Regulatory effects and potential therapeutic implications of alarin in depression, and arguments on its receptor

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Alarin is a pleiotropic peptide involved in a multitude of putative biological activities, notably, it has a regulatory effect on depression-like behaviors. Although further elucidating research is needed, animal-based cumulative evidence has shown the antidepressant-like effects of alarin. In light of its regulatory role in depression, alarin could be used as a promising antidepressant in future treatment for depression. Nevertheless, the available information is still insufficient and the therapeutic relevance of alarin in depression is still of concern. Moreover, a plethora of studies have reported that the actions of alarin, including antidepressant activities, are mediated by a separate yet unidentified receptor, highlighting the need for more extensive research. This review focuses on the current understanding of the regulatory effects and future therapeutic relevance of alarin on depression, and the arguments on its receptors.

KEYWORDS
depression, antidepressant effect, alarin receptor, therapeutic implication, alarin

Abbreviations: 4E-BP1, 4E-binding protein 1; ACTH, adrenocorticotropic hormone; BDNF, brain-derived neurotrophic factor; CNS, central nervous system; CORT, corticosterone; CREB, cAMP response element-binding protein; CRH, corticotropin-releasing hormone; DA, dopamine; ERK, extracellular signal-regulated kinase; GALP, galanin-like peptide; GaIR, galanin receptor; GMAP, galanin message-associated peptide; GPCR, G protein-coupled receptor; GR, glucocorticoid receptor; HPA, hypothalamus–pituitary–adrenal gland; ICV, intracerebroventricular; IL, interleukin; MAOIs, monoamine oxidase inhibitors; MI, myocardial infarction; MR, mineralocorticoid receptor; mTOR, mammalian target of rapamycin; NEP, norepinephrine; NMDA, N-methyl-D-aspartate; NRIs, combined serotonin-NEP reuptake inhibitors; SSRIs, selective serotonin reuptake inhibitors; TCAs, tricyclic antidepressants; TNF, tumor necrosis factor; TrkB, tropomyosin-related kinase B; UCMS, unpredictable chronic mild stress.
Introduction

Depression is a common mental illness that presents with an abnormal lowering of mood (feeling sad, irritable, and empty), a loss of pleasure or interest in activities, feelings of excessive guilt or low self-esteem, and feelings of hopelessness about the future, anxiety, sleep disturbances, changes in appetite or body weight, feeling tired, thoughts of suicide, sexual dysfunction, and cognitive impairments (1). It is one of the most widespread health threats to humans, affecting about 4.4% of the global population (2). It is also a leading cause of disability worldwide, contributing significantly to health, social, and economic problems (3, 4). Although the pathogenesis of depression is not well elucidated, the interplay between genetic and psychosocial predisposing factors has been suggested to be involved (5, 6). Numerous medications for the treatment of depression are currently available, but most of them are associated with low therapeutic response rates as well as adverse effects, prompting researchers to look for a novel treatment option (7). In recent years, several neuropeptides, such as the galanin neuropeptide family, have been discovered to have promising therapeutic relevance in depressive mood disorders (8, 9). Galanin, the ancestral peptide of the galanin family, has been shown to be implicated in depression and to have therapeutic value (10). Alarin, another member of the galanin family, has also recently been demonstrated to play a pivotal regulatory role in depression-like behaviors (11).

Alarin is a 25-amino acid containing biologically active peptide from the family of galanin, which was discovered for the first time 16 years ago in the gangliocytes of human neuroblastic tumors (12). It is a splice variant of galanin-like peptide (GALP) that results from alternative splicing of the GALP gene through exon three exclusion (13). Its name was termed after its N-terminal alanine and C-terminal serine (12). In healthy individuals, the normal plasma concentration of alarin is 0.37 ±0.10 µg/L (14). Alarin is broadly distributed in the central nervous system (CNS) and different peripheral tissues of various species like skin, eyes, the gastrointestinal tract, and the endocrine glands.

According to numerous rodent studies, the alarin peptide is distributed in a wide range of nuclei in the brain of mice and rats, including the mitral cell layer of the olfactory bulb, the accessory olfactory bulb, the medial preoptic area, the amygdala, and the bed nucleus of the stria terminalis (15–18). Besides, alarin is markedly expressed in a number of nuclei in the hypothalamus, such as paraventricular nucleus (PVN), dorsomedial nucleus (DMN), and the arcuate nucleus (ARC), ventromedial nucleus (VMN), and lateral hypothalamus (15, 16, 19). Moreover, this neuropeptide is expressed in the locus coeruleus (LC), and locus subcoeruleus of the midbrain and hindbrains, as well as the trigeminal complex, the ventral cochlear nucleus, the facial nucleus, and the epithelial layer of the plexus chorois in the brain (15, 19). Based on a clinical study by Eberhard et al. alarin is present in a variety of nuclei in human brain as demonstrated from medium to high-intensity alarin-like immunoreactivity (LI) in all choroid plexus tumors, in the majority of ependymomas, and the minority of astrocytomas, meningiomas, and tumors of the cranial nerves (18). However, clinical studies demonstrating the tissue distribution of alarin in human are limited, which warrants further research to identify the difference in the expression of alarin and its receptor among various species.

Alarin is a pleiotropic peptide with multifaceted biological functions (15, 16, 19, 20). It is involved in a series of important physiological effects, like the regulation of energy homeostasis, feeding behavior, body weight, and body temperature. Central injection of alarin into PVN has been shown to increase acute food intake and body weight in animals (15, 16, 19, 20). Alarin also has thermoregulatory effects that elicit significant fever-like hypermetabolic /hyperthermic thermoregulatory responses in rats (21). Additionally, this peptide plays a critical function in glucose metabolism by stimulating insulin-mediated glucose uptake in a variety of tissues in rodents (16, 20, 22). Moreover, alarin has a reproductive function that promotes the secretion of sex hormones such as luteinizing hormone (LH) (15, 20). In addition, this peptide exhibits antiedema, anti-inflammatory, and vasoconstrictive effects to maintain normal eye and skin health, antifibrinolytic activity, and antioxidant properties to prevent cardiac fibrosis, as well as antimicrobial activity against gram-negative bacteria (23–25). More importantly, it has been demonstrated that alarin exhibits antidepressant-like activity in various animal studies (11, 26–28). Although alarin has a variety of biological activities, the specific receptors that mediate these actions are still unknown. This review primarily points out the current understanding regarding the regulatory role of alarin in depression and its future potential therapeutic implications. Besides, the article provides a synopsis of the arguments concerning the alarin-specific receptor.

