The molecular mechanism of chronic stress affecting the occurrence and development of breast cancer and potential drug therapy

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ABSTRACT
According to the 2020 data released by the International Agency for Research on Cancer, breast cancer has surpassed lung cancer as the world’s most newly diagnosed first-time cancer. Compared with patients with other types of cancer, those with breast cancer experience greater mental stress and more severe psychological impacts because of the life-threatening diagnosis, physical changes, treatment side effects, and family and social life dysfunctions. These usually manifest as anxiety, depression, nervousness, and insomnia, all of which elicit stress responses. Particularly under chronic stress, the continuous release of neurotransmitters from the neuroendocrine system can have a highly profound impact on the occurrence and prognosis of breast cancer. However, because of the complex mechanisms underlying chronic stress and the variability in individual tolerance, evidence of the role of chronic stress in the occurrence and evolution of breast cancer remains unclear. This article reviewed previous research on the correlation between chronic stress and the occurrence and development of breast cancer, particularly the molecular mechanism through which chronic stress promotes breast cancer via a variety of neurotransmitters and neurotransmitter systems.

Abbreviations: ACH, acetylcholine; ADR-α1, α1 adrenergic receptors; ADR-α2, α2 adrenergic receptors; AC, adenylate cyclase; α7-nAChr, α7 nicotinic acetylcholine receptors; BCScs, breast cancer stem cells; BDNF, brain-derived neurotrophic factor; β-AR/ADR-β, β-adrenergic receptor; β-END, β-endorphin; CCK, cholecystokinin; CNS, central nervous system; CCL2, CC chemokine ligand; CCR2, CC chemokine receptor; CUS, chronic unpredictable stress; CTCs, circulating tumor cells; CNS, chronic stress response; CSC, cancer stem cells; CBM, cognitive-behavioral stress management; CTNNB1, catenin β1; cAMP, cyclic adenosine monophosphate; DA, dopamine; DR, dopamine receptors; DAG, diacylglycerol; EPAC, exchange protein directly activated by cAMP; EGR1, early growth response gene-1; EGFR, epidermal growth factor receptor; ERK1/2, extracellular signal-regulated kinase 1/2; EMT, epithelial–mesenchymal transformation; E, epinephrine; FAK, focal adhesion kinase; GPCR, G-protein-coupled receptor; GABAB1e, GABA-B1e receptor; GAD, glutamic acid decarboxylase; GABA, α-aminobutyric acid; GC, glucocorticoids; HPA, hypothalamic–pituitary–adrenal axis; HSP, heat shock proteins; HA, histamine; IRS, insulin receptor substrate; IGFR-1, insulin-like growth factor-1 receptor; IP3, inositol triphosphate; JNK, c-Jun NH2-terminal kinase; LDHA, lactate dehydrogenase A; mACHRs, muscarinic acetylcholine receptors; MAPK, mitogen-activated protein kinase; MRI, medical and psychological intervention; mTOR, mammalian target of rapamycin; mGLUT, glutamic acid; NTS, neurotensin; NTs, neurotensin receptor; nAChR, nicotinic acetylcholine receptor; NE, norepinephrine; NPY, neuropeptide Y; PRL, lactotroph; PKC, protein kinase C; PRL, prolactin; PKA, protein kinase A; PLC, phospholipase C; PROP, propranolol; FOUSFI, FOU domain class 5 transcription factor 1; PMN, premetastatic niche; SSADH, succinate semialdehyde dehydrogenase; SNS, sympathetic nervous system; TrkB, tyrosine kinase receptor B; TPH1, tryptophan hydroxylase 1; VP, vasopressin; VEGF, vascular endothelial growth factor; hT, 5-hydroxytryptamine.

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Introduction

According to the 2020 data released by the International Agency for Research on Cancer of the World Health Organization, the incidence of new diagnoses of breast cancer has surpassed that of lung cancer for the first time. The prevention and treatment of breast cancer remains a critical research topic across the world. Presently, breast cancer is mostly regarded as a physical disease by modern medicine and its neuropsychological factors are ignored. Since the biopsychosocial medicine model was proposed by the American scholar Engel in 1977, emotions have been considered as another vital sign along with body temperature, pulse, breathing, and blood pressure. Compared with patients with other types of cancers, those with breast cancer have a significantly higher incidence of depression due to the side effects of radiotherapy and chemotherapy, mastectomy, menopause, and other factors. In such emotionally loaded microenvironment, these factors may even induce breast cancer recurrence and metastasis, thus directly affecting the prognosis and survival rate of patients with breast cancer.

An increasingly larger number of problems arise with therapies that target breast cancer cells; therefore, future research may shift from a single-purpose biomedical model to a more socially integrated model of medicine.

With the change in medical strategy, people have gradually realized that the development of breast cancer is not only related to physiological factors but also closely related to the stress caused by personal neuropsychological factors and social environment. Stress responses are a series of nonspecific adaptive responses that are produced when the body senses that the homeostatic balance is threatened or altered. Based on the duration and intensity of the stress, it can be classified as acute and chronic stress. Acute stress has a positive effect on the body. Conversely, chronic stress indicates that the body has been in a state of “overdrive” for a long period of time, which can cause serious complications and even promote tumor occurrence and metastasis. Common stressors are categorized as physical (various physical, chemical, and biological stimuli), psychological (conflict, frustration, hatred, fear, etc.), and social (professional competition, work burden, etc.) [10,71].

Research on the relationship between psychosocial factors and breast cancer has been increasing. In this review, we discuss the biological pathways and mechanisms of chronic psychological stress in breast cancer. We explore the effect of the continuous secretion of certain neurotransmitters on the occurrence and development of breast cancer, and we summarize the potential drugs or blockers that influence the neuroendocrine system for the treatment of breast cancer, further clarify the mechanism of neurotransmitter action, and suggest novel strategies to develop drugs against breast cancer.

Relationship between chronic stress, the neuroendocrine system, and breast cancer

In today’s fast-paced society, an individual experiences a high degree of personal stress, which usually manifests as anxiety, depression, tension, and insomnia. In addition to these, patients with breast cancer face changes in their body image, side effects of treatment, and family and social factors that lead to mental stress and psychological distress, all of which lead to chronic psychological stress [88]. Currently, the role of chronic psychological stress in tumor progression is one of the prominent topics in oncology research [76]. Chronic stress refers to the continuous, nonspecific, adaptive response of the body caused by the activation of the classical hypothalamic–pituitary–adrenal (HPA) axis [160]. Chronic psychological stress has been confirmed to be involved in various physiological and pathological processes of the human body and has been shown to promote disease progression by weakening the mental or physical health of the patient. In particular, the evolution of malignant tumors and the remodeling of the tumor microenvironment have severe adverse effects [10].

Increasing evidence has shown that chronic psychological stress can induce the occurrence and prognosis of breast cancer through the neuroendocrine system [126]. The ancient Greek physician Galen pointed out that psychology and physiology are connected and that organic diseases of the body can be cured by psychotherapy [141]. Nerves mainly transmit information from the central nervous system (CNS) to organs and tissues. Sensory nerves, motor nerves, and autonomic nerves (sympathetic and parasympathetic) form a neural network directly or indirectly connected to the neuroendocrine system, thus enabling functional communication between various parts of the body [56]. Other physical stressors, such as surgical stress and cold stress, have also been shown to promote the growth of breast cancer cells [16, 64]. Regardless of the stress source, the nonspecific response of the body is mainly produced by the HPA axis and sympathetic nervous system (SNS). The HPA axis primarily mediates the secretion of glucocorticooids (GCs), whereas SNS mainly mediates the secretion of catecholamines such as epinephrine (E) and norepinephrine (NE) [22]. SNS rapidly improves the body’s alertness and resilience during stress, which enables the body to cope with stress [34]; however, the continuous activation of the SNS under chronic stress significantly elevates the expression levels of GC, E, NE, and other neurotransmitters in the tumor microenvironment, thereby promoting the cell growth and metastasis of breast cancer [75] (Fig. 1).

