Abstract. Breast cancer is the most common malignant neoplasm in women worldwide, and the treatment regimens currently available are far from optimal. Targeted therapy, based on molecular typing of breast cancer, is the most precise form of treatment, and CXC chemokine receptor 2 (CXCR2) is one of the molecular markers used in targeted therapies. As a member of the seven transmembrane G-protein-coupled receptor family, CXCR2 and its associated ligands have been increasingly implicated in tumor-associated processes. These processes include proliferation, angiogenesis, invasion, metastasis, chemoresistance, and stemness and phenotypic maintenance of cancer stem cells. Thus, the inhibition of CXCR2 or its downstream signaling pathways could significantly attenuate tumor progression. Therefore, studies on the biological functions of CXCR2 and its association with neoplasia may help improve the prognosis of breast cancer. Furthermore, the targeting of CXCR2 could supplement the present clinical approaches of breast cancer treatment strategies. The present review discusses the structures and mechanisms of CXCR2 and its ligands. Additionally, the contribution of CXCR2 to the development of breast cancer and its potential therapeutic benefits are also discussed.

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1. Introduction

Breast cancer is an aggressive malignancy, and is a major threat to the health of women worldwide. According to the 2018 Global Cancer Statistics, breast cancer is the second leading cause of cancer-associated mortality worldwide, following lung cancer. Furthermore, there were ~2.1 million new cases in 2018 worldwide, and these accounted for approximately a quarter of the total number of female patients with cancer (1). In China, breast cancer is the fifth leading cause of cancer-associated mortality, according to statistics collected by the China National Cancer Center in 2018 (2). On the therapy options for breast cancer, molecular typing is a critical basis. Based on the expression of estrogen receptor (ER), progesterone receptor (PR), human epidermal growth factor receptor 2 (HER2) and Ki-67, there are five primary molecular subtypes of breast cancer: Luminal A, luminal B, triple-negative/basal like, HER2-enriched and normal-like (3). Currently, molecular typing-based integrative treatment regimens consist of surgery, radiotherapy, chemotherapy, endocrine therapy and targeted therapy, which have improved treatment efficacy, including improving overall survival (OS) and progression-free survival (4,5). However, most patients with cancer still experience drug resistance, recurrence and metastasis (6). In recent years, CXC chemokine receptor 2 (CXCR2) has emerged as a critical functional receptor. CXCR2 serves an important role in various aspects of breast cancer, including the diverse range of pathological processes associated with tumor progression (7). Combined analysis demonstrated that patients with solid tumors and elevated CXCR2 expression had poorer prognosis, including OS, recurrence-free survival and disease-free survival (8). The present review summarizes the biological roles of CXCR2 in breast cancer and systematically examines the pathways and mechanisms of CXCR2 associated with the initiation and development of breast cancer, as well as the potential therapeutic value of an anti-CXCR2 treatment.
2. Structure and interactions of the chemokine CXC and its receptor CXCR2

Chemokines are small (6-14 kDa), secreted peptides that mediate the migration of leukocytes to inflammation and secondary lymphoid organs. Chemokines are also essential for other pathophysiological processes, including infectious diseases, asthma, and atherosclerosis (9-11). Based on the position of the four cysteine (Cys) residues at the N-terminus, the chemokines are divided into four subtypes: CXC (α, CC (β), XC (γ) and CX3C (δ)) (12). To date, 50 chemokines and 20 chemokine receptors have been identified; the majority of chemokines belong to the CC and CXC subgroups (13). CXC is further classified into ELR+ and ELR- CXC chemokines, based on the glutamate-leucine-arginine (Glu-Leu-Arg, ELR) sequence that occurs before the first Cys at the N-terminus (14).