Role of alarin in depression and its potential therapeutic implications

Overview of the pathophysiology of depression

Depression is a seriously debilitating disorder that is thought to develop as a result of a constellation of genetic and psychosocial factors (5, 6). Numerous genomic studies indicate that depression is a heritable psychiatric disorder associated with a complex genetic architecture and several genetic variants (29). Although there is no solid evidence elucidating the pathophysiological mechanism underlying depression, a number of neurobiological hypotheses have been postulated in the past few decades, including the monoamine hypothesis, the neurotrophic hypothesis, the GABAergic
hypothesis, the glutamatergic hypothesis, and the alteration of
the hypothalamic-pituitary-adrenal (HPA) axis (30–34). The
monoamine hypothesis, which entails the catecholamine
hypothesis and the indolamine hypothesis, posits that the
underlying pathophysiological basis of depression is a deficiency
of the neurotransmitters serotonin, norepinephrine (NEP), or
dopamine (DA) produced by the serotonergic, noradrenergic,
and dopaminergic neurons in the monoaminergic systems,
respectively (35). The catecholamine hypothesis is based on the
findings that monoamine oxidase inhibitors (MAOIs) and
tricyclic antidepressants (TCAs), which boost the levels and
transmission of catecholamines, such as NEP and DA, exhibit
antidepressant effects. This indicates that the depletion of
catecholamines in the brain is linked to the pathomechanism
of depression. Additionally, dysregulation of catecholamine
transmission is postulated to be involved in depression (30, 35– 38). The indolamine hypothesis, on the other hand, proposes a
decrease in the activity of serotonin [or 5-hydroxytryptamine
(5-HT)] in the brain as well as a deficiency in pre-synaptic and
post-synaptic 5-HT receptors, particularly the 5-HT1A auto-
receptor, which is a potential predisposing factor to depression
(39–42).

According to the neurotrophic hypothesis, another
pathophysiological hypothesis of depression, the
downregulation of neurotrophins like brain-derived
neurotrophic factor (BDNF) is related to persistent stress
and depression. Numerous studies indicate that decreased
hippocampal and cortical BDNF levels are significantly
correlated with stress-induced depressive-like behaviors
(43–45). Moreover, the GABAergic hypothesis suggests that
the pathogenesis of depression is associated with reduced
brain concentration of the inhibitory neurotransmitter γ-
aminobutyric acid (GABA), as well as changes in its receptors
(GABA-A receptors) mediating GABAergic inhibition in
the prefrontal and occipital cortex (31, 32). GABAergic
transmission, which is crucial for the regulation of hippocampal
neurogenesis and neural maturation, plays a prominent role in
the brain's control of stress and represents a major vulnerability
factor in depressive mood disorders. Persistent stress is thought
to result in synaptic loss and changes in the interneuronal
plasticity in circuits underlying affective and cognitive
processes that contributes to the occurrence of depression.
Recent reports have shown stress-induced alterations in the
structure, connectivity, and neurochemistry of interneurons in
depressed individuals. In particular, stress-induced plasticity
of GABAergic inhibition occurs via changes in GABA synthesis,
release, and the expression of specific GABA,A-receptor subunits
has been shown to be involved in the development of depression
(46–49). A significant reduction in the concentration of GABA
has been observed in the cortex, hypothalamus, and olfactory
bulb as a result of chronic stress (50). Several lines of evidence
indicate that chronic stress reduces GABAergic synaptic
transmission in the ventral hippocampus due to the selective
loss of hippocampal neurons, and leads to depression (33, 34).

On the other hand, the glutamatergic hypothesis argues
that abnormalities in glutamatergic neurotransmission or
glutamatergic toxicity are implicated in the biological
mechanisms underlying depression. The glutamatergic
hypothesis is based on the observation of quick and large
antidepressant effects from blocking an ionotropic glutamate
receptor called the glutamate N-methyl-D-aspartate (NMDA)
receptor (51–53). Moreover, alteration of the HPA axis has been
demonstrated to participate in the development of depression.
According to this theory, the pathophysiology of depressive
disorders is fundamentally linked to increased glucocorticoid
secretion and an overactive HPA axis (54–57).

Regulatory effects of alarin in depression

Accumulating evidence shows that some members of
the galanin neuropeptide family are involved in mediating
behavioral functions linked to stress and anxiety, suggesting
their potential involvement in mood disorders like depression
(58). Galanin, which is abundantly distributed in the
hypothalamus and amygdala, is linked to stress response
and depression as it is co-expressed with and modulates NEP
and serotonin systems (59–63). Therapeutically, antagonists
of galanin receptor (GalR) sub-type 1 (GalR1) and GalR3, as
well as agonists of GalR2 have been established to have an
antidepressant-like effect by enhancing the activities of selective
serotonin reuptake inhibitors (SSRIs) and promoting the
availability of transcription factors (64, 65). Importantly, recent
research has reported that alarin, which is also localized in the
brain areas linked with depression, like the medial amygdala
and hypothalamus, has been demonstrated to have regulatory
effects on depression-like behaviors in various animal models
(11, 25–27, 66).

Wang and coworkers revealed for the first time that alarin
exhibits potent antidepressant-like effects in both the acute
stress model and the unpredictable chronic mild stress (UCMS)
depression-like model. The study additionally demonstrated
that alarin is distributed in the hypothalamus and LC, which
are recognized as critical regions in mood-related disorders
(26). Zhuang et al. consistently showed that central injection
of alarin significantly ameliorates depression-like behaviors in
the UCMS mouse model (28). Subsequently, the antidepressant
properties of alarin were further confirmed by several other
studies (11, 27, 66). However, the antidepressant-like effect of
alin has been shown to be dose-limited, with no remarkable
effect at higher doses (at 2.0 nmol or higher). Although the
mechanism by which alarin produces the antidepressant effect
is not yet fully understood, a number of potential mechanisms
have recently been reported. Here, some putative mechanisms

of alarin-induced antidepressant actions are briefly discussed in two separate sections: the TrkB-mTOR signaling pathway and the hypothalamus–pituitary–adrenal (HPA) axis.