Neurotransmitters are chemical substances that transmit messages among neurons or between neurons and effectors. The role of neurotransmitters in breast cancer is complex and varied. Based on their chemical structures, neurotransmitters can be classified as biogenic amines, amino acids, peptides, etc. Biogenic amine neurotransmitters include dopamine (DA), NE, and serotonin (5-HT, also known as 5-hydroxytryptamine). Amino acid neurotransmitters include γ-amino butyric acid (GABA), glycine, glutamic acid (mGluR), histamine (HA), and acetycholine (ACH). Peptide neurotransmitters include endogenous opioid peptide, neurovasopressin, cholecystokinin, and neuro peptide Y (NPY). This paper summarized the mechanisms regulating the development of breast cancer using the abovementioned classification framework. Obtaining mechanistic insights into the differential biological responses of different neurotransmitters under chronic stress allows us to understand into how drugs can be used to block the neuroendocrine system in breast cancer treatment.

Biogenic amine neurotransmitters regulate the pathological mechanism of breast cancer

α-epinephrine and breast cancer

Surgery, environment, and psychosocial stress factors activate a series of signal transduction pathways in the human body, triggering the stress response in the autonomic nervous system. Adrenergic receptors are the principal SNS mediators that regulate organ function, maintain systemic homeostasis in a resting state, and trigger the body’s “fight or flight” response under stress. Following the activation and excitation of adrenergic receptors, the sympathetic nerves release E and NE that initiate various intracellular signaling cascades [2, 29]. There is increasing evidence demonstrating that α-adrenergic receptors play an important role in breast cancer cell proliferation, migration, and invasion [145, 153] (Fig. 2A). Despite evidence that the application of adrenergic receptor antagonists can slow down cancer progression and
reduce patient mortality, there are still some controversies about whether they are absolutely beneficial for patients with cancer. For example, people pay more attention to psychological stress and use the restraint stress model or the chronic unpredictable stress model to study how changes in environmental temperature affect tumor progression in ways other than the through the immune system. As another example, the studies mentioned below cannot rule out the influence of other hormones involved in the complex neuroendocrine response under stress on tumor growth. However, it is certain that adrenaline signals are the key effector mediators that play a role in stress and tumors driving.

**Classification and biological function of receptors**

ADR-α is coupled to Gq protein through the phospholipase C/inositol triphosphate/Ca^{2+} pathway, and the diacylglycerol/protein kinase C (PKC) pathway plays a role. ADR-α2 coupled with Gi protein inhibits adenylate cyclase (AC) and decreases the intracellular levels of cyclic adenosine monophosphate (cAMP) [51, 101]. ADR-α is found in arteries and capillaries and helps regulate blood supply to many organs, including the breasts.

**Promotion of proliferation**

ADR-α2 is a key regulator of human breast cancer cell proliferation. Some researchers have studied its mechanism, mainly involving the mitogen-activated protein kinase (MAPK) signaling pathway. In the breast cancer cell lines MCF-7 and T47D, activation of ADR-α2 increases prolactin (PRL) levels and activates STAT5, extracellular signal-regulated kinase (ERK1/2), and Akt [152]. ERK1/2 is one of the key signaling proteins of the Ras/Raf/MAPK signaling pathway, and it participates in various functions, including the regulation of carcinogenesis and the progression of tumor cells. In addition, PRL can activate ERK1/2 and Akt through Src, and ADR-α2 can stimulate ERK phosphorylation through Src-independent pathways [1].

**β-epinephrine and breast cancer**

Chronic stress can activate β-adrenergic receptor (ADR-β), which plays an important role in growth, invasion, metastasis, angiogenesis, immunosuppression, etc. [39]. It can also reduce the therapeutic effect of anticancer drugs, which negatively affects the prognosis of patients with breast cancer. Evidence has shown that the application of ADR-β antagonists can slow down the progression of breast cancer, reduce the mortality rates of patients, and improve their prognosis. Such antagonists are expected to become biomarkers for diagnosis, monitoring the treatment effect, and determining the prognosis of breast cancer [107]. Ultimately, there exists a link between stress, ADR-β, and breast cancer. The following section focuses on the key mediators involved in ADR-β signaling implicated in the stress response and the progression of breast cancer.

**Classification and biological function of receptors**

ADR-β is distributed in the liver, kidney, lung, brain, breast, lymphatic tissue, and vasculature [62]. β1, β2, and β3 receptors bind to Gs proteins and, when stimulated by adrenaline, activate AC to produce cAMP, a second messenger. cAMP production reactivates its two main downstream target molecules: protein kinase A (PKA) or exchange protein directly activated by cAMP (EPAC) [72]. The production and activation of cAMP and its downstream effectors lead to an array of signaling pathways that affect the biological response as indicated in Fig. 2B. ADR-β signaling can also regulate the biological activities of fibroblasts, epithelial cells, and vascular endothelial cells, which are related to cancer cells, especially lymphatic and immune cells [75, 84].

**Induced stem cell-like properties**

Compared with common breast cancer cells, breast cancer stem cells show enhanced tumorigenesis, invasion, and migration abilities, which is considered essential for the proliferation, invasion, and other malignant biological characteristics of breast cancer cells. Cui et al. used NOD/SCID mice to establish a stress model and found that ADR-p2 promotes glucose metabolism balance in breast cancer cells, shifting it toward glycolysis, and also promotes the expression of lactate dehydrogenase A under chronic stress. The product of this is lactic acid that promotes glucose metabolism balance in breast cancer cells, shifting it toward glycolysis, and also promotes the expression of lactate dehydrogenase A under chronic stress.

**Chemical resistance**

Research has shown that ADR-α can affect chemotherapy drugs used to treat breast cancer. Su et al. [139] demonstrated that the activation of α-adrenergic signaling leads to the upregulation of MDR1. Increased MDR1 expression increases the resistance of MCF-7 cells to paclitaxel. When MCF-7 cells were implanted subcutaneously in severe combined immunodeficiency (SCID) mice, the stress condition led to the upregulation of MDR1 expression.
Fig. 2. Biogenic amine neurotransmitters regulate the pathological mechanism of breast cancer

A. ADR-α1 is coupled to Gq protein and exerts physiological effects via the PLC/IP3/Ca\(^{2+}\) and DAG/PKC pathways. ADR-α2 coupled with Gi protein inhibits AC expression and decreases the intracellular cAMP level. ADR-α2 stimulates ERK phosphorylation via the Ras/Raf/MAPK signaling pathway and Src-independent pathway to promote breast cancer cell proliferation.

B. Upon stimulation by adrenaline, ADR-β and Gs protein conjugate to activate AC, resulting in the production of the second messenger cAMP, which then activates its two downstream target molecules PKA or EPAC, leading to biological effects.

C. Under chronic stress, ADR-β2 promotes glucose homeostasis in breast cancer cells to shift toward glycolysis and promotes LDHA expression. The resultant weakly acidic microenvironment enhances the deubiquitination of MYC by USP28, improves the stability of MYC, and activates the promoter of Slug, ultimately promoting the stem cell-like properties of breast cancer.

D. The increased expression of CCL2 and CCR2 in chronically stressed mice results in the accumulation of macrophages in the lungs, which ultimately leads to a significant increase in the metastasis and colonization of circulating breast cancer cells in the lungs.

