress-mediated growth impairment, assay

selection, chemical prioritization, integrated approaches to testing and assessment (IATA), and future quantitative AOP development for oxidative and mitochondrial toxicity.

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AI disclosure

Artificial intelligence (AI) tools were used to support literature prioritization, review and AOP-Wiki page preparation in this work. AOP-helpFinder was used for automated literature mining, and ChatGPT (OpenAI) was used as an auxiliary tool for title and abstract screening, extraction of study metadata, and identification of potential weight-of-evidence indicators. AI-assisted outputs were used only to organize and prioritize information and were verified against the original sources by the authors before inclusion. Additional AI assistance was used for formatting, copy-editing, citation cross-checking, and harmonization of the AOP-Wiki pages. All scientific interpretations, weight-of-evidence judgments, final wording, and conclusions were determined and approved by the authors, who take full responsibility for the content and integrity of the work.

AOP Development Strategy

Context

ROS are continuously formed during aerobic metabolism and can also be generated in response to environmental stressors. At controlled levels, ROS participate in redox signaling, whereas excessive ROS can disturb redox homeostasis and initiate oxidative stress (Schieber and Chandel, 2014; Sies et al., 2017). Proteins are major targets of oxidative attack because amino acid side chains, thiol groups, metal centers, and prosthetic groups can undergo oxidative modification. Protein oxidation can reduce enzyme activity, alter protein-protein interactions, impair folding, increase aggregation, disrupt degradation by proteasomal and lysosomal systems, and contribute to cellular dysfunction (Dalle-Donne et al., 2006).

AOP 333 was developed to represent the protein oxidation and cell injury/death-driven linear route within the broader ROS-growth AOP network. This route was selected because protein oxidation is a well-established consequence of oxidative stress and because mitochondrial proteins are central determinants of OXPHOS coupling and ATP production. Oxidative modification of respiratory-chain proteins, ATP synthase subunits, mitochondrial carriers, or proteins involved in maintaining mitochondrial membrane potential can reduce respiratory efficiency and ATP synthesis (Murphy, 2009; Nicholls and Ferguson, 2013; Sokolov et al., 2019). ATP depletion is an established contributor to loss of cell viability because cell survival depends on ATP-dependent ion gradients, membrane repair, stress responses, proteostasis, and execution of regulated cell death pathways. Severe ATP depletion can shift cellular outcomes toward irreversible injury or necrosis, whereas less severe depletion may permit apoptosis or adaptive responses (Leist et al., 1997; Bonora et al., 2012).

The AOP was designed to maximize reuse of existing AOP-Wiki content. AOP 478 was reviewed because it provides a curated AOP-Wiki context for oxidative stress downstream of free radical generation and includes oxidative molecular damage, including modified proteins, as a relevant consequence of oxidative stress (AOP-Wiki, 2026a). AOP 263 was used to anchor the downstream mitochondrial bioenergetic segment because it provides an OECD-published, well-supported module connecting decreased coupling of OXPHOS with decreased ATP pool and decreased growth-related outcomes (AOP-Wiki, 2026b; OECD, 2022; Song and Villeneuve, 2021). AOPs 12, 13, 17, 38, and 48 were reviewed because they reuse the KE 'Increase, Cell injury/death' and provide evidence that cell injury/death is a generic, reusable cellular response to diverse upstream perturbations. In particular, AOP 17 describes oxidative stress-related developmental neurotoxicity that includes cell injury/death, AOP 38 uses cell injury/death as an early response to protein alkylation in liver fibrosis, and AOP 48 includes mitochondrial dysfunction leading to cell injury/death in an excitotoxicity context (AOP-Wiki, 2026e-g).

Strategy

AOP 333 was developed using the principles described in OECD AOP guidance, including modular description of KEs and KERs, evidence evaluation using biological plausibility, empirical support, essentiality, and quantitative understanding, and clear description of the biological domain of applicability (OECD, 2018, 2021). The development approach combined reuse of existing AOP-Wiki content, targeted literature review, and an AI-human hybrid evidence workflow. The objective was to define a focused linear AOP within the broader ROS-growth AOP network rather than to create an isolated de novo pathway.

The evidence search began with development of event-specific search terms for each KE, including KE names, synonyms, endpoint terms, assay names, stressor terms, taxa, and species names. These terms were used in AOP-helpFinder to search PubMed for co-occurrence of KEs and related mechanistic concepts (Carvaillo et al., 2019; Jornod et al., 2022). AOP-helpFinder outputs, including PMIDs, titles, abstracts, and matched KE terms, were

exported and subjected to overlap analysis to remove redundant hits and filter taxa- or endpoint-irrelevant literature.

The second phase used ChatGPT (OpenAI, San Francisco, CA, USA)-assisted screening to prioritize abstracts and full-text records. The LLM was used as an auxiliary tool to extract study metadata, including stressor, species, biological system, dose or concentration, and exposure time; to classify evidence type, including biological plausibility, empirical support, and essentiality; and to identify weight-of-evidence indicators, including dose-response concordance, temporal concordance, incidence concordance, and intervention or rescue evidence. Studies were categorized as high, medium, or low priority. High-priority studies were retrieved for full-text review, medium-priority studies were reserved as supporting evidence, and low-priority studies were documented but not carried forward for detailed curation.

The final phase consisted of manual expert review and curation. Expert review verified LLM outputs against the full text, extracted evidence into KER evidence tables, and assigned weight-of-evidence calls for biological plausibility, empirical support, essentiality, and quantitative understanding. Targeted manual searches were performed to fill gaps for protein oxidation, mitochondrial bioenergetics, ATP depletion, cell injury/death, and growth outcomes. Studies were prioritized when they measured two or more KEs in the same biological system, reported dose and time information, or supported temporal, dose-response, incidence, or intervention concordance. Mechanistic reviews and OECD reports were used to support biological plausibility where relationships are widely established, whereas primary experimental studies were prioritized for empirical support where available.

Summary of the AOP

Events

Molecular Initiating Events (MIE), Key Events (KE), Adverse Outcomes (AO)

Sequence	Type	Event ID	Title	Short name
	MIE	1115	Increase, Reactive oxygen species	Increase, ROS
	KE	1392	Increase, Oxidative Stress	Increase, Oxidative Stress
	KE	1767	Increase, Protein oxidation	Increase, Protein oxidation
	KE	1446	Decrease, Coupling of oxidative phosphorylation	Decrease, Coupling of OXPHOS
	KE	1771	Decrease, Adenosine triphosphate pool	Decrease, ATP pool
	KE	55	Increase, Cell injury/death	Cell injury/death
	AO	1521	Decrease, Growth	Decrease, Growth

Key Event Relationships

Upstream Event	Relationship Type	Downstream Event	Evidence	Quantitative Understanding
Increase, Reactive oxygen species	adjacent	Increase, Oxidative Stress	High	Moderate
Increase, Oxidative Stress	adjacent	Increase, Protein oxidation	High	Moderate
Increase, Protein oxidation	adjacent	Decrease, Coupling of oxidative phosphorylation	Moderate	Low
Decrease, Coupling of oxidative phosphorylation	adjacent	Decrease, Adenosine triphosphate pool	High	High
Decrease, Adenosine triphosphate pool	adjacent	Increase, Cell injury/death	High	Moderate
Increase, Cell injury/death	adjacent	Decrease, Growth	High	Moderate

Stressors

Name	Evidence
Hydrogen peroxide	
tert-Butyl hydroperoxide	
Paraquat	
Heavy metals (cadmium, lead, copper, iron, nickel)	
Silver	
Silver nanoparticles	

Name	Evidence
Ionizing Radiation	
Ultraviolet B radiation	

Overall Assessment of the AOP

The overall weight of evidence supporting AOP 333 is considered moderate. Biological plausibility is high for all six KERs in the pathway. The mechanistic connections between oxidative stress, protein oxidation, impaired mitochondrial OXPHOS coupling, ATP depletion, cell injury/death, and decreased growth are individually well supported, and the AOP draws on conserved biological processes broadly applicable across aerobic eukaryotes. The central OXPHOS-to-ATP segment is directly associated with OECD-endorsed AOP 263, providing strong mechanistic and quantitative support for this portion of the pathway (OECD, 2022; Song and Villeneuve, 2021). The cell injury/death KE (Event 55) is a widely reused and modular AOP-Wiki element present in endorsed AOPs 12, 13, 17, 38, and 48, reinforcing the credibility of its use as a downstream consequence of severe energetic failure (AOP-Wiki, 2026a-e). Empirical support is high for the ROS-to-oxidative-stress and oxidative-stress-to-protein-oxidation relationships and moderate for the protein-oxidation-to-OXPHOS transition, where supporting evidence is observational and cross-stressor rather than from controlled selective-inhibition studies. The ATP-depletion-to-cell-death and cell-death-to-growth KERs have moderate empirical support. Essentiality is high for the OXPHOS-to-ATP relationship and moderate for the remaining KERs. Quantitative understanding is strongest for the OXPHOS-to-ATP KER and low to moderate elsewhere, reflecting the difficulty of predicting organism-level growth outcomes from upstream molecular damage endpoints. The main uncertainties are the causal versus correlational character of the protein oxidation-OXPHOS association, the ATP threshold dependence of cell death mode and severity, and the multifactorial nature of organismal growth as an apical endpoint. AOP 333 is currently most suitable for qualitative and semi-quantitative use in mechanistic interpretation, hazard identification, and support for integrated testing and assessment strategies (OECD, 2018; Becker et al., 2015).

Domain of Applicability

Life Stage Applicability

Life Stage	Evidence
All life stages	Moderate

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
humans	Homo sapiens	Moderate	NCBI
mammals	mammals	Moderate	NCBI
fish	fish	Moderate	NCBI
crustaceans	Daphnia magna	Moderate	NCBI
green algae	Ulva compressa	Moderate	NCBI

Sex Applicability

Sex	Evidence
Unspecific	Moderate

The domain of applicability for AOP 333 is broad across aerobic eukaryotic organisms in which ROS generation, oxidative stress responses, protein oxidation, mitochondrial oxidative phosphorylation, ATP-dependent homeostasis, cell injury/death, and growth are biologically relevant. The AOP is most applicable to biological contexts in which increased ROS is sufficient to induce oxidative protein damage and where mitochondrial ATP production is important for cellular survival and growth.

The stressor domain includes direct ROS generators, redox-cycling chemicals, metals, nanoparticles, mitochondrial toxicants, hypoxia-reoxygenation, radiation, and inflammatory or pathogen-related stressors. Because the MIE is defined operationally as increased ROS rather than as a stressor-specific molecular interaction, AOP 333 should be applied with attention to evidence that the stressor actually induces oxidative stress and protein oxidation in the biological context under evaluation.

Essentiality of the Key Events

Essentiality is evaluated for the overall AOP based on whether preventing or modifying upstream KERs changes downstream KERs or the AO. Direct essentiality evidence is strongest for the OXPHOS to ATP relationship and for ATP dependence of cell viability. Essentiality for protein oxidation is biologically plausible but less directly demonstrated because selective prevention of protein oxidation without altering other oxidative stress processes is experimentally difficult.

Key event	Essentiality	Rationale	Experimental manipulation evidence (KE knock-out / inhibition / rescue)	Uncertainties
Event 1115: Reactive oxygen species, increased	Moderate	ROS are causally linked to oxidative stress because oxidative stress occurs when oxidant formation exceeds antioxidant capacity. Antioxidant and radical-scavenging interventions can reduce oxidative stress and downstream oxidative damage in many systems, supporting the importance of ROS as an upstream driver (Schieber and Chandel, 2014; Sies et al., 2017).	Indirect (stop/attenuation): antioxidant and ROS-scavenger pre-treatment reduces oxidative stress and downstream damage across oxidative-stress models (Schieber and Chandel, 2014; Sies et al., 2017). No selective single-source ROS knock-out is available.	ROS can also function in physiological signaling at low levels; oxidative stress can be sustained by altered antioxidant capacity even when a specific ROS source is removed.
Event 1392: Oxidative stress, increased	Moderate to high	Antioxidant interventions commonly reduce downstream oxidative damage and cell injury. Oxidative stress downstream of radical generation is represented in AOP 478 (AOP-Wiki, 2026a; Schieber and Chandel, 2014; Sies et al., 2017).	Indirect: modulation of antioxidant capacity alters progression to oxidative macromolecular damage; oxidative stress is the curated hub KE in endorsed AOP 478 (AOP-Wiki, 2026a; Carrothers et al., 2025).	Oxidative stress can be adaptive at low levels and harmful at higher intensity or duration.

Event 1767: Protein oxidation, increased	Moderate	Protein carbonylation and other oxidative modifications impair protein function and can contribute to mitochondrial dysfunction (Dalle-Donne et al., 2006). Cadmium-induced protein carbonylation and actin glutathionylation were reduced by oxidase/NOS inhibitors in mussel hemocytes (Canesi et al., 2010).	Direct (partial): cadmium-induced protein carbonylation and actin glutathionylation reduced by oxidase/NOS inhibitors in mussel hemocytes (Canesi et al., 2010); GSTA4 silencing raised mitochondrial protein carbonylation and target knockdown reduced respiration (Curtis et al., 2012).	Selective rescue of protein oxidation alone is uncommon; protein oxidation can be both causal and a marker of broader damage.
Event 1446: Coupling of OXPHOS, decreased	High	This KE is associated with AOP 263. Recovery of OXPHOS coupling can restore mitochondrial function and ATP production, supporting essentiality of this KE for downstream ATP depletion (AOP-Wiki, 2026b; OECD, 2022; Song and Villeneuve, 2021).	Direct (rescue): removal of uncouplers or restoration of coupling recovers mitochondrial membrane potential and ATP in the endorsed AOP 263 module (AOP-Wiki, 2026b; OECD, 2022; Song and Villeneuve, 2021).	Mild uncoupling may sometimes be adaptive and reduce ROS production.
Event 1771: ATP pool, decreased	High	ATP depletion is directly linked to loss of cell viability and can influence the mode of cell death. Intracellular ATP concentration can act as a switch between apoptosis and necrosis (Leist et al., 1997).	Indirect: ATP-restoration experiments reduce downstream injury/proliferation deficits; central KE in endorsed AOP 263 (Leist et al., 1997; Nicotera et al., 1998; OECD, 2022).	Cells may compensate through glycolysis or altered energy allocation.
Event 55: Cell injury/death, increased	Moderate	Cell injury/death is a shared KE used in AOPs 12, 13, 17, 38, and 48. Loss of viable cells provides a plausible mechanism for reduced growth and tissue function (AOP-Wiki, 2026c-g).	Indirect: ATP restoration/maintenance reduces injury in some systems, indicating energy-status dependence (Leist et al., 1997; Nicotera et al., 1998); widely reused modular KE (AOPs 12, 13, 17, 38, 48).	Growth can also decline through reduced proliferation, altered cell size, or developmental delay without overt cell death.

Event 1521: Growth, decreased (AO)	Not applicable (AO)	Growth is the adverse outcome and a regulatory-relevant endpoint across multiple taxa. AOP 263 provides precedent for decreased growth as an AO downstream of mitochondrial bioenergetic impairment (AOP-Wiki, 2026b; OECD, 2022; Song and Villeneuve, 2021).	As the adverse outcome, essentiality is assessed for upstream KEs; AOP 263 provides precedent for decreased growth as an AO downstream of these modules (OECD, 2022; Song and Villeneuve, 2021).	Growth is integrative and may arise from multiple mechanisms.
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Weight of Evidence Summary

Evidence assessment is organized by KER. Calls follow OECD weight-of-evidence considerations for biological plausibility, empirical support, and quantitative understanding (OECD, 2018, 2021).

Biological plausibility of KERs

KER	Biological plausibility call	Rationale
2009: ROS increase leads to oxidative stress increase	High	Oxidative stress reflects an imbalance between oxidant production and antioxidant capacity, and ROS are primary oxidant species in cellular redox biology (Schieber and Chandel, 2014; Sies et al., 2017). AOP 478 supports oxidative stress downstream of free radical generation (AOP-Wiki, 2026a).
3632: oxidative stress increase leads to protein oxidation increase	High	ROS and related oxidants can modify amino acid side chains, thiols, metal centers, and prosthetic groups, producing carbonylated, glutathionylated, misfolded, aggregated, or degraded proteins (Dalle-Donne et al., 2006; Sies et al., 2017).
3633: protein oxidation increase leads to decreased coupling of OXPHOS	Moderate to high	Mitochondrial OXPHOS depends on intact electron transport complexes, ATP synthase, metabolite carriers, and membrane-associated protein assemblies. Oxidative modification of these proteins can impair electron transfer, proton pumping, membrane potential, and ATP synthesis efficiency (Murphy, 2009; Nicholls and Ferguson, 2013; Sokolov et al., 2019).
2203: decreased coupling of OXPHOS leads to decreased ATP pool	High	This relationship is associated with AOP 263. OXPHOS coupling is a major determinant of ATP production in aerobic eukaryotic cells; reduced coupling lowers ATP synthesis efficiency (AOP-Wiki, 2026b; OECD, 2022; Song and Villeneuve, 2021).
2768: decreased ATP pool leads to increased cell injury/death	High	ATP is required for survival, ion homeostasis, membrane repair, proteostasis, and regulated death processes. Severe ATP depletion can switch cellular outcomes toward necrosis or irreversible injury (Leist et al., 1997; Bonora et al., 2012).

2767: increased cell injury/death leads to decreased growth	High	Growth depends on viable cell number, tissue integrity, and biomass accumulation. Increased cell injury/death reduces the cellular basis for growth and is reused across AOPs 12, 13, 17, 38, and 48 (AOP-Wiki, 2026c-g; Conlon and Raff, 1999).
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Empirical support for KERs

KER	Empirical support call	Rationale	Inconsistencies or evidence gaps
2009: ROS increase leads to oxidative stress increase	High	Paraquat increased ROS and antioxidant enzyme responses in <i>Chlorella vulgaris</i> (Qian et al., 2009). Infection-induced ROS coincided with antioxidant and inflammatory responses in golden pompano (Gao et al., 2022). AOP 478 reports evidence linking free radical generation/energy deposition to oxidative stress (AOP-Wiki, 2026a).	ROS is transient and often measured indirectly; oxidative stress biomarkers vary by assay and taxa.
3632: oxidative stress increase leads to protein oxidation increase	High	Oxidative stressors increase protein carbonyls or related protein oxidation endpoints. Cadmium and hydrogen peroxide increased protein carbonylation and redox modification in <i>Chlamydomonas</i> systems (Zaffagnini et al., 2012). Cadmium induced protein carbonylation and actin glutathionylation in mussel hemocytes (Canesi et al., 2010). Thermal stress in zebrafish increased protein carbonyls with antioxidant responses (Tseng et al., 2011).	Protein oxidation endpoints are heterogeneous; some studies measure total carbonyls whereas others identify specific oxidized proteins.
3633: protein oxidation increase leads to decreased coupling of OXPHOS	Moderate	Evidence links oxidative protein damage or mitochondrial proteome modification with altered mitochondrial function. Age-associated oxidative changes in zebrafish were associated with changes in mitochondrial oxidative status and aconitase activity (Almáida-Pagán et al., 2014). Hypoxia-reoxygenation altered mitochondrial proteome and bioenergetics in <i>Crassostrea gigas</i> (Sokolov et al., 2019).	Many studies measure correlation rather than direct causation; protein oxidation may occur alongside lipid peroxidation or other mitochondrial damage.
2203: decreased coupling of OXPHOS leads to decreased ATP pool	High	AOP 263 reports strong evidence for this KER (AOP-Wiki, 2026b; OECD, 2022; Song and Villeneuve, 2021). Cadmium exposure in oysters reduced state 3 respiration and affected mitochondrial bioenergetics (Sokolova et al., 2005).	Compensatory glycolysis and altered metabolic demand can obscure total ATP changes.
2768: decreased ATP pool leads to increased cell injury/death	Moderate to high	ATP depletion and cell death are linked in multiple systems. Intracellular ATP concentration influences the decision between apoptosis and necrosis (Leist et al., 1997). Calcium electroporation caused dose-dependent ATP depletion and cancer cell death (Hansen et al., 2015).	ATP assays may reflect both energy state and cell number; direct temporal separation of ATP depletion from cell death is needed.

2767: increased cell injury/death leads to decreased growth	Moderate	Cell injury/death is reused as a KE in several established AOPs (AOP-Wiki, 2026c-g). Methanol-exposed mouse embryos showed growth reduction and elevated cell death (Abbott et al., 1995). In bivalves, cadmium and temperature interactions caused cellular energy disruption, mortality, and reduced condition/growth-related outcomes (Cherkasov et al., 2006).	Growth can be reduced by mechanisms other than cell death; direct dose/time concordance between cell death and growth is not always measured.
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Inconsistencies and uncertainties

The main uncertainty for AOP 333 is the quantitative strength and directionality of the protein oxidation to OXPHOS coupling relationship. Protein oxidation can impair mitochondrial enzymes and respiratory complexes, but mitochondrial dysfunction can also enhance ROS generation and thereby increase protein oxidation. AOP 333 represents one biologically plausible and empirically supported direction within a broader feedback-prone network. Another uncertainty is that ATP depletion can lead to different cellular outcomes depending on severity and duration; moderate depletion may reduce proliferation or activate adaptive stress responses, whereas severe depletion promotes cell injury/death. Finally, growth is a multifactorial endpoint. Increased cell injury/death is an important contributor to impaired growth, but decreased growth can also arise through reduced proliferation, altered cell size, altered energy allocation, endocrine signaling, or developmental delay without overt cell death.

Quantitative Consideration

Quantitative understanding varies across the AOP. The relationship between OXPHOS coupling and ATP production has the strongest quantitative foundation, while the relationships linking oxidative stress to protein oxidation and cell injury/death to organismal growth are more often qualitative or semi-quantitative.

