Biological role of interleukin 1 receptor antagonist isoforms

William P Arend, Carla J Guthridge

Abstract
The interleukin 1 receptor antagonist (IL1Ra) family of molecules now includes one secreted isoform (sIL1Ra) and three intracellular isoforms (icIL1Ra1, 2, and 3). Extensive evidence indicates that the sole biological function of sIL1Ra seems to be to competitively inhibit IL1 binding to cell-surface receptors. Although intracellular IL1Ra1 may be released from keratinocytes under some conditions, the intracellular isoforms of IL1Ra may carry out additional as yet poorly defined roles inside cells. Maintenance of a balance between IL1 and IL1Ra is important in preventing the development or progression of inflammatory disease in certain organs. Both the secreted and intracellular isoforms of IL1Ra contribute to maintenance of this balance. An allelic polymorphism in intron 2 of the IL1Ra gene (IL1RN*2) predisposes to the development or severity of a variety of human diseases largely of epithelial cell origin. Both the impaired production of IL1Ra and the overproduction of IL1β are related to the presence of this allele. Restoration of the balance between IL1Ra and IL1 through a variety of approaches is a therapeutic goal in specific chronic inflammatory diseases.

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The interleukin 1 receptor antagonist was originally described as a secreted molecule (sIL1Ra) from monocytes and macrophages. The primary function of sIL1Ra seems to be to competitively inhibit the binding of IL1 to cell surface receptors. Three intracellular isoforms of this molecule have now been described. The biological role of the intracellular isoforms of IL1Ra remains unclear. Although these molecules may be released from cells to function as receptor antagonists, the intracellular isoforms of IL1Ra may carry out additional roles inside cells. This review will summarise characteristics of the described IL1Ra isoforms, the role of these molecules in biology, and reported disease associations of allelic polymorphisms of the IL1Ra gene.

Characteristics of IL1Ra isoforms
The early description of various IL1 inhibitory activities in biological fluids or cell supernatants, and the purification, cDNA cloning, and expression of the IL1Ra molecule, have been reviewed in detail. The original 17 kDa isoform of IL1Ra is secreted as variably glycosylated species of 22–25 kDa from monocytes, macrophages, neutrophils, hepatocytes, microglial cells, and other cells. sIL1Ra binds to types I and II IL1 receptors (IL1RI and IL1RII) on plasma membranes of a variety of cells with an avidity near equal to the two agonists, IL1α and IL1β, yet fails to induce receptor internalisation or discernible intracellular responses. Target cells exhibit full biological responses to occupancy of only three to five receptors per cell by IL1 agonists, and most cells express a relatively large number of the biologically active IL1RI. Because of this extreme sensitivity to IL1 stimulation and a large number of free receptors per cell, 100-fold excess or greater amounts of IL1Ra must be present to effectively inhibit cell stimulation.

The extended IL1Ra gene (IL1RN) is present on the long arm of human chromosome 2 at band 2q14, adjacent to the genes for IL1α and IL1β. The first described intracellular isoform of IL1Ra (icIL1Ra1) is created by an alternative transcriptional splice of an upstream exon into the amino terminus of sIL1Ra, creating a 18 kDa protein that lacks a signal peptide. icIL1Ra1 is a major protein in keratinocytes and other epithelial cells, and is also produced with delayed kinetics by monocytes and macrophages. A cDNA for a second intracellular isoform of IL1Ra (icIL1Ra2) was cloned from human neutrophils, and contained a 63 bp sequence inserted between the first and second exons for icIL1Ra1. This cDNA was present in fibroblasts, keratinocytes, and human myelomonocytic cells, although the predicted 25 kDa protein has not yet been described as a natural product in any cell. Thus, it remains unknown whether icIL1Ra2 exists as a protein in vivo. A third intracellular isoform of IL1Ra (icIL1Ra3) was recently described as a 16 kDa product of predominately the sIL1Ra mRNA, created by alternative translational initiation. icIL1Ra3 is a major protein in hepatocytes and neutrophils, and also is present in smaller amounts in monocytes, macrophages, and keratinocytes.

The binding characteristics of each isoform of IL1Ra have been examined with IL1R in both solid phase and soluble forms. Recombinant icIL1Ra1 and sIL1Ra exhibited equivalent binding to immobilised IL1RI, whereas icIL1Ra3 bound fourfold to fivefold less avidly. The biological activities of these isoforms of IL1Ra exhibited similar characteristics in the murine thymocyte assay, with sIL1Ra and icIL1Ra1 showing equivalent inhibition of IL1 stimulation and icIL1Ra3 being fourfold less active. Recombinant icIL1Ra1 and icIL1Ra2 exhibited similar patterns of
Table 1 Animal models of disease treated with recombinant IL1Ra