E. Trastuzumab resistance-dependent PI3K/Akt/mTOR pathway is controlled by catecholamine-induced ADR-β2 activation.
4T1 cells to construct a mouse chronic stress model of breast cancer and found that after its activation, ADR-β2 regulated hexokinase-2 expression in cancer cells and promoted glucose metabolism in tumors [73]. In addition, ADR-β2 stimulation in osteoblasts promoted the adhesion of breast cancer cells to bone marrow endothelial cells in an H-1-β- and selectin-dependent manner [35].

Promotion of migration and invasion
Changes in neuroendocrine responses in the body during chronic stress can result in a “premetastatic microenvironment” in the distal organs during the premetastatic tumor stage, making it more conducive to the survival of tumor cells. In other words, the premetastatic niche (PMN) is another important factor in the promotion of tumor metastasis. ADR-β overexpression in human breast cancer cells induces an invasive phenotype, whereas significantly reduced ADR-β expression reduces the effect of stress on metastasis in vivo [119]. The final site that tumor cells colonize and whether the tumor can survive in distant organs, far away from the primary focus, is affected by the microenvironment of the distant organs. The formation of PMN and survival of metastatic cells are closely related to the recruitment of macrophages in PMN. Chen et al. [28] used the ADR-β agonist isoproterenol in a mouse model of chronic unpredictable stress response before metastasis and showed that the number of monocytes in the peripheral blood, spleen, and bone marrow was significantly increased after isoproterenol administration. Furthermore, the expressions of CC chemokine ligand 2 and monocyte/macrophage chemokine receptor 2 in lung tissues were upregulated, leading to the accumulation of macrophages in the lung. The result was a significant increase in the metastatic colonization of tumor cells in the lungs of the breast cancer mouse model. In addition, Le et al. [78] found that chronic stress can regulate the COX2-PEG2 inflammatory pathway related to tumor-associated macrophages through ADR-β signal transduction of inflammatory cells, increase the expression of vascular endothelial growth factor (VEGF) and the density of lymphatic vessels in the primary tumor of breast cancer, induce stable expansion of lymphatic vessels draining the primary tumor, and increase lymph flow, thereby enhancing the ability of tumor cells to migrate and spread through the lymphatic vessels (Fig. 2D).

Chemical resistance
In addition to promoting the proliferation, migration, and invasion of breast cancer cells, ADR-β2 reverses chemical resistance to the breast cancer drug Herceptin (trastuzumab). The therapeutic antibody Herceptin is currently a first-line drug for the clinical treatment of breast cancer; it can specifically target breast cancer cells overexpressing HER2. However, drug resistance has become a critical problem, restricting the clinical efficacy of the drug. ADR-β2 and HER2 constitute a positive feedback loop in human breast cancer cells, affecting the biological behavior of breast cancer cells, suggesting that ADR-β2 activation is involved in trastuzumab resistance [83]. Further studies have found that in patients with breast cancer with HER2 overexpression, ADR-β2 expression is negatively correlated with the response to trastuzumab. ADR-β2 upregulates the expression of miR-21 and muc1 by activating STAT3 and the PI3K/Akt pathway. It also inhibits miR-199a/b-3p expression and induces the activation of mammalian target of rapamycin (mTOR). Therefore, the trastuzumab resistance-dependent PI3K/Akt/mTOR pathway is controlled by catecholamine-induced β2-AR activation (Fig. 2E). Furthermore, the ADR-β2 antagonist propranolol (PROP) not only enhances the antitumor activities of trastuzumab but also resensitizes resistant cells to trastuzumab. The next research direction needs to accurately assess the expression of ADR-β subtypes in tumor cells of patients with breast cancer and selectively use β-receptor blockers. Only prospective, multicenter clinical studies could provide reliable evidence for β-blockers to become a new adjuvant treatment for patients with breast cancer.

Dopamine and breast cancer
DA is a catecholamine neurotransmitter present in the mammalian neuroendocrine system [114]. Chronic stress causes the overexpression of DRs. The DA imbalance theory, which explains the mechanism of stress, is widely accepted [89]. Under chronic stress, rats continue to be in a state of depression; DA levels in the prefrontal lobe, striatum, hippocampus, and other brain regions are decreased; and the synaptic transmission function of dopaminergic neurons is significantly downregulated [93]. Imaging studies in patients with severe depression showed abnormal DR1 binding [23]. Long-term stress stimulation has been reported to significantly increase DR1 levels in the rat limbic system by 29%, which decreases after antidepressant treatment [102]. Selective DR1 agonists can reverse the decrease in DA levels caused by chronic unpredictable stress and can ameliorate behavioral despair in mice [42,111]. Abnormal function of the DA system can cause diseases of the central nervous system, such as schizophrenia and Parkinson’s disease. Numerous recent studies have found that the DR signaling pathway is closely related to the occurrence, development, and prognosis of breast cancer. DRs may become new targets for the targeted therapy of tumors. For example, the DR antagonist methiliprazine has been explored in basic research and clinical practice of pituitary tumors [9]. DA agonists such as bromocriptine and carergeline are first-line agents for the treatment of prolactinoma, and some researchers are currently researching the application of DA agonists for the treatment of breast cancer [37].

Overexpression of DRs can induce breast cancer. In 2003, Comings et al. concluded that the D2 receptor gene is a risk factor for breast cancer (P = 0.018) in a study on the genetic risk factors of breast cancer. Studies have found that individuals with chronic stress are susceptible to several cancers, including breast cancer. Interestingly, the expression levels of DRD2–DRD4 in patients with breast cancer are elevated [108]. The expression levels of DRD2–DRD4 in patients with breast cancer without psychological intervention were significantly higher than those in patients who received the intervention [3]. Bakhhtou et al. [12] found a difference in gene expression between patients with breast cancer and healthy people and found that DRD2 and DRD4 were overexpressed in peripheral blood mononuclear cells of patients at stage II and III breast cancer. At the same time, cellular experiments revealed that DRD2 agonists could induce apoptosis of breast cancer cells. Sachols et al. [117] found that pluripotent stem cells in healthy subjects did not express DRs, whereas CSCs expressed all five DRs, suggesting that DRs are a potential biomarker for screening CSCs and a potential target for triple-negative breast cancer therapy. However, many questions about DRs and tumors need to be further studied, especially the specific molecular mechanism of the antitumor effect of compounds through the DR signaling pathway. The important role of DR agonists and antagonists in tumors has been receiving increased attention from researchers. However, the role of DRs in food intake and glucose metabolism is still unclear. Future research can focus on tumor metabolism and discover the role of dopamine in other tumors. Based on current research results, it is expected to play a clinical guiding role.

Noradrenaline and breast cancer
Evidence suggests that the tumor-driven effect of β-adrenergic signaling is mainly dependent on local sympathetic ending-derived NE [36]. When SNS is activated by chronic stress, adrenaline and NE are first secreted by the adrenal medulla and distributed through the bloodstream to the tumor site. Sympathetic endings entering tumor tissue can release NE, which interacts directly with tumor cells [143]. A study on breast cancer demonstrated that mice with restraint stress exhibit rapid tumor growth and increased tumor innervation. Antagonism of the nicotinic ACh receptor (NACHR) in the peripheral nervous system with hexamethonium bromide completely eliminated cell growth. However, removal of the adrenal gland had no significant effect
on tumor growth and nerve count increase induced by restraint stress. This result confirms the dominant role of NE derived from local nerve fibers [7]. The author also proposed a feedforward circuit innervated by tumors: under restraint stress, the brain source is upregulated by activating the β3-AR/cAMP/Epac/c-Jun NH2-terminal kinase (JNK) pathway. The expression of brain-derived neurotrophic factor increases tumor tissue innervation through the action of tyrosine kinase receptor B. This new growth of nerve endings can release NE to promote tumor growth, thereby forming a feedforward circuit. In addition, the depressive state increases the level of NE, increases the phosphorylation of p38MAPK, and increases the malignancy of breast cancer cells [119]. NE activates the Akt signaling pathway under anxiety and promotes the development of breast cancer [147].