The coding sequence of the CXCR2 gene is located at 2q34-35 and contains three exons and two introns (15). CXCR2 is a G protein-coupled receptor that contains seven transmembrane regions, an extracellular N-terminus and an intracellular C-terminus (16). The N-terminus, the fourth transmembrane domain and the second extracellular loop are prerequisites for ligand binding and specificity, and determine the rate of receptor internalization (17). The C-terminus region is involved in receptor phosphorylation, internalization and G protein coupling. Only ligand monomers activate CXCR2, which interact via a two-site, two-step model. This model involves the binding of the N-terminal domain of CXCR2 with the N-loop and core domain of ligands at site 1. At site 2, CXCR2 activation is conferred by the insertion of the N-terminus signal domain of the ligands into the orthosteric pocket of the receptor (18). Interleukin-8 (IL-8), also termed CXCL8, is a CXCR2 ligand and was the first CXC chemokine derived from the medium of lipopolysaccharide and polyhydroxyalkanoate-stimulated human monocytes (19). In humans, CXCR2 is also known as IL-8 receptor B and interacts with ELR+CXC chemokines with high affinity. These chemokines include GRO-α/CXCL1, GRO-β/CXCL2, GRO-γ/CXCL3, ENA-78/CXCL5, GCP-2/CXCL6, NAP-2/CXCL7 and IL-8/CXCL8, which mediate angiogenesis (20).

3. Signaling pathways of CXCR2 activation

CXCR2 possesses no kinase activity. In addition to being coupled to G proteins, it binds to other proteins, such as G protein coupled receptor kinase 2/6, β-arrestin1/2, adaptor protein-2, protein phosphatase 2A and vasodilator-stimulated phosphoprotein. This enables CXCR2 to mediate different signaling cascades in breast cancer (21). G proteins are heterotrimeric protein complexes that are comprised of three subunits, known as α, β and γ, which are inactive in their resting state. Upon binding of the ligands to CXCR2, CXCR2 physically couples to the G protein (22). CXCR2 becomes subsequently activated and the guanosine diphosphate linked to the Gαs subunit of the G protein complex is converted to GTP. This transformation causes Gαs to dissociate from the receptor and Gβγ, leading to the activation of several downstream signaling pathways.

As presented in Fig. 1, the three main pathways activated by CXCR2 are the phosphatidylinositol-3 kinase (PI3K)/Akt pathway, the phospholipase C (PLC)/protein kinase C (PKC) pathway and the Ras/Raf/extracellular signal related kinases (ERK1/2) pathway (23-25). The PI3K/Akt pathway is the primary downstream signaling cascade mediated by PI3K, and protein kinase B (PKB) plays an essential role in this pathway. PKB is the oncogene product of the retrovirus AktB, which is also known as Akt. Akt activates IkB kinase (IKK), which phosphorylates IkB to expose the nuclear localization signals of NF-κB. This allows NF-κB subunits to translocate to the nucleus (26). The PI3K/Akt pathway is one of the most commonly altered pathways in human malignant tumors, and it is critical for cell survival, motility and angiogenesis (27). The phosphorylation of CXCR2 also results in the activation of mitogen-activated protein kinases (MAPK) signaling, which includes ERK. Ras activates ERK through Raf and MEK, and phosphorylated ERK translocates from the cytoplasm to the nucleus. This mediates the transcriptional activation of c-Jun, c-Fos, Elk-1, AP1 and ATF, which participate in various biological functions, such as cell proliferation and differentiation, morphology maintenance, cytoskeleton construction, apoptosis and tumorigenesis (28,29). The activation of PLC induces another signaling transduction pathway that generates two secondary messengers, inositol triphosphate and diacylglycerol. This leads to calcium mobilization from the endoplasmic reticulum and activates PKC. Subsequently, c-Jun N-terminal kinase influences cell apoptosis by mediating the activity of p53 and Bcl-2 (30,31). Additionally, the release of the Gαs subunit from the G protein trimer inhibits the conversion of ATP into cyclic AMP by adenylate cyclase, and this decreases the intracellular levels of cyclic AMP (21). Furthermore, CXCR2 signaling could trigger the pathways of focal adhesion kinase (FAK), Rho, Rac, the Janus kinase/signal transducer and the activator of the transcription pathways (32-34).