KER	Quantitative understanding call	Rationale
2009: ROS increase to oxidative stress increase	Low to moderate	ROS measurements are reactive, transient, and assay-dependent. Quantitative relationships can be defined within a specific assay, but generalizable prediction across taxa and stressors remains limited (Sies et al., 2017).
3632: oxidative stress increase to protein oxidation increase	Moderate	Protein carbonyl assays and redox proteomics provide quantitative measures of protein oxidation, but response-response relationships are not broadly generalizable across stressors or taxa (Dalle-Donne et al., 2006).
3633: protein oxidation increase to decreased OXPHOS coupling	Low to moderate	Specific oxidation of mitochondrial proteins can be associated with altered mitochondrial function, but predictive quantitative models are not yet established across taxa or stressors (Sokolov et al., 2019).
2203: decreased OXPHOS coupling to decreased ATP pool	High	AOP 263 reports strong quantitative understanding, supported by bioenergetic theory and experimental response-response relationships (AOP-Wiki, 2026b; OECD, 2022; Song and Villeneuve, 2021).
2768: decreased ATP pool to increased cell injury/death	Moderate	ATP thresholds influence the type and severity of cell death, and quantitative relationships are reported in defined systems, but thresholds vary by cell type and exposure condition (Leist et al., 1997; Hansen et al., 2015).

2767: increased cell injury/death to decreased growth	Low to moderate	Quantitative linkage between cell loss and organismal growth is plausible and can be modeled in defined systems, but empirical cross-taxa response-response relationships remain limited (Conlon and Raff, 1999).
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BMD/POD-anchored concordance

The following benchmark-dose/point-of-departure (BMD/POD) concordance table anchors AOP 333 to quantitative cross-KE ordering, in line with Handbook section 4C. The multiomics point-of-departure (moPOD) dataset for gamma-irradiated *Daphnia magna* (Song et al., 2023) provides POD magnitudes for increased ROS, decreased ATP, decreased OXPHOS coupling, and cell death, demonstrating the expected upstream-to-downstream POD ordering (more sensitive PODs upstream). The moPOD is presented as POD magnitude evidence, not as a causal re-ordering of KEs. The Lemna minor EDR50 range provides a whole-pathway apical anchor in an aquatic primary producer.

Key event (functional category)	POD metric	POD value (mGy/h)	POD ordering	Source
KE 1115: ROS, increased (mROS)	moPOD (multiomics POD)	0.4	1 (most sensitive)	Song et al., 2023
KE 1771: ATP pool, decreased	moPOD	2.5	2	Song et al., 2023
KE 1446: OXPHOS coupling, decreased (UPS/OXPHOS module)	moPOD	42.3	3	Song et al., 2023
KE 55: Cell injury/death (apoptosis)	moPOD	42.3	3 (least sensitive)	Song et al., 2023
Upstream KE chain → growth (Lemna minor, gamma)	EDR50 (growth)	31.5–54.8 (mGy/h)	whole-pathway apical	Xie et al., 2018, 2019, 2022

Considerations for Potential Applications of the AOP (optional)

AOP 333 can support mechanistic interpretation of growth impairment caused by oxidative stressors that induce protein oxidation, mitochondrial bioenergetic dysfunction, ATP depletion, and cell injury/death. The AOP is particularly relevant for hazard identification and chemical prioritization when evidence indicates increased ROS or oxidative stress together with protein carbonylation, redox proteomic signatures, mitochondrial membrane potential changes, reduced respiratory control, ATP depletion, cytotoxicity, or growth inhibition. The AOP may also support IATA development by linking upstream NAM endpoints, such as ROS assays, oxidative stress biomarkers, protein carbonyl assays, redox proteomics, mitochondrial membrane potential, oxygen consumption rate, ATP content, cytotoxicity assays, and organismal growth measurements.

AOP 333 can support chemical grouping and read-across for stressors that share evidence of oxidative protein damage, mitochondrial bioenergetic impairment, and ATP-associated cell injury. Because oxidative stress and protein oxidation are not chemical-specific, this AOP should not be used as a stand-alone basis for regulatory decisions. Instead, it should be applied as part of a weight-of-evidence framework that considers stressor mode of action, exposure context, assay specificity, taxonomic relevance, and concordance across multiple KEs. The AOP also highlights method-development needs, particularly standardized assays for protein oxidation, OXPHOS coupling, ATP depletion, and cell injury/death endpoints that can be connected quantitatively to apical growth outcomes.

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Appendix 1

List of MIEs in this AOP

Event: 1115: Increase, Reactive oxygen species

Short Name: Increase, ROS

Event Component

Process	Object	Action
reactive oxygen species biosynthetic process	reactive oxygen species	increased

AOPs Including This Key Event

AOP ID and Name	Event Type
Aop:186 - unknown MIE leading to renal failure and mortality	KeyEvent
Aop:213 - Inhibition of fatty acid beta oxidation leading to nonalcoholic steatohepatitis (NASH)	KeyEvent
Aop:303 - Frustrated phagocytosis-induced lung cancer	KeyEvent
Aop:383 - Inhibition of Angiotensin-converting enzyme 2 leading to liver fibrosis	KeyEvent
Aop:382 - Angiotensin II type 1 receptor (AT1R) agonism leading to lung fibrosis	KeyEvent
Aop:384 - Hyperactivation of ACE/Ang-II/AT1R axis leading to chronic kidney disease	KeyEvent

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AOP ID and Name	Event Type
Aop:396 - Deposition of ionizing energy leads to population decline via impaired meiosis	KeyEvent
Aop:409 - Frustrated phagocytosis leads to malignant mesothelioma	KeyEvent
Aop:413 - Oxidation and antagonism of reduced glutathione leading to mortality via acute renal failure	KeyEvent
Aop:416 - Aryl hydrocarbon receptor activation leading to lung cancer through IL-6 toxicity pathway	KeyEvent
Aop:418 - Aryl hydrocarbon receptor activation leading to impaired lung function through AHR-ARNT toxicity pathway	KeyEvent
Aop:386 - Deposition of ionizing energy leading to population decline via inhibition of photosynthesis	KeyEvent
Aop:387 - Deposition of ionising energy leading to population decline via mitochondrial dysfunction	KeyEvent
Aop:319 - Binding to ACE2 leading to lung fibrosis	KeyEvent
Aop:451 - Interaction with lung resident cell membrane components leads to lung cancer	KeyEvent
Aop:476 - Adverse Outcome Pathways diagram related to PBDEs associated male reproductive toxicity	MolecularInitiatingEvent
Aop:492 - Glutathione conjugation leading to reproductive dysfunction via oxidative stress	KeyEvent
Aop:497 - ERα inactivation alters mitochondrial functions and insulin signalling in skeletal muscle and leads to insulin resistance and metabolic syndrome	KeyEvent
Aop:500 - Activation of MEK-ERK1/2 leads to deficits in learning and cognition via ROS and apoptosis	KeyEvent
Aop:505 - Reactive Oxygen Species (ROS) formation leads to cancer via inflammation pathway	MolecularInitiatingEvent
Aop:513 - Reactive Oxygen (ROS) formation leads to cancer via Peroxisome proliferation-activated receptor (PPAR) pathway	MolecularInitiatingEvent
Aop:521 - Essential element imbalance leads to reproductive failure via oxidative stress	KeyEvent
Aop:540 - Oxidative Stress in the Fish Ovary Leads to Reproductive Impairment via Reduced Vitellogenin Production	MolecularInitiatingEvent
Aop:462 - Activation of reactive oxygen species leading the atherosclerosis	MolecularInitiatingEvent
Aop:299 - Deposition of energy leading to population decline via DNA oxidation and follicular atresia	KeyEvent
Aop:311 - Deposition of energy leading to population decline via DNA oxidation and oocyte apoptosis	KeyEvent
Aop:331 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and cell death	MolecularInitiatingEvent
Aop:327 - Excessive reactive oxygen species production leading to mortality (1)	MolecularInitiatingEvent
Aop:328 - Excessive reactive oxygen species production leading to mortality (2)	MolecularInitiatingEvent
Aop:329 - Excessive reactive oxygen species production leading to mortality (3)	MolecularInitiatingEvent
Aop:330 - Excessive reactive oxygen species production leading to mortality (4)	MolecularInitiatingEvent
Aop:26 - Calcium-mediated neuronal ROS production and energy imbalance	KeyEvent
Aop:534 - Succinate dehydrogenase (SDH) inhibition leads to oxidative stress	KeyEvent
Aop:273 - Mitochondrial complex inhibition leading to liver injury	KeyEvent
Aop:488 - Increased reactive oxygen species production leading to decreased cognitive function	MolecularInitiatingEvent
Aop:298 - Increase in reactive oxygen species (ROS) leading to human treatment-resistant gastric cancer	MolecularInitiatingEvent
Aop:27 - Cholestatic Liver Injury induced by Inhibition of the Bile Salt Export Pump (ABCB11)	KeyEvent
Aop:511 - The AOP framework on ROS-mediated oxidative stress induced vascular disrupting effects	MolecularInitiatingEvent
Aop:207 - NADPH oxidase and P38 MAPK activation leading to reproductive failure in Caenorhabditis elegans	KeyEvent
Aop:423 - Toxicological mechanisms of hepatocyte apoptosis through the PARP1 dependent cell death pathway	MolecularInitiatingEvent
Aop:481 - AOPs of amorphous silica nanoparticles: ROS-mediated oxidative stress increased respiratory dysfunction and diseases.	MolecularInitiatingEvent
Aop:282 - Adverse outcome pathway on photochemical toxicity initiated by light exposure	MolecularInitiatingEvent
Aop:569 - Decreased DNA methylation of FAM50B/PTCHD3 leading to IQ loss of children via PI3K-Akt pathway	KeyEvent
Aop:595 - Emerging OPFRS reproductive outcome pathway	MolecularInitiatingEvent
Aop:596 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	MolecularInitiatingEvent
Aop:598 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and reduced cell proliferation	MolecularInitiatingEvent
Aop:599 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and cell injury/death	MolecularInitiatingEvent

AOP ID and Name	Event Type
Aop:600 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell growth	MolecularInitiatingEvent
Aop:601 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell proliferation	MolecularInitiatingEvent
Aop:602 - Excessive reactive oxygen species leading to growth inhibition via oxidative DNA damage	MolecularInitiatingEvent
Aop:603 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell cycle disruption	MolecularInitiatingEvent
Aop:608 - Thyroid Hormone Excess Leading to Reduced, Swimming Performance via Hypomyelination	KeyEvent
Aop:613 - Peroxisome proliferator-activated receptor alpha activation leading to early life stage mortality via increased reactive oxygen species production	KeyEvent
Aop:622 - Calcineurin inhibitor induced nephrotoxicity leading to kidney failure	KeyEvent
Aop:636 - Increase in reactive oxygen species (ROS) leading to human amyotrophic lateral sclerosis (ALS)	MolecularInitiatingEvent
Aop:638 - Co-exposure to microplastics and cadmium leading to progression from NAFLD to liver tumorigenesis	MolecularInitiatingEvent
Aop:472 - DNA adduct formation leading to kidney failure	KeyEvent
Aop:324 - Increase in reactive oxygen species leading to growth inhibition via oxidative DNA damage and cell cycle disruption	MolecularInitiatingEvent
Aop:325 - Increase in reactive oxygen species leading to growth inhibition via oxidative DNA damage and cell death	MolecularInitiatingEvent
Aop:326 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and decreased cell proliferation	MolecularInitiatingEvent
Aop:332 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and decreased cell proliferation	MolecularInitiatingEvent
Aop:333 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	MolecularInitiatingEvent
Aop:647 - ROS formation leading to growth inhibition modulated by the Keap1-Nrf2 antioxidant response	MolecularInitiatingEvent

Biological Context

Level of Biological Organization

Cellular

Cell term

Cell term

cell

Organ term

Organ term

organ

Domain of Applicability

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
Vertebrates	Vertebrates	High	NCBI
human	Homo sapiens	Moderate	NCBI
human and other cells in culture	human and other cells in culture	Moderate	NCBI
mouse	Mus musculus	Moderate	NCBI
crustaceans	Daphnia magna	High	NCBI
Lemna minor	Lemna minor	High	NCBI
zebrafish	Danio rerio	High	NCBI

Life Stage Applicability

Life Stage Evidence

Life Stage Evidence

All life stages High

Sex Applicability**Sex Evidence**

Unspecific High

Mixed High

ROS is a normal constituent found in all organisms, *lifestages*, and *sexes*.

Key Event Description

Biological State: increased reactive oxygen species (ROS)

Biological compartment: an entire cell -- may be cytosolic, may also enter organelles.

Reactive oxygen species (ROS) are O_2^- -derived molecules that can be both free radicals (e.g. superoxide, hydroxyl, peroxy, alkoxy) and non-radicals (hypochlorous acid, ozone and singlet oxygen) (Bedard and Krause 2007; Ozcan and Ogun 2015). ROS production occurs naturally in all kinds of tissues inside various cellular compartments, such as mitochondria and peroxisomes (Drew and Leeuwenburgh 2002; Ozcan and Ogun 2015). Furthermore, these molecules have an important function in the regulation of several biological processes – they might act as antimicrobial agents or triggers of animal gamete activation and capacitation (Goud et al. 2008; Parrish 2010; Bisht et al. 2017).

However, in environmental stress situations (exposure to radiation, chemicals, high temperatures) these molecules have its levels drastically increased, and overly interact with macromolecules, namely nucleic acids, proteins, carbohydrates and lipids, causing cell and tissue damage (Brieger et al. 2012; Ozcan and Ogun 2015).

Reactive oxygen species (ROS) refers to the chemical species superoxide, hydrogen peroxide, and their secondary reactive products. In the biological context, ROS are signaling molecules with important roles in cell energy metabolism, cell proliferation, and fate. Therefore, balancing ROS levels at the cellular and tissue level is an important part of many biological processes. Disbalance, mainly an increase in ROS levels, can cause cell dysfunction and irreversible cell damage.

ROS are produced from both exogenous stressors and normal endogenous cellular processes, such as the mitochondrial electron transport chain (ETC). Inhibition of the ETC can result in the accumulation of ROS. Exposure to chemicals, heavy metal ions, or ionizing radiation can also result in increased production of ROS. Chemicals and heavy metal ions can deplete cellular antioxidants reducing the cell's ability to control cellular ROS and resulting in the accumulation of ROS. Cellular antioxidants include glutathione (GSH), protein sulfhydryl groups, superoxide dismutase (SOD).

ROS are radicals, ions, or molecules that have a single unpaired electron in their outermost shell of electrons, which can be categorized into two groups: free oxygen radicals and non-radical ROS [Liou et al., 2010].

<Free oxygen radicals>

superoxide	$O_2^{\cdot -}$
hydroxyl radical	$\cdot OH$
nitric oxide	$NO\cdot$
organic radicals	$R\cdot$
peroxy radicals	$ROO\cdot$
alkoxy radicals	$RO\cdot$
thiyl radicals	$RS\cdot$
sulfonyl radicals	$ROS\cdot$
thiyl peroxy radicals	$RSOO\cdot$
disulfides	$RSSR$

<Non-radical ROS>

hydrogen peroxide	H_2O_2
singlet oxygen	1O_2
ozone/trioxygen	O_3
organic hydroperoxides	$ROOH$
hypochlorite	ClO^-
peroxynitrite	$ONOO^-$
nitrosoperoxy carbonate anion	$O=NOOCO_2^-$
nitrocarbonate anion	$O_2NOCO_2^-$
dinitrogen dioxide	N_2O_2
nitronium	NO_2^+
highly reactive lipid- or carbohydrate-derived carbonyl compounds	

Potential sources of ROS include NADPH oxidase, xanthine oxidase, mitochondria, nitric oxide synthase, cytochrome P450, lipoxygenase/cyclooxygenase, and monoamine oxidase [Granger et al., 2015]. ROS are generated through NADPH oxidases consisting of $p47^{phox}$ and $p67^{phox}$. ROS are generated through xanthine oxidase activation in sepsis [Ramos et al., 2018]. Arsenic produces ROS [Zhang et al., 2011]. Mitochondria-targeted paraquat and metformin mediate ROS production [Chowdhury et al., 2020]. ROS are generated by bleomycin [Lu et al., 2010]. Radiation induces dose-dependent ROS production [Ji et al., 2019].

ROS are generated in the course of cellular respiration, metabolism, cell signaling, and inflammation [Dickinson and Chang 2011; Egea et al. 2017]. Hydrogen peroxide is also made by the endoplasmic reticulum in the course of protein folding. Nitric oxide (NO) is produced at the highest levels by nitric oxide synthase in endothelial cells and phagocytes. NO production is one of the main mechanisms by which phagocytes kill bacteria [Wang et al., 2017]. The other species are produced by reactions with superoxide or peroxide, or by other free radicals or enzymes.

ROS activity is principally local. Most ROS have short half-lives, ranging from nano- to milliseconds, so diffusion is limited, while reactive nitrogen species (RNS) nitric oxide or peroxynitrite can survive long enough to diffuse across membranes [Calcerrada et al. 2011]. Consequently, local concentrations of ROS are much higher than average cellular concentrations, and signaling is typically controlled by colocalization with redox buffers [Dickinson and Chang 2011; Egea et al. 2017].

Although their existence is limited temporally and spatially, ROS interact with other ROS or with other nearby molecules to produce more ROS and participate in a feedback loop to amplify the ROS signal, which can increase RNS. Both ROS and RNS also move into neighboring cells, and ROS can increase intracellular ROS signaling in neighboring cells [Egea et al. 2017].

In the primary event, photoreactive chemicals are excited by the absorption of photon energy. The energy of the photoactivated chemicals transfer to oxygen and then generates the reactive oxygen species (ROS), including superoxide (O_2^-) via type I reaction and singlet oxygen (1O_2) via type II reaction, as principal intermediate species in phototoxic reaction [Foote, 1991, Onoue et al. , 2009].

How it is Measured or Detected

Photocolorimetric assays [Sharma et al. 2017; Griendling et al. 2016] or through commercial kits purchased from specialized companies.

Yuan, Yan, et al., (2013) described ROS monitoring by using H_2 -DCF-DA, a redox-sensitive fluorescent dye. Briefly, the harvested cells were incubated with H_2 -DCF-DA (50 μ mol/L final concentration) for 30 min in the dark at 37°C. After treatment, cells were immediately washed twice, re-suspended in PBS, and analyzed on a BD-FACS Aria flow cytometry. ROS generation was based on fluorescent intensity which was recorded by excitation at 504 nm and emission at 529 nm.

Lipid peroxidation (LPO) can be measured as an indicator of oxidative stress damage Yen, Cheng Chien, et al., (2013).

Chattopadhyay, Sukumar, et al. (2002) assayed the generation of free radicals within the cells and their extracellular release in the medium by addition of yellow NBT salt solution [Park et al., 1968]. Extracellular release of ROS converted NBT to a purple colored formazan. The cells were incubated with 100 ml of 1 mg/ml NBT solution for 1 h at 37 °C and the product formed was assayed at 550 nm in an Anthos 2001 plate reader. The observations of the 'cell-free system' were confirmed by cytological examination of parallel set of explants stained with chromogenic reactions for NO and ROS.

On the basis of the pathogenesis of drug-induced phototoxicity, a reactive oxygen species (ROS) assay was proposed to evaluate the phototoxic risk of chemicals. The ROS assay can monitor generation of ROS, such as singlet oxygen and superoxide, from photoirradiated chemicals, and the ROS data can be used to evaluate the photoreactivity of chemicals [Onoue et al. , 2014, Onoue et al. , 2013, Onoue and Tsuda, 2006]. The ROS assay is a recommended approach by guidelines to evaluate the phototoxic risk of chemicals [ICH, 2014, PCPC, 2014].

<Direct detection>

Many fluorescent compounds can be used to detect ROS, some of which are specific, and others are less specific.

□ ROS can be detected by fluorescent probes such as *p*-methoxy-phenol derivative [Ashoka et al., 2020].

□ Chemiluminescence analysis can detect the superoxide, where some probes have a wider range for detecting hydroxyl radical, hydrogen peroxide, and peroxynitrite [Fuloria et al., 2021].

□ ROS in the blood can be detected using superparamagnetic iron oxide nanoparticles (SPION)-based biosensor [Lee et al., 2020].

□ Hydrogen peroxide (H_2O_2) can be detected with a colorimetric probe, which reacts with H_2O_2 in a 1:1 stoichiometry to produce a bright pink colored product, followed by the detection with a standard colorimetric microplate reader with a filter in the 540-570 nm range.

□ The levels of ROS can be quantified using multiple-step amperometry using a stainless steel counter electrode and non-leak Ag|AgCl reference node [Flaherty et al., 2017].

□ Singlet oxygen can be measured by monitoring the bleaching of *p*-nitrosodimethylaniline at 440 nm using a spectrophotometer with imidazole as a selective acceptor of singlet oxygen [Onoue et al., 2014].

<Indirect Detection>

Alternative methods involve the detection of redox-dependent changes to cellular constituents such as proteins, DNA, lipids, or glutathione [Dickinson and Chang 2011; Wang et al. 2013; Griendling et al. 2016]. However, these methods cannot generally distinguish between the oxidative species behind the changes and cannot provide good resolution for the kinetics of oxidative activity.