Promotion of migration and invasion

NE released under chronic stress can promote the invasion and metastasis of breast cancer cells. Sloan et al. [125] constructed a stress model of mice using binding methods and found that chronic stress enhanced the metastasis of breast cancer cells to distant tissues (including lung and lymph nodes) but had little effect on the growth of the primary tumor. This stress-induced metastasis is mediated by promoting macrophage infiltration, and stress simultaneously induces the expression of premetastatic genes such as TGFβ, Arg1, and CSF1 in the primary tumor microenvironment. Stress also promotes the adhesion of tumor cells and vascular endothelial cells. Isoproterenol can induce integrin-mediated cell adhesion through the β2-AR/cAMP/Epac/Rap1 pathway [110]. Studies have shown that NE can induce the release of the chemokine GROs from vascular endothelium and promote β1-integrin-mediated adhesion of breast cancer cells to vascular endothelium [138]. Under chronic stress, NE can activate focal adhesion kinase through the ADRB2/Src signal axis, thereby inhibiting the anoikis of breast cancer cells [133]. As a result, a small portion of the surviving circulating tumor cells adheres to the vascular endothelial cells of the distant organs, escapes the circulatory system, and are transferred to specific remote organs. Yamazaki et al. [155] treated MDA-MB-231 breast cancer cells with NE and found that NE can induce the production of reactive oxygen species and promote the expression of heme oxygenase-1, MMP-2, and MMP-9 genes, thereby increasing the invasiveness of breast cancer cells.

Promotion of angiogenesis and lymphangiogenesis

NE can promote the formation of tumor blood vessels and lymph vessels. Chen et al. found that VEGF expression in breast cancer cells can be upregulated by NE and that the culture supernatant of breast cancer cells treated with NE promoted the formation of a capillary-like network of endothelial cells. In a study on cocultured endothelial and breast cancer cells, NE was found to activate the Notch pathway of endothelial cells and significantly upregulate the expression of Notch ligand Jagged-1 on breast cancer cells. The Notch pathway is closely related to the interaction of tumor cells and mesenchymal cells and angiogenesis. The upregulation of Jagged-1 caused by NE can be reversed by β2-AR, PKA, and mTOR inhibitors, indicating that β2-AR/PKA/mTOR pathway is involved in this process.

5-hydroxytryptamine and breast cancer

One of the causes of chronic stress is the decreased release of 5-HT in the CNS, which reduces the content of the synaptic cleft. 5-hydroxytryptamine (5-HT) was first isolated from serum by author Rapport, and is more commonly known as serotonin [130]. 5-HT is present in a large amount in the digestive system and a small amount in the CNS. It participates in the regulation of mood and cognition, as well as the regulation of the functions of the gastrointestinal tract, such as movement, secretion, and immunity [60,94]. Studies have confirmed that 5-HT levels in the CNS of patients with depression are reduced. The 5-HT level was also significantly reduced in a chronic restraint stress mouse model [46,118]. 5-HT6 receptor antagonists are closely related to the CNS and are also involved in the regulation of neurotransmitters, such as ACH, DA, and GABA, playing an important role in alleviating anxiety and depression [61]. Furthermore, the HT3 receptor is an important target for controlling digestive disorders such as anorexia and bulimia. Activation of the 5-HT3 receptor can increase intracellular Ca2+ levels, promote the release of neurotransmitters and neuropeptides, and stimulate central and peripheral neurons, leading to vomiting and inflammation. 5-HT3 receptor antagonists can effectively counter eating disorders and vomiting caused by early chemotherapy and radiotherapy [164]. Therefore, 5-HT3 receptor antagonists can be clinically developed to inhibit dopaminergic neurons and promote the increase of adrenocorticotropic hormone levels to relieve stress response.

In the mammalian glands, 5-HT produced by the prolactin-induced expression of the 5-HT biosynthesis gene tryptophan hydroxylase 1 induces lactation-to-involution switching via 5-HT7 receptor-dependent tight junction disruption [66]. Although 5-HT has an inhibitory effect on the growth of normal breast epithelial cells, it stimulates the proliferation of breast cancer cells at submicromolar concentrations. Although the mechanism by which 5-HT affects the malignant phenotype of breast cancer remains unclear, the angiogenic effect of 5-HT suggests that it stimulates the proliferation and invasion of cancer cells, which is necessary for cancer progression [55,57]. Additionally, in the triple-negative breast cancer cell line MDA-MB-231, 5-HT was shown to promote cell invasion and proliferation through the 5-HT7/7 receptor and had a stronger proliferation effect in the first stage of metastasis than that after local infiltration [50]. Depression often promotes the dysregulation of the immune system of patients with cancer and accelerates tumor recurrence and metastasis. Compared with the nondepressive group, patients with breast cancer with depression had higher 5-HT levels, higher 5-HT1 expression, and lower overall survival rates [85].

Amino acid neurotransmitters regulate the pathological mechanism of breast cancer

γ-aminobutyric acid and breast cancer

Inhibitory neurotransmitter GABA is involved in important life activities such as human growth and development and nitrogen balance. Recent studies have reported that GABA can regulate tumor cell proliferation, migration, differentiation, and apoptosis [113]. The decrease in GABA levels in the cerebrospinal fluid and plasma of patients with depressive disorder results in weakening of the inhibitory effect on central neuron activity, leading to symptoms such as anxiety and insomnia, and further acceleration of the occurrence of chronic stress [14,43].

GABA receptors can be categorized into ion channels and metabotropic receptors according to their sensitivity to agonists and antagonists. Among them, GABA_A and GABA_B receptors are ion channels, whereas GABA_A receptors are metabotropic, belonging to the G-protein-coupled receptor (GPCR) family [20]. GABA_A comprises two subunits, GABA_A1 and GABA_A2. The prominent feature of GABA_A1 is its molecular diversity; to date, 14 splicing isomers ranging from GABA_A1a to GABA_A1h have been identified. GABA_A1e (GB1e) was shown to promote the proliferation and migration of MCF-7 cells and protect cells from apoptosis caused by Tm and Tg that are stressed by the endoplasmic reticulum [68]. GB1e can also significantly promote the cloning and microsphere formation of MCF-7 cells and the expression of the breast cancer stem cell marker molecule CD44. Molecular experiments showed that GB1e may promote the enhancement of cancer stem cell characteristics by activating the MAPK and ERK1/2 signaling pathways [20]. This is the first exploration of the function of GABA_A1e in tumors and the first to establish a connection between GABA_A receptors and the tumor microenvironment and tumor stem cells.

ABAT, a key enzyme in GABA metabolism, catalyzes the degradation
of GABA into succinate hemialdehyde and 1-glutamic acid, thereby contributing to the reduction of GABA levels [67]. Based on an analysis of a breast cancer metabolomics database, we found that tumor tissues with low ABAT expression had high GABA content and that the promoter of ABAT contained four conservative Snail-binding site E-box sequences (CAGGTG). Accordingly, we hypothesized that due to the change in the copy number of the ABAT genome and Snail inhibition, the ABAT expression level in basal-like breast cancer was decreased, leading to an increase in the levels of its metabolic substrate GABA and activation of related signaling pathways through GABA, thereby promoting the occurrence and development of breast cancer. In addition, GABA-mediated activation of the Ca\(^{2+}\)-NFAT1 axis downregulated ABAT expression in basal-cell-like breast cancer, thereby promoting the occurrence and metastasis of breast cancer [31] (Fig. 3A). Gina et al. silenced GABRP in two BLBC cell lines, HCC1187 and HCC70, and found that BLBC-related cytoteratin expression decreased, such as the expressions of KRT75, KRT68, KRT14, and KRT17. Simultaneously, the silencing of GABRP expression reduced ERK1/2 phosphorylation in the two cell lines and ultimately inhibited tumorigenic potential and migration in vitro [127]. GABRA3 activated the Akt pathway to promote breast cancer cell migration, invasion, and metastasis [53].