These pathways act as modulators in breast cancer cell metabolism, survival, proliferation, apoptosis, angiogenesis, transduction and motility. These form a positive feedback loop to enhance CXCR2 functionality by upregulating the expression of cytokines and chemokines, such as CXCL8 (19).

4. Biological characteristics of CXCR2 in breast cancer

CXCR2 and breast cancer growth. Tumorigenesis is a process that involves multiple genes, several steps and various signaling pathways, including the complex regulatory network of chemokines and their receptors (35,36). Studies have found that the CXCL8-CXCR2 axis is closely associated with multiple stages of breast cancer growth. These include the regulation of transcription of CDK inhibitors p21Cip1, p27Kip1 and p57Kip2, and the induction of tumor cell proliferation, differentiation, stress response and apoptosis (15,20,37). Paradoxically, CXCL8 enhances the immune system response and increases its ability to execute anti-tumor effects, whereas it also transforms the tumor microenvironment to promote tumor growth. Compared with healthy volunteers, patients with breast cancer have elevated levels of CXCL8, and the severity of this overexpression is positively correlated with disease stage (38). The levels of CXCR2 in malignant tissues are higher than those in benign and normal tissues (39). Polymorphisms in the CXCL8 and CXCR2 genes also suggest that elevated CXCL8 and CXCR2 expression may be risk factors of breast cancer (38).
However, the biological significance of CXCR2 in cancer cell proliferation remains controversial. Shao et al. (40) performed small interfering RNA-mediated knockdown of endogenous CXCL8 that upregulated p27Kip1 and downregulated cyclin D1. The decreased Akt phosphorylation and NF-κB activation resulted in reduced cell proliferation in both MDA-MB-231 and BT549 breast cancer cell lines. This indicated that CXCL8 and CCL2 overexpression enhances tumor proliferation (40). By contrast, other studies have shown that the overexpression of CXCR2 induces premature senescence, and silencing of CXCR2 prolongs cell passage via p53, NF-κB or C/EBPβ-associated pathways (39,41). Overall, several studies have reported that CXCR2 is a tumor-stimulating receptor that could be exploited as a marker of poor prognosis in a variety of cancer types. Thus, inhibiting CXCR2 production may promote cancer cell apoptosis (42,43). Therefore, CXCR2 may have different functions in normal, precancerous and tumor cells and requires further investigation.

In the tumor microenvironment, breast cancer growth in both autocrine and paracrine manners are regulated by CXCR2 and its ligands produced by stromal cells (44). Furthermore, neutrophils, myeloid cells and bone marrow-derived suppressor cells express CXCR2 and assist in tumor cell proliferation (44). Following the entry of neutrophils into the tumor site, an increase in cytokine secretion contributes to the production of an inflammatory microenvironment (45). Additionally, bone marrow-derived suppressor cells differentiate into M2-type macrophages, which facilitate cancer cell growth (46). Previous studies have demonstrated the knockout of the CXCR2 gene in host cells to inhibit tumor growth and increased tumor cell apoptosis (47-49).

CXCR2 and breast cancer angiogenesis. Once tumors exceed 1-2 mm in diameter, angiogenesis is initiated for growth and metastasis (50,51). CXCR2 affects angiogenesis in breast cancer primarily by interacting with CXCL8 and CXCL1, however the specific mechanism is yet to be determined (52-54).

Addison et al. (53) detected the expression of CXCR2 using a CXCR2 antibody in human microvascular endothelial cells and confirmed that the chemotaxis of ELR+CXC chemokine-mediated microvascular endothelial cell was obstructed, and was sensitive to pertussis toxins (53). Studies in CXCR2-deficient mice indicated that CXCL8 is the strongest ligand for CXCR2, and is mediated by the activation of the ELR+CXC chemokine (52). In cancer cells, CXCL8 and vascular endothelial growth factor (VEGF) cooperate to establish and expand tumor neovascularization. Furthermore, glucose deprivation and endoplasmic reticulum stress

