Review

The Role of Cyclooxygenase-2 in Colorectal Cancer

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Abstract

Colorectal cancer is the third common cancer in this world, accounting for more than 1 million cases each year. However, detailed etiology and mechanism of colorectal cancer have not been fully understood. For example, cyclooxygenase-2 (COX-2) and its product prostaglandin E2 (PGE2) have been closely linked to its occurrence, progression and prognosis. However, the mechanisms on how COX-2 and PGE2 mediate the pathogenesis of colorectal cancer are obscure. In this review, we have summarized recent advances in studies of pathogenesis and control in colorectal cancer to assist further advances in the research for the cure of the cancer. In addition, the knowledge gained may also guide the audiences for reduction of the risk and control of this deadly disease.

Key words: cancer, colon, cyclooxygenase-2, invasion, prostaglandin

Introduction

Colorectal cancer (CRC) is a common fatal cancer in developed countries such as USA [1, 2], accounting for more than 1 million cases each year [3] and 700,000 deaths [4]. Most recently, it is reported that colorectal cancer ranks fourth in the world's deadly cancer, accounting approximately 900,000 deaths yearly, due to increase of aging populations, obesity and lack of exercise [5]. In fact, colorectal cancer (CRC) is the second most frequent deadly cancer in the United State. The proposed causative factors include but not limited to genetic mutations and chronic inflammation. Until recently, it is still not known what causes CRC because of complexity of the etiology of CRC.

Etiologically, colon cancer is caused by spread of colorectal cancer cells to other parts of the body. The causative factors include genetic alterations, overexpression of Cytooxgenase-2 (COX-2), smoking, drinking of alcohol, harmful diet and lack of physical exercises [6-8]. Among those ricking factors, the occurrence of the cancer is closely linked to overexpression of COX-2, which has been noted in most of CRC [9, 10]. Increased levels of COX-2 mRNA and protein are found in the great majority of colorectal adenocarcinomas compared with levels in adjacent histologically normal mucosa [9]. A causal role for COX-2 in colorectal carcinogenesis is demonstrated in vivo in a murine model, and the biologic effects of upregulation of the enzyme are mediated predominantly through increased PGE2 production [11]. Therefore, many researchers have been trying to use COX-2 inhibitors such as nonsteroidal anti-inflammatory drugs (NSAID) and COX-2 inhibitors (COXIB) to control this deadly disease.

In this review, we summarize recent advances in understanding of COX-2 signaling in etiology of CRC. We also try to renew our interest in prevention and control of colorectal cancer by NSAID and COXIB.

COXs

COXs are important regulators of angiogenesis, inflammation and carcinogenesis. COXs are located at luminal side in the endoplasmic reticulum and...
associated with the nuclear envelope [12], containing three isoforms, that is, COX-1, COX-2 and COX-3 [13]. COX-1 is a housekeeping enzyme to meet the basic requirement for prostaglandins (PGs) [13, 14]. COX-3 is a variant of COX-1 mainly within central nervous system [15, 16]. In contrast, COX-2 is an inducible isofrom [17] in normal tissue such as colorectal, kidney, reproductive organs and stomach [18, 19]. However in carcinogenesis, COX-2 can be constantly upregulated [17, 20], for example, adenocarcinoma, squamous cell carcinoma, cholangiocarcinoma, endometrial carcinoma and hepatocellular carcinoma [21, 22].

Many factors, for example, DCA, IL-1β and LPS might promote expression of COX-2 moderately in normal fibroblasts (NFs), but profoundly in cancer-associated fibroblasts (CAFs) [23]. Our results have clearly demonstrated that COX-2 is enhanced by DCA, HGF and IL-1β [24-27]. As a result, Prostaglandin E₂ (PGE₂) production is greatly promoted, and such promotion increases proliferation and invasiveness of epithelial cancer cells [25, 27, 28]. Nevertheless, COX-2 inhibitors such as NS398 may decrease proliferation and invasiveness of colorectal cancer cells by overexpression of COX-2 and its product PGE₂ [25, 27, 28].

**Stromal Cells in Colorectal Carcinogenesis**

Stromal cells, for example, fibroblasts actively participate in carcinogenesis [29]. We have reported that fibroblasts from the stromal compartment play a pivotal role in COX-2 signalling and carcinogenesis [25-27, 30]. As shown previously, cancer-associated fibroblasts (CAFs) may promote epithelial ovarian cancer [31]. Cytokines, for example, IL-1β, Tumor Necrosis Factor-α (TNF-α) and other compounds, for instance, deoxycholic acid (DCA) stimulates COX-2 expression, which enhances PGE₂ production in colorectal fibroblasts [32-38]. In addition, COX-2 expression and PGE₂ production in CAFs from biopsies of colorectal cancer tissues are much greater than those from normal fibroblasts (NFs) [33]. Therefore, we should focus on the mechanism how COX-2 expression and PGE₂ production is mediated and how such findings are linked to progression and invasion of colorectal cancers.