The tropomyosin-related kinase B-mammalian target of rapamycin signaling pathways

According to several animal-based studies, alarin has been found to alleviate depression by targeting various components in the TrkB-mTOR signaling pathways as discussed below. The putative antidepressant mechanism of alarin using this pathway is further illustrated in Figure 1.

Tropomyosin-related kinase B receptor (TrkB): Alarin has been found to exert antidepressant-like effects by influencing tropomyosin-related kinase B (TrkB) which alters the downstream signaling pathways. TrkB is a member of the neurotrophic tyrosine kinase (NTRK) receptor family encoded by the NTRK2 gene. TrkB has a high level of expression in the brain and serves as a common upstream receptor for a number of ligands such as the brain-derived neurotrophic factor (BDNF), neurotrophin 3 (NT3), and neurotrophin 4 (NT4) (67, 68). This receptor is crucial for neural cell differentiation, survival, and proliferation (69). Current research has shown that defective TrkB is closely linked with the pathophysiology of depression. This is evident from the significantly reduced TrkB expression in the prefrontal cortex and hippocampus of postmortem human subjects with major depression (70–72). Hence, alarin is postulated to have an effect similar to that of antidepressants by targeting TrkB receptor-mediated activation of downstream cascades, especially extracellular signal-regulated kinase (ERK) and protein kinase B (AKT) signaling pathways (11, 26, 27). TrkB receptor-mediated Ras-ERK and PI3K-AKT intracellular signaling pathways are involved in neuronal survival and neurogenesis and result in antidepressant-like effects (73). While the TrkB-activated Ras-ERK pathway is involved in cell proliferation, differentiation, apoptosis, and synaptic plasticity, the PI3K-AKT pathway is involved in the regulation of cell growth, survival, proliferation, and movement (74–77). Alarin is thus demonstrated to have antidepressant properties in UCMS-treated mice, likely by increasing the expression of AKT and ERK in some areas of the brain, including the prefrontal cortex, hippocampus, olfactory bulb, and hypothalamus. Despite the putative role of TrkB in mediating the antidepressant activities of alarin, additional research is needed to determine whether alarin directly acts on TrkB or whether TrkB itself is an unidentified receptor of alarin.

Mammalian target of rapamycin (mTOR) and downstream components: The mTOR is another target of alarin to achieve its antidepressant-like effects via ERK and AKT pathway activation. mTOR is a widely expressed serine/threonine kinase that belongs to the phosphatidylinositol 3-kinase (PI3K)-related kinase protein family and is a key player in controlling protein translation (78). It is a target of the ERK/AKT pathway that integrates signals from neuronal activity, growth factors, and nutrient levels to regulate the initiation of protein translation via its major downstream components, namely ribosomal protein S6 kinase (p70S6K) and eukaryotic initiation factor 4E-binding protein 1 (4E-BP1) (78, 79). Activation of mTOR phosphorylates p70S6K and 4E-BP and induces protein synthesis to regulate many integrated physiological functions of the nervous system, including neuronal development, synaptic plasticity, memory storage, and cognition (78, 80, 81). On the other hand, chronic stress disrupts mTOR signaling and markedly decreases synaptic proteins such as post-synaptic density 95 (PSD-95) and pre-synaptic protein synapsin I, indicating its role in the pathophysiology of depression (79, 82). The loss of synaptic proteins that play an important role in synaptic function (or plasticity) commonly results in impaired synaptic quantity and function, inducing neuronal changes, and increasing vulnerability to depression (83–85). The antidepressant-like effect of alarin may therefore be mediated by reactivating the mTOR signaling pathway, which will enhance synaptic protein synthesis and synaptic plasticity while reversing synaptic protein loss and impaired function (27). Increased synaptic expression of PSD-95 and synapsin I, by improving the development, maturation, and function of synapses, relieves depressive symptoms (82, 86–89). Intracerebroventricular (ICV) injection of alarin has also been proven to improve depression-like behaviors through the alarin-mediated restoration of UCMS-induced reduction of phospho-mTOR and phospho-4E-BP1 in the prefrontal cortex, hippocampus, hypothalamus, and olfactory bulb. Besides, alarin was observed to reverse UCMS-induced downregulations of p70S6K, PSD-95, and synapsin I expression in these brain regions (27). Additionally, alarin has been shown to affect the activity of the cAMP response element-binding protein (CREB), a downstream messenger of ERK and AKT. Acute alarin treatment has been reported to markedly increase phosphorylated CREB levels in the prefrontal cortex of stressed animals to basal levels of vehicle-treated mice, but no significant changes in CREB levels have been observed in other brain areas, such as the hippocampus, the olfactory bulb, and the hypothalamus (11). Several previous studies have indicated that rapid, long-lasting, potent antidepressant effects in both depressed patients and rodent models can be produced by increasing the activity of mTOR and the downstream signaling components that contribute to synaptic protein synthesis and synaptic plasticity (86, 90). This shows that alarin has pharmacological properties resembling those of antidepressant drugs and may be used as a treatment option for depression.