Figure 1. Structure, major signaling cascades and receptor recycling of CXCR2. CXCR2 belongs to the G protein-coupled receptor family that possesses seven transmembrane structures. It contains an extracellular N-terminus, an intracellular C-terminus, three extracellular loops and three cytoplasmic loops. Following ligand binding, CXCR2 physically couples to the G protein. Subsequently, CXCR2 is activated and the GDP linked to the Gα subunit of the G protein complex is converted to GTP. The Gα subunit coupled to the inner cell membrane dissociates from CXCR2 and the Gβγ subunits. Several downstream pathways are induced, and the main three signaling pathways are via PI3K/Akt, PLC/PKC and Ras/Raf/ERK1/2. Moreover, the Gα subunit inhibits adenylate cyclase activity and decreases the efficiency of ATP conversion to cAMP. GRK phosphorylates the C-terminus of the receptor, and mediates the desensitization and endocytosis of the receptor via β-arrestin recruitment of endocytic components. AP-2 also regulates CXCR2 internalization and sequestration. Internalized CXCR2 is subjected to degradation by lysosomes, or recycled to the outer membrane surface. These pathways modulate cell metabolism, survival, proliferation, apoptosis, angiogenesis, transduction and motility. As a positive feedback loop to enhance CXCR2 functionality is formed by upregulating the expression of cytokines and chemokines. CXCR2, CXC chemokine receptor 2; PI3K, phosphatidylinositol-3 kinase; PLC, phospholipase C; PKC, protein kinase C; ERK, extracellular signal related kinase; cAMP, cyclic AMP; GRK, G-protein coupled receptor kinase; GPCR, G-protein coupled receptor; GDP, guanosine disphosphate; AC, adenylate cyclase.
effectively induce the upregulation of CXCL8 (55). CXCL8 and VEGF are regulated by distinct pathways in different cell lines. MDA-MB-231 cells mainly activates the MAPK-ERK pathway, and the activity of the PI3K/Akt pathway is increased in GIH01A cells. Both signaling pathways are activated in MDA-MB-468 and Hs578T cell lines (56). CXCL8 generated by endothelial cells binds to CXCR2 to mediate interactions between CXCR2 and VEGFR receptor 2 (VEGFR2). This includes the transactivation of VEGFR2 via Src kinase-mediated receptor phosphorylation, which is required for CXCL8 to induce endothelial cell permeability (56). The CXCL8-CXCR2 axis also induces VEGF transcription and stimulates VEGFR2 activation through the NF-κB pathway in endothelial cells (57). Moreover, the CXCL8-CXCR2 axis activates the expression of EGFR to mediate endothelial cell migration and capillary formation (58). It also elevates integrin αvβ3 levels, which serve a key role in endothelial cell survival and cancer cell migration during tumor angiogenesis (59).

Another study revealed that the expression of CXCL8 in ER+ cells was lower than that in ER− cells, and exogenous ERα substantially interfered with CXCL8 expression. This suggests that the inactivation of ERα and upregulation of CXCL8 could promote angiogenesis in human breast cancer (60). The silencing of CXCR2 further indicated the importance of CXCL8-mediated angiogenesis. Nannuru et al (61) analyzed the microvessel density of primary tumor sections, and found that silencing CXCR2 in C666 cells considerably decreased tumor angiogenesis compared with the control group.

Furthermore, thrombin stimulates tumors to secrete CXCL1 in endothelial cells, which reinforces tumor angiogenesis. Thus, thrombin-induced angiogenesis could be perturbed by the CXCL1 antibody (54). In 4T1 cells, shRNA-knockdown of CXCL1 impeded tumor growth and angiogenesis (54).