Acetylcholine and breast cancer

Classification and biological function of receptors

Cholinergic receptors are divided into muscarinic ACh receptors and nAChRs, which mediate the production of the neurotransmitter ACH. In mammals, nAChRs are divided into neuronal and muscle nAChRs [30]. Neuronal nAChRs can be divided into α and β subunits (α2, α3, α4, and α6) and β subunits (β2, β3, and β4) can form heteropentamers, and α7 and α9 subunits can form homopentamers [140]. Obviously, the different structures of α7 and α9 lead to completely different pharmacological functions. But their structures are so similar that they are difficult to distinguish (Table 1). As a specific antagonist of neuronal nAChRs, α-conotoxin has a strong ability to distinguish between different α and β subunit interfaces of neuronal nAChR subtypes. Therefore, α-conotoxin is a valuable molecular probe for studying the fine structure and physiological and pharmacological functions of various nAChR subtypes, and it is also a source of new drugs for the treatment of nAChR-related diseases [79]. The nAChRs are closely associated with the occurrence and development of diseases, and the different nAChR subtypes determine the different pathological and physiological process (Fig. 3B).

Overexpression of α7 and α9-nAChR

Mounting evidence has shown that the overexpression of ACHRs mediates the proliferation, apoptosis, angiogenesis, and epithelial–mesenchymal transformation of breast cancer cells [103]. Members of the nAChR family, α7-nAChR and α9-nAChR, have been found to be expressed in both MCF-7 and MDA-MB-468 cells [30]. Aldehyde dehydrogenase (ALDH), which is responsible for deoxidation in cells, has been used as a specific marker of stem cells in recent years. It has been used to identify the enrichment of CSC subsets and the determination of poor prognosis in patients with breast cancer, which is mainly associated with increased metastasis and chemotherapy resistance [121]. Nicotine, an N-choline receptor agonist, can induce human MCF-7 breast cancer cells to develop resistance to the anticancer drug Adriamycin. Additionally, nicotine promotes the migration of breast cancer by activating PKC, and the number of ALDH-positive CSCs in MCF-7 cells is dependent on the concentration of nicotine [74]. The proliferation of ALDH-positive cells in MCF-7 cells treated with the α7-nAChR specific agonist PHA 543,613 was increased, and this effect was blocked by the N-choline antagonist α-BTX [59]. In vivo studies have shown that nicotine may promote the growth of solid tumors by inducing cancer cell proliferation, adhesion, and angiogenesis.

Higher expression of α9-nAChR mRNA was detected in ER (-) tumor tissues than in ER (+) tumor tissues. Mouse transplant tumor models and the NNK human mammary epithelial cell line MCF-10A can undergo carcinization on exposure to nicotine [81]. Further studies showed that NNK could not only upregulate the phosphorylation level of ERK1/2 and stimulate the expression of genes encoding tumor-related factors but also downregulate the expression of the tumor suppressor gene CDKN2A, thereby inducing cancer [52]. Small RNA interference experiments confirmed that the proliferation of breast cancer cells induced by nicotine or NNK can be inhibited by silencing the expression of α9-nAChR [49]. In contrast, α9-nAChR overexpression can stimulate the proliferation of breast cancer cells [151]. It was demonstrated that nicotine and NNK regulate the pathological processes of cancer by inducing the expression of genes related to cell process or metabolism, and this is closely related to the activation or high expression of nAChR.

Glutamate and breast cancer

mGluR is a major excitatory neurotransmitter in the mammalian neuroendocrine system. Strong or chronic repeated stress can induce excessive and persistently activated excitation, resulting in increased free radical generation, causing pathological damage through oxidative stress [47]. Increasing evidence proves that mGluR is involved in the development of breast cancer [161]. Xiao et al. constructed an mGluR4 eukaryotic expression plasmid capable of high expression in MCF-7 cells. Immunofluorescence results showed that mGluR4 was localized in the cell membrane and cytoplasm. CGK-8 experiments showed that high mGluR4 expression significantly promoted the proliferation of breast cancer cells [26]. Speyer et al. [135] discussed the potential role of mGluR1 in mediating endothelial cell (EC) proliferation and tumor-induced angiogenesis. Using mGluR1-specific inhibitors or shRNA silencing, the growth of EC and the formation of stromal tubes was shown to be dependent on the mGluR1 signaling pathway. Additionally, in mouse models of breast cancer, the loss of mGluR1 activity led to decreased angiogenesis. We think mGluR1 acts as a pro-angiogenic factor and a mediator of tumor progression in breast cancer. Therefore, researchers suggest that mGluR1 is a potential new molecular target for anti-angiogenesis therapy in breast cancer. Although the mechanism of mGluR in the nervous system has been studied in detail, its role in the occurrence and development of breast cancer remains to be elucidated.

Histamine and breast cancer

HA is involved in the pathological process of physiological allergies and various diseases as a central neurotransmitter [90]. Martinel et al. applied restraint-cold stress to stimulate rats to determine the HA level in the heart, lungs, liver, kidneys, and stomach. The results showed that the HA levels in the heart in both the stress and the control groups were not significantly different [90]. The HA level showed a significant increase in the stress group but remained unchanged in the control group.
Amino acid neurotransmitters regulate the pathological mechanism of breast cancer

A. GABA_A and GABA_C receptors are ionotropic channels, whereas GABA_B receptors are metabotropic. ABAT is a key enzyme in GABA metabolism that catalyzes the degradation of GABA to produce succinic semialdehyde and L-glutamic acid, thereby contributing to reducing GABA levels. The downregulation of ABAT expression in basal cell-like breast cancer may be due to GABA-mediated activation of the Ca^{2+}-NFAT1 axis. In addition, GABAB1e (GB1e) may significantly enhance the cloning and microsphere formation ability of MCF-7 cells through endoplasmic reticulum stress and further activates the MAPK/AKT/ERK signaling pathway to promote breast cancer cell proliferation, migration, and invasion.

B. Neuronal nAChRs are categorized as heteropentamers and homopentamers. Their structures and pharmacological functions are different. When activated by an agonist, the influx of Na^+ and Ca^{2+} and the efflux of K^+ occurs, regulating the expressions of MAPK, PKC, VEGF, etc. These proteins are closely associated with the occurrence and development of breast cancer.
Under various stress conditions, the level of released HA increases, and, through its receptors, it participates in the regulation of the autonomic nervous system; cardiovascular activities; behavior; and secretion of ACTH, vasopressin, PR, β-endorphin, and NE [91]. HA expression and biosynthesis have been detected in breast cancer cells. Breast cancer cells treated with the H4R agonist clobenpropit, VUF8430, and JNJ28610244 showed enhanced cell migration and invasion. The increased levels of ERK1/2 phosphorylation and TGF-β1 were detected using western blotting, the increased expression of Smad2/3-positive nuclei was detected using indirect immunofluorescence, and the formation of tumors was detected using a mammosphere assay [48]. Increased histidine decarboxylase gene expression is associated with improved relapse-free and overall survival among patients with breast cancer. HA treatment (5 mg kg⁻¹) in 4T1 tumor-bearing mice reduced tumor growth and increased apoptosis. Although no immunomodulatory effects were observed in wild-type mice, significant correlations between tumor weight and cytotoxic lymphocyte infiltration were detected in H4R knockout mice. H4R is differentially modulated by agonists and antagonists impacting tumor growth and immunity in 4T1 tumor-bearing mice [100]. For this reason, HD could be a potential biomarker for breast cancer.

