Peroxisomes, oxidative stress, and inflammation

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INTRODUCTION

Peroxisomes are essential organelles of human cells. In this article, we review peroxisome biology; summarizing how the organelle is formed, how it functions, and what happens when these processes are compromised. In addition, we describe an emergent link between the organelle and cellular aging pathways. In the latter analysis, we connect peroxisome function with the generation and destruction of specific inflammatory mediators and speculate on the organelle's involvement in initiating and progressing human disease.

PEROXISOME FUNCTION

Degradation
Peroxisomes synthesize and degrade a wide variety of cellular compounds. Through α- and β-oxidations, specific long-chain, very-long-chain, and 3-methyl-branched-chain fatty acids are degraded. These processes may occur entirely within the organelle or may involve participation of other organelles - e.g., mitochondria. The notion that peroxisomes shuttle metabolites for continued processing and/or anaplerotic metabolism is part of an emergent theme for the organelle; specifically, that it is integrated into a variously interacting endomembrane system responsible for a number of critical cellular processes.

The peroxisome's handling of hydrogen peroxide, a reactive oxygen species produced by oxidative reactions occurring within the organelle, also bears on this point. Under most conditions, hydrogen peroxide is produced and immediately processed by the organelle's resident enzymes.
Inflammatory drugs, including aspirin, are popularly seen as a major strategy to combat these processes. Specific pathways involving the presence of peroxisomes - organelle function is linked to the inflammatory response. Peroxisomes contain enzymes which contribute to the synthesis of critical cellular constituents including bile acids, ether phospholipids, and docosahexaenoic acids, among others. Bile acids, derived from cholesterol, are important molecules involved in digestion through their ability to emulsify fats. Ether phospholipids, including plasmalogens, represent a vital class of membrane protective molecules, found throughout cells of the body. Docosahexaenoic acids, peroxisomally produced omega-3 fatty acids, are the pivotal precursors of resolvins (“resolution-phase interaction products”), maresins (“macrophage mediator in resolving inflammation”), and protectins (formerly called “neuroprotectins”). These molecules possess potent anti-inflammatory, inflammatory resolving, and immunoregulatory activities. Importantly, conversion of docosahexaenoic acids to these biologically active mediators is accelerated by non-steroidal anti-inflammatory drugs, including aspirin, which inhibits the cyclooxygenase-2 enzyme.

**PEROXISOME FORMATION**

**Biogenesis**

From a functional perspective, the peroxisome is clearly a major player in cellular metabolism and a key component of organismal physiology. As to how it acquires these capacities, the answer lies in a magnificently choreographed series of biochemical processes which bring about its biogenesis.

Defining the origins of the peroxisome membrane has taken some time - many years in fact. Growth and division of existing organelles gained considerable early support until evidence was obtained that the endoplasmic reticulum was also providing membrane. The current consensus is that both processes contribute to peroxisome membrane growth and proliferation. Once assembled, the peroxisome membrane acquires additional membrane proteins including those constituting the import machinery. Although still not described in complete detail, this apparatus is known to consist of the following components: soluble receptors which recognize peroxisomal targeting signals on nascent proteins/enzymes; docking proteins which serve to concentrate and direct the receptor-ligand complex at the organelle membrane; and several molecules involved in facilitating the translocation process and recycling essential components for additional rounds of import.

Mechanisms are in place to recycle unneeded, damaged, or aged import factors, as well as to degrade the entire organelle when appropriate.
Redox balance

Many of the enzymes imported by peroxisomes oxidize substrates and produce hydrogen peroxide as a metabolic by-product. This hydrogen peroxide is normally processed to water and oxygen by catalase or other organellar peroxidases, thus maintaining oxidative balance. In recent years, several circumstances have been described in which the balance is upset and peroxisomes begin to produce excess hydrogen peroxide and related downstream reactive oxygen species. These include certain disease states in which catalase is either not produced or is unstable, as well as situations in which the enzyme is inactivated or mislocalized. Certain xenobiotics/environmental toxins appear to be able to inhibit activity of the enzyme, and aging cells are progressively less able to correctly compartmentalize the critical antioxidant enzyme.