**CXCR2 and breast cancer metastasis.** Metastasis is a basic biological characteristic of malignant neoplasia. Distant metastasis confers breast cancer a worse prognosis, with the five-year survival rate of 27% in the United States between 2008 and 2014, whereas the five-year survival rate of the localized stage was of 99% (62). Metastasis occurs predominantly through the lymphatic system, blood, direct infiltration and planting. This process is extremely complex, dynamic and continuous, and contains several independent processes. For example, when tumors metastasize via the blood circulation, the cancer cells proliferate within the primary lesion and form new blood vessels. Subsequently, the aggressive cells detach from the primary tumor. This is followed by epithelial-mesenchymal transition (EMT), which results in cells with properties similar to interstitial cells. Epithelial cells subjected to EMT, lose their intercellular connections and polarity, and experience changes in cell morphology and increased migration capacity. Tumor cells grow into the surrounding interstitial space by infiltration, and are in close contact with local capillaries and lymphatic endothelial cells. Once cells have invaded this interstitial space, they subsequently penetrate the walls of the vessel and protrude into the lumen. This enables transport to the target tissue, where the cells then proliferate in the matrix to form a new secondary tumor (63,64).

CXCR2 is involved in the migration, invasion and metastasis of breast cancer cells in various ways (61). A number of studies have investigated the effects of CXCL8 on cancer invasion and migration (65-68). High levels of CXCL8 expression promote angiogenesis and attract neutrophils to release enzymes involved in tissue remodeling and tumor formation. Thus, ectopic expression of CXCL8, stimulated by IL-8 and TNF-α, could exacerbate the metastatic potential of breast cancer (65). A CpG island located upstream of the CXCL8 promoter in the highly metastatic cell lines MDA-231 and MDA-345, results in the upregulation of CXCL8 expression (66). However, the loss of CXCL8 production in MDA-MB-231 and BT549 cells consequently attenuates migration and invasion, which may be due to decreased integrin β3 transcription (40). COX-2-mediated CXCL8 production in ER-negative breast cancer cells promotes osteoclast formation and bone metastasis (67). Compared with patients with no metastasis, those with breast cancer and bone metastases demonstrate increased CXCL8 levels. Furthermore, there is a significant positive correlation between plasma CXCL8 levels and bone resorption (68). In addition, CXCL8 regulates the actin cytoskeleton through Ca2+-activated PLC-dependent PKC and Rho-GTPase (69). Additionally, CXCL8 also promotes cell migration through the stimulation of the intracellular Akt pathway (27). Triple-negative breast cancer (TNBC) cells secrete CXCL8, which activates CXCR2 in tumor-associated fibroblasts and tumor-associated macrophages. This results in STAT3 phosphorylation and upregulation of CXCL8 transcription and translation. The binding of CXCL8 to CXCR2 in cancer cells facilitates metastasis (70), which was also observed in MDA-MB-231 cells and xenograft mouse models with CXCR2-knockout, as well as decreased migration, compared with wild-type cells (70). These data indicate that the CXCR2-CXCL8 axis is multifaceted in tumor progression and metastasis, and renders cancer cells invasive.

Bone marrow-derived mesenchymal stem cells (MSCs) express CXCL1 and CXCL5 to recruit PyMT breast cancer cells and prompt the migration in a CXCR2-dependent manner in vitro. The CXCR2-inhibitory antibody, SB265610, substantially curbs the migration of cancer cells to MSC-conditioned media (71). Additionally, it was proposed that TNF-α-activated MSCs secrete CXCR2 ligands to recruit CXCR2+ neutrophils to tumor sites (72). Tumor-associated neutrophils stimulate metastasis through matrix metalloproteinases and other soluble molecules. A co-culture system, consisting of tumor cells and neutrophils, substantiated to higher expression levels of metastasis-associated genes in tumor cells, which identified an MSCs/neutrophil/tumor cell axis associated with cancer metastasis (72).

The silencing of CXCR2 in breast cancer cell lines using short hairpin RNA results in attenuated cell invasion. Moreover, when these shRNA-treated cells were transplanted into an orthotopic mouse xenograft model, spontaneous lung metastases were decreased by 40% compared with the control group (61). In addition, the loss of CXCR2 expression in stroma cells (neutrophils, macrophages and endothelial cells) in the tumor microenvironment also prevented cancer cell migration (47). It was hypothesized that CXCR2 is involved in the paracrine loop between tumor cells and its surroundings, which potentiates invasion and metastasis.

