Inflammation and oncogenesis: a vicious connection

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Abstract

Epidemiological and experimental data suggest a close connection between inflammation and tumorigenesis. Solid tumors are typically infiltrated with immune cells and inflammation impacts most, if not all, stages of tumorigenesis. Molecular and cellular pathways, which connect inflammation and cancer, have emerged as attractive targets for prevention and therapy. In this review we discuss general mechanisms and concepts of cancer promoting inflammation.

Keywords
cancer; inflammation; immunity; cytokines

Introduction

Most, if not all, solid tumors are infiltrated with immune and inflammatory cells. This can represent an ongoing anti-tumor response or be a sign of immune system subversion by the tumor for its own benefit. The first possibility has been addressed within the frame of the “tumor immunosurveillance” concept, proposed by Old, Schreiber and coworkers [1]. The immune system can play an anti-tumorigenic role in certain cases, especially in blood, chemically and virally induced cancers by eliminating pre-malignant as well as fully transformed cells. This process largely depends on altered immunogenic epitopes expressed by cancer cells as well as on stress, necrosis and other immunostimulatory signals, which help immune system to recognize tumor antigens as non-self. On the effector side, immunosurveillance relies on CD8+ cytotoxic T cells (CTLs) and natural killer (NK) cells as well as help from antigen presenting dendritic cells (DCs) and CD4+ Th1 cells [1].

However, cancer cells possess a great ability to mutate, evolve and rapidly grow. Hence, the cancer can easily outsmart the immune system through the growth of low-immunogenic or resistant clones or by directly subverting the anti-tumor immune response and use it for tumor promotion. Often referred to as “tumor escape” [2], this situation is very well illuminated by the fact that advanced tumors always exhibit a significant immune infiltrate but are rarely rejected. Tumors also may remain dormant for a long time, reflecting an “equilibrium” between tumor growth (immune dependent or independent) and immune destruction [3]. Thus, while
the host immune system may be engaged in early tumor detection and destruction, it has become increasingly evident that immune cells and inflammatory processes that either precede or are subsequent to cancer development play a pivotal pro-tumorigenic role [4,5]. Various immune cells, including T and B lymphocytes, macrophages, DC, neutrophils, NK cells, mast cells and other cell types are frequently found to be concentrated in tumors relative to the surrounding tissue [6–9]. Therefore, it appears that such cells are actively recruited in response to tumor-derived signals as a result of tumor selection and evolution. However, it is also plausible that such cells may be initially recruited into the tumor as a part of the anti-tumor response, but once present within the tumor microenvironment they are diverted towards pro-tumorigenic responses. For instance, myeloid cells, which can give rise to “M1” macrophages that produce IL-12 and other anti-tumorigenic products, differentiate within the tumor microenvironment into myeloid derived suppressor cells (MDSC) or “M2” macrophages that produce various immunosuppressive and pro-angiogenic molecules [10–12]. Similarly, various tumor promoting T cells, including Th2, Th17 and Treg cells, can be recruited or differentiated in situ within tumors, while cells important for anti-tumor responses, such as Th1 cells or CD8+ CTLs, are either underrepresented or functionally disarmed [13–16]. Importantly, there is no unequivocal correlation between the presence of a T cell infiltrate and tumor prognosis as for sporadic colon cancer it represents better prognosis [17,18], while in breast cancer an infiltrate with a high CD4+ to CD8+ ratio correlates with worse prognosis [19]. Furthermore, tumors can induce and perpetuate tumor-associated inflammation and use it to their own benefit [20].

What determines the overall contribution of inflammatory processes to tumor development?

1) First, many inflammatory mediators (for instance, cytokines) are also important growth and survival factors that stimulate the survival and proliferation of pre-malignant cells [21].

2) Inflammatory mediators often activate oncogenic transcription factors, such as NF-κB and STAT3 [22–24], whereas oncoproteins such as Ras and Myc can initiate inflammatory response [25,26].