**Peptide neurotransmitters regulate the pathological mechanism of breast cancer**

**NPY and breast cancer**

NPY is one of the most abundant neuropeptides and plays an important role in regulating emotions and stress-related behaviors [159]. Studies have shown that NPY is involved in stress response to varying degrees and can freely cross the blood–brain barrier [58]. The NPY system is significantly altered during chronic stress. Zambello et al. [158] found that the expressions of NPY mRNA in the hippocampus, Y1 receptor mRNA in the ventral and medial hypothalamus, and Y2 receptor mRNA in the ventral and medial hypothalamus were significantly lower in mice under chronic stress than in those in the control group. The anxiety and depression symptoms shown by the mice in the stress group were significantly increased compared with those of the mice in the control group. This suggests a functional connection between the hippocampus and hypothalamus NPY system and that chronic stress may damage the NPY system, indirectly leading to the occurrence of emotional problems [17].

Receptor autoradiography showed that 85% of breast cancer cases had NPY receptors. The NPY family includes three peptides: NPY, polypeptide YY, and pancreatic polypeptide; of these, NPY is more important in tumors [125]. NPY acts through four GPCRs: Y1, Y2, Y4, and Y5. Y1, Y2, and Y5 are associated with various aspects of tumorigenesis and angiogenesis [6]. NPY receptors are expressed in the colon, kidneys, adrenal gland, reproductive organs, testicles, and breasts [19]. Y1 is the dominant receptor in tumors, whereas Y2 is expressed preferentially in nontumor breast tissue. Reubi et al. proposed that oncogenic transformation may be conferred from Y2 to Y1 subtype [112]. Recent in vitro studies have implicated Y5 receptors in breast cancer growth and angiogenesis. Additionally, NPY inhibited the inflammatory response and enhanced the activity of MCF-7 cells in a hypoxic environment [40]. This is probably because NPY affects the inflammatory response, which in turn affects HIF-1 expression in tumor cells.

**Neurotensin and breast cancer**

Neurotensin (NTS) shows various central and peripheral effects by binding to NTS receptor (NTSR) [45]. There are three known subtypes of NTSR: NTSR1, NTSR2, and NTSR3. NTSR1, the high-affinity NTSR, has been well studied and is associated with the occurrence, development, proliferation, invasion, and prognosis of various malignant tumors [87, 97,98]. However, currently, there are only few studies on the role of NTSR1 in breast cancer. In this study, the relevant literature on the occurrence and development of breast cancer mediated by the NTS/NTSR1 signaling pathway in recent years has been reviewed. NTS and NTSR1 are explored as potential diagnostic markers and drug action targets for breast cancer treatment.

Somai et al. [131] found that NTSR1 and NST activation can promote the transcriptional activation of Bel-2 in breast cancer cells and promote the growth of tumor cells. Bakirtzi et al. [13,162] showed that binding of NTS and NTSR1 promoted the expression of interleukin-8, which was dependent on NF-κB, and NF-κB activated the insulin-like growth factor-1 receptor (IGF-1R)-mediated PI3K/Akt pathway. This activation promoted the expression of NF-κB-dependent related genes so that epithelial cells could evade apoptosis, thereby leading to cell proliferation. Studies on insulin receptor substrates (IRSs) confirmed that the combination of IRSs and IGF-1R does play an important role in the proliferation and metastasis of breast cancer. Souazé et al. [134] found that NTS and NTSR1 were expressed in MDA-MB-231 cells. Exogenous NT can enhance the movement and invasion of cancer cells after activating NTR1. Watters et al. found that NTSR is the target gene of estradiol and that estradiol can increase the transcription of NTSR in the hypothalamic arcuate nucleus neuroendocrine nucleus and preoptic region. MAPK signals activated by NTS can also significantly increase the HSP27 phosphorylation at site 82. The overexpression of heat shock proteins in patients with breast cancer promoted the proliferation and invasion of cancer cells, which is a marker of poor prognosis for malignant breast cancer and is related to the degree of tumor invasion [8]. High HSP27 expression in breast cancer can affect the sensitivity of tumor cells to hormone therapy [45,54,150]. In addition, MAPK signals
activated by NTS can also significantly increase the phosphorylation of HSP27 at site 82 [54] (Fig. 4). NTS can also express early growth response gene-1 (EGR-1), and the promoter region of epidermal growth factor receptor (EGFR) has an EGR-1 binding site [163]. Thus, EGFR expression is promoted, which is closely related to breast cancer occurrence. EGFR is a benzimidazole compound. At present, there are clinically specific antibodies and small-molecule kinase inhibitors against EGFR receptors used in breast cancer treatment. Some studies have pointed out that there are EGFR mutants in breast cancer, and these mutants are produced due to the deletion, mutation, and rearrangement of EGFR. The deletion of certain domains of EGFR leads to the destruction of receptor downregulation mechanisms, activation of abnormal signal transduction pathways, and inhibition of cell apoptosis. The EGFR ligand also has a great influence on intracellular signal transduction. Coexpression of the two often indicates a poor tumor prognosis. In addition, KRAS protein is downstream of the EGFR signaling pathway. Therefore, KRAS gene testing can screen those patients with breast cancer for whom EGFR-targeted therapeutic drugs are effective and realize the individualized treatment of patients with breast cancer.

Thus, NTSR1 can serve as an important target for breast cancer diagnosis and treatment, but many of its mechanisms have not yet been fully determined. For example, it is not understood if other molecules

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**Fig. 4.** The NTS/NTSR1 signaling pathway mediates the occurrence and development of breast cancer.

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are involved in the invasion and metastasis of NTS–NTSR1 mediated by ECM degrading enzymes. In various NTSR1-positive malignant tumors, such as pancreatic and breast cancers, NTS usually regulates the growth and proliferation of tumor cells through an autocrine mechanism. Therefore, it has been proposed that the stability of NTS mediated by MMPs may be a new regulation mode of tumor cells [24]. Then, the question arises whether the elevated MMP level in the process of NTS-NTSR1 promoting breast cancer invasion and metastasis causes the degradation of NTS and terminate this signal. Studies have found that the regulation of NTS stability by MMP in tumor cells may be more complicated and that the regulation of NTS by MMP may be related to the degree of dependence of tumor cells on sex hormones [44,124]. Then, is the role of MMPs positive or negative and what is the relationship between MMPs and the regulation of estrogen signaling? These questions need to be further investigated. **Table 2** summarizes the roles and molecular mechanisms of the above neurotransmitters in promoting breast cancer during chronic stress.
Fig. 5. Structures of other chemicals expected to reverse or interfere in the occurrence and development of breast cancer which listed in Tables III to VII. (1) Fenoldopam; (2) L-Stepholidine; (3) A77636; (4) C17; (5) Thioridazine; (6) Sulpiride; (7) Bromocriptine; (8) Doxazosin; (9) Rauwolscine; (10) Yohimbine; (11) Tramadol; (12) Propranolol; (13) Quercetin 3-O-glucuronide; (14) ICI 118,551; (15) Atenolol; (16) Isoproterenol; (17) SR269970; (18) 4-DAMP; (19) B3-1113; (20) Cevimeline; (21) Garcinol; (22) Darifenacin; (23) Luteolin; (24) EGCG; (25) Quercetin; (26) YM 298,198; (27) BAY36-7620; (28) YM 298,198; (29) Riluzole; (30) OUP-186; (31) Clobenpropit; (32) JNJ7777120; (33) Scorpion venom; (34) JMV449; (35) SR-48,692.
Fig. 5. (continued).
Fig. 5. (continued).
models; the inhibitory effect was obvious when combined with dexamethasone. Dexamethasone and the combined treatment could significantly reduce the expression of MMP2 in xenograft tumor while increasing the expression of NTSR1. This preliminarily explains the relationship between antipsychotics and breast cancer (95, 116). For example, the antipsychotic drug thioridazine significantly promoted apoptosis of drug-resistant and metastatic breast cancer through the antagonism of the DRD2 receptor (145). Therefore, researchers believe that doxazosin can be a potential drug for breast cancer treatment. The following section summarizes the research progress of DR-α2 antagonists in the treatment of breast cancer (Table 4).