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<th>Disease Model</th>
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<td>Collagen-induced arthritis in mice</td>
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<td>Streptococcal cell wall induced arthritis in rats</td>
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<td>Immune complex induced arthritis in mice</td>
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<td>Antigen-induced arthritis in rabbits</td>
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<td>Septic shock in rabbits, baboons, rats, and mice</td>
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<td>Bacterial meningitis in rabbits</td>
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<td>Ischaemic brain injury in rats</td>
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<td>Experimental allergic encephalomyelitis in rats</td>
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<td>Streptococcal cell wall induced colitis in rats</td>
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<td>Experimental shigellosis in rabbits</td>
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<td>Immune complex induced colitis in rabbits</td>
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<td>Acetic acid-induced colitis in rats</td>
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<td>Lipopolysaccharide-induced pleurisy in rabbits</td>
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<td>Monocrotaline-induced pulmonary hypertension in rats</td>
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<td>Allergic-induced late asthmatic reaction in guinea pigs</td>
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<td>Bleomycin- or silica-induced pulmonary fibrosis in mice</td>
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<td>Immune complex-induced lung injury in rats</td>
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<td>Ischaemic/reperfusion lung injury in rats</td>
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<td>Crescentic glomerulonephritis in rats</td>
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<td>Anti-glomerular basement membrane antibody induced glomerulonephritis in rats</td>
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<td>Pre-term delivery in mice induced by IL1</td>
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<td>Osteoclast formation and bone resorption in ovariecotised mice and rats</td>
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<td>Hepatic fibrosis induced by dimethylhydrosamine in rats</td>
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<td>Post-cardiac transplant coronary arteriopathy in piglets</td>
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<td>Graft versus host disease in mice</td>
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<td>Streptozotocin-induced diabetes in mice</td>
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have not been established, but conceivably they could lead to increased IL1 production. However, these important observations indicate that adequate levels of IL1Ra are necessary to counter the potentially injurious effects of IL1 in tissues where other mechanisms may possibly stimulate the local production of IL1.

The circulating levels of IL1Ra in human diseases, and the local production of this anti-inflammatory cytokine in various tissues, have been examined in some detail.13 sIL-1Ra is produced by hepatocytes as an acute phase protein and increased levels have been described in the circulation of patients with a wide variety of inflammatory, infectious, and neoplastic diseases, as well as following surgery.25 However, studies in many human diseases indicate that the level of production of all isoforms of IL1Ra in local tissues may not be sufficient to effectively block the inflammatory consequences of local IL1 production. These observations suggest that mechanisms regulating IL1Ra production by macrophages and other cells in diseased tissues may be inadequate. Whether sIL1Ra in the circulation, produced as an acute phase protein, diffuses into tissues and inhibits IL1 effects is not known.

**Role of icIL1Ra in biology**

The biological effects of the intracellular isoforms of IL1Ra have been less thoroughly investigated. As summarised above, icIL1Ra1 and icIL1Ra2 bind equally as well as sIL1Ra to IL1RI and inhibit the stimulatory effects of IL1. In contrast, icIL1Ra3, which predominates in neutrophils and hepatocytes, is a weak inhibitor of receptor binding of IL1.26 Although the intracellular isoforms of IL1Ra are synthesised in the cytoplasm and appear not to be transported to the nucleus, keratinocytes may release icIL1Ra1 under some conditions. In addition to secretion of icIL1Ra1 by an unknown leaderless pathway, keratinocytes may express icIL1Ra1 on their plasma membrane with release from that location as well.28 Although keratinocytes possess small amounts of the active IL1RI, the biologically inactive IL1RII predominates on these cells, particularly after activation.27 IL1RII are cleaved from expressing cells, avidly bind IL1β, and function in the cell microenvironment as IL1 inhibitors. Thus, although keratinocytes are capable of releasing small amounts of icIL1Ra1, the functional importance of this cytokine as an extra-cellular receptor blocker in the skin has not been established.

Intracellular IL1Ra1 also may block IL1 induced gene expression inside cells without release to the exterior. Human ovarian and cervical cancer cells expressing icIL1Ra1 exhibited a decrease in IL1 stimulated GRO and IL8 mRNA expression in comparison with cell lines lacking icIL1Ra1.29 Furthermore, expression of icIL1Ra1 in non-producing cells, using a retroviral expression vector, led to a pattern of response analogous to that seen with cells spontaneously producing icIL1Ra1. This effect of icIL1Ra1 was not mediated at the level of transcription but appeared to involve alterations in GRO mRNA stability or degradation. In other studies, fibroblasts from patients with systemic sclerosis expressed higher levels of icIL1Ra1 in vitro than did fibroblasts from normal controls, after stimulation with IL1β or TNFα.30 Additional studies indicated that intracellular proIL1α, and not mature IL1α released into the cell microenvironment, induced the production of icIL1Ra1 by the fibroblasts. Whether the high levels of production of icIL1Ra1 by skin fibroblasts were related to the abnormal functional characteristics of these cells was not determined.