3) Tumor-associated inflammation can suppress anti-tumor immune response and divert tumor specific immune cells from being anti-tumorigenic to become pro-tumorigenic.

4) Inflammation can stimulate tumor angiogenesis.

5) Inflammation can stimulate tumor invasiveness and metastatic dissemination [27].

Types of tumor-promoting inflammation

Several types of inflammation, which differ by cause, mechanism, outcome and intensity exist [28], and all of them potentially can promote cancer development and progression. How tumor-promoting inflammation is induced? First, repetitive injury and infections can result in a chronic inflammatory response, for instance infection with Helicobacter pylori or Hepatitis C virus (HCV) cause gastritis, ulcers and hepatitis, eventually leading to gastric or liver cancer, respectively [21]. Infection with Bacteroides sp. facilitates tumor development in spontaneous intestinal cancers [29]. Chronic inflammation can also be induced by environmental exposure or dietary/metabolic factors. Particulate material and other components of tobacco smoke trigger injury and chronic lung inflammation, thereby increasing the risk of lung cancer [30]. Obesity is a risk-factor for liver cancer development and obesity-associated inflammation may serve as a critical driving force for liver cancer promotion [31–34]. Several types of autoimmunity may also contribute tumor development, for example inflammatory bowel disease increases the risk of colitis-associated cancer (CAC) and Celiac disease is a risk factor for lymphoma [35–37]. However, not all chronic inflammatory diseases increase cancer risk equally. Ulcerative colitis imposes much greater risk for CAC than Crohn’s disease, and rheumatoid arthritis does not increase cancer risk at all. Most likely, chronic inflammation needs to act synergistically with carcinogen exposure and tissue injury and repair.
Even cancers that evolve without underlying chronic inflammation exhibit tumor associated inflammation and contain inflammatory infiltrates [26]. Oncogene activation (as shown for Ras and Myc), or cell senescence induced by DNA damage or oncogene activation, can enhance the transcription of pro-inflammatory genes, coding for cytokines and chemokines [25,38–40]. The last but not least, type of inflammatory response associated with cancer is therapy-induced inflammation. Since most of the cancer cells are resistant to apoptosis, their death induced by chemo- or radiotherapy is necrotic in nature and proteins released by necrotic are potent inducers of inflammation [41]. A similar inflammatory response can be triggered by hypoxia and nutrient deprivation, which result in the necrotic death of cells at the core of large tumors. Two outcomes, which are not mutually exclusive, can be induced by this type of inflammation. First, activation of the immune system by products of necrotic cells can enhance the efficiency of tumor antigen presentation resulting in anti-tumor immunity that helps the host to eradicate the remaining tumor [42,43]. Second, inflammatory mediators released by dying cancer cells can lead to activation of tumor infiltrating macrophages that produce cytokines that eventually activate oncogenic transcription factors in remaining cancer cells that stimulate their survival and proliferation [23,44]. The balance between therapy-induced tumor eradication and regrowth is very delicate and is likely to depend on the extent of therapy induced cell death, the type of cancer being treated and the inflammatory microenvironment associated with the tumor.

**Does inflammation induce tumorigenesis?**

Epidemiological, pharmacological and genetic evidences provide a solid support that inflammation can increase cancer risk and can promote tumor progression [26]. However, it remains to be determined whether chronic inflammation can cause tumor-initiating genetic alterations or can only act in conjunction with carcinogen exposure. In the case CAC, it was suggested that chronic inflammation and colonic injury can directly cause DNA alterations [45,46]. However, chronic inflammation and loss of protective mucus can also increase intestinal permeability for environmental toxins and mutagens, which induce mutations in stem cells that give rise to cancer [47]. Furthermore, inflammation can stimulate the proliferation of cells that harbor oncogenic mutations induced by carcinogens rather than induce mutations themselves. Nonetheless, inflammation can result in the production of reactive oxygen and nitrogen species (ROS and RNI) by immune cells as well as immune mediated stimulation of ROS production in pre-malignant cells, induction of “mutagenic” enzymes such as activation-induced cytidine deaminase (AID) [48,49] and inactivation of DNA damage gatekeeper pathways such as the mismatch repair response (MMR) [10,45] or p53 [50,51]. Inflammation may also lead to epigenetic modifications including DNA and histone methylation that eventually lead to silencing of tumor suppressor loci [52–55]. In summary, it remains to be fully established whether inflammatory alone can result in tumor initiation (Figure 1) but as described below, there is ample evidence that inflammation is tumor promoting.