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List of Key Events in the AOP

[Event: 1392: Increase, Oxidative Stress](#)

Short Name: Increase, Oxidative Stress

Event Component

Process	Object	Action
oxidative stress		increased

AOPs Including This Key Event

AOP ID and Name	Event Type
Aop:220 - Cyp2E1 Activation Leading to Liver Cancer	KeyEvent

AOP333

AOP ID and Name	Event Type
Aop:17 - Binding of electrophilic chemicals to SH(thiol)-group of proteins and /or to seleno-proteins involved in protection against oxidative stress during brain development leads to impairment of learning and memory	KeyEvent
Aop:284 - Binding of electrophilic chemicals to SH(thiol)-group of proteins and /or to seleno-proteins involved in protection against oxidative stress leads to chronic kidney disease	KeyEvent
Aop:377 - Dysregulated prolonged Toll Like Receptor 9 (TLR9) activation leading to Multi Organ Failure involving Acute Respiratory Distress Syndrome (ARDS)	KeyEvent
Aop:411 - Oxidative stress Leading to Decreased Lung Function	MolecularInitiatingEvent
Aop:424 - Oxidative stress Leading to Decreased Lung Function via CFTR dysfunction	MolecularInitiatingEvent
Aop:425 - Oxidative Stress Leading to Decreased Lung Function via Decreased FOXJ1	MolecularInitiatingEvent
Aop:429 - A cholesterol/glucose dysmetabolism initiated Tau-driven AOP toward memory loss (AO) in sporadic Alzheimer's Disease with plausible MIE's plug-ins for environmental neurotoxicants	KeyEvent
Aop:452 - Adverse outcome pathway of PM-induced respiratory toxicity	KeyEvent
Aop:464 - Calcium overload in dopaminergic neurons of the substantia nigra leading to parkinsonian motor deficits	KeyEvent
Aop:470 - Deposition of energy leads to abnormal vascular remodeling	KeyEvent
Aop:478 - Deposition of energy leading to occurrence of cataracts	KeyEvent
Aop:479 - Mitochondrial complexes inhibition leading to left ventricular function decrease via increased myocardial oxidative stress	KeyEvent
Aop:481 - AOPs of amorphous silica nanoparticles: ROS-mediated oxidative stress increased respiratory dysfunction and diseases.	KeyEvent
Aop:482 - Deposition of energy leading to occurrence of bone loss	KeyEvent
Aop:483 - Deposition of Energy Leading to Learning and Memory Impairment	KeyEvent
Aop:505 - Reactive Oxygen Species (ROS) formation leads to cancer via inflammation pathway	KeyEvent
Aop:521 - Essential element imbalance leads to reproductive failure via oxidative stress	KeyEvent
Aop:26 - Calcium-mediated neuronal ROS production and energy imbalance	AdverseOutcome
Aop:488 - Increased reactive oxygen species production leading to decreased cognitive function	KeyEvent
Aop:396 - Deposition of ionizing energy leads to population decline via impaired meiosis	KeyEvent
Aop:437 - Inhibition of mitochondrial electron transport chain (ETC) complexes leading to kidney toxicity	KeyEvent
Aop:535 - Binding and activation of GPER leading to learning and memory impairments	KeyEvent
Aop:171 - Chronic cytotoxicity of the serous membrane leading to pleural/peritoneal mesotheliomas in the rat.	KeyEvent
Aop:138 - Organic anion transporter (OAT1) inhibition leading to renal failure and mortality	KeyEvent
Aop:177 - Cyclooxygenase 1 (COX1) inhibition leading to renal failure and mortality	KeyEvent
Aop:186 - unknown MIE leading to renal failure and mortality	KeyEvent
Aop:200 - Estrogen receptor activation leading to breast cancer	KeyEvent
Aop:444 - Ionizing radiation leads to reduced reproduction in Eisenia fetida via reduced spermatogenesis and cocoon hatchability	KeyEvent
Aop:447 - Kidney failure induced by inhibition of mitochondrial electron transfer chain through apoptosis, inflammation and oxidative stress pathways	KeyEvent
Aop:476 - Adverse Outcome Pathways diagram related to PBDEs associated male reproductive toxicity	KeyEvent
Aop:497 - ERα inactivation alters mitochondrial functions and insulin signalling in skeletal muscle and leads to insulin resistance and metabolic syndrome	KeyEvent
Aop:457 - Succinate dehydrogenase inhibition leading to increased insulin resistance through reduction in circulating thyroxine	KeyEvent
Aop:459 - AhR activation in the thyroid leading to Subsequent Adverse Neurodevelopmental Outcomes in Mammals	KeyEvent
Aop:507 - Nrf2 inhibition leading to vascular disrupting effects via inflammation pathway	KeyEvent
Aop:509 - Nrf2 inhibition leading to vascular disrupting effects through activating apoptosis signal pathway and mitochondrial dysfunction	KeyEvent
Aop:510 - Demethylation of PPAR promotor leading to vascular disrupting effects	KeyEvent
Aop:511 - The AOP framework on ROS-mediated oxidative stress induced vascular disrupting effects	KeyEvent
Aop:538 - Adverse outcome pathway of PFAS-induced vascular disrupting effects via activating oxidative stress related pathways	KeyEvent

AOP ID and Name	Event Type
Aop:260 - CYP2E1 activation and formation of protein adducts leading to neurodegeneration	KeyEvent
Aop:450 - Inhibition of AChE and activation of CYP2E1 leading to sensory axonal peripheral neuropathy and mortality	KeyEvent
Aop:501 - Excessive iron accumulation leading to neurological disorders	KeyEvent
Aop:540 - Oxidative Stress in the Fish Ovary Leads to Reproductive Impairment via Reduced Vitellogenin Production	KeyEvent
Aop:471 - Neuron defect induced early behavioral change	KeyEvent
Aop:31 - Oxidation of iron in hemoglobin leading to hematotoxicity	KeyEvent
Aop:534 - Succinate dehydrogenase (SDH) inhibition leads to oxidative stress	AdverseOutcome
Aop:462 - Activation of reactive oxygen species leading the atherosclerosis	KeyEvent
Aop:331 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and cell death	KeyEvent
Aop:595 - Emerging OPFRS reproductive outcome pathway	KeyEvent
Aop:596 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	KeyEvent
Aop:598 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and reduced cell proliferation	KeyEvent
Aop:599 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and cell injury/death	KeyEvent
Aop:600 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell growth	KeyEvent
Aop:601 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell proliferation	KeyEvent
Aop:602 - Excessive reactive oxygen species leading to growth inhibition via oxidative DNA damage	KeyEvent
Aop:603 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell cycle disruption	KeyEvent
Aop:608 - Thyroid Hormone Excess Leading to Reduced, Swimming Performance via Hypomyelination	KeyEvent
Aop:616 - organic UV filter and its Photoproducts reproductive toxicity pathways	KeyEvent
Aop:622 - Calcineurin inhibitor induced nephrotoxicity leading to kidney failure	KeyEvent
Aop:625 - Increased 11β-Hydroxysteroid dehydrogenase type 1 activity leading to MASLD progression via insulin resistance-associated oxidative stress	KeyEvent
Aop:628 - Increased 11β-Hydroxysteroid dehydrogenase type 1 activity leading to MASLD progression via lipogenesis-associated oxidative stress	KeyEvent
Aop:472 - DNA adduct formation leading to kidney failure	KeyEvent
Aop:642 - Intestinal FXR inhibition leading to steatohepatitis via gut-liver axis dysregulation	KeyEvent
Aop:324 - Increase in reactive oxygen species leading to growth inhibition via oxidative DNA damage and cell cycle disruption	KeyEvent
Aop:325 - Increase in reactive oxygen species leading to growth inhibition via oxidative DNA damage and cell death	KeyEvent
Aop:326 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and decreased cell proliferation	KeyEvent
Aop:332 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and decreased cell proliferation	KeyEvent
Aop:333 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	KeyEvent
Aop:645 - Binding and activation of AhR lead to cardiovascular aging	KeyEvent
Aop:646 - Binding and activation of AhR/PPARγ lead to lipid metabolism disorders	KeyEvent
Aop:647 - ROS formation leading to growth inhibition modulated by the Keap1-Nrf2 antioxidant response	KeyEvent

Stressors

Name

Acetaminophen
Chloroform
furan

Name

Platinum
Aluminum
Cadmium
Mercury
Uranium
Arsenic
Silver
Manganese
Nickel
Zinc
nanoparticles

Biological Context**Level of Biological Organization**

Molecular

Domain of Applicability**Taxonomic Applicability**

Term	Scientific Term	Evidence	Links
rodents	rodents	High	NCBI
Homo sapiens	Homo sapiens	High	NCBI

Life Stage Applicability

Life Stage	Evidence
All life stages	High

Sex Applicability

Sex	Evidence
Mixed	High

Taxonomic applicability: Occurrence of oxidative stress is not species specific.

Life stage applicability: Occurrence of oxidative stress is not life stage specific.

Sex applicability: Occurrence of oxidative stress is not sex specific.

Evidence for perturbation by prototypic stressor: There is evidence of the increase of oxidative stress following perturbation from a variety of stressors including exposure to ionizing radiation and altered gravity (Bai et al., 2020; Ungvari et al., 2013; Zhang et al., 2009).

Key Event Description

Oxidative stress is defined as an imbalance in the production of reactive oxygen species (ROS) and antioxidant defenses. High levels of oxidizing free radicals can be very damaging to cells and molecules within the cell. As a result, the cell has important defense mechanisms to protect itself from ROS. For example, Nrf2 is a transcription factor and master regulator of the oxidative stress response. During periods of oxidative stress, Nrf2-dependent changes in gene expression are important in regaining cellular homeostasis (Nguyen, et al., 2009) and can be used as indicators of the presence of oxidative stress in the cell.

In addition to the directly damaging actions of ROS, cellular oxidative stress also changes cellular activities on a molecular level. Redox sensitive proteins have altered physiology in the presence and absence of ROS, which is caused by the oxidation of sulfhydryls to disulfides on neighboring amino acids (Antelmann & Hellmann 2011). Importantly Keap1, the negative regulator of Nrf2, is regulated in this manner (Itoh, et al. 2010).

ROS also undermine the mitochondrial defense system from oxidative damage. The antioxidant systems consist of superoxide dismutase, catalase, glutathione peroxidase and glutathione reductase, as well as antioxidants such as α -tocopherol and ubiquinol, or antioxidant vitamins and minerals including vitamin E, C, carotene, lutein, zeaxanthin, selenium, and zinc (Fletcher, 2010). The enzymes, vitamins and minerals catalyze the conversion of ROS to non-toxic molecules such as water and O₂. However, these antioxidant systems are not perfect and endogenous metabolic processes and/or exogenous oxidative influences can trigger cumulative oxidative injuries to the mitochondria, causing a decline in their functionality and efficiency, which further promotes cellular oxidative stress (Balasubramanian, 2000; Ganea & Harding, 2006; Guo et al., 2013; Karimi et al., 2017).

However, an emerging viewpoint suggests that ROS-induced modifications may not be as detrimental as previously thought, but rather contribute to signaling processes (Foyer et al., 2017).

Sources of ROS Production

Direct Sources: Direct sources involve the deposition of energy onto water molecules, breaking them into active radical species. When ionizing radiation hits water, it breaks it into hydrogen (H*) and hydroxyl (OH*) radicals by destroying its bonds. The hydrogen will create hydroxyl free radicals (HO2*) if oxygen is available, which can then react with another of itself to form hydrogen peroxide (H2O2) and more O2 (Elgazzar and Kazem, 2015). Antioxidant mechanisms are also affected by radiation, with catalase (CAT) and peroxidase (POD) levels rising as a result of exposure (Seen et al. 2018; Ahmad et al. 2021).

Indirect Sources: An indirect source of ROS is the mitochondria, which is one of the primary producers in eukaryotic cells (Powers et al., 2008). As much as 2% of the electrons that should be going through the electron transport chain in the mitochondria escape, allowing them an opportunity to interact with surrounding structures. Electron-oxygen reactions result in free radical production, including the formation of hydrogen peroxide (H2O2) (Zhao et al., 2019). The electron transport chain, which also creates ROS, is activated by free adenosine diphosphate (ADP), O2, and inorganic phosphate (Pi) (Hargreaves et al. 2020; Raimondi et al. 2020; Vargas-Mendoza et al. 2021). The first and third complexes of the transport chain are the most relevant to mammalian ROS production (Raimondi et al., 2020). The mitochondria has its own set of DNA and it is a prime target of oxidative damage (Guo et al., 2013). ROS is also produced through nicotinamide adenine dinucleotide phosphate oxidase (Nox) stimulation, an event commenced by angiotensin II, a product/effector of the renin-angiotensin system (Nguyen Dinh Cat et al. 2013; Forrester et al. 2018). Other ROS producers include xanthine oxidase, immune cells (macrophage, neutrophils, monocytes, and eosinophils), phospholipase A2 (PLA2), monoamine oxidase (MAO), and carbon-based nanomaterials (Powers et al. 2008; Jacobsen et al. 2008; Vargas-Mendoza et al. 2021).

How it is Measured or Detected

Oxidative Stress: Direct measurement of ROS is difficult because ROS are unstable. The presence of ROS can be assayed indirectly by measurement of cellular antioxidants, or by ROS-dependent cellular damage. Listed below are common methods for detecting the KE, however there may be other comparable methods that are not listed

- Detection of ROS by chemiluminescence (<https://www.sciencedirect.com/science/article/abs/pii/S0165993606001683>)
- Detection of ROS by chemiluminescence is also described in OECD TG 495 to assess phototoxic potential.
- Glutathione (GSH) depletion. GSH can be measured by assaying the ratio of reduced to oxidized glutathione (GSH:GSSG) using a commercially available kit (e.g., <http://www.abcam.com/gshgssg-ratio-detection-assay-kit-fluorometric-green-ab138881.html>).
- TBARS. Oxidative damage to lipids can be measured by assaying for lipid peroxidation using TBARS (thiobarbituric acid reactive substances) using a commercially available kit.
- 8-oxo-dG. Oxidative damage to nucleic acids can be assayed by measuring 8-oxo-dG adducts (for which there are a number of ELISA based commercially available kits), or HPLC, described in Chepelev et al. (Chepelev, et al. 2015).

Molecular Biology: Nrf2. Nrf2's transcriptional activity is controlled post-translationally by oxidation of Keap1. Assay for Nrf2 activity include:

- Immunohistochemistry for increases in Nrf2 protein levels and translocation into the nucleus Western blot for increased Nrf2 protein levels
- Western blot of cytoplasmic and nuclear fractions to observe translocation of Nrf2 protein from the cytoplasm to the nucleus qPCR of Nrf2 target genes (e.g., Nqo1, Hmox-1, Gcl, Gst, Prx, TrxR, Srxn), or by commercially available pathway-based qPCR array (e.g., oxidative stress array from SABiosciences)
- Whole transcriptome profiling by microarray or RNA-seq followed by pathway analysis (in IPA, DAVID, metacore, etc.) for enrichment of the Nrf2 oxidative stress response pathway (e.g., Jackson et al. 2014)
- OECD TG422D describes an ARE-Nrf2 Luciferase test method

In general, there are a variety of commercially available colorimetric or fluorescent kits for detecting Nrf2 activation.

Assay Type & Measured Content	Description	Dose Range Studied	Assay Characteristics (Length/Ease of use/Accuracy)
ROS Formation in the Mitochondria assay (Shaki et al., 2012)	"The mitochondrial ROS measurement was performed flow cytometry using DCFH-DA. Briefly, isolated kidney mitochondria were incubated with UA (0, 50, 100 and 200 μ M) in respiration buffer containing (0.32 mM sucrose, 10mM Tris, 20 mM Mops, 50 μ M EGTA, 0.5 mM MgCl ₂ , 0.1 mM KH ₂ PO ₄ and 5 mM sodium succinate) [32]. In the interval times of 5, 30 and 60 min following the UA addition, a sample was taken and DCFH-DA was added (final concentration, 10 μ M) to mitochondria and was then incubated for 10 min. Uranyl acetate-induced ROS generation in isolated kidney mitochondria were determined through the flow cytometry (Partec, Deutschland) equipped with a 488-nm argon ion laser and supplied with the Flomax software and the signals were obtained using a 530-nm bandpass filter (FL-1 channel). Each determination is based on the mean fluorescence intensity of 15,000 counts."	0, 50, 100 and 200 μ M of Uranyl Acetate	Long/ Easy High accuracy
Mitochondrial Antioxidant Content Assay Measuring GSH content (Shaki et al., 2012)	"GSH content was determined using DTNB as the indicator and spectrophotometer method for the isolated mitochondria. The mitochondrial fractions (0.5 mg protein/ml) were incubated with various concentrations of uranyl acetate for 1 h at 30 °C and then 0.1 ml of mitochondrial fractions was added into 0.1 mol/l of phosphate buffers and 0.04% DTNB in a total volume of 3.0 ml (pH 7.4). The developed yellow color was read at 412 nm on a spectrophotometer (UV-1601 PC, Shimadzu, Japan). GSH content was expressed as μ g/mg protein."	0, 50, 100, or 200 μ M Uranyl Acetate	

H2O2 Production Assay Measuring H2O2 Production in isolated mitochondria (Heyno et al., 2008)	“Effect of CdCl2 and antimycin A (AA) on H2O2 production in isolated mitochondria from potato. H2O2 production was measured as scopoletin oxidation. Mitochondria were incubated for 30 min in the measuring buffer (see the Materials and Methods) containing 0.5 mM succinate as an electron donor and 0.2 μM mesoxalonnitrile 3-chlorophenylhydrazon (CCCP) as an uncoupler, 10 U horseradish peroxidase and 5 μM scopoletin.”	0, 10, 30 μM Cd2+ 2 μM antimycin A	
Flow Cytometry ROS & Cell Viability (Kruiderig et al., 1997)	“For determination of ROS, samples taken at the indicated time points were directly transferred to FACScan tubes. Dih123 (10 mM, final concentration) was added and cells were incubated at 37°C in a humidified atmosphere (95% air/5% CO2) for 10 min. At t 5 9, propidium iodide (10 mM, final concentration) was added, and cells were analyzed by flow cytometry at 60 ml/min. Nonfluorescent Dih123 is cleaved by ROS to fluorescent R123 and detected by the FL1 detector as described above for Dc (Van de Water 1995)”“For determination of ROS, samples taken at the indicated time points were directly transferred to FACScan tubes. Dih123 (10 mM, final concentration) was added and cells were incubated at 37°C in a humidified atmosphere (95% air/5% CO2) for 10 min. At t 5 9, propidium iodide (10 mM, final concentration) was added, and cells were analyzed by flow cytometry at 60 ml/min. Nonfluorescent Dih123 is cleaved by ROS to fluorescent R123 and detected by the FL1 detector as described above for Dc (Van de Water 1995)”		Strong/easy medium
DCFH-DA Assay Detection of hydrogen peroxide production (Yuan et al., 2016)	Intracellular ROS production was measured using DCFH-DA as a probe. Hydrogen peroxide oxidizes DCFH to DCF. The probe is hydrolyzed intracellularly to DCFH carboxylate anion. No direct reaction with H2O2 to form fluorescent production.	0-400 μM	Long/ Easy High accuracy
H2-DCF-DA Assay Detection of superoxide production (Thiebault et al., 2007)	This dye is a stable nonpolar compound which diffuses readily into the cells and yields H2-DCF. Intracellular OH or ONOO- react with H2-DCF when cells contain peroxides, to form the highly fluorescent compound DCF, which effluxes the cell. Fluorescence intensity of DCF is measured using a fluorescence spectrophotometer.	0-600 μM	Long/ Easy High accuracy
CM-H2DCFDA Assay (Eruslanov & Kusmartsev, 2009)	The dye (CM-H2DCFDA) diffuses into the cell and is cleaved by esterases, the thiol reactive chlormethyl group reacts with intracellular glutathione which can be detected using flow cytometry.		Long/Easy/ High Accuracy

Method of Measurement	References	Description	OECD-Approved Assay
Chemiluminescence	(Lu, C. et al., 2006; Griending, K. K., et al., 2016)	ROS can induce electron transitions in molecules, leading to electronically excited products. When the electrons transition back to ground state, chemiluminescence is emitted and can be measured. Reagents such as luminol and lucigenin are commonly used to amplify the signal.	No
Spectrophotometry	(Griending, K. K., et al., 2016)	NO has a short half-life. However, if it has been reduced to nitrite (NO2-), stable azocompounds can be formed via the Griess Reaction, and further measured by spectrophotometry.	No
Direct or Spin Trapping-Based electron paramagnetic resonance (EPR) Spectroscopy	(Griending, K. K., et al., 2016)	The unpaired electrons (free radicals) found in ROS can be detected with EPR and is known as electron paramagnetic resonance. A variety of spin traps can be used.	No
Nitroblue Tetrazolium Assay	(Griending, K. K., et al., 2016)	The Nitroblue Tetrazolium assay is used to measure O2.- levels. O2.- reduces nitroblue tetrazolium (a yellow dye) to formazan (a blue dye), and can be measured at 620 nm.	No
Fluorescence analysis of dihydroethidium (DHE) or Hydrocyans	(Griending, K. K., et al., 2016)	Fluorescence analysis of DHE is used to measure O2.- levels. O2.- is reduced to O2 as DHE is oxidized to 2-hydroxyethidium, and this reaction can be measured by fluorescence. Similarly, hydrocyans can be oxidized by any ROS, and measured via fluorescence.	No
Amplex Red Assay	(Griending, K. K., et al., 2016)	Fluorescence analysis to measure extramitochondrial or extracellular H2O2 levels. In the presence of horseradish peroxidase and H2O2, Amplex Red is oxidized to resorufin, a fluorescent molecule measurable by plate reader.	No
Dichlorodihydrofluorescein Diacetate (DCFH-DA)	(Griending, K. K., et al., 2016)	An indirect fluorescence analysis to measure intracellular H2O2 levels. H2O2 interacts with peroxidase or heme proteins, which further react with DCFH, oxidizing it to dichlorofluorescein (DCF), a fluorescent product.	No

HyPer Probe	(Griendling, K. K., et al., 2016)	Fluorescent measurement of intracellular H ₂ O ₂ levels. HyPer is a genetically encoded fluorescent sensor that can be used for in vivo and in situ imaging.	No
Cytochrome c Reduction Assay	(Griendling, K. K., et al., 2016)	The cytochrome c reduction assay is used to measure O ₂ •– levels. O ₂ •– is reduced to O ₂ as ferricytochrome c is oxidized to ferrocyanochrome c, and this reaction can be measured by an absorbance increase at 550 nm.	No
Proton-electron double-resonance imaging (PEDRI)	(Griendling, K. K., et al., 2016)	The redox state of tissue is detected through nuclear magnetic resonance/magnetic resonance imaging, with the use of a nitroxide spin probe or biradical molecule.	No
Glutathione (GSH) depletion	(Biesemann, N. et al., 2018)	A downstream target of the Nrf2 pathway is involved in GSH synthesis. As an indication of oxidation status, GSH can be measured by assaying the ratio of reduced to oxidized glutathione (GSH:GSSG) using a commercially available kit (e.g., http://www.abcam.com/gshgssg-ratio-detection-assay-kit-fluorometric-green-ab138881.html).	No
Thiobarbituric acid reactive substances (TBARS)	(Griendling, K. K., et al., 2016)	Oxidative damage to lipids can be measured by assaying for lipid peroxidation with TBARS using a commercially available kit.	No
Protein oxidation (carbonylation)	(Azimzadeh et al., 2017; Azimzadeh et al., 2015; Ping et al., 2020)	Can be determined with ELISA or a commercial assay kit. Protein oxidation can indicate the level of oxidative stress.	No
Seahorse XFp Analyzer	Leung et al. 2018	The Seahorse XFp Analyzer provides information on mitochondrial function, oxidative stress, and metabolic dysfunction of viable cells by measuring respiration (oxygen consumption rate; OCR) and extracellular pH (extracellular acidification rate; ECAR).	No