Brain-derived neurotrophic factor (BDNF): Alarin is also postulated to exhibit antidepressant effects through the upregulation of the expression of neurotrophin, particularly BDNF (26). BDNF is a dimeric protein in the neurotrophin family, which is thoroughly distributed in the brain, more abundantly in the cerebral cortex, hypothalamus, hippocampus, striatum, basal forebrain, and cerebellum (91). It promotes the growth, development, neuroprotection, and survival of neurons,
A schematic illustration of the putative antidepressant mechanism of alarin targeting the TrkB-mTOR signaling pathway and BDNF secretion. In this pathway, alarin may exert antidepressant-like effects by interacting with TrkB, possibly as a ligand, and stimulating downward signaling components like ERK and AKT. Besides, alarin may affect the activity of CREB by enhancing its phosphorylation. It also exhibits antidepressant effects by increasing mTOR, 4E-BP1, p70S6K, PSD-95, and synapsin I. Moreover, alarin enhances the secretion of natural antidepressant peptide (BDNF) through a pathway that is not yet well defined. The synthesis and release of BDNF from the transcription process activates TrkB receptors and the downstream mTOR signaling pathway, which coincides with the therapeutic response. BDNF-or alarin-activated mTOR triggers the translational machinery, increasing synaptic protein (PSD-95 and synapsin I) synthesis involved in synaptogenesis to confer an antidepressant effect. BDNF, brain-derived neurotrophic factor; CREB, cAMP response element-binding protein; CRE, cAMP response element; ERK, extracellular signal-regulated kinase; mTOR, mammalian target of rapamycin; PI3K, phosphatidylinositol 3-kinase; p70S6K, ribosomal protein S6 kinase; PKA, protein kinase A; PSD-95, post-synaptic density 95; 4E-BP1, eukaryotic initiation factor 4E-binding protein 1; TrkB, tropomyosin-related kinase B.

FIGURE 1

as well as regulates neuronal apoptosis and increases synaptic plasticity. This ultimately prevents neuronal degeneration and death, which reveals the antidepressant effect of BDNF (67, 92). BDNF is suggested to exhibit a principal antidepressant role by fostering synaptic plasticity by acting as a ligand of the TrkB receptor, stimulating mTOR to induce synaptic protein synthesis (93–95). Therefore, alarin can potentially alleviate depression by up-expressing BDNF, as evidenced by the increased expression of BDNF mRNA in the prefrontal cortex and hippocampus of UCMS-treated mice after alarin administration (26). This action of alarin is similar to the effects of most antidepressants, which upregulate BDNF expression in rodent brain and human blood. The increase in endogenous BDNF, in turn, attenuates the symptoms of depression by reversing neuronal atrophy and promoting cell proliferation in the hippocampus and prefrontal cortex (96, 97). However, this alarin-induced BDNF
upregulation is mediated by an unidentified receptor, even though it probably belongs to the GPCR receptor family.

The hypothalamic–pituitary–adrenal axis

Moreover, alarin regulates depression by controlling adrenocortical hormone secretion and the interaction of HPA axis organs (Figure 2). Like galanin, alarin plays a significant role in the regulation of depression-like behaviors via its antagonist activity against the hyperactivity of the HPA axis (93, 98). The abnormality of the HPA system, which is essential in stress responses and homeostasis maintenance, is involved in the pathophysiology of depression (57, 99, 100). It has been proposed that hyperactivity of the HPA axis and disruption of the stress feedback mechanism are common pathophysiological features of major depressive disorder in humans. A large amount of data suggests that aberrant stimulation of the HPA axis in depression activates the hormonal cascade in the brain (57). The hypothalamus secretes corticotropin-releasing hormone (CRH) in response to psychological stress in cortical brain regions. CRH stimulates the anterior pituitary gland to release corticotropin or adrenocorticotropic hormone (ACTH), which in turn triggers glucocorticoid secretion from the adrenal cortex to the plasma (91). In normal circumstances, the HPA axis is regulated to prevent it from becoming overactive by mineralocorticoid (MR) and glucocorticoid (GR) receptors. However, an imbalance in the MR/GR ratio, mainly a decrease in limbic GR receptor function and increased functional activity of the MR system, is seen in stress-related conditions like depression (101–103). Chronic stress in major depressive disorder induces neuronal degeneration in the hippocampus, leading to suppression and impairment of GRs that inhibit the negative feedback of glucocorticoids (104, 105). This disturbed stress feedback mechanism further activates the HPA axis and leads to the over-secretion of glucocorticoid levels (106–108).

Alarin may therefore have a crucial role in ameliorating depression by acting as a hormone regulator that downregulates the activity of the HPA axis. Available data shows that ICV injection of alarin in UCMS mice is associated with a decline in CRH expression levels in the hypothalamus, and a decrease in serum levels of CRH, ACTH, and corticosteroids [cortisol in humans and corticosterone (CORT) in animals] released from the HPA axis (26). Another study also found that alarin negatively affects the expression of enzymes involved in steroidogenesis, such as cholesterol desmolase, 11β-hydroxylase, and aldosterone synthase, in the adrenal glands. In addition, although it has no appreciable effect on blood concentrations of aldosterone and CORT, alarin substantially reduces blood hormone levels derived from the HPA axis, such as CRH, ACTH, and pro-opiomelanocortin (POMC) (66). This effect of alarin resembles those of other antidepressants that reduce the elevated levels of ACTH, cortisol, and CORT in depressed subjects (106–108). Collectively, alarin-mediated normalization of the serum levels of the key hormones of the HPA axis alleviates depression symptoms and exhibits an antidepressant-like effect.

Furthermore, the antidepressant-like properties of alarin may be mediated by restoring altered levels of the pro-inflammatory and anti-inflammatory cytokine, thereby reducing activation of the HPA axis (28). Based on numerous prior animal and human studies, inflammatory cytokines play an essential role in the pathophysiology of depression (109–112). Clinical data suggest that pro-inflammatory cytokines such as interleukin (IL)-1α, tumor necrosis factor (TNF)-α, and IL-6 disrupt GRs, activate the HPA axis, impair the central serotonin system, and consequently induce depressive symptoms (113). In contrast, anti-inflammatory cytokines are associated with reduced hyperactivity of the HPA axis and depressive symptomatology (28). Here, alarin reverses the UCMS-induced rise in pro-inflammatory cytokines (TNF-α and IL-6) and a decrease in the levels of anti-inflammatory cytokine (IL-10) in the blood, prefrontal cortex, hippocampus, and hypothalamus. In addition, alarin restores the UCMS-induced decrease in GR expression in these brain areas, which ultimately reduces HPA axis overactivity and depression (28).