### ADR-β2 antagonists

The high expression of ADR-β in some breast cancer cell types affects the recurrence and metastasis of breast cancer, suggesting that blocking the ADR-β pathway is an effective strategy to slow down the progression of breast cancer (105). A meta-analysis by Wang et al., a retrospective analysis by Zhao et al., a systematic analysis of cohort studies of breast cancer by Childers et al., and a clinical study analysis by Choy evaluated the relationship between ADR-β antagonists and breast cancer prognosis.
Table 3

| Subtype | Drug | Material basis | Mechanism | Effect | Model | References |
|---------|------|----------------|-----------|--------|-------|------------|
| DR agonists | Fenoldopam | 1 | Suppressed cell invasion and induced cell death | Inhibited the G protein-coupled receptor (GPR) pathway | MDA-MB-231 cells BALB/c mice | [88] |
| DR agonists | Lisinopril | 2 | Suppressed cell invasion and induced cell death | Inhibited the G protein-coupled receptor (GPR) pathway | MDA-MB-231 cells 4T1 cells BALB/c mice | [156] |
| DR agonists | Lisinopril | 3 | Suppressed cell invasion and induced cell death | Inhibited the G protein-coupled receptor (GPR) pathway | MDA-MB-231 cells 4T1 cells BALB/c mice | [156] |
| DR agonists | Lisinopril | 4 | Suppressed cell invasion and induced cell death | Inhibited the G protein-coupled receptor (GPR) pathway | MDA-MB-231 cells 4T1 cells BALB/c mice | [156] |
| DR antagonist | Bromocriptine | 5 | Activated DRD2 | Inhibited cell invasion and induced cell death | MCF-7 cells MDA-MB-231 cells MCF-7 cells BALB/c mice | [109] |
| DR antagonist | Sulpiride | 6 | Activated DRD2 | Inhibited cell invasion and induced cell death | MCF-7 cells MDA-MB-231 cells MCF-7 cells BALB/c mice | [157] |

The results showed that BBS significantly reduced the risk of recurrence and distant metastasis, but most of these studies involved small samples and the follow-up time was relatively short; thus, larger studies are needed to further explore the relationship between BBS and breast cancer prognosis.

At present, PROP, labetalol, metoprolol, bisoprolol, and atenolol are commonly used clinical ADR-β antagonists in the treatment of heart diseases such as hypertension and arrhythmia. Recent studies have suggested a correlation between ADR-β antagonists and the prognosis of breast cancer. PROP is well tolerated and has high safety. It has been clinically used to treat infantile hemangioma, and a phase II clinical trial of its use for the treatment of breast cancer is underway. PROP is used in combination with chemotherapy drugs to prevent cardiotoxicity caused by anthracyclines and inhibit breast cancer metastasis induced by SNS activation caused by postoperative stress [80]. PORP can inhibit the infiltration of macrophages in the tumor parenchyma to a large extent, indicating that ADR-β signaling plays an important role in stress-mediated tumor metastasis. PROP can also inhibit glucose metabolism and uptake in 4T1 breast cancer cells. By inhibiting the ADR-β signal, PROP can reduce the Ki-67 proliferation index of breast cancer cells, reduce the expression of various cyclins, and enhance the expression of the tumor suppressor gene p53 protein in the cells. It has also been shown to induce apoptosis of breast cancer cells and significantly inhibit tumor growth in patients with advanced breast cancer [96]. Rivero et al. demonstrated that β-blockers can effectively reduce the number of lung metastases and tumor size of triple-negative breast cancer cells by simulating an MDA-MB-231 cell metastasis model in NSG-immunodeficient mice. The following section summarizes the research progress of ADR-β2 antagonists in the treatment of breast cancer (Table 5).

Other receptor antagonists

The following section summarizes the research progress of other neurotransmitter receptors antagonists in the treatment of breast cancer (Table 6).

Nonpharmacological interventions to reverse chronic stress

Tumor treatment has already entered the era of comprehensive treatment, and exploratory research on stress management strategies is extremely challenging. Studies have suggested that adding medical and psychological intervention and cognitive-behavioral stress management to the treatment of patients with cancer can reduce the patients’ stress level and improve their quality of life, thereby improving clinical efficacy [15,146]. Benign stress often brings beneficial effects, and it is no exception for malignant tumors. The elevated levels of adrenaline and IL-6 induced by exercise can lead to tumor growth through the mobilization and redistribution of NK cells [65,104]. Results from mouse models also verified that the benign stress of pancreatic cancer, lung cancer, and melanoma lung metastasis can stabilize the expression levels of CCR5 and NKG2D in NK cells through the activation of sympathetic nerves, thereby enhancing their antitumor effects [132]. Clinical studies have shown that behaviors such as mindfulness, meditation, and yoga are effective in reducing chronic stress and improving the quality of life of patients with cancer [11,115,122]. In short, a healthy lifestyle, a positive life attitude, and professional stress management guidance are essential for the prevention and treatment of tumors.

The COVID-19 pandemic continues to affect our lives and has led to several inconveniences for patients with breast cancer; this has made them prone to negative emotions, such as anxiety and depression, which affects the effectiveness of treatment. Strengthening the psychological care of patients undergoing chemotherapy after breast cancer surgery can relieve their distress and help them successfully complete the course of chemotherapy. One of the consequences of the COVID-19 pandemic is the need to prevent infections and delay in breast cancer screenings. The
Canadian Association of Breast Imaging/Canadian Association of Radiologists has developed safety guidelines for breast imaging and concluded that delaying breast cancer screening should be avoided if possible even during the COVID-19 pandemic.

**Summary and prospects**

In the personalized treatment of patients with breast cancer, the emergence of chronic stress is always accompanied by neurological disorders and disorders that affect the prognosis of patients to varying degrees. Independently existing psychological problems are unrelated to the technical problems involved in the treatment process. Existing treatments for breast cancer may not reduce the continuous stress faced by patients due to the psychological effects. Considering that the mechanism of chronic stress promoting the occurrence and evolution of breast cancer has so far not been systematically and comprehensively revealed, this article reviews the research on the correlation between chronic stress and the occurrence and development of breast cancer. The molecular mechanism by which chronic stress promotes breast cancer development through related neurotransmitters progress in elucidating potential drugs or blockers in the treatment of breast cancer have been emphasized. Novel ideas for the prevention and treatment of breast cancer in clinical practice, and the theoretical basis for the use of classical blockers in the treatment of breast cancer have been discussed. One interesting finding of this review is that drugs used to treat mental illness are implicated in playing an important role in the treatment of breast cancer, thereby expanding the new uses of classical drugs (Fig. 6).