**Inflammation and tumor growth**

Tumor growth (often called tumor promotion) is the sum total of malignant cell proliferation vs. malignant cell death. Both processes are strongly impacted by inflammation and inflammatory cytokines produced by tumor infiltrating immune cells, such as IL-6 and TNF-α, can serve as mitogens and survival factors for pre-malignant and fully established cancer cells (Figure 1). Inflammation also contributes to the induction of angiogenesis, which is critical for supplying the growing tumor with necessary nutrients and oxygen [56,57].

Much of the growth stimulating cross-talk between immune and malignant cells is mediated by cytokines that activate the oncogenic transcription factors NF-κB and STAT3 (reviewed in [22]). Activation of either NF-κB or STAT3 is found in over 50% of all cancers [23,44,58] and
is a pre-requisite for the expression of a variety of target genes important for tumorigenesis, including anti-apoptotic genes (c-IAP, Bcl-xL, Bcl-2, c-FLIP), proliferative genes (Cyclins, c-Myc), stress-response genes (SOD2, ferritin heavy chain, hsp70), chemokines and pro-angiogenic molecules (VEGF, bFGF, CXCL12) [22,59–61]. On the other hand, in immune cells, NF-κB and STAT3 are instrumental for the production of pro-inflammatory cytokines, which mediate NF-κB and STAT3 activation in cancer cells, including IL-1, TNF, IL-6 and IL-23.

In models of inflammation-associated colon and liver cancers, ablation of IKKβ, a protein kinase required for NF-κB activation in myeloid cells, reduces the production of various pro-inflammatory cytokines including IL-1, IL-6, TNF and IL-12/IL-23 and results in decreased tumor multiplicity and size [62,63]. The tumor promoting action of these cytokines is mostly mediated by NF-κB (e.g. TNF and IL-1) or STAT3 (e.g. IL-6, IL-11) activation in premalignant enterocytes and hepatocytes [64–70]. Other relevant STAT3 activators in tumor cells can include members of the IL-6 family, such as OSM, LIF, CNTF and IL-27, cytokines of the IL-10 family (IL-22, IL-19 and others), and growth factors such as EGF [71–74]. NF-κB in both immune cells and cancer cells can also be activated by signals from Toll-like or NOD-like receptors or by inflammatory cytokines such as IL-1 or TNF. It was also suggested that IL-17 can drive NF-κB activation [75] and can indirectly cause STAT3 activation by stimulating IL-6 production [15]. STAT3 itself was shown to prolong NF-κB activation and nuclear retention [68], thereby providing a basis for the cross-talk between these important oncogenic pathways driven by inflammatory signals. Importantly, however, NF-κB and STAT3 are not classical oncogenes, at least not in solid tumors, as they are not subject to direct mutational activation.