**CXCR2 and drug resistance in breast cancer.** Drug resistance is one of the challenges of advanced breast cancer therapy.
Tumors gradually become less responsive to chemoradiotherapy during treatment and therefore, the mortality rate of breast cancer remains high. As a consequence of drug resistance, there are few recognized therapeutic strategies (73,74).

A wealth of evidence has shown that malignant cells that survive primary chemoradiotherapy express higher levels of CXCR2 ligands (65,75). Additionally, high levels of CXCL1, CXCL3, CXCL5, CXCL6, CXCL7 and CXCL8 are observed in drug-resistant breast cancer cells, which diminishes the effectiveness of other medical interventions (7.76). Shi et al (76) revealed that the multidrug-resistant human breast cancer cell line, MCF-7/R, has higher expression levels of CXCL6 and CXCL8 compared with the sensitive control cell line MCF-7/S. The inhibition of CXCL6 and CXCL8 with antibodies enhances the sensitivity to paclitaxel and doxorubicin treatment in MCF-7/R cells. The inhibition of CXCL6 and CXCL8 expression reverses the chemoresistance of MCF-7/S cells to doxorubicin (76). Similarly, other studies demonstrated that the inhibition of CXCL8 in MDA-MB-231 and BT549 cells improved the efficacy of chemotherapy (40). The inhibition of IL-8 or CXCR2 was shown to prevent the paclitaxel-induced autocrine inflammatory feed-forward loop (77). Sharma et al (75) used Cl66-wt, 4T1-wt, Cl66sh-CXCR2 and 4T1sh-CXCR2 cells that expressed varying levels of CXCR2, to assess the effect of CXCR2 levels on chemosensitivity. Furthermore, it was reported that the silencing of CXCR2 could increase the cytotoxicity of paclitaxel and doxorubicin, and intensify the antitumor activity of these drugs (75).

Regarding the intracellular mechanism of CXCR2-mediated drug resistance, previous studies suggested that AKT regulates chemical resistance through a broad anti-apoptotic molecule, such as PED (78-80). Patients with breast cancer and Akt phosphorylation at serine 473 are more sensitive to paclitaxel treatment (81). The overexpression of COX2 results in chemical resistance to breast cancer by generating prostaglandin H2 and activating NF-κB (82), which was also observed in a study by Xu et al (39). These results suggest that CXCR2-dependent regulatory pathways are crucial for cancer cells to maintain chemoresistance, and provide evidence that supports the targeting of the CXCR2 axis as an adjunctive therapy to prevent drug resistance.

**CXCR2 and breast cancer stem cells.** Cancer stem cells (CSCs) were first identified in breast cancer (83), and the presence of CSCs may be responsible for the high level of heterogeneity observed in this disease (84). Breast cancer stem cells (BCSCs) are resistant to the effects of chemoradiotherapy and endocrine therapy due to their self-renewal and differentiation potential. This allows BCSCs to initiate and maintain tumor growth, invasion, metastasis, drug resistance and recurrence (85). The importance of CSCs was demonstrated in several cancer types (86). Trastuzumab prolongs survival in patients with HER2-positive breast cancer treatment, which is in part due to its ability to limit CSCs (87). Neoadjuvant lapatinib was shown to suppress BCSCs in HER2-positive tumors and its combination with lapatinib prolongs the time to progression in patients with trastuzumab resistance (88). CXCL8 transmits signals through CXCR2, which induces EMT (89). EMT is a process that regulates invasion and metastasis, and enables cells to obtain stem cell characteristics (36). Singh et al (90) demonstrated a positive correlation between the level of CXCL8 in metastatic pleural and peritoneal effusion and the ability of these tumor cells to form mammospheres in vitro. The inhibition of CXCR2 using a small molecule CXCR1/2 antagonist, SCH563705, abolished the impact of CXCL8 expression. This was quantified by enzyme-linked immunosorbent assay (90). The suppression of CXCR2 expression limits tumor spheroid formation and aldelfluor-positive rate in breast cancer cells, and increases the efficacy of anti-HER2 therapy in HER2-positive patients (91). Furthermore, CXCR2 regulates the activity of CSCs through both HER2-dependent and HER2-independent pathways. One such pathway is through FAK, which is also associated with BCSC maintenance (92). In vivo studies have shown that MSCs are recruited to the tumor site, co-localized with CSCs in the microenvironment and secreted cytokines. This increases the number of CSCs present in the tumor, thereby accelerating tumor growth (93). Liu et al (93) suggested that this effect is initiated by CXCL6 secreted by cancer cells, and is maintained by CXCL6 ligands (GCP-2 and NAP-2) that are secreted by cancer cells and stromal cells. In addition, cancer cell-derived CXCL1 induces the expression of CXCL8 and GRO-α in MSCs, which promotes the formation and maintenance of CSCs (94). The inhibition of CXCR2 or downstream pathways can decrease the number and activity of CSCs in vitro and in xenografts, and increase the efficacy of docetaxel to decrease tumor volume (95). Recently, studies have identified CXCR2 as a novel BCSC biomarker of only TNBC. CXCR2 was observed to be co-expressed with CSC-associated proteins, such as NANOG and SOX2 (96,97). Furthermore, 4T1 cells that express CXCR2 have characteristics of CSCs, including a low proportion of CXCR2-positive cells (~1%), hypoxia, elevated expression of CSC-associated mRNA, increased tumor spherule formation, tumor xenografts, resistance to radiotherapy and chemotherapy (96).