**PGEs and Their Receptors**

COX-2 is an enzyme regulating PGE₂ within our body [39]. Prolonged PGE₂ increase is usually a sign of inflammation, cancer genesis and spread. COX-2 mediates biosynthesis and release of prostaglandins using arachidonic acid (AA) as the substrate [39]. In other words, this enzyme first converts arachidonic acid into prostaglandin G₂ and prostaglandin H₂, and then synthesizes prostaglandin D₂, E₂, F₂α, F and thromboxane A₂, exerts their actions through the cognate G-protein-coupled receptors (GPCRs) [39].

Prostaglandins are active lipid compounds which have multiple hormone functions to participate in inflammation and progression of colon cancer [40, 41].

Prostaglandin signaling is involved in the progression of many diseases including chronic diseases such as cancer, suggesting prostaglandins are indeed associated with regulation of both acute and chronic inflammation [9]. The main form of prostaglandin involved in colorectal cancer is PGE₂. PGE₂ can act on the receptors, for example, EP₁, EP₂, EP₃, and EP₄ to induce PGE₂ signal cascade, leading to changes of intracellular calcium, cAMP and some inflammatory factors. As a result, physiological or pathological processes follow [23, 42]. Recent investigations support that PGE₂ may enhance progression of colorectal cancer [41, 43, 44], and EP₄ is a therapeutic target for cancer therapy [45, 46]. COX-2 derived PGE₂ can also contribute to tumor development through several mechanisms including inhibition of apoptosis. However, the mechanisms by which PGE₂ regulates apoptosis are still largely unknown. The EP₂ and EP₄ receptors mediate their activities through cAMP production. Suppression of apoptosis has been seen in intestinal cells by cAMP through the induction of the IAP family member-inhibitor of apoptosis 2 (IAP-2) [47, 48]. Therefore, it is wise to investigate the hypothesis that anti-apoptotic effects of PGE₂ are mediated through cAMP, which results in the induction of the IAP family member c-IAP2.

**The cellular distribution of COX-2 and PGE₂**

The sub-epithelial tissue lies immediately underneath the epithelial layer. The mesenchyme of a tissue lies immediately underneath the epithelium. That is, fibroblasts are the predominant cells of the sub-epithelial layer. Interestingly, the cellular site of COX-2 upregulation in the earliest intestinal cancer tissue is sub-epithelial, and not epithelial.

In fact, recent studies have demonstrated that fibroblasts play a far more varied role than you anticipate. The cells express many receptors for cytokines and hormones and modulate intestinal secretory responses to inflammatory mediators by releasing PGE₂.

The sub-epithelial site of inducible COX-2 expression in early murine intestinal adenomas conflicts with alternative evidence that, rather, upregulation of this enzyme occurs in colorectal...
epithelial cells [49]. To reconcile the issue, we hypothesize that in normal and premalignant colorectal tissue, mesenchymal cells are the principal source of COX-2 expression. Nevertheless, once malignant transformation has occurred, the enzyme is also expressed in the epithelium. The question, then, arises of which mesenchymal cells might express COX-2 in nonmalignant colorectal tissue.

**COX-2 signaling is a marker for tumorigenesis**

COX-2 is a short-lived rate-limiting enzyme [19, 50]. It converts arachidonic acid (AA) to prostaglandins (PGs) and thromboxanes [51]. A major PG product of COX-2 is PGE$_2$, which may regulate angiogenesis, immunity and tumorigenesis [52-54]. In carcinogenesis, COX-2 may be uncontrollably elevated, transcriptionally or post-transcriptionally [55, 56]. Therefore, increase of COX-2 expression is a marker for tumor diagnosis, which is associated with patients' survival rate [17, 57-60]. The significant roles of COX2 signaling has closely been correlated various types of cancer as reported [13, 18, 61-63]. Recent reviews have summarized recent advances in the role of COX-2 and prostaglandin E$_2$ in the pathogenesis of colorectal cancer [18, 61, 62]. A report suggests that PGE$_2$ signaling mediates chronic inflammation in the colorectal microenvironment [86]. Cytokines and other compounds such as DCA, IL-1β, tumor necrosis factor α (TNF-α) and lipopolysaccharide (LPS) may promote expression of COX-2 mRNA and protein in human colorectal fibroblasts, profoundly in cancer-associated fibroblasts (CAF) [25-27]. When stimulated with the pro-inflammatory cytokines interleukin (IL)-1β or TNF-α, orbital fibroblasts express high levels of COX-2 and synthesized correspondingly high levels of PGE$_2$ [87]. Powell and colleagues described a population of specialized sub-epithelial “myofibroblasts” with pleiotropic capabilities, including the ability to modulate intestinal secretory responses to inflammatory mediators by releasing PGE$_2$ [88]. We have noted that IL-1β or TNF-α promotes production of PGE$_2$ by as much as 25-fold in human colorectal fibroblasts we obtained from colonoscopies [30]. Equivalent or greater increases in COX-2 mRNA and more profoundly expressed in the adjacent normal tissue compared to cancer tissue (p < 0.002) [80], suggesting that the role of COX-2 appears to be involved in the early stages of progression of the oncogenic mechanism of CRC, initially affecting the host (tumor microenvironment) of the tumor in normal cells and later tumor epithelial cells of the tumor itself. The authors suggest that treatment should be given to prevent CRC rather than to suppress this progression especially to colorectal tumors in which the expression was found to be greatest.