We should pay more attention to the psychological state of patients and put forth scientific and reasonable stress management strategies. We advocate timely and targeted psychological interventions for patients with breast cancer with depressive symptoms. For example, meditation, exercise, and cognitive-behavioral therapy may become a feasible method for the treatment of patients with cancer and help to improve the psychological state of patients. Paying attention to the clinical manifestations of depressive symptoms in patients with breast cancer and improving the effectiveness of antitumor treatment for patients with breast cancer will undoubtedly provide novel therapeutic strategies for breast cancer treatment. We also want to attract the readers’ attention to the fact that drugs used to treat mental illness are implicated in playing an important role in the treatment of breast cancer, thereby expanding the new uses of classical drugs.

There are still many unsolved problems regarding the role of neurotransmitters in tumor genesis and development: (1) Are the mechanisms of neurotransmitters in different types of breast cancer the same? (2) How do neurotransmitters participate in the regulation of tumor cells through other cells (such as fibroblasts, endothelial cells, and immune cells) in the tumor microenvironment? (3) Glucocorticoids are often clinically used to reduce the side effects of chemotherapy in patients with cancer. Will this have an adverse effect on cancer treatment? (4) Is there any connection between the inflammatory response and neurotransmitters in tumors? The effect of the nervous system on breast cancer is a non-negligible factor, and it is evident that different neurotransmitters have different mechanisms of action in different types of breast cancer, ultimately adding to the complexity of this relationship. This will be the focus of our future studies.

**Table 4**

| Subtype | Drug         | Material basis | Model          | Effect                             | Mechanism                                      | References |
|---------|--------------|----------------|----------------|------------------------------------|-----------------------------------------------|------------|
| ADR-α1 antagonists | Doxazosin | 8              | MDA-MB-231      | Induced apoptosis                  | Blocked EGFR and NF-κB signaling              | [63]       |
|         | Doxazosin    | 8              | MCF-7           |                                    |                                               |            |
|         | Doxazosin    | 8              | MCF72A          |                                    |                                               |            |
|         | Rauwolscine  | 9              | MCF4-L5         | Inhibited the proliferation        |                                               | [145]      |
|         | Rauwolscine  | 9              | Mouse tumor C4-HD |                                    |                                               |            |
|         | Yohimbine    | 10             | MCF-7           | Reversed cell proliferation        | Enhanced the pump function of P-glycoprotein and conferred resistance of MCF-7 cells to paclitaxel | [139]      |
|         | Yohimbine    | 10             | SCID mice       |                                    |                                               |            |
|         | Tramadol     | 11             | MDA-MB-231 BALB/c mice | Inhibited proliferation, migration, and invasion | Reduced ERR expression                        | [153]      |

**Table 5**

| Subtype | Drug         | Material basis | Breast cancer cell line | Effect                                      | Mechanism                                      | References |
|---------|--------------|----------------|-------------------------|---------------------------------------------|-----------------------------------------------|------------|
| Natural products that inhibit ADR-β2 | Quercetin-3-O-glucuronide (Q3G) | 13              | MDA-MB-231               | Suppressed cell invasion and MMP-9 induction and inhibited the binding of [3H]-NA to ADR-β2 | Suppressed ROS generation, cAMP and RAS activation, phosphorylation of ERK1/2, and HMOX1, MMP2, and MMP9 gene expression ERK1/2 and COX-2 phosphorylation levels were significantly decreased after β-blocker treatment. | [155] |
| ADR-β2 antagonists | Propranolol | 12             | MDA-MB-435               | Inhibited viability of cells, arrested cell cycle progression at G0/G1 and S phase, and induced cell apoptosis |                                               | [154]      |
|         | Propranolol  | 12             | MDA-MB-231               |                                    |                                               |            |
|         | ICI 118,551  | 14             | MDA-MB-48               |                                    |                                               |            |
|         | Atenolol     | 15             | MDA-MB-453               | Inhibited cell proliferation              |                                               | [21]       |
|         | Isoproterenol | 16            | MDA-MB-231 ZR-75         | Inhibited cell proliferation              | Inhibited DNA synthesis                       | [129]      |

**CRediT authorship contribution statement**

Hui-min Liu: Writing – review & editing, Visualization, Writing – original draft. Le-le Ma: Writing – review & editing, Visualization. Chunyu Li: Investigation, Writing – review & editing. Bo Cao: Writing – review & editing, Visualization. Yifang Jiang: Investigation, Writing –
Table 6
Other receptor antagonists.

| Subtype                       | Drug                  | Material basis | Model         | Effect                                      | Mechanism                                                                 | References         |
|-------------------------------|-----------------------|----------------|---------------|---------------------------------------------|---------------------------------------------------------------------------|--------------------|
| 5-HT antagonists              | SB269970              | 17 T47D        | MCF-7         | Antiproliferation and anti-invasion         | Gα-activated cAMP and Gβγ-activated kinase signaling during invasion and Gβγ-activated PI3K/Akt signaling during proliferation | [50]               |
|                               | BJ-1113               | 19 MDA-MB-231  | MCF-7         | The inhibitory effect of BJ-1113 against MDA-MB-231 tumor growth was greater than that of SB269970, a 5-HT7 receptor antagonist |                                                              |                    |
| mAChR antagonists             | Cevimeline            | 20 MDA-MB-231  | MCF-7         | Inhibited cell proliferation               | PLC activation                                                             | [30]               |
|                               | 4-DAMP                | 18 MDA-MB-231  | MCF-7         | Inhibited cell proliferation               | PLC activation                                                             | [69,120,123]       |
|                               | Darifenacin           | 22 MDA-MB-231  | MCF-7         | Inhibited cell proliferation               | PLC activation                                                             | [86,99,106]        |
| nAChR antagonists             | Epigallocatechin-3-gallate (EGCG) | 24 MCF-7 | | Inhibited cell proliferation               | Blocking α9-nAChR signaling pathway                                       | [144]              |
|                               | Garcinol              | 21 MDA-MB-231  | MCF-7         | Inhibited cell proliferation               | Downregulate the expression of α9-nAChR and cyclin D3 protein            | [27]               |
| mGluR1-specific antagonists   | Riluzole              | 29 MDA-MB-231  | MCF-7         | Inhibited cell proliferation in a dose-response manner in all EC types | Inhibition of angiogenesis                                                  | [41,136]          |
|                               | YM 298,198            | 28 MCF-7       |               | Inhibited cell proliferation in all the cell types, except HUVEC | Inhibition of angiogenesis                                                  | [135]              |
| H3R antagonists               | OUP-186               | 30 MDA-MB-231  | MCF-7         | Induced breast cancer cell death            | Activated caspase-3/7                                                      | [142]              |
|                               | Clobenpropit          | 31 MDA-MB-231  | MCF-7         | Cell death was only slightly induced        | Activated caspase-3/7                                                      |                    |
|                               | JNJ7777120            | 32 MDA-MB-231  | MCF-7         | Inhibited cell migration and invasion       | Reduced E-cadherin, cytoplasmic and nuclear β-catenin, and nuclear Slug and an increase in vimentin and α-smooth muscle actin expression | [48]               |
| Peptide neurotransmitter      | Scorpion venom        | 33 MDA-MB-231  | MCF-7         | Reduced motility and invasion in breast cancer cells | Inhibited MMP activity; upregulation of p53 and downregulation of Bcl-xl and BID protein expression by modulating signaling proteins Erk1/2 and STAT3 and DNA damage in breast and colorectal cancer cell lines | [4,5]             |
| agonist                       | JMV449                | 34 MCF-7       |               | Promoted apoptosis                          | MAPK activation caused the expression of B-cell lymphoma-2 (BCL-2)       | [131]              |
| NTS antagonist                | SR48692               | 35             |               |                                            |                                                                          | [77]               |

In summary, we highlight in the Fig. 5 which are the potential targets for drug development.

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Hui-min Liu: Writing – review & editing.
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Declaration of Competing Interest

No author has an actual or perceived conflict of interest with the contents of this article.
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