Molecular Biology: Nrf2. Nrf2's transcriptional activity is controlled post-translationally by oxidation of Keap1. Assays for Nrf2 activity include:

Method of Measurement	References	Description	OECD-Approved Assay
Immunohistochemistry	(Amsen, D., de Visser, K. E., and Town, T., 2009)	Immunohistochemistry for increases in Nrf2 protein levels and translocation into the nucleus	No
qPCR	(Forlenza et al., 2012)	qPCR of Nrf2 target genes (e.g., Nqo1, Hmox-1, Gcl, Gst, Prx, TrxR, Srxn), or by commercially available pathway-based qPCR array (e.g., oxidative stress array from SABiosciences)	No
Whole transcriptome profiling via microarray or via RNA-seq followed by a pathway analysis	(Jackson, A. F. et al., 2014)	Whole transcriptome profiling by microarray or RNA-seq followed by pathway analysis (in IPA, DAVID, metacore, etc.) for enrichment of the Nrf2 oxidative stress response pathway	No

Update - non-endorsed (by You Song, 02/09/2026)

ToxCast/Tox21 assays (invitrodb v4.3): ATG_NRF2_ARE_CIS; Tox21_ARE_BLA_Agonist_ratio; TOX21_ARE_KS_LUC_Agonist; LTEA_HepaRG_NFE2L2; LTEA_HepaRG_GCLC; LTEA_HepaRG_CAT

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Event: 1767: Increase, Protein oxidation

Short Name: Increase, Protein oxidation

Event Component

Process	Object	Action
protein oxidation		increased

AOPs Including This Key Event

AOP ID and Name	Event Type
Aop:327 - Excessive reactive oxygen species production leading to mortality (1)	KeyEvent
Aop:596 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	KeyEvent
Aop:598 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and reduced cell proliferation	KeyEvent
Aop:599 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and cell injury/death	KeyEvent
Aop:600 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell growth	KeyEvent
Aop:601 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell proliferation	KeyEvent
Aop:603 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell cycle disruption	KeyEvent
Aop:332 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and decreased cell proliferation	KeyEvent
Aop:333 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	KeyEvent
Aop:647 - ROS formation leading to growth inhibition modulated by the Keap1-Nrf2 antioxidant response	KeyEvent

Biological Context

Level of Biological Organization

Molecular

Cell term

Cell term

cell

Organ term

Organ term

organ

Domain of Applicability

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
humans	Homo sapiens	High	NCBI
mammals	mammals	High	NCBI
fish	fish	High	NCBI
crustaceans	Daphnia magna	Moderate	NCBI
green algae	Ulva compressa	Moderate	NCBI

Life Stage Applicability

Life Stage	Evidence
All life stages	Moderate

Sex Applicability

Sex	Evidence
Unspecific	Moderate

The biological domain of applicability for this KE is broad because proteins are universal biological macromolecules and many amino-acid residues are susceptible to oxidative modification. The KE is applicable wherever proteins are exposed to oxidants and where oxidative modification can be measured. It is therefore relevant across unicellular algae, invertebrates, fish, mammals, plants and human-derived cell systems. The evidence base is strongest in mammalian toxicology and biomedical studies, but ecotoxicological evidence supports relevance in algae, fish, mollusks and crustaceans.

The KE is not intrinsically limited by life stage or sex. However, the magnitude and toxicological importance of protein oxidation may be modified by antioxidant capacity, proteasomal and lysosomal degradation capacity, protein turnover, metal ion availability, oxygen availability, temperature, inflammatory status, nutritional status, mitochondrial activity, and exposure duration. Tissues or cells with high metabolic demand, high mitochondrial density, high inflammatory activity, or low proteostatic reserve may be especially susceptible.

Within the ROS-growth AOP network, this KE functions as a molecular damage event linking oxidative stress to downstream impairment of mitochondrial function and cellular injury. Nevertheless, the KE should remain modular. It may be reused in any AOP in which increased oxidative modification of proteins is measured or inferred as a discrete biological state, regardless of whether the downstream effect is impaired oxidative phosphorylation, cell death, altered signaling, immune dysfunction, neurotoxicity, growth inhibition or another adverse outcome.

Key Event Description

Protein oxidation refers to an increase in oxidative modification of proteins relative to an appropriate control state. Proteins are abundant and chemically diverse macromolecules that contain amino-acid side chains and peptide backbones susceptible to attack by ROS and related oxidants. Oxidation can lead to formation of protein carbonyls, oxidation of sulfur-containing amino acids such as cysteine and methionine, nitration or hydroxylation of aromatic residues, disulfide formation, S-glutathionylation, fragmentation, cross-linking, aggregation, altered folding and changes in enzymatic or structural function (Stadtman and Levine, 2003; Dalle-Donne et al., 2006; Fedorova et al., 2014).

The KE is defined by the observed or measured increase in oxidatively modified proteins rather than by a particular upstream stressor or downstream consequence. Protein oxidation can be reversible or irreversible depending on the chemical modification. Reversible thiol oxidation, disulfide formation, S-glutathionylation and methionine oxidation may participate in redox signaling and adaptive regulation, whereas irreversible carbonylation, backbone cleavage and protein aggregation are more commonly associated with protein dysfunction, proteostatic burden and cellular injury (Stadtman and Levine, 2003; Dalle-Donne et al., 2006; Reichmann et al., 2018).

Within oxidative stress AOPs, protein oxidation is an important molecular damage KE because it links redox imbalance to functional impairment of enzymes, structural proteins, signaling proteins and organelle proteins. In the ROS-growth AOP network, oxidation of mitochondrial respiratory proteins, cytoskeletal proteins or metabolic enzymes may contribute to decreased coupling of oxidative phosphorylation, impaired ATP production, altered cell cycle regulation, increased cell injury/death and reduced growth. However, these downstream consequences should be described on separate KER and AOP pages so that KE 1767 remains modular and reusable.

How it is Measured or Detected

Protein oxidation can be measured using biochemical, immunochemical, fluorescence-based and proteomic approaches. No single method captures all forms of protein oxidation. Protein carbonylation is one of the most widely used and relatively stable indicators of oxidative protein damage, but other modifications such as methionine sulfoxide, cysteine oxidation, nitrotyrosine, S-glutathionylation and protein cross-linking may be more appropriate in particular biological contexts. Confidence is highest when the method directly detects a defined oxidized protein modification or oxidized peptide, and lower when broad assays are used without complementary specificity checks.

Measurement approach	Endpoint measured	Representative method names	Scientific confidence and limitations
Protein carbonyl assays	Protein carbonyl groups formed by direct oxidation or by adduction of reactive carbonyl species	DNPH derivatization with spectrophotometry, ELISA, immunoblotting or OxyBlot; hydrazide-based probes	Widely used, relatively stable and broadly accepted as a marker of protein oxidation. DNPH methods are sensitive but do not identify individual proteins unless combined with immunoblotting or proteomics. Carbonyls may arise from direct oxidation or from secondary reactions with lipid peroxidation products (Levine et al., 1990; Dalle-Donne et al., 2006; Fedorova et al., 2014).
Redox proteomics	Oxidized proteins or oxidized amino-acid residues at protein or peptide level	2D gel electrophoresis plus anti-DNP immunoblotting; LC-MS/MS redox proteomics; carbonyl-reactive enrichment workflows	High mechanistic value because it can identify protein targets and modification sites. Requires careful sample handling, derivatization or enrichment, and appropriate bioinformatic analysis (McDonagh et al., 2005; Fedorova et al., 2014; Butterfield and Dalle-Donne, 2014).
Thiol oxidation assays	Oxidation state of protein thiols and disulfides	Biotin-switch methods; maleimide labeling; redox Western blot; differential alkylation; thiol redox proteomics	Useful for reversible cysteine oxidation and redox signaling. Interpretation depends on preservation of redox state during sampling and on whether reversible signaling events or irreversible damage are being assessed (Dalle-Donne et al., 2006; Reichmann et al., 2018).
S-glutathionylation assays	Protein S-glutathionylation as a reversible thiol redox modification	Anti-glutathione immunoblotting; redox proteomics; mass spectrometry	Mechanistically informative for redox regulation and oxidative stress responses. It may represent adaptive regulation rather than irreversible damage and should be interpreted in biological context (Dailianis et al., 2009; Zaffagnini et al., 2012).
Nitrotyrosine and other specific oxidized residue assays	Specific oxidized or nitrated amino-acid residues	Anti-nitrotyrosine immunoassays; LC-MS/MS; targeted proteomics	Provides higher chemical specificity for particular oxidant pathways, such as peroxynitrite-associated nitration, but does not capture all protein oxidation. Best used when the expected chemistry is known.
Advanced oxidation protein products and aggregate assays	Bulk oxidized protein products, cross-linked proteins or protein aggregates	AOPP assays; aggregate detection; protein insolubility assays	Useful for broad screening of oxidative protein burden but less specific than defined chemical or proteomic measurements. Should be interpreted as supportive evidence, especially when combined with carbonyl or mass-spectrometric endpoints.

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Event: 1446: Decrease, Coupling of oxidative phosphorylation

Short Name: Decrease, Coupling of OXPHOS

Event Component

Process	Object	Action
proton binding	mitochondrion	increased
oxidative phosphorylation uncoupler activity	mitochondrion	increased
regulation of mitochondrial membrane potential	mitochondrion	decreased

AOPs Including This Key Event

AOP ID and Name	Event Type
Aop:267 - Uncoupling of oxidative phosphorylation leading to growth inhibition via glucose depletion	MolecularInitiatingEvent
Aop:263 - Uncoupling of oxidative phosphorylation leading to growth inhibition via decreased cell proliferation	MolecularInitiatingEvent
Aop:264 - Uncoupling of oxidative phosphorylation leading to growth inhibition via ATP depletion associated cell death	MolecularInitiatingEvent
Aop:265 - Uncoupling of oxidative phosphorylation leading to growth inhibition via increased cytosolic calcium	MolecularInitiatingEvent

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AOP ID and Name	Event Type
Aop:266 - Uncoupling of oxidative phosphorylation leading to growth inhibition via decreased Na-K ATPase activity	MolecularInitiatingEvent
Aop:268 - Uncoupling of oxidative phosphorylation leading to growth inhibition via mitochondrial swelling	MolecularInitiatingEvent
Aop:534 - Succinate dehydrogenase (SDH) inhibition leads to oxidative stress	KeyEvent
Aop:331 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and cell death	KeyEvent
Aop:596 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	KeyEvent
Aop:598 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and reduced cell proliferation	KeyEvent
Aop:612 - Peroxisome proliferator-activated receptor alpha activation leading to early life stage mortality via reduced adenosine triphosphate	KeyEvent
Aop:613 - Peroxisome proliferator-activated receptor alpha activation leading to early life stage mortality via increased reactive oxygen species production	KeyEvent
Aop:326 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and decreased cell proliferation	KeyEvent
Aop:332 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and decreased cell proliferation	KeyEvent
Aop:333 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	KeyEvent

Stressors

Name

2,4-Dinitrophenol
 Carbonyl cyanide-p-trifluoromethoxyphenylhydrazone
 Carbonyl cyanide m-chlorophenyl hydrazone
 Pentachlorophenol
 Triclosan
 Emodin
 Malonoben

Biological Context

Level of Biological Organization

Cellular

Cell term

Cell term

cell

Domain of Applicability

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
zebrafish	Danio rerio	High	NCBI
human	Homo sapiens	High	NCBI
mouse	Mus musculus	High	NCBI
rat	Rattus norvegicus	High	NCBI
Lemna minor	Lemna minor	High	NCBI

Life Stage Applicability

Life Stage	Evidence
Embryo	High
Juvenile	High

Life Stage		Evidence
Adult, reproductively mature		Moderate
Sex Applicability		
Sex	Evidence	
Unspecific	High	
Taxonomic applicability domain		
This key event is in general considered applicable to most eukaryotes, as the mitochondrion and oxidative phosphorylation are highly conserved (Roger 2017).		
Life stage applicability domain		
This key event is considered applicable to all life stages, as ATP synthesis by oxidative phosphorylation is an essential biological process for most living organisms.		
Sex applicability domain		
This key event is considered sex-unspecific, as both males and females use oxidative phosphorylation as a main process to generate ATP.		
Key Event Description		
Decreased coupling of oxidative phosphorylation (OXPHOS), or uncoupling of OXPHOS, describes dissipation of protonmotive force (PMF) across the inner mitochondrial membrane (IMM) by environmental stressors. In eukaryotes, the mitochondrial electron transport chain mediates a series of redox reactions to create a PMF across the IMM. The PMF is used as energy to drive adenosine triphosphate (ATP) synthesis through phosphorylation of adenosine diphosphate (ADP). These processes are coupled and referred to as OXPHOS. A number of chemicals can dissipate the PMF, leading to uncoupling of OXPHOS. This key event describes the main outcome of the interactions between an uncoupler and the transmembrane PMF. An uncoupler can bind to a proton in the mitochondrial inter membrane space, transport the proton to the matrix side of the IMM, release the proton and move back to the inter membrane space. These processes are repeated until the transmembrane PMF is dissipated. This KE is therefore a lumped term of these processes and represents the final consequence of the interactions.		
How it is Measured or Detected		
Uncoupling of oxidative phosphorylation can be indicated by reduced mitochondrial membrane potential, increased proton leak and/or increased oxygen consumption rate.		
- Mitochondrial membrane potential can be determined using ToxCast high-throughput screening bioassays such as “APR_HepG2_MitoMembPot”, “APR_Hepat_MitoFxnI”, and “APR_Mitochondrial_membrane_potential”, and the Tox21 high-throughput screening assay “tox21-mitotox-p1”. - Mitochondrial membrane potential can also be measured using commercially available fluorescent probes such as TMRM (tetramethylrhodamine, methyl ester, perchlorate), TMRE (tetramethylrhodamine, ethyl ester, perchlorate) and JC-1 (Perry 2011). - Proton leak and oxygen consumption rate can be measured using a high-resolution respirometry (Affourtit 2018) or a Seahorse XF analyzer (Divakaruni 2014).		
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Event: 1771: Decrease, Adenosine triphosphate pool

Short Name: Decrease, ATP pool

Event Component

Process	Object	Action
ATP biosynthetic process	ATP	decreased

AOPs Including This Key Event

AOP ID and Name	Event Type
Aop:328 - Excessive reactive oxygen species production leading to mortality (2)	KeyEvent
Aop:329 - Excessive reactive oxygen species production leading to mortality (3)	KeyEvent
Aop:264 - Uncoupling of oxidative phosphorylation leading to growth inhibition via ATP depletion associated cell death	KeyEvent
Aop:263 - Uncoupling of oxidative phosphorylation leading to growth inhibition via decreased cell proliferation	KeyEvent
Aop:290 - Mitochondrial ATP synthase antagonism leading to growth inhibition (1)	KeyEvent
Aop:291 - Mitochondrial ATP synthase antagonism leading to growth inhibition (2)	KeyEvent
Aop:286 - Mitochondrial complex III antagonism leading to growth inhibition (1)	KeyEvent
Aop:287 - Mitochondrial complex III antagonism leading to growth inhibition (2)	KeyEvent
Aop:266 - Uncoupling of oxidative phosphorylation leading to growth inhibition via decreased Na-K ATPase activity	KeyEvent
Aop:331 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and cell death	KeyEvent
Aop:596 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	KeyEvent
Aop:598 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and reduced cell proliferation	KeyEvent
Aop:599 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and cell injury/death	KeyEvent
Aop:600 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell growth	KeyEvent
Aop:601 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell proliferation	KeyEvent

AOP ID and Name		Event Type
Aop:612 - Peroxisome proliferator-activated receptor alpha activation leading to early life stage mortality via reduced adenosine triphosphate		KeyEvent
Aop:326 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and decreased cell proliferation		KeyEvent
Aop:332 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and decreased cell proliferation		KeyEvent
Aop:333 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death		KeyEvent
Stressors		
Name		
Carbonyl cyanide-p-trifluoromethoxyphenylhydrazone		
Carbonyl cyanide m-chlorophenyl hydrazone		
2,4-Dinitrophenol		
Malonoben		
Pentachlorophenol		
Triclosan		
Emodin		
Biological Context		
Level of Biological Organization		
Cellular		
Cell term		
Cell term		
cell		
Domain of Applicability		
Taxonomic Applicability		
Term	Scientific Term	Evidence Links
zebrafish	Danio rerio	High NCBI
human	Homo sapiens	High NCBI
rat	Rattus norvegicus	High NCBI
mouse	Mus musculus	High NCBI
Lemna minor	Lemna minor	High NCBI
Life Stage Applicability		
Life Stage	Evidence	
Embryo	High	
Juvenile	High	
Adult, reproductively mature	Moderate	
Sex Applicability		
Sex	Evidence	
Unspecific	High	
Taxonomic applicability domain		
This key event is in general considered applicable to all eukaryotes utilizing ATP as a direct source of energy and signaling molecule.		
Life stage applicability domain		
This key event is considered applicable to all life stages, as all developmental stages require energy supply to maintain necessary physiological processes.		

Sex applicability domain

This key event is considered sex-unspecific, as both males and females use ATP as an essential energy molecule.

Key Event Description

Decreased adenosine triphosphate (ATP) pool describes the loss of balance between ATP synthesis and ATP consumption, leading to reduced total ATP. As a primary form of biological energy, ATP is used by many biological processes (Bonora 2012). Decrease in ATP level normally attributes to metabolic disorders in major ATP synthetic pathways, such as mitochondrial oxidative phosphorylation, fatty acid β -oxidation, glycolysis and plant photophosphorylation.

How it is Measured or Detected

-The ATP pool in cells or tissue can be quantified using a well-established ATP bioluminescent assay (Lemasters 1978; Wibom 1990). Assay principles: ATP can react with luciferase and luciferin from firefly and the luminescence emitted from the reaction is proportional to the ATP concentration:



-ToxCast high-throughput screening bioassays, such as "NCCT_HEK293T_CellTiterGLO" and "NIS_HEK293T_CTG_Cytotoxicity" can be used to measure this KE.

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Event: 55: Increase, Cell injury/death**Short Name: Cell injury/death****Event Component****Process Object Action**

cell		increased
death		

AOPs Including This Key Event

AOP ID and Name	Event Type
Aop:48 - Binding of agonists to ionotropic glutamate receptors in adult brain causes excitotoxicity that mediates neuronal cell death, contributing to learning and memory impairment.	KeyEvent
Aop:13 - Chronic binding of antagonist to N-methyl-D-aspartate receptors (NMDARs) during brain development induces impairment of learning and memory abilities	KeyEvent
Aop:38 - Protein Alkylation leading to Liver Fibrosis	KeyEvent
Aop:12 - Chronic binding of antagonist to N-methyl-D-aspartate receptors (NMDARs) during brain development leads to neurodegeneration with impairment in learning and memory in aging	KeyEvent
Aop:144 - Endocytic lysosomal uptake leading to liver fibrosis	KeyEvent
Aop:17 - Binding of electrophilic chemicals to SH(thiol)-group of proteins and /or to seleno-proteins involved in protection against oxidative stress during brain development leads to impairment of learning and memory	KeyEvent
Aop:278 - IKK complex inhibition leading to liver injury	KeyEvent
Aop:281 - Acetylcholinesterase Inhibition Leading to Neurodegeneration	KeyEvent
Aop:273 - Mitochondrial complex inhibition leading to liver injury	KeyEvent
Aop:377 - Dysregulated prolonged Toll Like Receptor 9 (TLR9) activation leading to Multi Organ Failure involving Acute Respiratory Distress Syndrome (ARDS)	KeyEvent
Aop:265 - Uncoupling of oxidative phosphorylation leading to growth inhibition via increased cytosolic calcium	KeyEvent
Aop:264 - Uncoupling of oxidative phosphorylation leading to growth inhibition via ATP depletion associated cell death	KeyEvent

AOP ID and Name	Event Type
Aop:266 - Uncoupling of oxidative phosphorylation leading to growth inhibition via decreased Na-K ATPase activity	KeyEvent
Aop:268 - Uncoupling of oxidative phosphorylation leading to growth inhibition via mitochondrial swelling	KeyEvent
Aop:479 - Mitochondrial complexes inhibition leading to left ventricular function decrease via increased myocardial oxidative stress	KeyEvent
Aop:490 - Co-activation of IP3R and RyR leads to reduced IQ and increased socio-economic burden through non-cholinergic mechanisms	KeyEvent
Aop:494 - AhR activation leading to liver fibrosis	KeyEvent
Aop:530 - Endocytotic lysosomal uptake leads to intestinal barrier disruption	KeyEvent
Aop:331 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and cell death	KeyEvent
Aop:596 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	KeyEvent
Aop:599 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and cell injury/death	KeyEvent
Aop:624 - Increased 11β-Hydroxysteroid dehydrogenase type 1 activity leading to MASLD progression via insulin resistance-associated mitochondrial dysfunction	KeyEvent
Aop:625 - Increased 11β-Hydroxysteroid dehydrogenase type 1 activity leading to MASLD progression via insulin resistance-associated oxidative stress	KeyEvent
Aop:626 - Increased 11β-Hydroxysteroid dehydrogenase type 1 activity leading to MASLD progression via insulin resistance-associated endoplasmic reticulum stress	KeyEvent
Aop:627 - Increased 11β-Hydroxysteroid dehydrogenase type 1 activity leading to MASLD progression via lipogenesis-associated mitochondrial dysfunction	KeyEvent
Aop:628 - Increased 11β-Hydroxysteroid dehydrogenase type 1 activity leading to MASLD progression via lipogenesis-associated oxidative stress	KeyEvent
Aop:629 - Increased 11β-Hydroxysteroid dehydrogenase type 1 activity leading to MASLD progression via lipogenesis-associated endoplasmic reticulum stress	KeyEvent
Aop:325 - Increase in reactive oxygen species leading to growth inhibition via oxidative DNA damage and cell death	KeyEvent
Aop:333 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	KeyEvent
Aop:647 - ROS formation leading to growth inhibition modulated by the Keap1-Nrf2 antioxidant response	KeyEvent

Biological Context

Level of Biological Organization

Cellular

Cell term

Cell term

eukaryotic cell

Domain of Applicability

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
human	Homo sapiens	High	NCBI
human and other cells in culture	human and other cells in culture	High	NCBI
Rattus norvegicus	Rattus norvegicus	High	NCBI
mouse	Mus musculus	High	NCBI

Life Stage Applicability

Life Stage Evidence

All life stages

Sex Applicability

Sex Evidence

Unspecific

Cell death is an universal event occurring in cells of any species (Fink and Cookson,2005).