Potential therapeutic implication of alarin in depression

Based on existing theories on the etiology of depression, many antidepressant drugs have been discovered and utilized in the treatment of depression for a long time. A variety of antidepressant medications targeting these hypothetical pathomechanisms are currently under consideration for the treatment of depression (114). Nonetheless, inhibitors of monoamine transporters like SSRIs, NEP reuptake inhibitors (NRIs), and combined serotonin-NEP reuptake inhibitors (SNRIs) are nowadays noted as the most widely prescribed antidepressant medications (7). These medications attenuate a postulated deficiency of monoamine transmission by raising extrasynaptic NEP and/or serotonin levels (115). However, currently available antidepressant drugs raise concerns about their efficacy and adverse effects, increasing the necessity of developing appropriate antidepressants with new mechanisms. Hence, researchers are urged to work diligently on developing novel therapeutic drugs with efficient antidepressant effects and fewer side effects, as well as seeking better targets for antidepressant therapies to achieve and maintain remission.

Currently, numerous neuropeptides, including the galanin neuropeptide family and their receptors, are attracting researchers’ interest to use them as novel therapeutic targets for depressive disorders. This is because such neuropeptides colocalize with the neurotransmitters NEP, 5-HT, and DA in some brain areas and mediate anxiety and stress-related behavioral functions (8, 9). Hence, designing drugs that enhance peptidergic transmission, regulating the activity
FIGURE 2
A hypothetical antidepressant activity of alarin through the regulation of the HPA-axis. Alarin relieves depression by acting as a hormone regulator that downregulates the activity of the HPA axis (CRH, ACTH, and Corticosteroids), and an enzyme inhibitor that decreases the expression of enzymes involved in steroidogenesis (desmolase, 11β-hydroxylase, and aldosterone synthase), thus lowering corticosteroid hormone synthesis. Besides, alarin alleviates depression by reducing HPA-axis overactivity resulting from increased pro-inflammatory cytokines (IL-1α, TNF-α, and IL-6) and decreased anti-inflammatory cytokines (IL-10). Alarin also has antidepressant properties due to its ability to restore hippocampal volume loss, GR resistance/downregulation, prefrontal cortex degradation, and amygdala hyperactivity. Arrow ends (→) indicate a stimulatory effect, while negative signs (–) show an inhibitory effect. ACTH, adrenocorticotropic hormone; CORT, corticosterone; CRH, corticotropin-releasing hormone; GR, glucocorticoid receptor; HPA, hypothalamic-pituitary-adrenal; IL, interleukin; MR, mineralocorticoid receptor; TNF, tumor necrosis factor.

and functions of the co-expressing neurons, may serve as a promising treatment strategy for depression. Importantly, the parent peptide of the galanin family, galanin, has been reported to have remarkable therapeutic effects in depressed patients. The interaction of the serotonergic system and neuropeptides is a key aspect of galanin’s potential for treating major depression in rodents (10, 116, 117).

More interestingly, a large number of studies involving a variety of experimental rodent models have recently revealed that alarin exhibits potent antidepressant-like effects by employing several mechanisms. It has neuroprotective effects and prevent the occurrence of depression by upregulating the expression of TrkB, BDNF, and mTOR, as well as by downregulating the activity of the HPA axis (11, 26–28). As a consequence, it may become possible to derive therapeutic benefits from this new knowledge. Given its growing antidepressant role, alarin could be a peptide of the galanin family, in addition to galanin, that may be taken into consideration in the therapeutic arena of depression. Nowadays, it attracts researchers’ interest toward its potential implications in clinical practice as a treatment option for depression. But clinical studies regarding the pharmacological properties of alarin are still inadequate, warranting the need for further extensive clinical research. A better understanding of the antidepressant mechanisms of alarin may therefore make it possible to use it as an alternative treatment strategy for depression in clinical settings in the future. Overall, alarin has an intriguingly enormous potential for use in neuroscience research and novel drug development for depression, albeit its particular receptor remains to be determined.

Arguments on alarin receptor

A large amount of data has demonstrated that the various physiological effects of the galanin family, such as galanin, the galanin message-associated peptide (GMAP), GALP, and spexin, are generally mediated by three subtypes of GalRs in the G-protein-coupled receptors (GPCR) family, namely GAIr1, Galr2, and Galr3 (118). However, several receptor binding studies unveiled that alarin does not bind to any of the three GalRs (119). A study by Santic et al. reported that synthetic
human alarin could not bind to membrane preparations of human neuroblasts expressing GalR. It has low receptor binding affinity for GalRs, with a dissociation constant (Ki) > 1,000 for GALR1, Ki > 1,000 for GALR2, and Ki > 10⁶ for GALR3 (13). Another study by Boughton et al. on hypothalamic explants further strengthened that alarin has no detectable affinity toward any of the three GalR subtypes (20). The inability of alarin to function via GalRs is proposed to be the result of the exclusion of exon three, leading to a loss of the GalR-binding domain necessary for the ability of other members of the galanin family to bind to GalRs (13). Thus, as alarin does not share any similarity with galanin and is not able to compete with radio-labeled galanin for known GalRs, alarin actions are unlikely to be mediated via such receptors (13, 15, 20).

Instead, the biological actions of alarin seem to be mediated via a distinct unknown receptor different from GalRs. In addition, evidence regarding the physiologic and pharmacologic characteristics of the alarin receptor is still insufficient. However, it has been indicated that alarin appears to mediate its effects through numerous, yet unidentified alarin-specific receptors other than GalRs (13, 19, 20). Fraley et al. demonstrated that unlike GALP, which is mediated by GalRs and is associated with motor deficits after ICV injection, central injection of alarin in mice is linked to the absence of motor impairments, supporting that alarin may act via a separate receptor (120, 121). It is also supported by another study that demonstrates that the alarin antagonist (the so-called Ala6-25Cys), but not the GalR blockers (M35 or C7), reverses the attenuating effects of alarin on myocardial infarction (MI)-induced cardiac fibrosis. Furthermore, this truncated alarin peptide (Ala6-25Cys) that missed the first five amino acids of the full-length alarin restores the inhibitory effects of alarin on angiotensin II-induced cardiac fibrosis (24, 122-124). Ala6-25Cys also antagonizes the acute orexigenic effects of centrally administered alarin (13, 19). This highlights that alarin produces biological effects that are unlikely to be mediated by the known GalRs, but rather by a specific receptor that has not yet been determined (24). But this unidentified alarin-specific receptor may or may not be other subtypes of GalRs that have not yet been known.

