A Novel Anti-cancer Strategy Targeting CD44-positive Cancer Stem Cells

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Introduction

Despite progress in cancer therapy, cancer continues to be a common and lethal disease worldwide. Recent advances in cancer stem cell (CSC) research have indicated that CSCs, a subpopulation of cancer cells that can self-renew and differentiate into various cell types, are responsible for tumor initiation, recurrence, and metastasis [1,2]. The existence of CSCs is considered to be a cause of drug-resistant cancers, since they eliminate sources of biochemical stress. Numerous studies have been performed to isolate CSCs from other cancer cells, and several significant markers have been identified for some cancers. Importantly, the expression of several CSC markers has been reported to enhance tumor progression and maintain a malignant phenotype.

CD44 is one of the most famous CSC markers for gastrointestinal cancer. Several studies have demonstrated that CD44 enhances the malignant phenotype. High expression levels of CD44 are significantly correlated with both tumor size and invasiveness [3,4]. The expression of a splice variant of CD44 (CD44v) is frequently associated with the progression of colon and gastric cancers [5-7]. The multifunctional roles of CD44 in the microenvironment and in the regulation of cancer stemness, as well as the demonstrated prognostic value of CD44 for some cancers, suggest that targeting CD44 could be a promising approach with the potential to eliminate CSCs.

In CSCs, reactive oxygen species (ROS) are kept at low levels. ROS play an important role in the formation of a proliferation-permissive intracellular environment and the preservation of a self-renewal capacity [8,9]. In cancer tissues, a high ROS level is counteracted by antioxidant enzymes that are produced by cancer cells or inflammatory cells [10]. Although ROS produce oxidative stress in most cancer cells, CSCs possess enhanced mechanisms for protecting against such stress induced by ROS. One of the main mechanisms is glutathione metabolism. CD44v isoforms interact with and stabilize xCT (SLC7A11), a subunit of the cystine/glutamate antiporter known as system xc(\textsuperscript{−}). The expression of xCT on the cell surface is essential for the uptake of cystine, which is required for intracellular glutathione (GSH; a major antioxidant), and thereby potentiates the ability of cancer cells to increase glutathione synthesis and defend themselves against ROS [11-13]. This mechanism is reportedly associated with not only drug resistance, but also the promotion and growth of tumors. The ablation of CD44 induced the loss of xCT from the cell surface and suppressed tumor growth in a transgenic mouse model of gastric cancer [12].

Despite accumulating evidence regarding CD44-targeting strategies for CSCs, the promised agents have not yet been established [14,15]. 5-Aminosalicylic acid (5-ASA), the most commonly used anti-inflammatory medication for inflammatory bowel diseases (IBD), has been reported to reduce the risk of CRC significantly [16,17]. Since chronic inflammation is a key factor in carcinogenesis, the effect of 5-ASA therapy on reducing the risk of CRC may be mediated by a reduction in mucosal inflammation. Recent studies have revealed that sulfasalazine, one of the 5-ASA drugs, might have an even more important role. Sulfasalazine consists of sulfapyridine and the active moiety 5-ASA that are released in the gut. Intact sulfasalazine, but not its metabolites, was found to inhibit the cystine/glutamate transporter (Figure 1) [18]. As a result, GSH synthesis can be efficiently suppressed by the absence of intracellular cystine [19]. Several studies have also shown the function of sulfasalazine as a specific inhibitor of xCT-mediated cystine transport and its ability to inhibit the growth, invasion, and metastasis of several types of cancer [11,18,20]. We have recently shown that in a mouse experiment, sulfasalazine selectively induced apoptosis in CD44v-expressing head and neck cancer cells and promoted the sensitivity of tumors to anti-cancer drugs [21]. Additionally, in a gastric cancer mouse model, sulfasalazine treatment suppressed the development of precancerous lesions and cancer [22]. Therefore, sulfasalazine might be a promising agent targeting CD44-positive CSCs for the prevention and treatment of CSCs.

Figure 1: Scheme for CD44-targeting therapy using sulfasalazine (modified from Reference 12); System xc(\textsuperscript{−}) is a cystine/glutamate exchange transporter composed of a light-chain subunit (xCT) and a heavy-chain subunit (CD98hc); CD44v maintains a high level of interaction with and stabilizes xCT (SLC7A11), a subunit of the cystine/glutamate antiporter known as system xc(\textsuperscript{−}). The expression of xCT on the cell surface is essential for the uptake of cystine, which is required for intracellular glutathione (GSH; a major antioxidant), and thereby potentiates the ability of cancer cells to increase glutathione synthesis and defend themselves against ROS [11-13]. This mechanism is reportedly associated with not only drug resistance, but also the growth and proliferation of tumors. The ablation of CD44 induced the loss of xCT from the cell surface and suppressed tumor growth in a transgenic mouse model of gastric cancer [12].

Based on accumulated evidence, we conducted a retrospective review of the records of ulcerative colitis (UC)-associated CRC patients who underwent curative colorectal surgery [23]. We then compared the patients' clinicopathological findings according to the length of sulfasalazine administration. As a result, long-term sulfasalazine administration was shown to reduce proliferative CD44v9-positive...
cells in cancer and increase the degree of adenocarcinoma differentiation. Furthermore, an in vitro assay revealed that sulfasalazine promoted the expression of epithelial differentiation markers (E-cadherin and CDX2) and inhibited the proliferation of CD44-positive cancer cells. These findings indicate the importance of CD44v9-positive cells in UC-associated cancer progression and differentiation, suggesting that sulfasalazine may serve as a novel therapeutic agent that targets CD44v9-positive cells. There were some limitations in this study: a small number of patients, not considering the dose intensity of sulfasalazine, and the limitation of patients to UC-associated cancer patients. Therefore, further study is needed to investigate whether sulfasalazine affects sporadic cancers as well.

Consistent with previous observations, CD44 is considered to be an ideal target for the development of clinical therapeutics against CSCs. However, only a few clinical trials targeting CD44 have been conducted so far, and none of these trials have resulted in an effective treatment [14]. The limitations demonstrated in previous clinical trials examining anti-CD44 therapies highlight the need for the development of improved anti-CD44 reagents, such as high-affinity anti-CD44v peptides, to replace highly immunogenic antibodies [24] as well as the use of CSC-specific CD44v peptides for cancer vaccines based on promising CD44 DNA vaccine studies in animal models [25,26].

Conclusion

In conclusion, targeting CD44 is a promising approach with the potential to eliminate CSCs. Better understanding of the fundamental characteristics of CD44 and improvements in drug development targeting CD44 should bring new hope to patients with life-threatening cancers.

References