Key Event Description

Two types of cell death can be distinguished by morphological features, although it is likely that these are two ends of a spectrum with possible intermediate forms. Apoptosis involves shrinkage, nuclear disassembly, and fragmentation of the cell into discrete bodies with intact plasma membranes. These are rapidly phagocytosed by neighbouring cells. An important feature of apoptosis is the requirement for adenosine triphosphate (ATP) to initiate the execution phase. In contrast, necrotic cell death is characterized by cell swelling and lysis. This is usually a consequence of profound loss of mitochondrial function and resultant ATP depletion, leading to loss of ion homeostasis, including volume regulation, and increased intracellular Ca^{2+} . The latter activates a number of nonspecific hydrolases (i.e., proteases, nucleases, and phospholipases) as well as calcium dependent kinases. Activation of calpain I, the Ca^{2+} -dependent cysteine protease cleaves the death-promoting Bcl-2 family members Bid and Bax which translocate to mitochondrial membranes, resulting in release of truncated apoptosis-inducing factor (tAIF), cytochrome c and endonuclease in the case of Bid and cytochrome c in the case of Bax. tAIF translocates to cell nuclei, and together with cyclophilin A and phosphorylated histone H2AX (γH2AX) is responsible for DNA cleavage, a feature of programmed necrosis. Activated calpain I has also been shown to cleave the plasma membrane Na^{+} - Ca^{2+} exchanger, which leads to build-up of intracellular Ca^{2+} , which is the source of additional increased intracellular Ca^{2+} . Cytochrome c in cellular apoptosis is a component of the apoptosome.

DNA damage activates nuclear poly(ADP-ribose) polymerase-1 (PARP-1), a DNA repair enzyme. PARP-1 forms poly(ADP-ribose) polymers, to repair DNA, but when DNA damage is extensive, PAR accumulates, exits cell nuclei and travels to mitochondrial membranes, where it, like calpain I, is involved in AIF release from mitochondria. A fundamental distinction between necrosis and apoptosis is the loss of plasma membrane integrity; this is integral to the former but not the latter. As a consequence, lytic release of cellular constituents promotes a local inflammatory reaction, whereas the rapid removal of apoptotic bodies minimizes such a reaction. The distinction between the two modes of death is easily accomplished in vitro but not in vivo. Thus, although claims that certain drugs induce apoptosis have been made, these are relatively unconvincing. DNA fragmentation can occur in necrosis, leading to positive TUNEL staining (see explanation below). Conversely, when apoptosis is massive, it can exceed the capacity for rapid phagocytosis, resulting in the eventual appearance of secondary necrosis.

Two alternative pathways - either extrinsic (receptor-mediated) or intrinsic (mitochondria-mediated) - lead to apoptotic cell death. The initiation of cell death begins either at the plasma membrane with the binding of TNF or FasL to their cognate receptors or within the cell. The latter is due to the occurrence of intracellular stress in the form of biochemical events such as oxidative stress, redox changes, covalent binding, lipid peroxidation, and consequent functional effects on mitochondria, endoplasmic reticulum, microtubules, cytoskeleton, or DNA. The intrinsic mitochondrial pathway involves the initiator, caspase-9, which, when activated, forms an "apoptosome" in the cytosol, together with cytochrome c, which translocates from mitochondria, Apaf-1 and dATP. The apoptosome activates caspase-3, the central effector caspase, which in turn activates downstream factors that are responsible for the apoptotic death of a cell (Fujikawa, 2015). Intracellular stress either directly affects mitochondria or can lead to effects on other organelles, which then send signals to the mitochondria to recruit participation in the death process (Fujikawa, 2015; Malhi et al., 2010). Constitutively expressed nitric oxide synthase (nNOS) is a Ca^{2+} -dependent cytosolic enzyme that forms nitric oxide (NO) from L-arginine, and NO reacts with the free radical such as superoxide (O_2^-) to form the very toxic free radical peroxynitrite (ONOO^-). Free radicals such as ONOO^- , O_2^- and hydroxyl radical (OH^-) damage cellular membranes and intracellular proteins, enzymes and DNA (Fujikawa, 2015; Malhi et al., 2010; Kaplowitz, 2002; Kroemer et al., 2009).

How it is Measured or Detected

Necrosis:

Lactate dehydrogenase (LDH) is a soluble cytoplasmic enzyme that is present in almost all cells and is released into extracellular space when the plasma membrane is damaged. To detect the leakage of LDH into cell culture medium, a tetrazolium salt is used in this assay. In the first step, LDH produces reduced nicotinamide adenine dinucleotide (NADH) when it catalyzes the oxidation of lactate to pyruvate. In the second step, a tetrazolium salt is converted to a colored formazan product using newly synthesized NADH in the presence of an electron acceptor. The amount of formazan product can be colorimetrically quantified by standard spectroscopy. Because of the linearity of the assay, it can be used to enumerate the percentage of necrotic cells in a sample (Chan et al., 2013).

The MTT assay is a colorimetric assay for assessing cell viability. NAD(P)H-dependent cellular oxidoreductase enzymes may reflect the number of viable cells present. These enzymes are capable of reducing the tetrazolium dye MTT 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide to its insoluble formazan, which has a purple color. Other closely related tetrazolium dyes include XTT, MTS and the WSTs. Tetrazolium dye assays can also be used to measure cytotoxicity (loss of viable cells) or cytostatic activity (shift from proliferation to quiescence) of potential medicinal agents and toxic materials. MTT assays are usually done in the dark since the MTT reagent is sensitive to light (Berridge et al., 2005).

Propidium iodide (PI) is an intercalating agent and a fluorescent molecule used to stain necrotic cells. It is cell membrane impermeant so it stains only those cells where the cell membrane is destroyed. When PI is bound to nucleic acids, the fluorescence excitation maximum is 535 nm and the emission maximum is 617 nm (Moore et al., 1998).

Alamar Blue (resazurin) is a fluorescent dye. The oxidized blue non fluorescent Alamar blue is reduced to a pink fluorescent dye in the medium by cell activity (O'Brien et al., 2000) (12).

Neutral red uptake, which is based on the ability of viable cells to incorporate and bind the supravital dye neutral red in lysosomes (Repetto et al., 2008) (13). Moreover, quantification of ATP, signaling the presence of metabolically active cells, can be performed (CellTiter-Glo; Promega).

ATP assay: Quantification of ATP, signaling the presence of metabolically active cells (CellTiter-Glo; Promega).

Apoptosis:

TUNEL is a common method for detecting DNA fragmentation that results from apoptotic signalling cascades. The assay relies on the presence of nicks in the DNA which can be identified by terminal deoxynucleotidyl transferase or TdT, an enzyme that will catalyze the addition of dUTPs that are secondarily labeled with a marker. It may also label cells that have suffered severe DNA damage.

Caspase activity assays measured by fluorescence. During apoptosis, mainly caspase-3 and -7 cleave PARP to yield an 85 kDa and a

25 kDa fragment. PARP cleavage is considered to be one of the classical characteristics of apoptosis. Antibodies to the 85 kDa fragment of cleaved PARP or to caspase-3 both serve as markers for apoptotic cells that can be monitored using immunofluorescence (Li, Peng et al., 2004).

Hoechst 33342 staining: Hoechst dyes are cell-permeable and bind to DNA in live or fixed cells. Therefore, these stains are often called supravital, which means that cells survive a treatment with these compounds. The stained, condensed or fragmented DNA is a marker of apoptosis (Loo, 2002; Kubbies and Rabinovitch, 1983).

Acridine Orange/Ethidium Bromide staining is used to visualize nuclear changes and apoptotic body formation that are characteristic of apoptosis. Cells are viewed under a fluorescence microscope and counted to quantify apoptosis.

Update - non-endorsed (by You Song, 02/09/2026)

ToxCast/Tox21 assays (invitrodb v4.3): APR_HepG2_CellLoss_*; ATG_XTT_Cytotoxicity; LTEA_HepaRG_LDH_cytotoxicity

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List of Adverse Outcomes in this AOP

Event: 1521: Decrease, Growth

Short Name: Decrease, Growth

Event Component

Process	Object	Action
growth	multicellular organism	decreased

AOPs Including This Key Event

AOP ID and Name	Event Type
Aop:263 - Uncoupling of oxidative phosphorylation leading to growth inhibition via decreased cell proliferation	AdverseOutcome
Aop:290 - Mitochondrial ATP synthase antagonism leading to growth inhibition (1)	AdverseOutcome
Aop:291 - Mitochondrial ATP synthase antagonism leading to growth inhibition (2)	AdverseOutcome
Aop:286 - Mitochondrial complex III antagonism leading to growth inhibition (1)	AdverseOutcome
Aop:287 - Mitochondrial complex III antagonism leading to growth inhibition (2)	AdverseOutcome
Aop:245 - Reduction in photophosphorylation leading to growth inhibition in aquatic plants	AdverseOutcome
Aop:265 - Uncoupling of oxidative phosphorylation leading to growth inhibition via increased cytosolic calcium	AdverseOutcome
Aop:264 - Uncoupling of oxidative phosphorylation leading to growth inhibition via ATP depletion associated cell death	AdverseOutcome
Aop:266 - Uncoupling of oxidative phosphorylation leading to growth inhibition via decreased Na-K ATPase activity	AdverseOutcome
Aop:267 - Uncoupling of oxidative phosphorylation leading to growth inhibition via glucose depletion	AdverseOutcome

AOP ID and Name	Event Type
Aop:268 - Uncoupling of oxidative phosphorylation leading to growth inhibition via mitochondrial swelling	AdverseOutcome
Aop:473 - Energy deposition from internalized Ra-226 decay lower oxygen binding capacity of hemocyanin	AdverseOutcome
Aop:331 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and cell death	AdverseOutcome
Aop:596 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	AdverseOutcome
Aop:598 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and reduced cell proliferation	AdverseOutcome
Aop:599 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and cell injury/death	AdverseOutcome
Aop:600 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell growth	AdverseOutcome
Aop:602 - Excessive reactive oxygen species leading to growth inhibition via oxidative DNA damage	AdverseOutcome
Aop:603 - Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell cycle disruption	AdverseOutcome
Aop:601 - Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell proliferation	AdverseOutcome
Aop:567 - Binding to plastoquinone B site leading to decreased population growth rate via photosystem II inhibition	AdverseOutcome
Aop:324 - Increase in reactive oxygen species leading to growth inhibition via oxidative DNA damage and cell cycle disruption	AdverseOutcome
Aop:325 - Increase in reactive oxygen species leading to growth inhibition via oxidative DNA damage and cell death	AdverseOutcome
Aop:326 - Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and decreased cell proliferation	AdverseOutcome
Aop:332 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and decreased cell proliferation	AdverseOutcome
Aop:333 - Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	AdverseOutcome
Aop:647 - ROS formation leading to growth inhibition modulated by the Keap1-Nrf2 antioxidant response	AdverseOutcome

Stressors

Name

2,4-Dinitrophenol
 Carbonyl cyanide-p-trifluoromethoxyphenylhydrazone
 Carbonyl cyanide m-chlorophenyl hydrazone
 Pentachlorophenol
 Triclosan
 Emodin
 Malonoben

Biological Context

Level of Biological Organization

Individual

Domain of Applicability

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
human	Homo sapiens	Moderate	NCBI
rat	Rattus norvegicus	Moderate	NCBI
mouse	Mus musculus	Moderate	NCBI
zebrafish	Danio rerio	High	NCBI
fathead minnow	Pimephales promelas	High	NCBI
Lemna minor	Lemna minor	High	NCBI

Term	Scientific Term	Evidence	Links
Daphnia magna	Daphnia magna	Moderate	NCBI
Life Stage Applicability			
Life Stage	Evidence		
Embryo	High		
Juvenile	High		
Sex Applicability			
Sex	Evidence		
Unspecific	High		
Taxonomic applicability domain			
This key event is in general applicable to all eukaryotes.			
Life stage applicability domain			
This key event is applicable to early life stages such as embryo and juvenile.			
Sex applicability domain			
This key event is sex-unspecific.			
Key Event Description			
Decreased growth refers to a reduction in size and/or weight of a tissue, organ or individual organism. Growth is normally controlled by growth factors and mainly achieved through cell proliferation (Conlon 1999).			
How it is Measured or Detected			
Growth can be indicated by measuring weight, length, total volume, and/or total area of a tissue, organ or individual organism.			
Revised version (non-endorsed, added by You Song, 02/09/2026)			
Growth can be measured as a change in size, mass, biomass, length, area, volume, or growth rate of a tissue, organ, or whole organism over a defined exposure period. The appropriate measurement depends on the biological system, life stage, and test species.			
At the organism level, growth is commonly assessed by measuring body weight, wet weight, dry weight, body length, standard length, total length, shell length, shoot height, root length, frond number, frond area, cell density, biomass, or relative growth rate. Growth inhibition is typically expressed as a statistically significant reduction relative to the control, or as an effect concentration causing a defined percentage reduction in growth, such as ECx, ErCx, EyCx, LOEC, NOEC, or benchmark concentration/dose values.			
For unicellular algae and cyanobacteria, growth is commonly quantified from changes in biomass over time. Biomass may be estimated using direct cell counts, electronic particle counters, microscopy, optical density, fluorescence, chlorophyll-related measurements, or other validated biomass proxies. Growth inhibition is commonly calculated from average specific growth rate and yield.			
For aquatic macrophytes such as Lemna spp., growth is typically quantified using frond number and at least one additional growth-related variable, such as total frond area, dry weight, fresh weight, or image-based area measurements. Digital imaging, flatbed scanning, stereomicroscopy, or automated image-analysis tools can be used to quantify frond area and related morphological endpoints.			
For terrestrial plants, growth can be measured using seedling emergence, survival, shoot height, root length, fresh or dry biomass, leaf area, and visual signs of phytotoxicity such as chlorosis, necrosis, deformation, or delayed development. Measurements may be made using rulers or calipers, analytical balances, scanners, digital imaging, or plant image-analysis software.			
For aquatic invertebrates, growth may be assessed using body length, body area, wet weight, dry weight, developmental stage, or size at defined time points. In small organisms such as Daphnia, body length is often measured from microscope or stereomicroscope images using calibrated image-analysis software.			
For fish and amphibians, growth is commonly assessed using wet weight, dry weight, standard length, total length, snout-vent length, condition factor, developmental stage, and specific growth rate. Measurements may be obtained using balances, calipers, digital imaging, or stereomicroscopy, depending on organism size and life stage.			
For mammals and other vertebrates in repeated-dose or reproductive/developmental toxicity studies, growth-related endpoints are typically evaluated using body weight, body-weight gain, food and water consumption, pup or offspring weight, litter growth, organ weight, and developmental landmarks. These endpoints are usually interpreted together with clinical observations, survival, pathology, and reproductive or developmental parameters.			
Because growth is an integrative apical endpoint, measurement should be performed over a biologically appropriate time window and interpreted in relation to survival, developmental delay, nutritional status, reproductive output, and general toxicity. Direct measurements of organismal size, biomass, or growth rate provide the strongest evidence for this key event, whereas indirect indicators such as altered morphology, reduced condition factor, or reduced organ weight may provide supporting evidence depending on biological context.			

Relevant OECD Test Guidelines include, but are not limited to:

- OECD TG 201: Freshwater Alga and Cyanobacteria, Growth Inhibition Test. This guideline directly assesses inhibition of algal or cyanobacterial growth, typically using average specific growth rate and yield as response variables.
- OECD TG 208: Terrestrial Plant Test: Seedling Emergence and Seedling Growth Test. This guideline assesses effects on seedling emergence and early growth of terrestrial plants, including biomass, shoot height, and visible phytotoxic effects.
- OECD TG 211: *Daphnia magna* Reproduction Test. Although the primary endpoint is reproductive output, the test can provide chronic organism-level information relevant to population performance, and growth-related observations such as body size may be reported as supporting endpoints when included in the study design.
- OECD TG 212: Fish, Short-term Toxicity Test on Embryo and Sac-Fry Stages. This guideline assesses toxicity during early fish development from fertilized egg to the end of the sac-fry stage. Growth-related developmental observations may support interpretation of early-life-stage impairment.
- OECD TG 215: Fish, Juvenile Growth Test. This guideline directly evaluates effects on juvenile fish growth. Effects are expressed using growth rate, and concentration-response analysis can be used to estimate concentrations causing a defined percentage change in growth rate.
- OECD TG 221: *Lemna* sp. Growth Inhibition Test. This guideline directly assesses effects on vegetative growth of *Lemna* spp. using average specific growth rate and yield, commonly based on frond number and additional variables such as frond area or biomass.
- OECD TG 228: Determination of Developmental Toxicity to Dipteran Dung Flies. This guideline is relevant where chemical exposure affects development and growth-related performance in terrestrial invertebrates.
- OECD TG 241: Larval Amphibian Growth and Development Assay. This guideline evaluates growth and development in amphibians from fertilization through the early juvenile period.
- OECD TG 407: Repeated Dose 28-day Oral Toxicity Study in Rodents. This guideline includes body weight and food/water consumption measurements as part of the assessment of repeated-dose systemic toxicity.
- OECD TG 408: Repeated Dose 90-day Oral Toxicity Study in Rodents. This guideline includes body weight and related clinical observations as part of subchronic systemic toxicity assessment.
- OECD TG 416: Two-Generation Reproduction Toxicity. This guideline evaluates effects on reproduction as well as growth and development of offspring across generations.
- OECD TG 422: Combined Repeated Dose Toxicity Study with the Reproduction/Developmental Toxicity Screening Test. This guideline includes body weight, food/water consumption, offspring observations and measurements, and developmental parameters.
- OECD TG 443: Extended One-Generation Reproductive Toxicity Study. This guideline evaluates reproductive and developmental effects, including health, growth, development, and function of offspring following pre- and postnatal exposure.
- OECD TG 453: Combined Chronic Toxicity/Carcinogenicity Studies. This guideline includes body weight and body-weight gain as part of the evaluation of chronic systemic toxicity and carcinogenicity.

Regulatory Significance of the AO

Growth is a regulatory relevant chronic toxicity endpoint for almost all organisms. Multiple OECD test guidelines have included growth either as a main endpoint of concern, or as an additional endpoint to be considered in the toxicity assessments. Relevant test guidelines include, but not only limited to:

- Test No. 201: Freshwater Alga and Cyanobacteria, Growth Inhibition Test
- Test No. 208: Terrestrial Plant Test: Seedling Emergence and Seedling Growth Test
- Test No. 211: *Daphnia magna* Reproduction Test
- Test No. 212: Fish, Short-term Toxicity Test on Embryo and Sac-Fry Stages
- Test No. 215: Fish, Juvenile Growth Test
- Test No. 221: *Lemna* sp. Growth Inhibition Test
- Test No. 228: Determination of Developmental Toxicity to Dipteran Dung Flies (*Scathophaga stercoraria* L. (Scathophagidae), *Musca autumnalis* De Geer (Muscidae))
- Test No. 241: The Larval Amphibian Growth and Development Assay (LAGDA)
- Test No. 407: Repeated Dose 28-day Oral Toxicity Study in Rodents
- Test No. 408: Repeated Dose 90-Day Oral Toxicity Study in Rodents
- Test No. 416: Two-Generation Reproduction Toxicity
- Test No. 422: Combined Repeated Dose Toxicity Study with the Reproduction/Developmental Toxicity Screening Test
- Test No. 443: Extended One-Generation Reproductive Toxicity Study
- Test No. 453: Combined Chronic Toxicity/Carcinogenicity Studies

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Appendix 2

List of Key Event Relationships in the AOP

List of Adjacent Key Event Relationships

Relationship: 2009: Increase, ROS leads to Increase, Oxidative Stress

AOPs Referencing Relationship

AOP Name	Adjacency	Weight of Evidence	Quantitative Understanding
Reactive Oxygen Species (ROS) formation leads to cancer via inflammation pathway	adjacent	High	Not Specified
Essential element imbalance leads to reproductive failure via oxidative stress	adjacent		
unknown MIE leading to renal failure and mortality	adjacent		
ERα inactivation alters mitochondrial functions and insulin signalling in skeletal muscle and leads to insulin resistance and metabolic syndrome	adjacent	High	
Oxidative Stress in the Fish Ovary Leads to Reproductive Impairment via Reduced Vitellogenin Production	adjacent	High	Low
Activation of reactive oxygen species leading the atherosclerosis	adjacent	High	
Deposition of ionizing energy leads to population decline via impaired meiosis	adjacent	High	Moderate
Calcium-mediated neuronal ROS production and energy imbalance	adjacent	High	
Succinate dehydrogenase (SDH) inhibition leads to oxidative stress	adjacent	High	High
The AOP framework on ROS-mediated oxidative stress induced vascular disrupting effects	adjacent	High	High
AOPs of amorphous silica nanoparticles: ROS-mediated oxidative stress increased respiratory dysfunction and diseases.	adjacent	High	High
Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and cell death	adjacent	High	Moderate
Emerging OPFRS reproductive outcome pathway	adjacent	High	High
Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	adjacent	High	
Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and cell injury/death	adjacent		
Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell growth	adjacent		
Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell proliferation	adjacent		
Excessive reactive oxygen species leading to growth inhibition via oxidative DNA damage	adjacent		
Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell cycle disruption	adjacent		
DNA adduct formation leading to kidney failure	adjacent	High	High
Increase in reactive oxygen species leading to growth inhibition via oxidative DNA damage and cell cycle disruption	adjacent	High	Moderate
Increase in reactive oxygen species leading to growth inhibition via oxidative DNA damage and cell death	adjacent	High	Moderate
Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and decreased cell proliferation	adjacent	High	Moderate
Increase in reactive oxygen species leading to growth inhibition via protein oxidation and decreased cell proliferation	adjacent	High	Moderate
Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	adjacent	High	Moderate
ROS formation leading to growth inhibition modulated by the Keap1-Nrf2 antioxidant response	adjacent	High	High

Evidence Supporting Applicability of this Relationship

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
human	Homo sapiens	High	NCBI
fish	fish	High	NCBI
crustaceans	Daphnia magna	High	NCBI
green algae	Ulva compressa	High	NCBI
mouse	Mus musculus	High	NCBI
rat	Rattus norvegicus	High	NCBI

Life Stage Applicability

Life Stage	Evidence
All life stages	High

Sex Applicability

Sex	Evidence
Unspecific	High

This KER is broadly applicable to aerobic eukaryotic systems in which ROS production and antioxidant buffering can be measured. The current AOP-Wiki relationship page identifies human, mouse and rat with high evidence, but the ROS-growth evidence base supports extension to algae, fish, crustaceans, mollusks and other organisms relevant to environmental toxicology (AOP-Wiki, 2026a). The relationship is expected to be conserved because it is based on redox chemistry and conserved antioxidant-defense systems rather than on a taxon-specific receptor or signaling pathway.