Under conditions in which peroxisomal reactive oxygen species amass, dramatic effects on cells are seen. For example, cellular proteins, lipids, and DNA are oxidatively damaged, organelle function is compromised and metabolism is slowed.

Genetic disease

Peroxisomes fail to form or are deficient in one or more of their constituent enzymes in a series of devastating genetic diseases described in ever greater detail over the past 30 years or so. The severity of the clinical manifestations reflects the extent of the organelle’s impaired function. Many affected children die within the first decade of life with deficits manifest in nearly all organ systems. To date, treatment approaches have largely been limited to palliative care. Advances in gene and/or protein therapies promise to improve clinical outcomes.

Oxidative stress, inflammation, and degenerative disease

Oxidative stress and inflammation are inextricably tied processes. Chronic inflammation is associated with elevated reactive oxygen species levels; anti-inflammatory cascades are linked to diminished reactive oxygen species concentrations. And the converse is true: elevated oxidative stress triggers inflammation, whereas redox balance inhibits the cellular response. Thus, oxidative stress and inflammation may be seen as both causes and consequences of cellular pathology. We suggest here that through the peroxisome’s role in cellular redox balance, as well as its ability to synthesize various anti-inflammatory molecules and degrade pro-inflammatory mediators, the organelle is part of a critical network controlling cell function and organismal well-being (Figure 1). What is surprising is that these vital roles for the organelle have been unappreciated for so long.

We have previously argued that peroxisomes function as important communication centers - integrating signals from various sources to alter their own metabolism as well as that of other organelles, and to initiate or inhibit cellular aging programs. A major redox-based interplay exists between peroxisomes and mitochondria, a relationship that warrants additional analysis. Several reports indicate that altering peroxisomal redox balance triggers...
oxidative stress in mitochondria - resulting in reactive oxygen species production, diminished membrane potential, and compromised organelle function [7,21]. Obviously, the cellular consequences of diminished mitochondrial function are profound. However, restoring peroxisomal redox balance - for example by supplementing (peroxisomal) catalase, renews mitochondrial [38]. Mitochondria repolarize and aging cells delay appearance of senescence markers. Increasing oxidative stress in peroxisomes is “progeric” on cells; eliminating the stress revives them. This approach of targeted antioxidant prophylaxis has also been successful in disease models. For example, in a human cell model for psoriasis, catalase supplementation reduces expression of the inflammatory cytokine, tumor necrosis factor α, that is thought to be a major initiator of the chronic inflammation seen in psoriatic tissue [23]. Similarly, in in vitro [23] and in vivo (Terlecky SR - unpublished) models of ischemia-reperfusion (heart attack), damage to cardiomyocytes and cardiac tissue is dramatically inhibited. In a rat cell model of Alzheimer’s disease, β-amyloid peptide-induced neuronal toxicity is significantly reduced (Terlecky SR - unpublished). Enhancing peroxisomal catalase also reduces inflammatory cytokine production in appropriately challenged human fibroblasts (Terlecky SR - unpublished). The evidence is mounting - peroxisome redox balance is a major determinant of cell stress and the presence or absence of cell pathology. Perhaps it is not surprising then that epidemiological studies suggest a very strong link exists between diminishing cellular catalase levels and the onset of degenerative disease [34].

CONCLUSION

The treatment of human degenerative disease requires, in our view, a distance from the reductionist and extremely focused approaches long applied by the scientific and medical communities. Rather, we suggest a broader attack, targeting oxidative stress, chronic inflammation, and the resultant pro-aging programs initiated in cells and tissues (Figure 2). There are many ways to approach this - the direction discussed in this article focuses on the peroxisome. The organelle plays a key role in controlling inflammation and maintaining oxidative balance in cells. By targeting peroxisomes - perhaps a number of devastating diseases could be more effectively treated or prevented. We suggest enhancing peroxisome function and maintaining the organelle’s redox balance by all means possible. Mitochondrial integrity/activity will be maintained or enhanced, oxidative stress will be reduced, inflammation will be held in check, and cellular pathology will be all but eliminated. Optimistic dreams? Perhaps. But as is often said in science, “That’s why you do the experiment”.

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