Editorial: Molecular Mechanisms Protecting against Tissue Injury

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The Editorial on the Research Topic

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In response to tissue injury acute inflammatory reactions occur that aim to restore homeostasis (Medzhitov, 2008). However, hampered resolution of inflammation can result in chronic inflammation and/or pathologic wound repair (Nathan and Ding, 2010). These conditions can result from excessive oxidative and inflammatory and/or overwhelmed adaptive response systems. They may also be triggered by a variety of other conditions including diabetes, infections, or aging. Targeted up-regulation of cytoprotective systems may be a therapeutic approach to ameliorate exacerbation of injury and prevent pathologic wound repair, fibrosis and/or cancer (Nathan, 2002). Such protective systems include various anti-oxidant, anti-inflammatory, anti-apoptotic molecules and also transporters or channels. In this Frontiers research topic various concepts how specific mechanisms can determine tissue damage or protection and their therapeutic potential in a number of pathological conditions and diseases are discussed.

Tissue damage control is important at different levels to maintain homeostasis of cells, tissues and whole organisms. For example, when immunological reactions are primarily directed against the cause of tissue damage (e.g., pathogenic microbes), excessive collateral damage may occur. In such cases, avoiding additional tissue damage would be more important than elimination of the disease-triggering stimulus (Soares, 2014; Soares et al., 2014). Severe tissue injury can lead to chronic inflammation, fibrosis, disturbed developmental changes and cancer (Reuter et al., 2010), which underscores the need to prevent tissue damage and/or to promote regeneration.

The coordinate immunological functions to maintain tissue homeostasis is orchestrated by a selected group of immune cells (Shalapour and Karin, 2015) such as dendritic cells (Mirzaee et al.) and regulatory T-cells (Lei et al.). Targeted modulation of these regulators may thus give control on the decisive machinery that determines immunity, inflammation, tissue remodeling, and cancer (Sutmuller et al., 2007). Infusion of regulatory T-cells facilitates tissue regeneration by preventing undesired immunological activity and by controlling resident non-immune tissue cells and forms an alternative strategy to dampen tissue injury (Lei et al.).

Mizumura and colleagues describe how autophagy may promote tissue damage or repair and novel developments in its regulation (Mizumura et al.). In particular, the role of selective autophagy in a variety of human diseases and the therapeutic potential of this system is discussed.

Surgery and other types of traumatic injury not only cause inflammatory injury, fibrosis, and scar formation (Brouwer et al., 2015), but are associated with the release of free heme (Wagener et al., 2003a). Heme is the prosthetic group of a number of physiologically important hemoproteins (e.g., hemoglobin, cytochromes or cyclooxygenase). However, when heme is not embedded in apoproteins which occurs in pathophysiological situations such as hemolysis or tissue injury, it can mediate or fuel oxidative, inflammatory, and fibrotic insults and may act as a danger signal...
it can protect tumor cells against immune surveillance (Was et al., 2010). Moreover, HO-1 has recently been suggested to be involved in transforming obesity to diabetes (Jais et al., 2014). Therefore, the effects of such cytoprotective systems appear to be critically dependent on cell-type and tissue-context-specific mechanisms.

Another review in this Frontiers topic by Horbach et al. addresses the role of the nuclear factors upstream stimulatory factor (USFs)-1 and -2 in the context of carcinogenesis and tissue injury. USFs have primarily been considered to be involved in the regulation of metabolism, but have also been shown to be intimately associated with tissue protection and the pathogenesis of cancer. Here, the complexity of USF-1 and -2 regulation by various kinases in carcinogenesis is discussed.

Finally, a feasible approach to afford targeted protection in various pathological conditions is to trigger defined protective pathways with safe plant components. For example, induction of anti-inflammatory pathways with herbal compounds and antioxidants suppressed expression of proinflammatory cytokines in human peripheral blood mononuclear cells (Spatuzza et al.), adipose cells (Zagotta et al.), and dendritic cells (Mirzaee et al.). Interestingly, many dietary and natural compounds have been demonstrated to activate nuclear factor erythroid 2-related factor (Nrf2), which in turn induces a number of protective enzymes, such as HO-1, and promote therapeutic effects in cardiovascular diseases (Barbagallo et al., 2013). However, when translating novel protective strategies from the bench to the clinic possible differences in experimental outcome between animal models, cell lines, healthy controls and patients need to be considered (Dorrestein et al., 2015).

Better insights into the observed differences between pre-conditioning and post-conditioning in relation to tissue repair could further deepen our understanding of these regulatory pathways. It appears likely that learning more about the molecular mechanisms protecting against tissue damage will enable the development of better strategies to prevent or ameliorate wound repair and promote healthy aging.

AUTHOR CONTRIBUTIONS

All authors listed, have made substantial, direct and intellectual contribution to the work, and approved it for publication.

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