In addition to CXCR2, CXCR1 was also found to be present on the surface of BCSCs, which facilitates the growth of BCSCs stimulated by tissue damage or inflammation (98). CXCR1 is also known as IL-8 receptor A, and is an active receptor selectively expressed by BCSCs (99). CXCR1 belongs to the same chemokine XCC receptor family as CXCR2, and CXCR1 has 75% sequence similarity with CXCR2 (100). It was proposed that CXCR1 has a prominent role in both the initiation and therapy of breast cancer (101). The inhibition of CXCR1 results in the depletion of BCSCs in vitro (102). CXCR1 inhibition combined with chemotherapy, results in the release of CXCL8 by dying bulk (non-CSC) tumor cells and binds with CXCR1 on the surface of BCSCs. As a result, the BCSCs are protected from apoptotic signals triggered by the Fas ligand (102). Several CXCR2 inhibitors, such as Reparixin, also block CXCR1. This further decreases the enrichment of BCSCs and prevents tumor recurrence (77,79).

The aforementioned results uncover the integral role played by CXCR2 signaling in the complex inflammatory cytokine response. Furthermore, the role of CXCR1 in maintaining the activity of BCSCs via autocrine or paracrine pathways, and its
effects on treatment efficacy and disease prognosis have been uncovered (87).

5. Preclinical and clinical evidence of targeting CXCR2 in breast cancer

Targeted therapy offers a unique opportunity to suppress the activity of key genes associated with tumorigenesis (103). CXCR2 and its ligands are influential targets involved in tumor growth and regulation (Fig. 2) (19). Additionally, preclinical studies on other malignant neoplasms have revealed that targeting CXCL8 alone, or in combination with other regimens, may provide an effective, novel option for cancer management (104-107).

Although CXCL8 is the most commonly studied ligand of CXCR2 in breast cancer, other CXCR2 receptor agonists, such as GRO-α, GRO-β, GRO-γ and CXCL5, are co-regulated with CXCL8. This co-regulation of CXCR2 ligands may limit the effectiveness of targeting CXCL8 alone (108), which could be circumvented by suppressing CXCR2 and its downstream signaling pathways. Blocking the function of CXCR2, by impeding ligand binding or subsequent pathway activation, essentially prevents the biological effects of multiple ELR+ CXC chemokines with a single treatment. This intervention may be more tumor-specific than the direct inhibition of cytokines, and could decrease the severity of adverse events.

