Enhanced 15-lipoxygenase activity and elevated eicosanoid production in kidney tumor microenvironment contribute to the inflammation and immune suppression

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Macrophage infiltration is a hallmark in the majority of solid tumors. Our studies demonstrated that macrophages that infiltrate human renal cell carcinoma (RCC) display markedly enhanced expression and activity of 15-lipoxygenase-2 (15-LOX2). Obtained data suggest that enhanced lipoxygenase activity in tumor-associated macrophages stimulates cancer inflammation and causes immune dysfunction.

Advanced, metastatic RCC disease is uniformly resistant to radiation and chemotherapy and it represents one of the most receptive cancers to immunotherapy. However, the clinical response is weakened, mostly due to tumor-induced immune suppression that represents a major obstacle for successful cancer immunotherapy. Multiple immunosuppressive mechanisms have been identified in RCC that propagate conditions interfering with the development and activation of an effective anti-tumor response. Tumor microenvironment, which encompasses immune, stromal and vascular cells, exerts profound immunosuppressive and tolerogenic activity on antigen presenting as well as T effector cells by secretion of bioactive proteins and lipids. Together, these factors favor conditions that allow tumors to escape immune recognition and foster proliferative and metastatic potential of tumor cells.

Growing body of evidence suggests that cancer progression is facilitated by enhanced production of eicosanoids that promote cancer inflammation, immune suppression, angiogenesis and proliferation of malignant cells. Eicosanoids are enzymatic products of arachidonic acid (AA) and can be formed by cyclooxygenase (COX), lipoxygenase (LOX) or cytochrome P-450 epoxygenase. Importantly, eicosanoids strongly contribute to the immunoregulatory mechanisms and may control inflammation through regulation of cytokine and chemokine production. Arachidonate products of cyclooxygenase and lipoxygenase pathways can be secreted in substantial amounts by both epithelial cancer cells and tumor-infiltrating inflammatory cells including tumor-associated macrophages and myeloid-derived suppressor cells. Numerous studies provided clinical and pharmacological evidence for involvement of COX2-PGE2 pathway in the pathogenesis of various cancers, including colon, lung, breast and bladder cancers. The other AA metabolizing enzymes (e.g., lipoxygenases) have drawn less attention for their potential roles in human tumor initiation and/or progression.

Lipoxygenases (LOXs) are iron-containing dioxygenases that catalyze formation of fatty acid hydroperoxides from polyunsaturated fatty acids. The specific bioactivities of LOXs include hydroperoxidase, leukotriene synthase, lipoxin synthase and hepxoylin synthase activities. Four major types of mammalian LOXs were described: 5-LOX, 8-LOX, 12-LOX and 15-LOX. In humans, two distinct subtypes of 15-LOX exist: 15-LOX1 (encoded by 15ALOX gene) and 15-LOX2 (encoded by 15ALOXB gene). Arachidonic acid (AA) is the major substrate for 15-LOX2, whereas 15-LOX1 prefers linoleic acid as substrate but could also metabolize AA. 15(S)-HETE is the major metabolite of AA formed by 15-LOX2 and 15-LOX1.

Recently we have demonstrated that progression of human renal cell carcinoma is associated with enhanced expression and activity of 15-LOX2. RCC tissues are frequently infiltrated with CD45+CD11b+CD68+HLA-DR+ tumor-associated macrophages (TAMs). TAMs seem to be central players in deregulated eicosanoid production observed in RCC because they secrete the highest levels of major 15-LOX2 arachidonate metabolite 15(S)-HETE and highly expressed 15-LOX2 at both gene and protein levels. Isolated TAMs produce substantial amounts of pro-inflammatory chemokine CCL2 and immunosuppressive cytokine...
IL-10. Co-incubation of TAMs with autologous T lymphocytes resulted in dramatic (4–10 fold) increase of IL-10 production. Remarkably, inhibition of lipoxygenase activity significantly reduced secretion of CCL2 and IL-10 by TAMs, and also prevented TAM-mediated induction of IL-10 in T cells. In addition, TAMs were able to induce tolerogenic transcription factor FOXP3 and inhibitory receptor CTLA-4 in T lymphocytes. However, the ability of TAMs to convert T cells into FOXP3+ T regulatory cells or induce CTLA-4 did not depend on 15-LOX2 activity. It would be important in further studies to delineate the mechanism by which TAMs induce FOXP3+ T regs in kidney cancer.

Cancer therapy and immunomodulatory therapy of RCC is less effective in treating large tumors partly due to immune suppression associated with advanced metastatic cancer disease. Our data suggest that tumor-associated macrophages play an important role in the development of immune suppression and T cell tolerance in human RCC through secretion of immunosuppressive factors and induction of immune unresponsiveness in T cells. We demonstrated that immune suppression in RCC is closely relates to deregulated eicosanoid metabolism in the tumor microenvironment. Figure 1 provides a schematic overview of the contribution 15-LOX2-expressing tumor-associated macrophages in immune dysfunction in patients with kidney cancer. This figure illustrates that macrophages infiltrating human RCC have a significant impact on cancer inflammation via CCL2-mediated recruitment of monocytes to the tumor site and inhibition of generation of anti-tumor immune response via secretion of IL-10 and also by conversion of T cells into T regs.

Our results suggest that manipulation of deregulated metabolism of arachidonic acid and reduction of eicosanoid levels in tumor microenvironment represents an attractive approach to regulate both cancer-related inflammation and immune suppression in human cancers including RCC. Inhibition of 15-LOX2 activity could potentially synergize with existing RCC immunotherapy such as IL-2, IFNα or dendritic-based therapy thereby enhancing its therapeutic effect and improving overall survival of cancer patients.
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