The interplay between NF-κB and STAT3 in colitis associated cancer: malignant cooperation between immune and cancer cells

The critical role of NF-κB in linking inflammation and tumorigenesis was first demonstrated in a mouse model of CAC [62]. Ablation of IKKβ in intestinal epithelial cells largely abolished the development of colonic adenomas. The pro-oncogenic role of NF-κB in CAC is most likely mediated through induction of anti-apoptotic proteins, particularly Bcl-XL [62,64,65]. NF-κB activation in epithelial cells has no effect on cell proliferation. By contrast, STAT3 in intestinal epithelial cells impacts both cell proliferation and cell survival [64,65]. Accordingly, STAT3 ablation in intestinal epithelial cells results in a dramatic decrease in both tumor induction and tumor growth, as well as reduced expression of Bcl-XL, and other pro-survival and tissue protective proteins and cell cycle regulators [64,65,74]. As mentioned above, NF-κB activation in myeloid cells also enhances tumor growth but the effect is mainly due to enhanced epithelial cell proliferation rather than survival [62]. This effect of NF-κB in myeloid cells is mediated through the production of TNF [76], IL-6 [64,77] and other cytokines.

Inflammation and metastasis

Ninety percent of cancer deaths are due to metastatic growth. Immune cells are present in all advanced tumors and specifically at the invasive front of the tumor and are involved in various forms of direct and indirect interactions with metastasizing cells and micrometastases [20, 27]. Indeed, the inflammatory microenvironment was found to influence several key stages of metastatic process [78] (Figure 1). The process of epithelial-mesenchymal transition (EMT), which is critical for metastasis, can be triggered by several cytokines, including TGFβ, IL-1, TNF-α and IL-6 [79–81] and may be a consequence of NF-κB and STAT3 activation [23] through induction of EMT regulators such as Snail, ZEB and Twist [80,82]. Inflammatory signals also regulate the production and activity of various proteases, which degrade extracellular matrix and facilitate invasion and extravasation of cancer cells [23,83].
Chemokines can directly stimulate the migration of malignant cells towards blood vessels [23,84,85], whereas cytokines such as TNF can increase vascular permeability [86]. Furthermore, cytokines are important for the survival, recruitment, colonization and regrowth of the metastatic seeds through the same mechanisms that affect the growth and survival of primary tumors [87–89].

Conclusions and perspective

The connection between inflammatory immune responses and tumorigenesis has been extensively investigated during the past decade and some of the underlying mechanisms have been elucidated. As a result, our view of the role played by the immune system in tumorigenesis has shifted from a strict anti-tumorigenic function to a more balanced view according to which the immune system, while having some negative effects on tumor growth at early stages of the tumorigenic process, has an overall tumor promoting effect. Chronic inflammation caused by infection, autoimmune disease and exposure to irritants as well as tumor associated inflammation, contribute to tumor promotion, progression and metastatic spread. Whereas in inflammation-associated cancer, inflammation can be viewed as a causative agent affecting either tumor initiation or early promotion, tumor-elicited inflammation acts as a late tumor promoter to enhance progression and metastasis. Many of the tumor promoting effects of inflammation depend on production of chemokines and cytokines and activation of the oncogenic transcription factors NF-κB and STAT3. Interference with the production, secretion and receptor binding by chemokines and cytokines, which have been confirmed to have pro-tumorigenic activities, represent new therapeutic approaches whose efficacy should be evaluated both as single agents and as adjuvants for chemo- and radio-therapy. Keeping in mind the reciprocal relationship between the anti-tumor (immunosurveillance) and pro-tumor (cancer promoting inflammation) arms of the immune system, it is also of importance to evaluate the therapeutic efficacy of agents that interfere with activation of pro-tumorigenic pathways in combination with agents or treatment that enhance anti-tumor immunity.

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References


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Inflammation is important at various stages of tumorigenesis. Inflammation may contribute to tumor initiation by inducing DNA damage through intermediates like ROS and via activation of epigenetic mechanisms, which lead to silencing of tumor suppressor genes. During tumor promotion, immune and inflammatory cells produce cytokines and chemokines, which facilitate cancer cell survival, proliferation and promote the angiogenic switch. This results in increased tumor growth. Cytokines and chemokines also induce further recruitment and differentiation of immune cells in the tumor microenvironment. At the tumor progression and metastasis stages, immune cells further contribute by production of cytokines and chemokines.
to increase cell survival, motility and invasiveness, as well as to promote epithelial mesenchymal transition (EMT).