The role of icIL1Ra1 in rhinovirus (RV) upper respiratory infections in humans was examined in recent studies. icIL1Ra1 protein was readily detected in the nasal washings of normal volunteers, with the levels markedly increasing after symptomatic RV infection.30 Moreover, the icIL1Ra1 production in vivo was prolonged and accompanied the clinical resolution of symptoms. Unstimulated normal respiratory epithelial cells contained icIL1Ra1 mRNA and protein, with release of the protein after RV infection in the absence of increased mRNA levels. These observations suggest that icIL1Ra1 may play an important part in the resolution of RV infections. This cytokine may be present constitutively in respiratory epithelial cells, as is true of keratinocytes, with release by unknown translational or post-translational mechanisms. However, these studies did not investigate any possible function of icIL1Ra1 inside epithelial cells.

In summary, the biological role of icIL1Ra1 remains unclear. Keratinocytes, or other epithelial cells, may apparently release this cytoplasmic protein under certain conditions, and it may function as a receptor blocker in the cell microenvironment. Some evidence suggests that this and other intracellular isoforms of IL1Ra may carry out additional functions inside cells. We are investigating this possibility using a number of experimental approaches.

**Allelic polymorphisms of the IL1Ra gene and disease**

Further evidence for the biological importance of a balance between IL1 and IL1Ra can be derived from recent studies on an allelic polymorphism of the IL1Ra gene. A polymorphism exists in intron 2 of IL1RN caused by two to six copies of an 86 bp tandem repeat.31 The A1 allele, containing four repeats, is the most common allele and is found in 73.6% of the population. Allele A2 (IL1RN*2) contains two repeats, is found in 21.4% of the population, and has been associated with a variety of human diseases, primarily of epithelial cells or tissues. An association of allele A2 with an increased prevalence of ulcerative colitis (UC), but not with Crohn’s disease, has been described in some but not all population groups (summarised by Tountas et al32). This association may not exist in Northern European ethnic groups, but it has been found in both in American-based Hispanic and Jewish populations.32 Other disease associations of IL1RN*2 include alopecia areata,33 lichen...
scleruous, psoriasis, multiple sclerosis, diabetic nephropathy, hypochlorhydria and gastric cancer, and susceptibility to severe sepsis.

A possible association with allele IL1RN*2 has also been studied in some rheumatic diseases. Systemic lupus erythematosus, particularly photosensitivity and discoid skin lesions, was observed to be increased in both white and Japanese populations carrying IL1RN*2. However, this allele may influence the severity of SLE rather than susceptibility. IL1RN*2 is also associated with more severe forms of Sjögren's syndrome, with the levels of IL1Ra protein being decreased in saliva but increased in serum. Lastly, the IL1Ra intron 2 polymorphism did not influence the susceptibility to or severity of rheumatoid arthritis.

The possible mechanisms of IL1RN*2 association with disease remain unclear. Steady state levels of IL1Ra mRNA in human keratinocytes, presumably representing icIL1Ra mRNA, were not related to the IL1RN allelic polymorphism. In early studies increased IL1Ra secretion, but not cell associated IL1Ra, as well as reduced IL1α production were observed in cytokine stimulated monocytes from IL1RN*2 normal donors. However, these results have not been substantiated in more recent studies where total IL1Ra production (both cell associated and secreted) was decreased in unstimulated monocytes or stimulated cells from both normal donors or UC patients carrying the IL1RN*2 allele. The possibility also exists that a gene linked to or associated with IL1RN*2 might explain the apparent disease association. Indeed, the presence of IL1RN*2 is associated with increased monocyte production of IL1β, apparently through strong linkage disequilibrium with the IL1β-31T diallelic polymorphism. Thus, more severe disease in people carrying allele 2 of the IL1Ra gene may be related to an imbalance between IL1Ra and IL1β, with both increased production of IL1β and possibly decreased IL1Ra.

**Summary**

IL1Ra production in many cells and tissues immediately follows that of IL1. Mature IL1 may actually stimulate production of sIL1Ra, and proIL1α inside fibroblasts may induce production of icIL1Ra1. The sole function of sIL1Ra in the cell microenvironment is to competitively inhibit receptor binding of IL1. Thus, the biological role of sIL1Ra may be to dampen or attenuate the potent biological consequences of IL1 both in normal physiology and in pathophysiological conditions.

icIL1Ra1 may also inhibit receptor binding of IL1 after release from epithelial cells, but the intracellular structural variants of IL1Ra may carry out additional roles inside cells not involving interaction with IL1 receptors. ProIL1α is synthesised in the cytoplasm followed by movement to the plasma membrane, or to the nucleus where the pro-piece may increase transcription. Whether icIL1Ra1 influences these intracellular effects of proIL1α has not been examined. However, the balance between IL1Ra and IL1 in some tissues, such as the joint and the vessel wall, may influence the relative propensity for the development of inflammatory disease. Whether this balance pertains only to sIL1Ra, or also includes intracellular isoforms of IL1Ra, may depend on the particular cell or tissue involved. The association of various inflammatory diseases, largely of epithelial tissues, with allele IL1RN*2 may be related to both decreased production of IL1Ra and to increased production of IL1β. Rather than the exogenous administration of recombinant sIL1Ra, or delivery of IL1Ra by gene therapy, stimulation or enhancement of endogenous IL1Ra production may be a more physiological approach to treatment of human disease.
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