**COX-2-promoted carcinogenesis is related to angiogenesis**

Suppression of COX-2 inhibits corneal neovascularization in colorectal cancer [81] by promoting production of angiogenic vascular endothelial growth factor (VEGF), a potent angiogenic growth factor [82]. In fact, overexpression of COX-2 promotes overexpression of VEGF, which induces tumor angiogenesis in Apc/COX-2 knockout mouse model [83]. Deletion of COX-2 gene may result in reduced growth in tumor xenografts and vascular density [84], probably via activation of Rac1 and Cdc42 [85]. In all, evidence indicates that COX-2 may induce uncontrollable angiogenesis in colorectal cancer.

**Cytokines and other compounds regulating COX-2 signaling**

Recent reviews have summarized recent advances in the role of COX-2 and prostaglandin E$_2$ in colorectal cancer. Nevertheless, once malignant transformation has occurred, the enzyme is also expressed in the epithelium. The question, then, arises of which mesenchymal cells might express COX-2 in nonmalignant colorectal tissue.
protein expression preceded the increases in PGE₂ synthesis. We also report that inducible COX-2 expression is substantially more robust in cancer-associated than normal colorectal fibroblasts [28]. IL-1β may stimulate expression of COX-2 and production of PGE₂ synthesis in cancer-associated fibroblasts by activating COX-2 promoter activities. The rate at which COX-2 mRNA decays can be dramatically retarded in vitro by PGE₂ [89]. We believe that this mechanism may be an important contributing factor to the enhanced PGE₂ synthesis of cancer-associated colorectal fibroblasts. Those reactions are probably through activation of protein kinase C (PKC) [90], suggesting that stromal cells also play an important role in colorectal carcinogenesis. Such activation may be through enhanced production of PGE₂ and as a result, mediating proliferation, invasion and apoptosis of colorectal cancer cells [25, 27]. Such activation of COX-2 is closely associated with increased proliferation and invasiveness in human colorectal epithelial cancer cells [25, 27, 28], because COX-2 inhibitor (NS398) or PKC inhibitor (Bisindoylmaleimide I, BIM or Staurosporine, STA) can dramatically down-regulate the proliferation and invasiveness of colorectal epithelial cancer cells [25, 27, 28]. Combination of PKC and COX-2 inhibitors can synergistically inhibit melanoma metastasis [91]. We have noted that IL-1β and TNF-α induce mRNA overexpression of COX-2 and promote production of PGE₂ in human colorectal fibroblasts, especially in CRC-associated strains [27, 92]. We have also noted that DCA strongly promotes COX-2 expression and PGE₂ production in colorectal cancer fibroblasts in vitro [25]. Inducible nitric oxide synthetases (iNOS) are also probably involved in the carcinogenesis of colorectal cancer because activation of iNOS by LPS is associated with activation of COX-2 signaling, and inhibition of iNOS by iNOS inhibitor 1400W or iNOS siRNA may nullify production of nitric oxide (NO) and PGE₂ [26]. NO may also promote mRNA production of COX-2 from colorectal cancer cells, for example, HCA7 and HCT116 [56]. We have reported that deoxycholic acid (DCA) also dramatically promotes COX-2 expression in colorectal cancer fibroblasts in vitro [25]. The expression and activity of iNOS and COX-2 may also be induced by LPS in those fibroblasts, resulting in increased NO production and COX-2 expression [26]. DCA is also a transcriptional activator of COX-2 in esophageal cancer cells [93].

**Nonsteroidal Anti-inflammatory drugs**

For decades, a significant progress is achieved to discoveries of effective drugs for CRC. One of those is nonsteroidal anti-inflammatory drugs (NSAID) which inhibit COX-2 [94-96]. NSAID includes aspirin, ibuprofen, naproxen, nimesulide and sulindac acid. Different NASID may act via different signaling pathways. For example, ibuprofen, indomethacin and naproxen can bind the activity site of COX-2, inhibit its activity reversibly, while aspirin acetylates the activity site of COX-2, attenuating its activity irreversibly. Some NSAID drugs for example, Aspirin, can facilitate the effect of COX-2 inhibitors for treatment of stage III colorectal cancer [97]. In fact, Aspirin may reduce colon cancer mortality in women by as much as 50 % [9, 98, 99]. Recently, a hybrid drug KSS19, a combination of NSAID refecoxib and cis-stilbene, has been found to be a potent COX-2 inhibitor, which inhibits colon cancer cell growth effectively [100].