Although it is not yet known, many scholars have put forth their hypotheses about the unidentified alarin receptor and its possible category of receptor family. Santic et al. hypothesize that the unknown alarin receptors most likely belong to the GPCR receptor superfamily and that alarin may therefore function similarly to other members of the galanin family by employing GPCR signaling pathways (13). This speculation is based on the fact that the vascular system expresses a significant number of GPCRs and the majority of neuropeptide receptors are members of the GPCR receptor superfamily (13, 119). On the other hand, other studies argue that the receptors for alarin may belong to the broad family of receptor tyrosine kinases (RTKs). This hypothesis stems from the observation that many peptide receptors, including receptors for epidermal growth factor, vascular endothelial growth factor, and insulin, belong to the RTK family, which suggests that the receptor for alarin could belong to the RTKs. Additionally, the involvement of TrkB, an RTK, in alarin-mediated antidepressant-like effects, supports the notion that alarin-specific receptors may be a member of RTKs (11, 70-72).

Furthermore, there is some other information regarding the undiscovered receptor of alarin. Few studies showed unidentified alarin receptor-expressing nuclei in the murine brain, as evidenced by induction of Fos-ir expression after ICV injection of alarin in numerous hypothalamic regions that regulate food intake, body weight, and endocrine hormone secretion (16). Additionally, the existing data indicated that the N-terminal region of the alarin peptide would be crucial for binding with its undiscovered receptor, based on the ground that the first few amino acids of alarin are conserved between rodents and primates (26). Generally, the unknown receptors of alarin pose another challenge in identifying them based on the latest effects in pharmacology and biology as well as the mechanisms underlying these effects. Therefore, unraveling the type of receptor, the physiological activity, and the mechanism of action of alarin remain active areas of future research. The alarin antagonist (Ala6-25Cys), which lacks the first five amino acids of alarin and acts as a specific inhibitor of alarin actions, has been indicated to be a useful tool that could have paramount importance in the search for alarin-specific receptors (19).

**Conclusion and future directives**

Depression is a mood disorder whose etiology has been explained by diverse proposed theories, but none of them has been able to adequately explain the nature of depression. Consequently, the effect of pharmacological modulation of some of the pathophysiological mechanisms of depression as an antidepressant therapy was disappointing. Besides, current widely used antidepressants that target monoamine transporters are reportedly associated with several limitations, like low therapeutic efficacy and side effects. Thus, discovering new targets and alternative antidepressants in depression treatment is required. Presently, multiple studies are ongoing to develop novel therapeutic strategies and replace the existing treatments for depression based on a more in-depth analysis of the behavioral and molecular mechanisms underpinning the disease.

Alarin, which is a 25-amino-acid long multifunctional neuropeptide, has recently gained recognition because its central administration is associated with the modulation of depressive-like behavior in animal models. Accumulated evidence shows that the expression of alarin in the brain, mainly in the hypothalamus and amygdala, is linked to its antidepressant-like effects. This reveals that alarin may potentially be used as a pharmacological agent for the treatment of depression, although
extensive elucidation of its physiological or pharmacological characteristics is required. Alarin might also be of utmost significance in the antidepressant application by centrally altering alarin expression and/or activities using a medication that able to cross the blood brain barrier and modifies alarin's effect. However, the details regarding antidepressant implications of alarin, including the effect of peripheral alarin administration and drugs that potentially modify the effect of alarin centrally, remain active area of research. Additionally, alarin may also be utilized in the future to treat depression that is resistant to standard antidepressant treatments, known as treatment-resistant depression. This could offer a solution to a major therapeutic challenge of treatment-resistant depression and greatly thrill mental health experts.

Nonetheless, data describing the alarin-specific receptors mediating the antidepressant effects and other biological activities are insufficient. A growing body of experimental evidence suggests that alarin is unlikely to bind to any known GalRs due to the absence of the GalR-binding domain. Besides, its particular receptor has not yet been determined, and this may open up a vast area of research to explore the unknown receptors for alarin. Overall, despite its promising therapeutic relevance, knowledge about the antidepressant effects of alarin is limited, so alarin is still in its infancy of use in the treatment of depression. This urges us to conduct more clinical research regarding the effects of alarin in depressed patients and its associated pharmacologic properties.

Author contributions

All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, article search, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agreed to be accountable for all aspects of the work.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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References