There are various orally active small-molecules that are non-competitive antagonists of CXCR2, including Repertaxin, SCH479833, SCH527123 and SCH563705. These antagonists exhibit anti-tumor properties in breast cancer (95), colorectal cancer (105) and melanoma (106) xenograft models. These properties include the obstruction of spontaneous hepatic metastasis of colon carcinomas (109). Moreover, the combination of paclitaxel treatment and the inhibition of CXCR2 slows breast tumor growth, attenuates tumor angiogenesis and lung metastasis, and diminishes the number of CSCs. Additionally,
targeting CXCR2 reduces drug resistance (75,110). Since CXCL8 is a prominent regulator of BCSC activity, a combined treatment of CXCR2 inhibitors and current HER2-targeted therapies is predicted to be an effective regimen to decrease CSC activity and increase survival in HER2-positive patients with breast cancer (90). A recent study in pancreatic ductal adenocarcinomas demonstrated that inhibition of CXCR2 predominantly in neutrophils/MDCs confers T cell entry to the tumor site (111). The inhibition of CXCR2 also enhances the sensitivity to anti-programmed death 1 therapy, which reinforces the endogenous anti-tumor immune response (111). Moreover, Uddin et al (98) revealed that the inhibition of proteasome significantly upregulates the levels of CXCL8 and its receptors CXCR1/2, through IKK. The inhibition of the induced CXCL8 expression or IKK activity may improve the effectiveness of proteasome inhibitors as a therapy for TNBC. Overall, CXCR2 and its ligands are potential therapeutic targets.

Due to the various functions of CXCL8 in homeostatic processes, the side effects of CXCR2 inhibitors cannot be ignored. Neutrophils are part of the immune surveillance system that monitors and destroys transformed cells, and CXCL8 exerts an anti-tumor effect by recruiting concentrated granulocytes (112). Therefore, the anti-tumor effects of CXCR2 inhibitors that influence neutrophil infiltration may unintentionally promote tumor growth. The solution to this dilemma depends on the new technologies of targeted drug therapies for cancer cells. In addition, CXCR2 inhibitors cause a marked decrease in the number of circulating neutrophils combined with chemotherapeutic agents, and this could cause a potentially synergistic increase in myelotoxicity (113).

Currently, CXCR2 antagonists are used in phase II clinical trials for chronic obstructive pulmonary disease (114), and obvious adverse reactions were reported in patients with severe asthma (113). Repertaxin was originally used to prevent CXCL8-induced reperfusion injury, and was the only clinically-tested CXCR2 inhibitor (21). Interfering with CXCR2 is particularly advantageous in breast cancer compared with other tumors. Phase I clinical trials have shown that Repertaxin, in combination with paclitaxel, is safe and well tolerated (99). Phase II studies on this combination are currently ongoing (99). The principle data on the efficacy and safety lay an essential foundation for the accelerated development of CXCR2 inhibitors as a treatment for breast cancer.

6. Conclusions and outlooks

Currently, controversies still exist in the field of CXCR2 research. One controversy is the association between CXCR2 and the expression of ER. Other controversies include the characterization of CXCR2 in cancer cell senescence, and its role in Akt-associated tumor metastasis (39,115-117). These controversies may be attributed to the limitations of the pathological tissue, cell type and animal model used.

The association between inflammation and tumor development is widely recognized, and the expression of many inflammatory cytokines in the tumor microenvironment has previously been reported (19). The poor efficacy of current breast cancer treatments may be associated with the molecular typing, and the function of CXCR2 in breast cancer. The anti-tumor effect of targeting CXCR2 was reflected in both in vitro and in vivo models of breast cancer, and further supports the development of CXCR2 inhibitors as a clinical therapy in breast cancer. Further exploration of whether CXCR2 inhibition plays a vital role in targeted therapies for breast cancer, based on molecular typing, is warranted.

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FG, LL and YW were significant contributors to the conception and design of this review article. LL, JW and YL were involved in the literature search. JW, YL and FG analyzed and summarized critically the existing knowledge obtained from the literature. FG and LL drafted the manuscript text. LW, FL, LL and FG designed and generated the figures. YW, LW and FL made substantial contributions to the manuscript revision regarding important intellectual content. All authors have read and approved the final manuscript.

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Patient consent for publication

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Competing interests

The authors declare that they have no competing interests.

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