The applicability domain should nevertheless be bounded by biological context and measurement feasibility. This KER is most relevant when the upstream KE is a measurable increase in ROS and the downstream KE is a measurable redox imbalance or antioxidant-response state rather than a distal oxidative damage endpoint alone. In organisms or compartments where ROS cannot be measured directly, evidence may rely on antioxidant-response or oxidative damage biomarkers, but these should be interpreted as indirect support. Applicability is strongest when ROS and oxidative stress endpoints are measured in the same system under the same exposure conditions.

Key Event Relationship Description

This KER describes the causal and predictive relationship by which an increase in reactive oxygen species leads to oxidative stress. ROS include superoxide, hydrogen peroxide, hydroxyl radical and secondary oxygen-derived reactive products. At low or transient levels, ROS can participate in normal cell signaling. However, when ROS production, flux or local concentration exceeds the capacity of enzymatic and non-enzymatic antioxidant defenses, the redox balance of the biological system shifts toward an oxidizing state, producing oxidative stress (Schieber and Chandel, 2014; Sies et al., 2017).

The downstream KE, oxidative stress, is not identical to increased ROS. Rather, it represents a systems-level imbalance between pro-oxidant pressure and antioxidant or repair capacity. The KER therefore depends not only on the magnitude of ROS increase, but also on the duration, localization and chemical identity of the ROS, the capacity of scavenging systems such as glutathione, superoxide dismutase, catalase and glutathione peroxidases, and the ability of the cell or organism to activate adaptive redox responses such as NRF2 signaling (Halliwell and Gutteridge, 2015; Griendling et al., 2016; Sies et al., 2017).

Within the ROS-growth AOP network, Relationship 2009 functions as a shared upstream KER. It connects the early measurable perturbation of increased ROS to the central hub event of oxidative stress, from which downstream AOP branches proceed through oxidative DNA damage, lipid peroxidation, protein oxidation, mitochondrial dysfunction, ATP depletion, altered cell proliferation, cell injury/death and decreased growth. This KER should remain modular and stressor-agnostic; stressor-specific mechanisms of ROS generation should be described in MIE or stressor sections where appropriate.

Evidence Supporting this KER

Biological Plausibility

Biological plausibility of Relationship 2009 is high. ROS are produced endogenously by mitochondrial electron transport, oxidase enzymes, peroxisomal reactions, photosynthetic electron transport and immune-cell oxidant systems, and they may also be generated by redox-cycling chemicals, metals, radiation and other stressors (Bedard and Krause, 2007; Murphy, 2009; Halliwell and Gutteridge, 2015). Oxidative stress is defined as a disturbance in the balance between oxidants and antioxidants in favor of oxidants, leading to disruption of redox signaling and/or molecular damage (Sies et al., 2017). Therefore, a sufficient increase in ROS has a direct mechanistic basis for causing oxidative stress when antioxidant and repair capacity are exceeded.

This relationship is also strongly supported by the known biology of antioxidant defenses. Superoxide dismutases convert superoxide to hydrogen peroxide; catalase, glutathione peroxidases and peroxiredoxins reduce hydrogen

peroxide and organic peroxides; and glutathione and thioredoxin systems maintain protein thiol redox balance. Increased ROS can consume these defenses, oxidize redox-sensitive proteins, activate NRF2-dependent antioxidant response pathways, and produce oxidative modification of lipids, proteins and nucleic acids (Schieber and Chandel, 2014; Griending et al., 2016; Sies et al., 2017).

Empirical Evidence

Empirical support for this KER is high. Numerous studies across taxa and stressor classes demonstrate concordant increases in ROS or ROS-generating conditions and oxidative stress endpoints. The strongest evidence comes from studies measuring both ROS and antioxidant-response or oxidative-stress biomarkers in the same biological system. Several examples from the ROS-growth concordance table are summarized below.

Biological system	Stressor	Exposure	Evidence for KE1115 (ROS increase)	Evidence for KE1392 (oxidative stress increase)	Concordance interpretation	Reference
Chlorella vulgaris	Paraquat	24 h; 0-1.0 uM	DCFH-DA fluorescence increased; LOEC for ROS approximately 0.5 uM paraquat.	SOD, POD and CAT activities increased at similar concentrations; antioxidant enzymes were approximately 3-5-fold above control at 0.5 uM.	Dose concordance supports ROS increase leading to oxidative stress in a photosynthetic eukaryote.	Qian et al. (2009)
Daphnia magna	Paraquat	48 h; 0.01-10 uM	ROS induction threshold reported around 0.1 uM paraquat.	SOD, CAT and GPx induction observed around 0.5 uM; TBARS increased around 1 uM.	ROS occurs at lower or similar concentrations than antioxidant and damage markers, supporting dose concordance.	Barata et al. (2005)
Trachinotus ovatus	Streptococcus agalactiae infection	0-120 h; 2×10^7 CFU/fish	ROS increased early, with maximum response around 6 h.	Antioxidant enzyme activities and antioxidant gene expression changed following the ROS response.	Temporal concordance supports ROS preceding redox-response activation during pathogen-induced oxidative stress.	Gao et al. (2022)
Mus musculus	Copper sulfate	42 days; 0-40 mg/kg bw	ROS increased at the lowest tested dose by day 42.	Antioxidant markers including SOD, GSH-related responses and oxidative stress/inflammatory indicators changed with exposure.	Concordant ROS and antioxidant-response changes support the relationship in mammals.	Jian et al. (2020)
Marine bivalves	Chlorothalonil	96 h; 0.1-10 ug/L	Stressor is thiol-reactive and associated with oxidative challenge; direct ROS was not the primary endpoint.	SOD, CAT and GPx activity changes and MDA/TBARS increases occurred in gill tissues.	Supports downstream oxidative stress following a stressor known to disturb redox balance; direct ROS evidence is weaker than in rows with ROS measurement.	Haque et al. (2019)

Biological system	Stressor	Exposure	Evidence for KE1115 (ROS increase)	Evidence for KE1392 (oxidative stress increase)	Concordance interpretation	Reference
Mya arenaria	Cyclic hypoxia/reoxygenation	3 weeks; repeated low oxygen exposure	Hypoxia/reoxygenation is a recognized ROS-generating condition in mitochondria.	Mitochondrial proton leak and oxidative stress-related bioenergetic changes were elevated under cyclic hypoxia.	Supports environmental modulation of ROS-associated oxidative stress and mitochondrial response.	Ouillon et al. (2021)

Uncertainties and Inconsistencies

The main uncertainties relate to measurement specificity and context dependence. ROS are chemically diverse and often short-lived, so different assays may detect different ROS species or generalized oxidant-dependent probe oxidation rather than a single ROS concentration. DCFH-DA and related probes are useful screening tools but can be influenced by peroxidases, metals, light, probe loading and cellular esterase activity (Wardman, 2007; Kalyanaraman et al., 2012). Consequently, apparent ROS increases must be interpreted with assay limitations in mind.

A second uncertainty is that ROS increases are not always adverse. Transient or localized ROS signals may activate adaptive stress responses and restore redox homeostasis without producing sustained oxidative stress. Conversely, oxidative stress may be inferred from antioxidant enzyme induction or oxidative damage biomarkers in studies where ROS were not directly measured. These cases support the KER less strongly than studies with direct, temporally resolved ROS measurements. Differences among taxa, life stages, tissues, exposure durations and antioxidant capacities may alter the threshold at which increased ROS becomes oxidative stress.

Quantitative Understanding of the Linkage

Quantitative understanding of this KER is low to moderate. The qualitative relationship is well established: oxidative stress occurs when ROS production or flux exceeds antioxidant and repair capacity. However, a universal quantitative threshold for ROS leading to oxidative stress cannot be defined because the relationship depends strongly on ROS species, subcellular localization, measurement method, antioxidant capacity, exposure duration, organism, cell type and co-stressors (Kalyanaraman et al., 2012; Griending et al., 2016; Sies et al., 2017).

Response-response relationship

Response-response information is available in specific systems. For example, in *Chlorella vulgaris* exposed to paraquat, ROS and antioxidant enzyme responses were observed at approximately 0.5 uM after 24 h, indicating local dose concordance between the upstream and downstream events (Qian et al., 2009). In *Daphnia magna* exposed to paraquat, ROS induction was reported at lower concentrations than antioxidant enzyme and TBARS responses, supporting an expected dose sequence in which ROS increases precede oxidative stress endpoints (Barata et al., 2005). These examples provide semi-quantitative support, but they cannot be generalized across all taxa or stressors.

Time-scale

The time scale of the KER can range from minutes to hours for ROS-sensitive signaling and antioxidant pathway activation, and from hours to days for measurable changes in antioxidant enzyme activities, glutathione status or oxidative damage biomarkers. In pathogen-exposed golden pompano, ROS increased early, followed by antioxidant enzyme and gene expression responses over subsequent hours to days, supporting temporal concordance (Gao et al., 2022).

Known modulating factors

Modulating factor	Details	Effect on the KER	Supporting evidence
Antioxidant capacity	Levels and activities of GSH, SOD, CAT, GPx, peroxiredoxins, thioredoxin systems and antioxidant vitamins.	Higher antioxidant capacity buffers ROS and raises the threshold for oxidative stress; depleted or impaired antioxidant systems lower the threshold.	Halliwell and Gutteridge (2015); Sies et al. (2017).

Modulating factor	Details	Effect on the KER	Supporting evidence
NRF2/ARE pathway activation	Induction of antioxidant and detoxification genes through NRF2-dependent signaling.	Adaptive NRF2 activation may reduce progression from increased ROS to sustained oxidative stress, but strong NRF2 activation can also serve as evidence that ROS has perturbed redox homeostasis.	Schieber and Chandel (2014); Sies et al. (2017); AOP-Wiki (2026c).
Subcellular localization of ROS	Mitochondria, chloroplasts, peroxisomes, membranes, nuclei and phagosomes differ in ROS production and local antioxidant buffering.	Localized ROS production can cause oxidative stress in a specific compartment even when whole-cell ROS measurements are modest.	Murphy (2009); Griendling et al. (2016).
Exposure duration and recovery time	Acute pulses, chronic low-level exposure and repeated stress can produce different redox outcomes.	Short pulses may be buffered or adaptive; sustained or repeated ROS elevations increase the probability of oxidative stress.	Sies et al. (2017); Ouillon et al. (2021).
Oxygen availability and hypoxia/reoxygenation	Oxygen tension affects mitochondrial electron transport and ROS formation.	Reoxygenation after hypoxia can increase mitochondrial ROS and enhance oxidative stress.	Ouillon et al. (2021).
Temperature and metabolic rate	Temperature and metabolic demand alter oxygen flux, mitochondrial activity and antioxidant capacity.	Higher metabolic activity or thermal stress can increase ROS formation and shift the balance toward oxidative stress.	Tseng et al. (2011).
Stressor chemistry	Redox cycling, metal-catalyzed reactions, radiation and mitochondrial inhibition generate ROS by different mechanisms.	Stressor type influences the ROS species, localization, time course and threshold for oxidative stress.	Bedard and Krause (2007); Murphy (2009); Qian et al. (2009); Gao et al. (2022).

Known Feedforward/Feedback loops influencing this KER

Known feedback and feedforward mechanisms influence the linkage. NRF2-dependent antioxidant responses can reduce ROS and restore homeostasis, whereas mitochondrial dysfunction, lipid peroxidation, inflammation and redox-sensitive signaling can amplify ROS generation and sustain oxidative stress. These feedbacks make the KER dynamic and nonlinear, particularly under chronic exposure or repeated stress.

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Relationship: 3632: Increase, Oxidative Stress leads to Increase, Protein oxidation

AOPs Referencing Relationship

AOP Name	Adjacency	Weight of Evidence	Quantitative Understanding
Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	adjacent	High	
Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and cell injury/death	adjacent		
Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell growth	adjacent		
Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and reduced cell proliferation	adjacent		

AOP Name	Adjacency	Weight of Evidence	Quantitative Understanding
Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell cycle disruption	adjacent		
Increase in reactive oxygen species leading to growth inhibition via protein oxidation and decreased cell proliferation	adjacent	High	Moderate
Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	adjacent	High	Moderate
ROS formation leading to growth inhibition modulated by the Keap1-Nrf2 antioxidant response	adjacent	High	Moderate

Evidence Supporting Applicability of this Relationship

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
humans	Homo sapiens	High	NCBI
mammals	mammals	High	NCBI
fish	fish	High	NCBI
crustaceans	Daphnia magna	High	NCBI
green algae	Ulva compressa	Moderate	NCBI

Life Stage Applicability

Life Stage	Evidence
All life stages	Moderate

Sex Applicability

Sex	Evidence
Unspecific	Moderate

This KER is broadly applicable to aerobic biological systems in which oxidative stress and protein oxidation can be measured. It is particularly relevant to tissues and cellular compartments exposed to high oxidant flux, including mitochondria, chloroplasts, peroxisomes, inflammatory cells, gill and digestive tissues, nervous tissues and rapidly metabolizing cells. The relationship is expected to be conserved because it is based on fundamental redox chemistry and protein chemistry rather than on a taxon-specific receptor or signaling pathway.

The KER should be applied with greatest confidence when upstream oxidative stress is assessed using direct or mechanistically interpretable redox endpoints and downstream protein oxidation is measured using specific markers such as protein carbonyls, oxidized thiols, methionine oxidation, AOPP, or redox proteomics. Applicability is weaker when protein oxidation is inferred only from broad stress responses or when oxidative stress and protein oxidation are not measured in the same biological context. Species, life stage and sex should be considered mainly as modifiers of sensitivity rather than determinants of whether the relationship can occur.

Key Event Relationship Description

This KER describes the relationship by which an increase in oxidative stress leads to increased protein oxidation. Oxidative stress represents a state in which oxidant generation or antioxidant depletion shifts the biological system toward a pro-oxidant condition. Under these conditions, reactive oxygen and nitrogen species, lipid-derived reactive aldehydes, metal-catalyzed oxidants and oxidized thiol/disulfide systems can modify proteins directly or indirectly. Protein oxidation includes irreversible modifications such as protein carbonyl formation, oxidation of aromatic and sulfur-containing amino acids, backbone fragmentation, crosslinking and aggregation, as well as reversible or regulatory modifications such as disulfide formation, S-glutathionylation, S-nitrosylation and other redox post-translational modifications (Stadtman and Levine, 2003; Dalle-Donne et al., 2006; Davies, 2016).

The relationship is biologically plausible because proteins are abundant cellular targets and many amino acid side chains react with oxidants or with secondary products of oxidative stress. Increased oxidative stress raises the probability that susceptible proteins will undergo oxidation, particularly when antioxidant defenses, reductive repair systems, proteasomal degradation or protein turnover cannot maintain proteostasis. The downstream KE therefore reflects a measurable biochemical consequence of the upstream oxidative-stress state.

Evidence Supporting this KER

Biological Plausibility

Biological plausibility of this KER is high. Oxidative stress produces or reflects oxidizing conditions that can modify proteins through multiple well-established chemical mechanisms. Hydroxyl radicals, peroxy radicals, singlet oxygen, hypochlorous acid, peroxynitrite and metal-catalyzed oxidants can oxidize amino acid side chains, while secondary products of lipid peroxidation, such as reactive aldehydes, can form protein adducts and carbonyl derivatives. These processes produce measurable protein carbonyls, oxidized methionine, oxidized cysteine residues, disulfides,

protein hydroperoxides, crosslinks and fragmented or aggregated proteins (Stadtman and Levine, 2003; Dalle-Donne et al., 2006; Davies, 2016).

The structural and functional relationship between the two KEs is direct. The upstream KE increases the oxidizing chemical environment, and the downstream KE is the covalent modification of protein targets under that oxidizing environment. Because proteins are abundant and essential for enzyme activity, signaling, structural integrity and energy metabolism, protein oxidation is a broadly expected consequence of oxidative stress when protective and repair mechanisms are insufficient.

Empirical Evidence

Empirical support for this KER is moderate to high. Multiple studies in diverse systems show that oxidative-stress conditions coincide with or precede increases in protein oxidation markers, especially protein carbonylation, oxidized thiols or glutathionylated proteins. The strongest evidence comes from experiments in which oxidative stress biomarkers and protein oxidation endpoints were measured in the same biological system and exposure context. However, the empirical evidence is not uniformly high because many studies measure protein oxidation alone as an oxidative damage endpoint, without direct upstream ROS or redox measurements in the same time course.

Biological system	Stressor or condition	Evidence relevant to KER	Interpretation
<i>Chlamydomonas reinhardtii</i>	Cadmium or hydrogen peroxide / oxidative stress conditions	Proteomic analyses identified protein carbonylation and redox modifications including glutathionylation of photosynthetic and metabolic proteins under oxidative stress conditions (Zaffagnini et al., 2012).	Supports occurrence of protein oxidation under oxidative-stress conditions in photosynthetic eukaryotes.
Zebrafish brain	Acute cold exposure	Protein carbonyls increased by 38% within 1 h after cold exposure, with increased antioxidant response markers over the same early time frame (Tseng et al., 2011).	Supports temporal association between oxidative stress response and protein oxidation in fish.
Freshwater fish <i>Channa punctata</i>	Deltamethrin, endosulfan and paraquat	Protein carbonyls were proposed and measured as biomarkers of exposure to oxidative-stress-inducing pesticides (Parvez and Raisuddin, 2005).	Supports stressor-induced protein oxidation in fish exposed to pro-oxidant pesticides.
<i>Mytilus galloprovincialis</i> hemocytes	Cadmium or 17 beta-estradiol	Redox parameters were altered by micromolar concentrations of stressors, consistent with oxidative stress and linked signaling processes in mussel hemocytes (Koutsogiannaki et al., 2014).	Supports relevance of oxidative stress/protein-oxidation processes in molluscan immune cells.
Mammalian / human cell systems	Hydrogen peroxide and related oxidants	Live-cell fluorescent detection approaches demonstrate oxidative stress-induced carbonylation of biomolecules, including proteins (Mukherjee et al., 2015).	Supports direct oxidative stress-induced carbonylation in mammalian cell systems.

Uncertainties and Inconsistencies

A major uncertainty is that protein oxidation comprises many different chemical modifications with different reversibility, biological consequences and measurement approaches. Protein carbonyls are widely used as relatively stable markers, but they represent only one subset of oxidative protein damage. Thiol oxidation, methionine oxidation and glutathionylation may be reversible or regulatory, while carbonylation and aggregation are often associated with irreversible damage. Therefore, different studies may use different operational definitions of protein oxidation, making quantitative comparison difficult (Dalle-Donne et al., 2006; Davies, 2016).

A second uncertainty is that oxidative stress is often inferred from antioxidant enzyme activity, glutathione status or damage endpoints rather than directly measured ROS flux. As a result, some empirical studies demonstrate co-occurrence of oxidative-stress markers and protein oxidation but cannot establish the exact sequence of events. Conversely, protein oxidation may arise secondarily from lipid peroxidation products, inflammation, metal-catalyzed reactions or impaired protein turnover, so the upstream oxidative-stress KE should be interpreted as a redox-state driver rather than a single chemical species. No strong contradictory evidence was identified for the general relationship that oxidative stress can increase protein oxidation.

Quantitative Understanding of the Linkage

Quantitative understanding of this KER is moderate. The qualitative biochemical relationship between oxidative stress and protein oxidation is well established, and response-response relationships exist in some experimental systems.

Response-response relationship

However, a general quantitative function predicting the magnitude of protein oxidation from a given oxidative-stress measurement has not been established across taxa, tissues, protein classes, stressors and assay methods. Quantitative prediction is complicated because the upstream KE can be measured by multiple endpoints, including ROS probes, glutathione status, antioxidant enzyme responses or pathway activation, while the downstream KE can be measured by protein carbonyls, oxidized thiols, methionine oxidation, glutathionylation, AOPP or redox proteomics.

Time-scale

The time scale of the linkage can range from minutes to days. Oxidation of susceptible amino acid residues may occur rapidly during an acute oxidant pulse, whereas accumulation of stable carbonylated proteins, protein aggregates or proteomic changes may require longer exposure or exceed the capacity of repair and degradation systems. In zebrafish exposed to acute cold stress, protein carbonylation increased within 1 h, showing that the downstream KE can occur rapidly in vivo under oxidative-stress conditions (Tseng et al., 2011). In *Chlamydomonas* and mammalian cell systems, protein oxidation and carbonylation are also detectable under defined pro-oxidant exposure conditions (Zaffagnini et al., 2012; Mukherjee et al., 2015).