Although COX-2 inhibitors are promising candidates for treatment of colorectal cancer, some concerns for treatment of colorectal cancer by COX inhibitors have been raised. For example, an elevated risk of myocardial infarction may be linked to its usage [101]. In addition, the extended use of nonselective NSAID is also associated with a number of pathological symptoms, for example, abdominal pain, dyspepsia, gastritis, gastrointestinal bleeding, nausea, and perforation of gastroduodenal ulcers [102]. Therefore, no major clinical trials of those inhibitors were successfully completed due to concerns of their adverse effect. Anyway, NSAID are effective in certain degrees for prevention and treatment of colorectal cancer. For example, a randomized trial demonstrates that NSAID are preventive for colorectal cancer on the patients with polyps [103, 104]. According to the results of large-scale trials, including the Adenomatous Polyp Prevention on Vioxx trial [105], the Adenoma Prevention with Celecoxib trial [104], the Prevention of Colorectal Sporadic Adenomatous Polyps trial [106] and colon polyp prevention trial [107], COXIB are effective for prevention of recurrence from sporadic colon cancer. Regular consumption of NSAID is also helpful for lowing the risk of colorectal, breast, lung and prostate cancer [108]. In all, it is still unknown how to prevent the potential risk when COX inhibitors are used for treatment of colorectal cancer.

To decrease the risk from COX inhibitors, many researchers have used low dose of COX inhibitors with other NSAID drugs that target other critical pathways in carcinogenesis. For example, combination of celecoxib with erlotinib (an EGFR tyrosine kinase inhibitor) is more effective to control polyp formation using an ApcMin/+ mice model and to inhibit cancer growth in a xenograft model [109]. Celecoxib with erlotinib treatment is more effective for treatment of the advanced non-small cell lung
cancer [110]. A 5-lipoxygenase inhibitor may inhibit resistant tumor cells to SC-236 (COX inhibitor) and tumor growth in a breast cancer animal model [111]. Combined treatment of celecoxib with peroxisome proliferators-activated receptor-γ agonist is better than either alone in a mouse breast cancer model [112]. Combination of aromatase inhibitors with celecoxib is better for patients suffering from metastatic breast cancer than either alone [113]. Therefore, we should reconsider the prospect of COX inhibitors for treatment of colon cancer.

**Perspective**

From previous work on certain types of cancer, COX-2 may be a key indicator to predict cancer prognosis. Along with the product PGE2, COX-2 is a major stimulator for progression of colorectal cancer. Up to now, the reports from basic and clinical investigations have shown that inhibition of PGE2 synthesis by specific COX-2 inhibitors, for example, nonsteroidal anti-inflammatory drugs (NSAIDs) may decrease the risk and improve prognosis of carcinogenesis of various types of cancer including colorectal cancer [64, 114-120]. Therefore, we need to strive forward to work out a protocol for successful use of COX-2 inhibitors in clinical applications to colorectal cancers and other types of cancer as well.

**Summary**

To summarize, the scientific basis for the current proposal, outlined above, is as follows: fibroblasts from mesenchymal (stromal) layer are the major target of cytokines such as IL1β and TNFα; fibroblasts are the predominant mesenchymal cells; fibroblasts from nonneoplastic colorectal tissue are a potent source of COX-2 expression, which is firmly established as an important factor in colorectal carcinogenesis. We believe that the investigation of COX-2 gene regulation is vital for control of this disease.

**Abbreviation**

CAF: cancer-associated fibroblast; COX: Cyclooxygenase; CRC: colorectal cancer; DCA: deoxycholic acid; EGF: epithelial growth factor; EP, FP, IP, TP and DP: prostanooid receptors; IL-1β: Interleukin 1β; iNOS: inducible nitric oxide synthase; LPS: lipopolysaccharide; NF: normal fibroblast; NO: nitric oxide; NSAID: nonsteroidal anti-inflammatory drug; p53: tumor protein p53; PG: prostaglandin; PKA: cAMP-dependent protein kinase; PKC: protein kinase C; TNF: tumor necrosis factor.

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**Author Contributions**

Juan Sheng, Hong Sun, Fubing Yu and Bo Li participated in collecting the information and drafting the manuscript. Yuan Zhang and Yingting Zhu and finalize the manuscript.

**Competing Interests**

The authors have declared that no competing interest exists.

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