ABSTRACT

Colorectal cancer (CRC) is one of the most life threatening disease with escalating mortality and morbidity. Some known causative factors include lifestyle, alcohol etc. but there are other aspects like long standing colitis, inhibition of apoptotic proteins, chemokines and their receptors and inflammasomes which increase the chances of CRC progression. In a clinical setting of diseases like colitis and Crohn’s disease, the risk of CRC varies on degree of inflammation, expression of chemokines and cytokines and other molecular alterations. Hence, the underlying molecular inductions determine the fate of CRC progression through extent of epithelial-mesenchymal transition, metastasis and invasive ability. The present viewpoint will showcase some of the latent and notable considerations in CRC progression.

KEYWORDS: Colon cancer; Apoptosis; Chemokines; Inflammasomes; Inflammation.

INTRODUCTION

Colorectal cancer (CRC) is a heterogeneous disease causing more than 4 million deaths. Apart from dietary factors, lifestyle and genetics, advancements from other ailments like inflammatory bowel disease are reasons for cause of CRC. Amongst the molecular pathogenesis, inflammation has a direct link to CRC. Not only by the assessment of mucosal biopsies revealed escalated levels of cell division and cell death, there were several conditions like epithelial mesenchymal transition (EMT) which had a higher role in cancer invasion rate or metastasis.

Cytokines which are categorized under small proteins play an important role in cell signaling. In carcinogenic conditions like colitis associated cancers, cytokines were said to have a significant influence. Factors like nuclear factor-kappa B (NF-kB), tumor necrosis factor (TNF) etc. regulate cell cycle, apoptosis, reactive oxygen species and mutations. Thus the present viewpoint will list few perplexing conditions, inflammatory modulators and receptors which would aid in CRC progression.

HYPOXIA - THE ENVIRONMENT FOR PROGRESSION

Hypoxia plays a huge role in the augmentation of solid tumors. During hypoxia, there is restricted oxygen supply which is regulated by hypoxia-inducible factor (HIF). HIF is not only involved in transcriptional activity of several genes, but its high levels are associated with poor prognosis of CRC. It was well evidenced that hypoxia promotes migration ability of cancer cells conditioning EMT. Cytokines such as tumor necrosis factor-α (TNF-α) were said to have direct relation with HIF-α levels enhancement via NF-kB activation. Additionally, Forkhead Box M1 (FOXM1) also advances EMT as FOXM1 promotes urokinase-type plasminogen activator receptor (uPAR) and matrix metalloproteinase 2 & 9.

In the condition of CRC, endoplasmic reticulum (ER) stress is down-regulated by hypoxic condition thereby increasing the aggression of metastasis. It was clearly observed in several solid tumors that FOXM1 transcrip-
tion factor was over expressed and ER stress was far reduced. 

The condition of hypoxia can be attenuated either by sensitizing the cancerous cells to ER stress or blocking uPAR pathway. Figure 1 depicts the role of hypoxia in regulating mammalian target of rapamycin (mTOR) in articulating vascular proliferation. It is evident that dysfunction in the PI3k/mTOR pathway leads to pathogenesis of CRC. The hypoxia inducing factors HIF-α and HIF-β trigger chemokines to initiate epithelial mesenchymal transition which is a lead step in metastasis.

CHEMOKINES AND THEIR RECEPTORS

Clinical evidences state that chemokine receptor 6 (CCR6) expressions was up-regulated in colorectal cancer. Chemokines in general can regulate the movement of tumor cells. CCR7, CCR9, C-X-C chemokine receptor type 1 (CXCR1), and C-X-C chemokine receptor type 2 (CXCR2) are also detected in tumor cells and their ligands can induce the chemotaxis of the corresponding receptor-expressing cells. These chemokines act as inducers of invasion within the tumor and also in regards to movement to other organs. When chemokine (c-c motif) ligand 20 (CCL20) triggered p130 phosphorylation, there was an enhanced migration and proliferation of cancer cells. The expression of chemokine receptor C-X-C chemokine receptor type 4 (CXCR4) in CRC metastasis was well established. Factors like high invasion rate, development of liver metastasis and colon carcinoma micro metastasis of liver were reported by CXCR4. Direct association of lymph node metastasis in CRC and CCR7 was observed in a study by Gunther et al. Not only does chemokine-chemokine receptor interaction supports metastasis in colon cancer, CCR6-CCL20 interaction was found to be involved in various inflammatory and immune disorders. CCR6 role in advanced colon cancer was clearly shown and also CCR6-CCL20 was reported to have involvement in EMT which resulted in poor outcome of colon cancer. CCR6 was described as independent risk factor in liver metastasis. On the other hand, CCL20 levels were higher in colon cancer patients with liver metastasis compared with controls without metastasis. In a study by Kawada and colleagues, amongst the colon cancer samples with CXCR3 expression, there was significantly high lymph node metastasis recorded. CXCR3 expression levels were directly proportional to poor prognosis which clearly indicates that CXCR3 activation was concomitant with colon cancer metastasis preferentially to the draining lymph nodes with poorer prognosis.

mTOR’s Role in Colon Cancer

Hyper activation of mechanistic target of rapamycin (mTOR) is due to signaling malfunctions upstream of mTOR in phosphatidylinositol-3-kinase (PI3K)/Akt/mTOR pathway. Mutagenesis in PI3Kα occurs in late tumorigenesis which is generally evident in more than 30% of colon cancers. mTORC1 inhibition had a role in improving regeneration capacity of intestinal stem cells but at the same time, abberant stimulation could trigger carcinogenesis. In humans, immunohistochemical studies on colorectal carcinoma samples revealed that mTORC1 signaling takes place in the early onset of tumorigenesis and plays a role in transforming normal cells to neoplastic. mTORC1 and mTORC2 were implicated in colorectal cancer biology to a greater extent.

