Myeloid-derived suppressor cells link inflammation to cancer

Dingzhi Wang¹ and Raymond N DuBois¹²³,*

¹Laboratory for Inflammation and Cancer; the Biodesign Institute at Arizona State University; Tempe, AZ USA; ²Department of Chemistry and Biochemistry; Arizona State University; Tempe, AZ USA; ³Department of Research and Division of Gastroenterology; Mayo Clinic; Scottsdale, AZ USA

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Our recent work has provided the first evidence that MDSCs promote chronic colonic inflammation and colitis-associated carcinogenesis. Our findings not only reveal a novel function of MDSCs in connecting inflammation to cancer, but also provide a rationale for developing effective therapeutic strategies to subvert inflammation- and tumor-induced immunosuppression.

Colorectal cancer (CRC) includes hereditary, sporadic, and colitis-associated CRC. Epidemiologic and experimental evidence strongly support the concept that chronic inflammation contributes to tumor initiation, progression, and metastasis. Indeed, ulcerative colitis, the most common form of inflammatory bowel disease (IBD), is associated with an increased risk for the development of CRC.¹ Since chronic inflammation is involved in immunosuppression,² one potential mechanistic explanation for the contribution of chronic inflammation to cancer is that it promotes tumor development via induction of immunosuppression, a protumorigenic environment by virtue of immune tolerance to tumor cells. This notion is supported by the results of our recent mouse model study of IBD-associated carcinomas in which myeloid-derived suppressor cells (MDSCs), recruited by chronic inflammation, promoted colitis-associated tumorigenesis and disease progression via suppression of antigenic CD8⁺ T-cell cytotoxicity.

MDSCs are a heterogeneous population of immature myeloid cells. In healthy individuals, immature myeloid cells differentiate into mature myeloid cells including macrophages, DCs, and granulocytes. However, this normal physiological process is interrupted in cancer patients. For example, the levels of MDSC in the blood are positively correlated with clinical cancer stage and metastatic tumor burden in mice and humans with colon cancers.⁴⁵ MDSCs have been shown to contribute to cancer immune evasion via suppressing T-cell activation, proliferation, trafficking, and viability, inhibiting natural killer (NK) cells, and promoting activation and expansion of Foxp3 positive regulatory T (Treg) cells.⁶ Although significant evidence demonstrates that MDSCs play key roles in tumor-induced immunosuppression, it remains unclear how MDSCs are recruited from the circulatory system to the colonic mucosa during chronic inflammation and tumorigenesis. Our recent work provides the first evidence showing that a chemokine receptor, C-X-C family receptor 2 (CXCR2) is required for infiltration of MDSCs from the circulatory system into inflamed colonic mucosa and colitis-associated tumors.

Previous studies have shown that several pro-inflammatory mediators such as IL-1β, IL-6, and PGE₂ induce MDSC accumulation and activation,⁷ suggesting that chronic inflammation might promote tumor initiation and progression by induction of immune suppression via MDSCs. However, so far, there has been no direct evidence demonstrating that MDSCs play a key role in connecting chronic inflammation to carcinogenesis. Our group was the first to report that colonic MDSCs recruited by chronic inflammation accelerate colitis-associated...
Moreover, our results showed that PGE₂ signaling induced expression of CXCR2 ligands in intestinal mucosa and tumors in vivo. These findings prompted us to postulate that PGE₂ might induce an infiltration of MDSCs into colonic mucosa and neoplasms through the induction of CXCR2 ligands. Our laboratory is currently testing this hypothesis.

Among prostaglandins, PGE₂ is the most abundant prostaglandin found in various types of human malignancies and is often associated with a poor prognosis. Research by our group and others has revealed that pro-inflammatory PGE₂ is a key inflammatory mediator and plays a predominant role in various hallmarks of cancer progression among most types of malignancy including in CRC. PGE₂ is produced from arachidonic acid by the action of cyclooxygenase enzymes (namely, COX-1 and COX-2) and prostaglandin E₂ synthases. COX-1 is constitutively expressed in most tissues and was thought to be a housekeeping enzyme responsible for maintaining basal prostanoid levels important for tissue homeostasis and platelet function. By contrast, COX-2 is an immediate-early response gene that is normally absent from most cells but is highly induced at sites of inflammation and tumors, particularly in IBD and CRC. Currently, the best agents for targeting the COX-2 enzyme are nonsteroidal anti-inflammatory drugs (NSAIDs). Both epidemiologic and clinical evidence has demonstrated that NSAIDs have beneficial effects on reducing the risk of developing CRC. NSAIDs (except aspirin) can increase the risk for cardiovascular complications. Our findings concerning the PGE₂-driven role of MDSCs in colorectal cancer might provide another potential mechanistic explanation for the well-characterized antitumor effects of NSAIDs.

In summary, our recent work has demonstrated that CXCR2 is required for the recruitment of MDSCs into the inflamed colonic mucosa and tumors suggest that PGE₂ might promote tumor formation and progression via recruiting MDSCs via a CXCR2-ligand-CXCR2 pathway (Fig. 1). These findings not only have extended our current knowledge of how chronic inflammation contributes to cancer, but also provided a rationale for the development of new therapeutic approaches to subvert tumor-induced immunosuppression via

**Figure 1.** A possible model for the role of PGE₂ in the regulation of tumor-induced immunosuppression. Pro-inflammatory prostaglandin E₂ (PGE₂), produced by tumor epithelial cells and tumor-associated stromal cells such as macrophages induces CXCR2 ligands, which in turn recruit CXCR2-expressing MDSCs into the tumor through the circulatory system. The recruited MDSCs promote tumor formation, growth, and progression primarily via suppressing CD8⁺ T-cell cytotoxicity against tumor cells.
CXCR2 antagonists and its neutralizing antibodies.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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