Known modulating factors

Modulating factor	Details	Influence on KER	Supporting evidence
Antioxidant and reductive capacity	Glutathione, thioredoxin, glutaredoxin, peroxiredoxins, catalase, superoxide dismutase and related systems.	Higher antioxidant/reductive capacity decreases the probability or magnitude of protein oxidation for a given oxidative challenge; depletion increases sensitivity.	Sies et al., 2017; Rouhier et al., 2015; Zaffagnini et al., 2012.
Metal availability	Iron, copper, cadmium and other redox-active or thiol-reactive metals.	Transition metals and thiol-reactive metals can promote site-specific oxidation, protein carbonylation or altered thiol redox state.	Stadtman and Levine, 2003; Parvez and Raisuddin, 2005; Koutsogiannaki et al., 2014.
Protein composition and localization	Proteins rich in cysteine, methionine, aromatic residues or metal-binding sites; mitochondrial, chloroplast and membrane-associated proteins.	Susceptible proteins and proteins located near ROS sources are more likely to undergo oxidation.	Davies, 2016; Dalle-Donne et al., 2006.
Proteostasis and repair capacity	Proteasome activity, autophagy, methionine sulfoxide reductases, thiol-disulfide exchange systems and protein turnover.	Efficient repair and degradation can reduce accumulation of oxidized proteins even when oxidative stress occurs.	Dalle-Donne et al., 2006; Davies, 2016.
Exposure duration and intensity	Acute versus chronic oxidative stress; pulse versus sustained oxidant generation.	Longer or more intense oxidative stress increases accumulation of stable oxidative protein damage, especially carbonyls and aggregates.	Mukherjee et al., 2015; Tseng et al., 2011.

Known Feedforward/Feedback loops influencing this KER

The linkage is expected to be nonlinear and threshold-dependent. Low or transient oxidative stress may lead to reversible redox signaling or repairable thiol modifications, whereas stronger or persistent oxidative stress is more likely to cause irreversible carbonylation, aggregation or loss of protein function. Quantitative evaluation is therefore strongest when upstream oxidative stress and downstream protein oxidation are measured in the same biological system across multiple exposure concentrations and time points.

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Relationship: 3633: Increase, Protein oxidation leads to Decrease, Coupling of OXPHOS

AOPs Referencing Relationship

AOP Name	Adjacency	Weight of Evidence	Quantitative Understanding
Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	adjacent		
Increase in reactive oxygen species leading to growth inhibition via protein oxidation and decreased cell proliferation	adjacent	Moderate	Low
Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	adjacent	Moderate	Low

Evidence Supporting Applicability of this Relationship

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
humans	Homo sapiens	Moderate	NCBI
mammals	mammals	Moderate	NCBI
fish	fish	Moderate	NCBI
crustaceans	Daphnia magna	Moderate	NCBI
green algae	Ulva compressa	Moderate	NCBI

Life Stage Applicability

Life Stage	Evidence
All life stages	Moderate

Sex Applicability

Sex	Evidence
Unspecific	Moderate

This KER is most applicable to aerobic eukaryotic cells and tissues in which mitochondria are important for ATP production and in which protein oxidation affects proteins involved in mitochondrial respiration, membrane potential, substrate transport or ATP synthesis. It is applicable across a broad range of taxa because the underlying chemistry of protein oxidation and the core architecture of OXPHOS are conserved. Applicability is strongest when the upstream KE is measured using specific protein oxidation endpoints, such as protein carbonyls, oxidized thiols, nitrated proteins, methionine oxidation, glutathionylation or redox proteomics, and when the downstream KE is measured using mechanistically informative mitochondrial endpoints such as membrane potential, oxygen consumption rate, respiratory control ratio, proton leak, ATP-linked respiration, or complex activity.

Confidence is lower when protein oxidation is measured only as a nonspecific bulk endpoint, when mitochondrial dysfunction is measured only as general cytotoxicity, or when the two KEs are not measured in the same biological system. The KER should also be interpreted cautiously under conditions where direct chemical uncoupling, lipid peroxidation, mitochondrial DNA damage, or generalized cell injury may be the dominant cause of decreased OXPHOS coupling.

Key Event Relationship Description

This KER describes the relationship by which increased protein oxidation leads to decreased coupling of oxidative phosphorylation. Protein oxidation refers to oxidative modification of protein amino acid residues or protein-associated cofactors, including carbonylation, thiol oxidation, methionine oxidation, tyrosine nitration, protein-peroxide formation, glutathionylation, and adduction by reactive aldehydes generated during lipid peroxidation. When such modifications affect mitochondrial proteins involved in electron transport, proton pumping, substrate transport, ATP synthase function, or maintenance of the inner mitochondrial membrane potential, OXPHOS efficiency can decline.

The downstream KE, decreased coupling of OXPHOS, describes a reduction in the efficiency with which electron transport and protonmotive force are coupled to ATP synthesis. AOP-Wiki Event 1446 describes this KE as dissipation or impairment of the protonmotive force across the inner mitochondrial membrane, measurable through decreased mitochondrial membrane potential, increased proton leak, altered oxygen consumption, or reduced respiratory control (AOP-Wiki, 2026c). Protein oxidation can contribute to this KE by impairing respiratory chain complexes, phosphate or nucleotide transporters, ATP synthase, redox cofactors, or mitochondrial membrane-associated proteins. This KER therefore links molecular damage to proteins with a cellular bioenergetic consequence.

The relationship is not intended to imply that all protein oxidation is adverse or that all oxidized proteins impair OXPHOS. Many reversible thiol modifications participate in redox regulation. The KER is most applicable when protein oxidation is persistent, extensive, affects mitochondrial or bioenergetic proteins, or exceeds cellular repair, reduction, and proteolytic capacity.

Evidence Supporting this KER**Biological Plausibility**

Biological plausibility of this KER is high. Proteins are major targets of oxidants because they are abundant, contain redox-active residues and cofactors, and often catalyze or participate in electron-transfer reactions. Reactive oxygen and nitrogen species, metal-catalyzed oxidants, lipid-derived aldehydes, and protein peroxides can modify cysteine, methionine, histidine, lysine, arginine, tyrosine and other residues, resulting in altered protein conformation, catalytic activity, complex assembly, stability, or degradation (Stadtman and Levine, 2003; Dalle-Donne et al., 2006; Davies, 2016).

Mitochondrial OXPHOS is particularly vulnerable to protein oxidation because it relies on multi-subunit protein complexes embedded in the inner mitochondrial membrane, iron-sulfur clusters, redox-active cofactors, substrate and nucleotide transporters, and maintenance of a protonmotive force. Oxidative modification of respiratory-chain subunits or transport proteins can reduce electron transfer, increase electron leak, impair proton pumping, alter substrate availability, or decrease membrane potential, thereby reducing coupling efficiency. Curtis et al. (2012) provided direct mechanistic evidence in 3T3-L1 adipocytes that increased carbonylation of mitochondrial proteins, including complex I-related proteins and transport proteins, was accompanied by decreased complex I activity, impaired respiration and reduced mitochondrial membrane potential. This provides a strong mechanistic bridge from the upstream KE to the downstream KE.

The relationship is also coherent with the broader OXPHOS AOP module. Decreased coupling of OXPHOS is a recognized measurable KE in AOP-Wiki and in the OECD-endorsed OXPHOS uncoupling leading to growth inhibition AOP. Although classical uncouplers act primarily through protonophoric mechanisms, oxidative damage to mitochondrial proteins provides an additional route to reduced coupling efficiency (AOP-Wiki, 2026c; OECD, 2022).

Empirical Evidence

Empirical support for this KER is moderate. The strongest empirical evidence comes from studies in which increased mitochondrial protein carbonylation or oxidative protein damage is measured together with reduced mitochondrial membrane potential, impaired respiration, decreased complex activity, or reduced coupling efficiency. However, many studies report either protein oxidation or mitochondrial dysfunction without measuring both KEs in a manner

that allows complete temporal, dose-response and incidence concordance assessment.

Evidence type	Summary of evidence	References
Direct mechanistic evidence in mammalian cells	GSTA4-silenced 3T3-L1 adipocytes displayed elevated carbonylation of mitochondrial proteins, including NADH dehydrogenase 1 alpha subcomplexes and phosphate carrier protein. Elevated protein carbonylation was accompanied by diminished complex I activity, impaired respiration, increased superoxide production and reduced mitochondrial membrane potential. Knockdown of selected carbonylation targets reduced basal and maximal respiration.	Curtis et al., 2012
Association in invertebrate life-history and mitochondrial function	Short-lived <i>Daphnia pulex</i> clones showed reduced complex I activity, increased oxidative damage and altered expression of ROS-scavenging enzymes. This supports an association between oxidative damage to cellular components and impaired mitochondrial respiratory function, although it does not isolate protein oxidation as the sole cause.	Ukhueduan et al., 2022
Bivalve hypoxia-reoxygenation evidence	Hypoxia-reoxygenation stress in the oyster <i>Crassostrea gigas</i> induced mitochondrial proteome and phosphoproteome shifts together with altered bioenergetic responses. This supports environmental relevance of oxidative/proteomic stress coupled to mitochondrial bioenergetic impairment, but direct protein oxidation-to-OXPHOS causality is not fully resolved.	Sokolov et al., 2019
Fish oxidative stress and mitochondrial response	Acute cold exposure in zebrafish brain induced oxidative stress responses and changes in uncoupling-protein/antioxidant mechanisms; protein carbonylation increased rapidly in the time course. This supports temporal feasibility of oxidative protein damage in relation to mitochondrial stress responses but does not provide a fully quantitative KER model.	Tseng et al., 2011
Photosynthetic eukaryote evidence	Large-scale redox proteomics in <i>Chlamydomonas reinhardtii</i> identified extensive protein glutathionylation under oxidative conditions, showing broad susceptibility of cellular proteins to redox modification. Evidence directly linking these modifications to decreased mitochondrial coupling in the same study is limited.	Zaffagnini et al., 2012

Uncertainties and Inconsistencies

A key uncertainty is that protein oxidation is chemically diverse. Reversible thiol oxidation and glutathionylation can act as regulatory or protective modifications, whereas carbonylation, nitration, aggregation or irreversible oxidation are more likely to be associated with functional impairment. As a result, the biological consequence of the upstream KE depends strongly on the specific protein target, modification type, dose, duration and cellular context.

A second uncertainty is that decreased coupling of OXPHOS may result from multiple upstream mechanisms, including lipid peroxidation, direct chemical uncoupling, mitochondrial DNA damage, calcium dysregulation, permeability transition, complex inhibition, or changes in mitochondrial dynamics. Protein oxidation may be causal, contributory, or secondary to these other mechanisms. Empirical support is strongest when oxidative modification of mitochondrial proteins is measured together with respiratory endpoints, but such studies remain relatively limited across ecotoxicological species.

Temporal concordance may also be difficult to establish. Protein oxidation of susceptible residues can occur within minutes to hours, but detectable impairment of OXPHOS coupling may require accumulation of damage, modification of key targets, or failure of repair and proteolytic systems. Conversely, mitochondrial dysfunction can increase ROS production and promote further protein oxidation, creating a feedforward loop that complicates the

assignment of a strictly unidirectional sequence.

Quantitative Understanding of the Linkage

Quantitative understanding of this KER is low to moderate. The qualitative linkage between oxidative modification of mitochondrial proteins and impaired mitochondrial coupling is well supported, and individual studies provide quantitative data on protein carbonylation, complex I activity, oxygen consumption and mitochondrial membrane potential. However, there is currently no generalizable mathematical model that predicts the magnitude of decreased OXPHOS coupling from a given amount of total protein oxidation across taxa, tissues, stressors and measurement platforms.

Response-response relationship

The response-response relationship is expected to be nonlinear and target-dependent. Total protein carbonyls or other bulk oxidation markers may correlate poorly with OXPHOS impairment if oxidation occurs mainly in proteins unrelated to mitochondrial respiration. Conversely, relatively small amounts of oxidation affecting key respiratory-chain subunits, ATP synthase, inner membrane transporters, or proteins required for maintenance of mitochondrial membrane potential may have substantial bioenergetic consequences. Curtis et al. (2012) provide a strong example of response-response evidence because elevated carbonylation of specific mitochondrial proteins was accompanied by reduced complex I activity, altered oxygen consumption and reduced membrane potential.

Time-scale

The time scale can range from minutes to days. Oxidation of susceptible mitochondrial protein residues may occur rapidly during an oxidant pulse, while measurable decreases in coupling efficiency may appear after sufficient oxidation of functionally important targets or after compensatory mechanisms are overwhelmed. In vivo studies of oxidative stress responses in fish show that protein oxidation can increase within hours under acute stress (Tseng et al., 2011), whereas environmentally relevant hypoxia-reoxygenation or chronic oxidative damage may alter mitochondrial proteome and function over longer time scales (Sokolov et al., 2019; Ukhueduan et al., 2022).

Known modulating factors

Modulating factor	Details	Influence on KER	Supporting evidence
Protein target and modification type	Carbonylation, nitration and irreversible oxidation of mitochondrial proteins are more likely to impair function than transient reversible thiol modifications.	Alters magnitude and probability of downstream OXPHOS impairment; oxidation of respiratory-chain subunits or transporters has higher expected impact.	Stadtman and Levine, 2003; Dalle-Donne et al., 2006; Davies, 2016; Curtis et al., 2012
Antioxidant and reductive repair capacity	Glutathione, thioredoxin, peroxiredoxins, methionine sulfoxide reductases and related systems can reverse or limit some oxidative protein modifications.	Higher antioxidant/repair capacity raises the threshold for downstream mitochondrial impairment.	Sies et al., 2017; Davies, 2016
Proteostasis capacity	Proteasomal and mitochondrial protein quality-control systems remove damaged proteins; reduced turnover permits accumulation.	Impaired proteostasis increases persistence of oxidized mitochondrial proteins and may increase downstream effect size.	Dalle-Donne et al., 2006; Davies, 2016
Mitochondrial abundance and energy demand	Cells with high mitochondrial density or high ATP demand may show stronger consequences of oxidation of OXPHOS proteins.	May increase sensitivity of downstream coupling endpoints to upstream protein oxidation.	Murphy, 2009; Nicholls and Ferguson, 2013

Exposure duration and intensity	Short transient oxidant pulses may cause reversible modification; persistent or high-intensity exposures can produce irreversible carbonylation and dysfunction.	Determines whether protein oxidation remains adaptive/regulatory or becomes damaging and functionally linked to OXPHOS impairment.	Davies, 2016; Curtis et al., 2012
Temperature, hypoxia-reoxygenation and oxygen availability	Environmental oxygen fluctuations and temperature stress affect ROS production, mitochondrial function and protein oxidation.	Can amplify oxidative modification and alter the timing and magnitude of downstream mitochondrial impairment.	Tseng et al., 2011; Sokolov et al., 2019

Known Feedforward/Feedback loops influencing this KER

A biologically important feedforward loop may occur because impairment of mitochondrial OXPHOS can increase electron leak and ROS production, which can further oxidize mitochondrial proteins. This loop can amplify the KER once mitochondrial protein oxidation begins to impair electron transport or membrane coupling. Negative feedback or adaptive responses may include activation of antioxidant pathways, increased protein turnover, mitophagy, mitochondrial biogenesis, and metabolic compensation through glycolysis. These feedback mechanisms are expected to influence the threshold and persistence of the downstream KE but are not yet sufficiently quantified for general application.

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Relationship: 2203: Decrease, Coupling of OXPHOS leads to Decrease, ATP pool

AOPs Referencing Relationship

AOP Name	Adjacency	Weight of Evidence	Quantitative Understanding
Uncoupling of oxidative phosphorylation leading to growth inhibition via decreased cell proliferation	adjacent	High	High
Uncoupling of oxidative phosphorylation leading to growth inhibition via ATP depletion associated cell death	adjacent	Moderate	Not Specified
Uncoupling of oxidative phosphorylation leading to growth inhibition via decreased Na-K ATPase activity	adjacent		
Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and cell death	adjacent	High	High
Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	adjacent		
Peroxisome proliferator-activated receptor alpha activation leading to early life stage mortality via reduced adenosine triphosphate	adjacent		
Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and decreased cell proliferation	adjacent	High	High
Increase in reactive oxygen species leading to growth inhibition via protein oxidation and decreased cell proliferation	adjacent	High	High
Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	adjacent	High	High

Evidence Supporting Applicability of this Relationship

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
zebrafish	Danio rerio	High	NCBI
human	Homo sapiens	High	NCBI
rat	Rattus norvegicus	High	NCBI
mouse	Mus musculus	High	NCBI

Life Stage Applicability

Life Stage	Evidence
Embryo	High
Juvenile	High

Sex Applicability

Sex	Evidence
Unspecific	High

Taxonomic applicability

Relationship 2203 is considered applicable to eukaryotes, as mitochondrial oxidative phosphorylation and ATP synthesis are highly conserved in these organisms. Uncoupling of oxidative phosphorylation leading to ATP depletion is a well-documented relationship in many taxa, such as human, rodents and fish.

Sex applicability

Relationship 2203 is considered applicable to all genders, as mitochondrial oxidative phosphorylation and ATP synthesis are fundamental biological processes and are not sex-specific.

Life-stage applicability

Relationship 2203 is considered applicable to all life-stages, as mitochondrial oxidative phosphorylation and ATP synthesis are essential energy production processes for maintaining basic biological activities.

Key Event Relationship Description

This key event relationship describes the dissipation of protonmotive force across the inner mitochondrial membrane by uncouplers (uncoupling of oxidative phosphorylation), leading to reduced total adenosine triphosphate (ATP) pool in cells or organisms.

Evidence Supporting this KER

The overall evidence supporting Relationship 2203 is considered high.

Biological Plausibility

The biological plausibility of Relationship 2203 is considered high.

Rationale: In eukaryotic cells, the major metabolic pathways responsible for ATP production are OXPHOS, citric acid (TCA) cycle, glycolysis and photosynthesis. Oxidative phosphorylation is much (theoretically 15-18 times) more efficient than the rest due to high energy derived from oxygen during aerobic respiration (Schmidt-Rohr 2020). As the ATP level is relatively balanced between production and consumption (Bonora 2012), ATP depletion is a plausible consequence of reduced ATP synthetic efficiency following uncoupling of OXPHOS.

Empirical Evidence

The empirical support of Relationship 2203 is considered high.

Rationale: The majority of relevant studies show good incidence, temporal and/or dose concordance in different organisms and cell types after exposure to known uncouplers, with relatively few exceptions.

Evidence:

- **Temporal concordance:** Exposure of zebrafish embryos to 0.5 μM of the classical uncoupler 2,4-DNP led to significantly uncoupling of OXPHOS after 21h, whereas significant reduction in ATP was only observed after 45h (Bestman 2015).
- **Dose concordance:** The uncoupler triclosan induced significant uncoupling of OXPHOS in zebrafish embryos at 15 μM , whereas higher (30 μM) concentration was required to caused significant ATP depletion (Shim 2016).
- **Dose concordance:** Exposure to 1 μM of the uncoupler CCCP led to 40% uncoupling of OXPHOS in rat RBL-2H3 cells, whereas the same magnitude of effect for ATP reduction required 1.6 μM of CCCP (Weatherly 2016).
- **Dose concordance:** Exposure to 10 μM of the uncoupler triclosan caused significant uncoupling of OXPHOS in rat RBL-2H3 cells, whereas significant reduction in ATP was observed at a higher concentration (30 μM) (Weatherly 2018).
- **Dose concordance:** Significant effect on uncoupling of OXPHOS required 2 μM FCCP, whereas a significant reduction in ATP required 20 μM FCCP in human RD cells (Kuruvilla 2003).
- **Incidence concordance:** In human colon cancer cells (SW480), exposure to 150 μM of the uncoupler flavanoid morin caused 60% reduction in MMP, whereas only around 35% decrease in ATP (Sithara 2017).
- **Incidence concordance:** Exposure of rat RBL-2H3 cells to 10 μM of the uncoupler triclosan led to 50% uncoupling of OXPHOS, whereas only 40% reduction in ATP (Weatherly 2016).
- **Incidence concordance:** Exposure to 5 μM of the uncoupler CCCP caused 71% uncoupling of OXPHOS, whereas only 64% reduction of ATP in human HL-60 cells (Sweet 1999).
- **Incidence concordance:** Exposure of human HeLa cells to 50 μM of the uncoupler CCCP for 1h led to 77% uncoupling of OXPHOS and 25% reduction in ATP (Koczor 2009).
- **Incidence concordance:** Exposure of the nematode *Caenorhabditis elegans* to 50 μM Arsenite for 1h led to approximately 45% uncoupling of OXPHOS and 20% reduction in ATP (Luz 2016).

Uncertainties and Inconsistencies

- A significant decrease followed by a significant increase in total ATP was observed in human RD cells during a 48h exposure to the uncoupler FCCP (Kuruvilla 2003), possibly due to the enhancement of other ATP synthetic pathways (e.g., glycolysis) as a compensatory action to impaired OXPHOS (Jose 2011)

Quantitative Understanding of the Linkage

The quantitative understanding of Relationship 2203 is high.

Rationale: Multiple mathematical models have been developed for describing the quantitative relationships between uncoupling of OXPHOS and ATP synthesis in vertebrates (Beard 2005; Schmitz 2011; Heiske 2017; Kubo 2020). These models, however, are highly complex metabolic or systems biological models and warrant further simplification to be used for this AOP.

Response-response relationship

A regression based quantitative response-response relationship between uncoupling of OXPHOS and ATP depletion was proposed for the crustacean *Daphnia magna* under UVB stress (Song 2020).

Known Feedforward/Feedback loops influencing this KER

- It is known that mild uncoupling of oxidative phosphorylation can enhance the activity of the mitochondrial electron transport chain to produce more ATP, and/or activate other ATP synthetic pathways (e.g., glycolysis) as a compensatory action to impaired OXPHOS (Jose 2011).