In the conditions of in vivo mTORC2 was down-regulated, there was a clear trend of reduction of proliferation and also there was reduction in the formation of tumor xenografts. EMT was also regulated by mTORC1 and mTORC2 and thus sequencing metastasis of CRC.

Targeting kinases like polo-kinase 1 which is highly expressed in proliferating cells during G2 and M phase of cell cycle was regarded as a potential target. Besides, induction of apoptosis via mTOR suppression is a strategic therapeutic intervention. BI2536 was one amongst the potent polo-kinase 1 inhibitor used in combination with NVP-BEZ235 which is a dual PI3K/mTOR inhibitor. These studies not only show the importance of epigenetic mechanisms in cancer signaling but also clearly denote the role of mTOR signaling in CRC progression.

Nod like receptors (Nlrp3) and their Role in CRC

Chronic inflammation is considered as a risk factor for the progression of CRC. Nod like receptor (Nlrp3) is a protein which assembles the inflammasome which is responsible for secretion of pro-inflammatory cytokines like IL-1β. This aids in cellular
functions like repair via immune cell activation and triggering pro-inflammatory cytokines. Inoperative NLR activation results in cell proliferation and tumorigenesis. Mice deficient of Nlrp3 were prone to dysplasia and tumor formation thus clearly stating the linkage between Nlrp3 deficiency and susceptibility to colitis associated carcinogenesis. Apoptosis associated speck like protein-1 (ASC) and caspase 1 are two major components of Nlrp3 which play a major role in inducing apoptosis of cancer cells. In experimental conditions of knocking out these two major components of Nlrp3, the mice developed tumorigenesis. Interleukin 18 (IL-18) which is secreted by Nlrp3 exerts extensive anti-tumor activity and aids in epithelium repair. The fact that IL-18 offers its protection against colitis and carcinogenicity was described in mice models where caspase 1 and Nlrp3 were knocked out and there was development of CRC. It was well defined that IL-18 promotes entereocyte proliferation to repair chemically induced injury of colonic epithelium and also inhibits hyperplasia during chronic stages of colitis. Also in the acute conditions of disease, IL-18 offers restoration of barrier integrity by controlled proliferation of stem cells at the base of the intestinal crypt which indirectly helps in intestinal homeostasis. In experimental conditions of azoxymthane (AOM) or dextran sulfate sodium (DSS) induced colitis, phosphorylated levels of STAT1 were reduced in the colon where the restoration is possible by IL-18. Nlrp3 inflammasome not only reduced the tumorigenesis but also suppressed liver colon cancer metastatic growth.

**SPINK1**

Serine peptidase inhibitor kazal type-1 (SPINK1) has been investigated for its role in multiple human carcinomas especially CRC. SPINK1 mutation was associated with pancreatitis. Epidermal growth factor receptor (EGFR) involves in intra-cellular message transduction. High expression of EGFR is also associated with poor prognosis. SPINK1 and epidermal growth factor (EGF) bind to EGFR and stimulate proliferation via mitogen-activated protein kinases (MAPK) which was recorded in pancreatic adenocarcinoma. In a study by Chen et al., the SPINK1 protein played a role in tumor proliferation and malignant progression in CRC through the EGFR pathway. SPINK1 was also used as a marker for predicting tumors in response to anti-EGFR treatment in CRC patients. Metallothionein expression in colon cancer was analogous with poor survival and SPINK1 caused CRC progression by down regulating metallothioneins expression.

**COX2**

The interest in cyclooxygenase (COX2) inhibitors as therapeutic interventions rose after its ability as an anti-carcinogenic agent with 40-50% reduction in CRC prevalence in non-steroidal anti-inflammatory drugs (NSAID) users. COX2 over expression has been reported in numerous colorectal adenomas and adenocarcinomas. The direct relation of COX2 with intestinal polyps. Ideal tumorigenesis mechanisms of COX2 are inhibition of apoptosis, increased effectiveness of invasiveness etc. It is noteworthy that though COX2 is preponderant to the cytoplasm of cancerous epithelial cells, it is negatively expressed in normal epithelium. In regards to the mucosa, CRC samples were found to have more messenger RNA (mRNA) levels than normal mucosa. In a clinical study involving 76 patients suffering with CRC, COX2 was conferred to advancement of the disease and decreased patient survival. The underlying mechanism of COX2 associated carcinogenicity is by attenuation of the mitochondrial apoptotic pathway distinguished by reduced cytochrome C release and diminished caspase activity. There was an increased expression of anti-apoptotic genes Bcl-2 recorded in cell lines like HCT-15. COX2 also was involved in death receptor (DR) pathway via DR expression in HCT-15 cells. Targeting CRC with COX2 inhibitors may not be a complete solution though NSAID users had less probability of CRC.

**CONCLUSION**

Biological mechanism underlying CRC have been markedly improved in recent years with respect to therapeutic interventions selecting target receptors, inflammasomes, cytokines and other chemokines. Finding the right agent for CRC treatment was achieved by inducing apoptosis, reducing EMT and decelerating proliferation and invasiveness. Treatment for CRC is not only designed via symptomatic approach but also targeting signaling mechanisms which drive the disease. Thus, this article highlights some of the facets in CRC which when targeted or inhibited can turn out to be a potential treatment escalating the quality of life (QoL) and improved prognosis.

**CONFLICTS OF INTEREST**

The authors declare that they have no conflicts of interest.

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