1. Cooper J. Diagnostic and statistical manual of mental disorders (4th edn, text revision)(DSM–IV–TR). Br J Psychiatry. (2001) 179:85.
2. Organization WH. Depression and other common mental disorders: global health estimates. Switzerland: World Health Organization (2017).
3. Wittchen H-U, Jacobi F. Size and burden of mental disorders in Europe—a critical review and appraisal of 27 studies. Eur Neuropsychopharmacol. (2005) 15:357–76. doi: 10.1016/j.euroneuro.2005.04.012
4. Murray CJ, Lopez AD. Global mortality, disability, and the contribution of risk factors: global burden of disease study. Lancet. (1997) 349:1436–42.
5. Fava M, Kendler KS. Major depressive disorder. Neuron. (2000) 28:335–41.
6. Kesler RC. The effects of stressful life events on depression. Ann Rev Psychol. (1997) 48:191–214.
7. Millan MJ. The role of monoamines in the actions of established and "novel" antidepressant agents: a critical review. Eur J Pharmacol. (2004) 500:371–84. doi: 10.1016/j.ejphar.2004.07.038
8. Hokfelt T, Barfod T, Bloom F. Neuropeptides: opportunities for drug discovery. Lancet Neurol. (2003) 2:463–72.
9. Holmes A, Heilig M, Rupniak NM, Steckler T, Griebel G. Neuropeptide systems as novel therapeutic targets for depression and anxiety disorders. Trends Pharmacol Sci. (2003) 24:580–8.
10. Ögren SO, Kuteeva E, Hokfelt T, Kehr J. Galanin receptor antagonists. CNS Drugs. (2006) 20:633–54.
11. Wang M, Zhou W, Zhou X, Zhuang F, Chen Q, Li M, et al. Antidepressant-like effects of alarin produced by activation of TrkB receptor signaling pathways in chronic stress mice. Behav Brain Res. (2015) 280:128–40. doi: 10.1016/j.bbr.2014.11.039
12. Santic R, Fenninger K, Graf K, Schneider R, Hauser-Kronberger C, Schilling FH, et al. Gangliocytes in neuroblastic tumors express alarin, a novel peptide derived by differential splicing of the galanin-like peptide gene. J Mol Neurosci. (2006) 29:145–52. doi: 10.1385/JMN:29:2:145
13. Santic R, Schmidhuber SM, Lang R, Rauch I, Voglas E, Eberhard N, et al. Alarin is a vasoactive peptide. Proc Natl Acad Sci USA. (2007) 104:10217–22.
14. Hu W, Fan X, Zhou B, Li L, Tian B, Fang X, et al. Circulating alarin concentrations are high in patients with type 2 diabetes and increased by glucagon-like peptide-1 receptor agonist treatment: an consort-compliant study. Medicine. (2019) 98:28. doi: 10.1097/MD.0000000000016428
15. Van Der Kolk N, Madison FN, Mohr M, Eberhard N, Koffer B, Fraley GS. Alarin stimulates food intake in male rats and LH secretion in castrated male rats. Neuropeptides. (2010) 44:333–40. doi: 10.1016/j.npep.2010.04.001
16. Fraley GS, Leathley E, Lundy N, Chheng E, King I, Koffer B. Effects of alarin on food intake, body weight and luteinizing hormone secretion in male mice. Neuropeptides. (2012) 46:99–104.
17. Eberhard N, Mayer C, Santic R, Navio RP, Wagner A, Bauer HC, et al. Distribution of alarin immunoreactivity in the mouse brain. J Mol Neurosci. (2012) 46:18–32. doi: 10.1007/s12031-011-9546-y
18. Eberhard N, Wei S, Reitsamer H, Koffer B. Expression of alarin in ependymoma and choroid plexus tumors. J Neuro Oncol. (2013) 114:165–71. doi: 10.1007/s11060-013-1177-4
19. Fraley GS, Leathley E, Nickols A, Gerometta E, Coombs E, Colton S, et al. Alarin 6-25Cys antagonizes alarin-specific effects on food intake and luteinizing hormone secretion. Neuropeptides. (2013) 47:37–41. doi: 10.1016/j.npep.2012.08.007
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Frontiers in Psychiatry 10 frontiersin.org

