Interleukin-1 and the Treatment of Auto-inflammatory Diseases

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Chronic inflammatory diseases fall into two categories: either “autoimmune” or “auto-inflammatory” [1]. Although nearly all autoimmune diseases have an inflammatory component, as in rheumatoid arthritis, in autoimmune diseases, the primary defect is the auto-reactive T-cell. The term “auto-inflammatory diseases” encompasses several local and systemic diseases due to monocyte dysfunction, each responsive to blocking interleukin-beta (IL-1β) or blocking the IL-1 receptor using the IL-1 receptor antagonist. The best example is Familial Mediterranean Fever since manifestations of this disease include fever, fatigue, painful inflamed serosal and synovial tissues, biochemical markers of systemic inflammation, and hematological responses of neutrophilia [2]. Other auto-inflammatory diseases are due to single amino acid mutations in the protein “cryopyrin” [3] and are called “cryopyrinopathies”; the same syndromes are also termed “cryopyrin associated periodic syndromes” (CAPS). Systemic inflammation can be lethal, and the best example is septic shock. However, septic shock represents exogenously induced inflammation or simply “exogenous inflammation.” Although many laboratory studies use bacterial products to stimulate IL-1β production, in the clinical sense, auto-inflammation is the result of IL-1-induced IL-1 or IL-1 induction of other cytokines such as IL-17. Either IL-1α or IL-1β can be the driving component in auto-inflammation. For example, in patients with neonatal onset multi-inflammatory disease (NOMID) treatment with the IL-1 receptor antagonist anakinra drives down not only the manifestations of the disease but also gene expression for IL-1α, IL-1β, TNFa, IL-6 and caspase-1 [4]. The most compelling evidence for the importance of IL-1 in inflammation comes from persons born with a mutation in the gene that codes for the IL-1 receptor antagonist. The syndrome is calledDIRA for deficiency of the IL-1 receptor antagonist [5,6]. At birth, these patients develop lethal inflammation with massive numbers of neutrophils in the skin and high levels of IL-17.

Although traditionally “auto-inflammatory diseases” are thought to be examples of rare diseases, in reality several common diseases such as type 2 diabetes, gout, dry eye disease, hydrenephrosis, suppurative, and heart failure are likely to be auto-inflammatory conditions. The common link is loss of the tight control over the processing and secretion of IL-1β from the activated monocyte. The processing and secretion of IL-1β is a function of the IL-1β/caspase-1 “inflammasome”, a complex of intracellular proteins that results in the secretion of IL-1β. IL-1 receptor blockade or neutralization results in a rapid and sustained reduction in disease severity; the responses are highly consistent with blockade of IL-1β activity in these diseases. The measurement of IL-6 in the circulation is the preferred method for assessing the severity of the inflammation in auto-inflammatory diseases, and the fall in serum IL-6 reveals the effectiveness of anti-IL-1 treatment. In cytokine biology, auto-inflammatory diseases are examples that reveal the causative role of a specific cytokine for disease severity. In addition to treating CAPS with IL-1 blockade, several trials are currently underway, including type 1 and pre-diabetes, post-myocardial infarction heart failure, osteoarthritis, gout and smoldering myeloma. CANTOS, the largest trial ever done in anti-cytokine therapeutics, with an enrollment of 17,200 patients, will test the hypothesis whether an antibody to IL-1β will reduce cardiovascular events in a high-risk type 2 diabetes cohort with a CRP > 2.0 mg/l, when on standard of therapy. In addition to blocking IL-1β, clinical trials reveal that monoclonal antibodies to IL-1α are also a therapeutic option in some diseases. IL-1α is expressed on platelets and also present as an active precursor in most epithelial cells. Thus, IL-1β as well as IL-1α emerge as master cytokines regulating diseases that were not appreciated as being due to endogenous inflammation.

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References


Bibliography

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