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Relationship: 2768: Decrease, ATP pool leads to Cell injury/death

AOPs Referencing Relationship

AOP Name	Adjacency	Weight of Evidence	Quantitative Understanding
Uncoupling of oxidative phosphorylation leading to growth inhibition via ATP depletion associated cell death	adjacent	Moderate	Not Specified
Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and cell death	adjacent	High	Moderate
Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	adjacent		
Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and cell injury/death	adjacent		
Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	adjacent	High	Moderate

Evidence Supporting Applicability of this Relationship

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
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Term	Scientific Term	Evidence	Links
humans	Homo sapiens	High	NCBI
mammals	mammals	High	NCBI
fish	fish	High	NCBI
crustaceans	Daphnia magna	High	NCBI
green algae	Ulva compressa	High	NCBI

Life Stage Applicability

Life Stage	Evidence
All life stages	Moderate

Sex Applicability

Sex	Evidence
Unspecific	Moderate

The biological domain of applicability is broad because ATP-dependent homeostasis is a conserved property of living cells. The KER is most directly applicable to eukaryotic cells and tissues in which mitochondrial and/or glycolytic ATP supply maintains cellular viability. It is particularly relevant to metabolically active tissues and developing organisms where energy demand is high. It is applicable to both sexes and to multiple life stages, although sensitivity may differ with developmental status, tissue type, temperature, oxygen availability, and metabolic reserve.

The chemical and stressor applicability domain includes stressors that reduce cellular ATP through mitochondrial inhibition, OXPHOS uncoupling, oxidative stress, membrane disruption, calcium overload, metabolic poisons, hypoxia or other mechanisms that impair ATP synthesis or increase ATP demand beyond compensatory capacity. In the ROS-growth AOP network, this KER is most relevant downstream of OXPHOS impairment caused by lipid peroxidation or protein oxidation, where energetic failure contributes to increased cell injury/death.

Key Event Relationship Description

This key event relationship describes the causal and predictive link by which a decrease in the cellular adenosine triphosphate (ATP) pool leads to increased cell injury and/or cell death. ATP is required to maintain ion gradients, plasma membrane integrity, mitochondrial homeostasis, macromolecular repair, vesicular trafficking, and regulated cell death programs. When ATP depletion is sufficiently severe or prolonged, energy-dependent adaptive and repair processes fail, calcium and sodium homeostasis are disrupted, mitochondrial permeability transition may be promoted, and cells may undergo apoptosis, necrosis, necroptosis-like injury or mixed forms of cell death depending on cellular context and residual ATP availability (Niemenen et al., 1994; Leist et al., 1997; Bonora et al., 2012).

The direction of this KER is from reduced ATP availability to increased cell injury/death. The KER is not intended to specify a single mode of cell death. Rather, it captures the general biological principle that loss of cellular energy supply increases the probability of irreversible cellular injury and death, with the exact death phenotype depending on cell type, severity of ATP depletion, duration of exposure, and availability of death-execution pathways.

Evidence Supporting this KER

The overall evidence supporting this KER is considered moderate to high. Biological plausibility is high because ATP is indispensable for cellular homeostasis and because severe ATP depletion is a well-established trigger of irreversible cell injury and death. Empirical support is moderate to high because multiple studies in mammalian cells, algae, aquatic organisms and cancer cell systems demonstrate concordance between ATP depletion and cell injury/death; however, the exact quantitative threshold varies substantially across biological systems and exposure conditions.

Biological Plausibility

Biological plausibility is high. ATP depletion compromises core cellular maintenance processes including ion pumping, membrane integrity, cytoskeletal dynamics, protein turnover, DNA repair, and mitochondrial function. When ATP supply falls below the level required for homeostasis, cells lose the ability to maintain electrochemical gradients and to execute energy-dependent adaptive responses. Severe energetic collapse promotes necrotic injury, while partial ATP depletion may permit regulated apoptotic execution depending on residual ATP availability and caspase competence (Niemenen et al., 1994; Leist et al., 1997; Nicotera et al., 1998; Zong and Thompson, 2006).

The mechanistic relationship is also supported by mitochondrial cell-death biology. ATP depletion often accompanies mitochondrial membrane depolarization, permeability transition, impaired oxidative phosphorylation, calcium dysregulation, and increased reactive oxygen species generation. These processes can amplify cellular injury and increase the probability of cell death (Kroemer et al., 1998; Green and Kroemer, 2004; Halestrap, 2009; Bonora et al., 2012).

Empirical Evidence

Empirical support is moderate to high. In mammalian systems, ATP depletion has been directly linked to cell killing after metabolic inhibition, and experimental work has shown that ATP depletion rather than mitochondrial depolarization can mediate hepatocyte death under some conditions (Niemenen et al., 1994). A widely cited study

demonstrated that intracellular ATP concentration influences whether cells die by apoptosis or necrosis, supporting both causality and phenotype dependence (Leist et al., 1997). Calcium electroporation studies provide dose-dependent evidence that ATP depletion is associated with reduced cancer cell survival and increased cell death (Hansen et al., 2015).

Evidence from environmental and ecotoxicological systems is consistent with this relationship. In *Chlamydomonas reinhardtii*, herbicide exposure produced ATP depletion and cell injury/death in a multiple-endpoint assay, demonstrating concordance between energetic disruption and cellular toxicity in an algal model (Nestler et al., 2012). In eastern oysters, cadmium exposure affected mitochondrial bioenergetics and was associated with cellular damage endpoints, supporting applicability of energetic failure to cell injury in aquatic invertebrates (Sokolova et al., 2005). In ROS-growth concordance data, mitochondrial toxicants and oxidative stressors including paraquat, rotenone, cadmium and hydrogen peroxide frequently produce decreased ATP or mitochondrial dysfunction together with cytotoxicity or tissue injury, although direct measurement of both KEs in the same study is not always available.

Evidence type	Summary	Representative references
Biological plausibility	ATP is required for ion homeostasis, membrane maintenance, repair, and regulated cell death execution; severe ATP depletion promotes irreversible cell injury/death.	Nieminen et al. 1994; Leist et al. 1997; Bonora et al. 2012
Temporal concordance	ATP depletion can occur rapidly after metabolic inhibition or mitochondrial impairment and precedes detectable loss of viability or death execution in several cell systems.	Nieminen et al. 1994; Hansen et al. 2015
Dose-response concordance	Increasing intensity of energetic perturbation or calcium electroporation increases ATP depletion and cell killing.	Hansen et al. 2015
Incidence concordance	Systems showing marked ATP depletion commonly show increased cytotoxicity, cell injury or cell death, although moderate ATP depletion may be compensated in some contexts.	Leist et al. 1997; Nestler et al. 2012; Sokolova et al. 2005
Essentiality / intervention	Experimental data indicate that ATP availability influences the form and occurrence of cell death; restoration or maintenance of energy status can reduce injury in some systems, but direct rescue evidence across taxa remains limited.	Leist et al. 1997; Nicotera et al. 1998

Uncertainties and Inconsistencies

The main uncertainty is that ATP depletion is not the only cause of cell injury/death. Cell death may also be initiated by DNA damage, receptor-mediated apoptosis, oxidative damage, calcium overload, lysosomal injury, proteotoxic stress or inflammatory signaling. Consequently, the presence of cell injury/death does not uniquely imply ATP depletion. The KER is strongest when ATP decline occurs before or at lower concentrations than cell death and when the upstream energetic perturbation is mechanistically established.

Another uncertainty concerns severity thresholds. Moderate ATP depletion may be reversible or may shift cells into cell-cycle arrest, reduced proliferation, or adaptive metabolic compensation rather than death. Conversely, very severe ATP depletion may prevent the energy-requiring execution of apoptosis and produce necrotic injury instead. Therefore, the downstream phenotype depends on the magnitude and duration of ATP depletion and on cellular metabolic reserve (Leist et al., 1997; Nicotera et al., 1998).

Empirical evidence across environmental species remains less dense than evidence from mammalian cell systems. Many ecotoxicological studies measure ATP, mitochondrial dysfunction, or cytotoxicity separately rather than measuring both KEs in the same time- and dose-resolved experiment. This limits the strength of concordance assessment across the full taxonomic applicability domain.

Quantitative Understanding of the Linkage

The quantitative understanding of this KER is considered moderate. Quantitative evidence supports a general response-response relationship in which larger or longer decreases in ATP increase the probability and severity of cell injury/death. However, a single universal threshold cannot be defined because ATP demand, ATP reserve, glycolytic capacity, cell type, death pathway, and exposure duration vary substantially among biological systems.

Several studies support threshold-like behavior. In hepatocytes, ATP depletion mediated killing after metabolic inhibition, supporting a causal threshold relationship between energetic collapse and cell death (Nieminen et al., 1994). Experiments in human T cells showed that intracellular ATP concentration can act as a switch influencing apoptotic versus necrotic death phenotypes (Leist et al., 1997). Calcium electroporation studies showed dose-dependent ATP depletion and reduced survival, supporting a quantitative relationship between the upstream energetic disturbance and the downstream cell death outcome (Hansen et al., 2015).

Response-response relationship

The expected response-response relationship is generally monotonic but non-linear. Small or transient ATP reductions may be tolerated or compensated. Larger reductions increase the probability of cell stress, impaired repair, loss of membrane integrity, and cell death. At extreme ATP depletion, necrotic injury is favored, whereas intermediate depletion may permit energy-dependent apoptosis depending on cell type and execution machinery (Leist et al., 1997; Nicotera et al., 1998).

Time-scale

The time scale of ATP depletion can range from minutes to hours following direct mitochondrial inhibition, uncoupling, metabolic inhibition, or membrane-disrupting interventions. Observable downstream cell injury/death may occur within hours to days depending on cell type, severity of ATP loss, and endpoint measured. In whole organisms, cell death may contribute to tissue injury or growth impairment over longer time frames.

Known modulating factors

Modulating factor	Details	Effect on this KER	References
Magnitude and duration of ATP depletion	Transient or moderate ATP depletion versus severe, sustained ATP depletion.	Severe and sustained ATP depletion increases probability of irreversible injury/death. Partial depletion may cause reversible stress or cell-cycle arrest.	Nieminen et al. 1994; Leist et al. 1997
Metabolic flexibility / glycolytic capacity	Ability to compensate for mitochondrial ATP loss by glycolysis or alternative ATP-generating pathways.	Higher metabolic flexibility may reduce sensitivity of the downstream cell death response.	Bonora et al. 2012; Zong and Thompson 2006
Cell type and proliferative/metabolic demand	Highly energy-demanding or poorly glycolytic cells may have lower tolerance to ATP depletion.	Alters threshold and time-scale for transition from ATP depletion to injury/death.	Bonora et al. 2012; Green and Kroemer 2004
Mitochondrial permeability transition and calcium homeostasis	Calcium overload and permeability transition can amplify ATP depletion and membrane failure.	Can accelerate progression to necrotic or mixed cell injury phenotypes.	Halestrap 2009; Nieminen et al. 1994
Apoptotic execution machinery	Caspase competence and residual ATP availability influence whether death is apoptotic or necrotic.	Determines cell death mode rather than the existence of injury/death per se.	Leist et al. 1997; Nicotera et al. 1998

Known Feedforward/Feedback loops influencing this KER

Feedback and feedforward processes may influence this linkage. ATP depletion can impair ion pumps, causing calcium dysregulation and mitochondrial permeability transition, which further suppresses ATP production and amplifies injury. Loss of mitochondrial function may also increase ROS generation, further damaging mitochondrial and cellular components. Conversely, glycolytic compensation and stress-response activation may temporarily buffer ATP depletion and delay cell death.

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Relationship: 2767: Cell injury/death leads to Decrease, Growth

AOPs Referencing Relationship

AOP Name	Adjacency	Weight of Evidence	Quantitative Understanding
Uncoupling of oxidative phosphorylation leading to growth inhibition via increased cytosolic calcium	adjacent	Moderate	Not Specified
Uncoupling of oxidative phosphorylation leading to growth inhibition via ATP depletion associated cell death	adjacent	Moderate	Not Specified
Uncoupling of oxidative phosphorylation leading to growth inhibition via decreased Na-K ATPase activity	adjacent		
Uncoupling of oxidative phosphorylation leading to growth inhibition via mitochondrial swelling	adjacent		
Increase in reactive oxygen species leading to growth inhibition via lipid peroxidation and cell death	adjacent	High	Moderate
Excessive reactive oxygen species leading to growth inhibition via protein oxidation and cell injury/death	adjacent		
Excessive reactive oxygen species leading to growth inhibition via fatty acid oxidation and cell injury/death	adjacent		
Increase in reactive oxygen species leading to growth inhibition via oxidative DNA damage and cell death	adjacent	High	Moderate
Increase in reactive oxygen species leading to growth inhibition via protein oxidation and cell death	adjacent	High	Moderate
ROS formation leading to growth inhibition modulated by the Keap1-Nrf2 antioxidant response	adjacent	High	High

Evidence Supporting Applicability of this Relationship

Taxonomic Applicability

Term	Scientific Term	Evidence	Links
humans	Homo sapiens	Moderate	NCBI
mammals	mammals	Moderate	NCBI
fish	fish	Moderate	NCBI
crustaceans	Daphnia magna	Moderate	NCBI
green algae	Ulva compressa	Moderate	NCBI

Life Stage Applicability

Life Stage	Evidence
All life stages	Moderate

Sex Applicability

Sex	Evidence
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Sex Evidence

Unspecific Moderate

The KER is applicable to biological systems in which growth depends on maintenance or expansion of viable cell number or biomass. This includes unicellular populations, developing embryos, juvenile organisms, growing tissues, and adult organisms in which tissue condition or somatic growth is assessed. Taxonomic applicability is broad across eukaryotes, but empirical support is strongest for algae, aquatic invertebrates, mollusks, fish, and mammalian embryo or cell models. The KER is not sex-specific, but sex, endocrine status, life stage, and environmental context may modulate sensitivity. The relationship is most relevant when cell injury/death is sufficiently extensive, sustained, or located in growth-relevant tissues. It is less predictive when growth is reduced by upstream mechanisms that suppress proliferation or metabolism without substantial cell death.

Key Event Relationship Description

This KER describes the causal and predictive relationship whereby an increase in cell injury and/or cell death leads to a decrease in growth. The upstream KE, cell injury/death, represents loss of cellular viability or severe cellular damage resulting in apoptosis, necrosis, or other forms of lethal cellular injury. The downstream KE, decreased growth, represents reduced accumulation of biomass, body size, length, cell density, tissue mass, or other growth-related endpoints at organ, organism, or population levels. The biological logic of the KER is that growth requires a positive balance between production of new cellular material and loss of existing cells. When cell injury/death is sufficiently frequent, persistent, or spatially distributed across growth-relevant tissues, net cell accumulation is reduced and tissue or organismal growth is impaired. In unicellular systems, increased cell death directly reduces viable cell density and biomass accumulation. In multicellular organisms, the relationship depends on the affected tissue, the ability to compensate through proliferation or regeneration, and the timing of injury relative to developmental or growth windows.

This relationship is not intended to imply that all decreases in growth are caused by cell death. Growth can also decrease through reduced cell proliferation, altered energy allocation, endocrine disruption, nutrient limitation, or developmental delay without overt lethality. Rather, the KER applies when increased cell injury/death is of sufficient magnitude or duration to reduce the viable cellular pool needed for growth or to damage growth-relevant tissues. Within the ROS-growth AOP network, this KER provides a terminal convergence relationship for pathways in which oxidative stress, DNA strand breaks, or ATP depletion produce cytotoxicity that contributes to reduced growth.

Evidence Supporting this KER**Biological Plausibility**

Overall call: High. Growth at the level of a tissue, organ, organism, or cell population depends on net accumulation of cells and cellular biomass. Increased cell death directly lowers the number of viable cells and can reduce tissue mass, disrupt morphogenesis, or impair the capacity for biomass accumulation. This relationship is strongly supported by developmental and cell-size control principles showing that final tissue and organism size depend on the balance among cell growth, cell division, and cell death (Conlon and Raff, 1999). In embryos and developing organisms, excessive cell death can reduce cell number available for organ formation and growth, whereas in unicellular populations and cell cultures, cytotoxicity directly reduces viable cell density. The KER is therefore mechanistically plausible across taxa, although the magnitude of growth impairment depends on the tissue affected, compensatory proliferation, regeneration, and exposure duration.

Empirical Evidence

Overall call: Moderate. Empirical support is moderate because multiple studies report concordance between cell injury/death and growth-related effects, but the evidence is heterogeneous and not always designed specifically to test this KER. In several systems, cell injury/death and growth inhibition are measured at different time points, and growth can be affected by mechanisms other than cell death. Nevertheless, the available data support the expected direction of effect across algae, fish embryos, mollusks, and mammalian embryo models.

Biological system	Stressor / context	Upstream evidence: cell injury/death	Downstream evidence: decreased growth	Concordance interpretation	Reference

Biological system	Stressor / context	Upstream evidence: cell injury/death	Downstream evidence: decreased growth	Concordance interpretation	Reference
<i>Chlamydomonas reinhardtii</i>	Paraquat	Loss of membrane integrity measured by SYTOX Green; cell death observed at approximately 0.5 μ M after 24 h.	Reduced cell density/growth after 72 h; growth LOEC approximately 0.1 μ M and EC50 approximately 0.26 μ M.	Partial temporal and endpoint concordance. Growth effects occurred at or below cytotoxicity thresholds, indicating that cell death contributes but is not the only driver of growth inhibition.	Jamers and De Coen, 2010
<i>Chlamydomonas reinhardtii</i>	Paraquat and herbicides	SYTOX Green cell death observed with paraquat; cell injury occurred alongside ATP depletion and other stress endpoints.	Assay system reported reduced growth/cell density and multiple mechanistic endpoints following herbicide exposure.	Supports association between cytotoxicity and reduced population growth, but includes multiple parallel mechanisms.	Nestler et al., 2012
Mouse and rat whole-embryo culture	Methanol	Cell death markedly elevated in embryos at growth-relevant concentrations.	Mouse and rat embryo growth reduction observed in exposed cultures.	Supports developmental concordance between increased embryonic cell death and growth impairment, with species differences in sensitivity.	Abbott et al., 1995
Eastern oyster, <i>Crassostrea virginica</i>	Cadmium and temperature interaction	Hemocyte mortality, lysosomal destabilization, and cellular energy disruption observed under cadmium stress.	Reduced condition index and increased mortality under combined cadmium and elevated temperature.	Supports linkage between cellular injury and reduced growth/condition, although growth is modified by temperature and energy budget effects.	Sokolova et al., 2005; Cherkasov et al., 2006
Fish embryos and juveniles	Rotenone	Histological lesions and tissue injury observed at low concentrations.	Developmental delay and growth-related impairment reported after short-term exposure.	Supports association between cellular/tissue injury and developmental growth impairment; direct measurement of cell death was limited.	Melo et al., 2015

Biological system	Stressor / context	Upstream evidence: cell injury/death	Downstream evidence: decreased growth	Concordance interpretation	Reference
Marine copepod, Paracyclops nana	Gamma radiation	Radiation induced oxidative stress and impaired survival/development.	Growth retardation and failure of nauplii to develop to adults observed.	Supports an adverse sequence from stress-induced cellular injury to growth retardation, although cell death was not always measured directly.	Won and Lee, 2014

Uncertainties and Inconsistencies

The main uncertainty is that decreased growth is an integrative endpoint and can arise through several mechanisms that do not require overt cell death. Reduced proliferation, ATP depletion, endocrine disruption, altered energy allocation, nutrient limitation, delayed development, or behavioral effects can all reduce growth. For this reason, cell injury/death should be interpreted as a sufficient but not always necessary contributor to decreased growth. A second uncertainty is that many studies measure cytotoxicity and growth at different times or in different tissues, which limits direct evaluation of temporal concordance. In some algal studies, growth inhibition occurs at lower concentrations than overt cell death, suggesting that non-lethal impairment of proliferation, photosynthesis, or energy metabolism may precede cell death. Conversely, mild or localized cell injury may be compensated by repair or proliferation and may not lead to measurable growth reduction. These uncertainties support a moderate, rather than high, empirical call for this KER.

Quantitative Understanding of the Linkage

Overall call: Low to moderate. Quantitative understanding is limited because the relationship between cell injury/death and growth depends on the proportion of cells affected, tissue location, developmental timing, compensatory proliferation, regenerative capacity, and organismal energy allocation. At a conceptual level, the linkage is quantitative: growth rate reflects the balance between biomass accumulation and biomass or cell loss, so increasing the frequency or magnitude of cell death should reduce net growth if cell replacement or compensatory growth is insufficient. However, few studies provide response-response models that predict growth reduction from a measured degree of cell injury/death across taxa or stressors.

Response-response relationship

In cell populations and unicellular organisms, the quantitative relationship can be relatively direct because viable cell density is part of the growth measurement. In multicellular organisms, the relationship is less direct because growth can continue despite localized cell death if compensatory proliferation or tissue repair occurs. Some data show concordance between cytotoxicity and growth inhibition, but these data are generally insufficient to define universal thresholds. Therefore, quantitative understanding should be considered low to moderate for broad AOP-Wiki application, with higher confidence possible for specific model systems where cell viability and growth rate are measured in the same assay and time course.

Known modulating factors

Modulating factor	Relevant details	Effect on the KER	Supporting references
Developmental stage	Embryonic and larval stages, rapid growth phases	Increases sensitivity because rapid tissue growth requires high net cell accumulation; cell death during development can disproportionately impair growth.	Abbott et al., 1995; Conlon and Raff, 1999
Tissue regenerative capacity	Capacity for compensatory proliferation or tissue repair	Reduces probability that cell death will translate into growth impairment when surviving cells can replace lost cells.	Conlon and Raff, 1999

Modulating factor	Relevant details	Effect on the KER	Supporting references
Exposure duration and timing	Acute versus chronic exposures; timing relative to growth window	Longer or developmentally timed exposures increase probability of growth effects from cell loss.	Jamers and De Coen, 2010; Melo et al., 2015
Energy and nutritional status	Energy budget, food availability, metabolic reserve	Can increase or decrease impact of cell death on growth by altering compensatory capacity and resource allocation.	Sokolova, 2013; Cherkasov et al., 2006
Environmental stressors	Temperature, oxygen availability, salinity, co-exposures	Can amplify cytotoxicity or reduce compensatory growth responses, modifying downstream growth effects.	Cherkasov et al., 2006; Won and Lee, 2014

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