study. unmedicated major depressive disorder patients: a single voxel 1H spectroscopy et al. Normal metabolite levels in the left dorsolateral prefrontal cortex of patients. resonance spectroscopy study of thalamus in treatment resistant depressive Arch General Psychiatry. 1017/S1461145704004080 monoaminergic neurons: a case for primary implication of the locus coeruleus. 12:2–19. Neuropsychopharmacology. network function in the hippocampus by activating nongenomic glucocorticoid receptors are associated with the antidepressant-like effect of 31:2857–66. doi: 10.1016/j.neuroscience.2001.01.002 33. Maggio J. Stress-induced plasticity of GABAergic inhibition. Front Cell Neurosci. (2014) 8:157. doi: 10.3389/fncel.2014.00157 34. Hu W, Zhang M, Czéh B, Flügge G, Zhang W. Stress impairs GABAergic and depression-like behaviour. J Neurosci. (2009) 35:1693–707. doi: 10.1016/j.neuroscience.2009.05.003 35. Schildkraut JJ. The catecholamine hypothesis of affective disorders: a review of theories and new developments. J Affect Dis. (2008) 114:1017–31. doi: 10.1016/j.ap prévu.142.9.1017 36. Berman RM, Cappiello A, Anand A, Oren DA, Heninger GR, Charney DS, et al. Antidepressant effects of ketamine in depressed patients. Nat Med. (2004) 10:575–82. doi: 10.1038/nm1195 37. Kuteeva E, Wardi T, Hökfelt T, Ogren S. Galanin enhances and alters synaptic density is associated with depression severity and network alterations. Nat Commun. (2019) 10:1–10. doi: 10.1038/s41467-019-09626-7 38. De Jong J, Linnoila M. Cerebrospinal fluid monoamine metabolites in treatment resistant depressive patients. Nat Neurosci. (2009) 12:2–19. Neuropsychopharmacology. glutocorticoid receptors are associated with the antidepressant-like effect of 1HmTOR signaling and synaptic plasticity. Behav Brain Res. (2016) 313:158–71. doi: 10.1016/j.bbr.2016.06.057 39. Åsberg M, Träskman L, Thorén P. 5-HIAA in the cerebrospinal fluid: a study of untreated depression and hippocampal volume loss. Am J Psychiatry. (2003) 160:1516–8. 40. Roy A, De Jong J, Linnola M. Cerebrospinal fluid monoamine metabolites and suicidal behavior in depressed patients: a 5-year follow-up study. Arch General Psychiatry. (1989) 46:609–12. 41. Maes M. The serotonin hypothesis of depression. Psychopharmacol Fourth Genet Preg. (1995) 1995:933–44. 42. Albert PR, Lemonde S. 5-HT1A receptors, gene expression, and depression: guilt by association. Neurosci. (2004) 10:575–93. doi: 10.1017/S0033875804426378 43. Martinowich K, Manji H, Lu B. New insights into BDNF function in depression and anxiety. Nat Neurosci. (2007) 10:1089–93. 44. Hasler G. Pathophysiology of depression: do we have any solid evidence of interest to clinicians? World Psychiatry. (2010) 9:155. doi: 10.1001/2005.5545.10.0b0298x 45. Sheline YI, Gado MH, Kramer HC. Untreated depression and hippocampal volume loss. Am J Psychiatry. (2003) 160:1516–8. 46. Duhan RS, Aghajanian GK, Sanacora G, Krystal JH. Synaptic plasticity and depression: new insights from stress and rapid-acting antidepressants. Nat Med. (2016) 22:338–49. 47. Holmes SE, Scheinot D, Fennema SJ, Nagamana M, Davis MT, Dell’Anno G, et al. Lower synaptic density is associated with depression severity and network alterations. Nat Commun. (2019) 10:1–10. doi: 10.1038/s41467-019-09626-7 48. Mody J, Maguire J. The reciprocal regulation of stress hormones and GABA receptors. Front Cell Neurosci. (2012) 6:4. doi: 10.3389/fncel.2012.00004 49. Acosta GR, Losada MEO, Rubio MC. Area-dependent changes in GABAergic function after acute and chronic cold stress. Neurosci Lett. (1993) 154:175–78. doi: 10.1016/0304-3940(93)90205-5 50. Trullas R, Skolnick P. Functional antagonists at the NMDA receptor complex exhibit antidepressant actions. Eur J Pharmacol. (1990) 185:1–10. 51. Brown ES, Varghese FP, McElwen BS. Association of depression with medical illness: does cortisol play a role? Biol Psychiatry. (2004) 55:1–9. 52. Hennings JM, Owashi T, Binder EB, Horstmann S, Menke A, Kloiber S, et al. Clinical characteristics and treatment outcome in a representative sample of depressed inpatients—findings from the Munich antidepressant response signature (MARS) project. J Psychiatric Res. (2009) 43:215–29. doi: 10.1016/j.jpsychires.2008.05.002 53. Holdboer F. Stress, hypercortisolism and corticosteroid receptors in depression: implications for therapy. J Affect Dis. (2001) 62:77–91. doi: 10.1016/S0165-0327(00)00352-9 54. Variante CM, Lightsman SL. The HPA axis in major depression: classical theories and new developments. Trends Neurosci. (2008) 31:464–8. doi: 10.1016/j.tins.2008.06.006 55. Kuteeva E, Hökfelt T, Wardi T, Ogren SO. Galanin, galanin receptor subtypes and depression-like behaviour. Cell Mol Life Sci CMLS. (2008) 65:1854–63. 56. Kuteeva E, Wardi T, Hökfelt T, Ogren SO. Galanin enhances and a galanin antagonist attenuates depression-like behaviour in the rat. Eur Neuropsychopharmacol. (2007) 17:64–9. doi: 10.1016/j.euroneuro.2006.03.003 57. McLaughlin PJ, Robinson JK. Galanin: involvement in behavior and neuroendocrinology, and therapeutic potential. Drug News Perspect. (2002) 15:647–55. doi: 10.1385/dnp.2002.15.8040200 58. Saar I, Labe J, Langel K, Runesson J, Webling K, Jarv J, et al. Novel systemically active galanin receptor 2 ligands in depression-like behaviour. J Neurochem. (2013) 127:114–23. doi: 10.1111/jnc.12274 59. Gendlich AL, Furazin T, Larm J. Distribution, regulation and role of hypothalamic galanin systems: renewed interest in a pleiotropic peptide family. Clin Exp Pharmacol Physiol. (2001) 28:100–10. doi: 10.1046/j.1440-1681.2001.03411.x 60. Davidson S, Lear M, Stanley L, Hing B, Baizan-Edge A, Herwig A, et al. Differential activity by polymorphic variants of a remote enhancer that supports galanin expression in the hypothalamus and amygdala: implications for obesity, depression and alcoholism. Neuropsychopharmacology. (2011) 36:2211–21. doi: 10.1038/npp.2011.9 61. Lu X, Sharkley L, Barlton T. The brain galanin receptors: targets for novel antidepressant drugs. CNS Neurol Dis Drug Targets. (2007) 6:183–92. 62. Lu X, Barr AM, Kinney JW, Sanna P, Conti B, Behrens MM, et al. A role for galanin in antidepressant actions with a focus on the dorsal raphe nucleus. Proc Natl Acad Sci. (2005) 102:874–9. doi: 10.1073/pnas.0408891102
89. Dwyer JM, Lepack AE, Duman RS. mTOR activation is required for the antidepressant effects of mGluR2/3 blockade. *Int J Neuropsychopharmacol.* (2012) 15:429–34. doi: 10.1016/j.i jnp.2011.07.002

90. Zheng W, Zhou Y-L, Liu W-J, Wang C-Y, Zhan Y-N, Li H-Q, et al. Rapid and longer-term antidepressant effects of repeated-dose intravenous ketamine for patients with unipolar and bipolar depression. *J Psychiatric Res.* (2018) 106:61–8. doi: 10.1016/j.jpsychires.2018.09.013

91. Masi G, Brovedani P. The hippocampus, neurotransf actors and depression. *CNS Drugs.* (2011) 25:913–31.

92. Calabrese F, Molteni R, Racagni G, Riva MA. Neurol nal plasticity: a link between stress and mood disorders. *Psychoneuroendocrinology.* (2009) 34:5208–16.

93. Picciotto MR, Brabant C, Einstein EB, Ramens HM, Neugebauer NM. Effects of galanin on monoaminergic systems and HPA axis: potential mechanisms underlying the effects of galanin on addiction-and stress-related behaviors. *Brain Res.* (2010) 1314:206–18. doi: 10.1016/j.brainres.2009.08.033

94. Pursolt RD, Le Pichon M, Jalife M. Depression: a new animal model sensitive to antidepressant treatments. *Nature.* (1977) 266:730–2.

95. Prut L, Belzung C. The open field as a paradigm to measure the effects of drugs on anxiety-like behaviors: a review. *Eur J Pharmacol.* (2003) 465:3–33. doi: 10.1016/s0014-2999(03)01272-x

96. Duman RS, Monteggia LM. A neurotrophic model for stress-related mood disorders. *Biol Psychiatry.* (2006) 59:1116–27.

97. Li N, He X, Zhang Y, Qi X, Li H, Zhu X, et al. Brain-derived neurotrophic factor signalling mediates antidepressant effects of lamotrigine. *Int J Neuropsychopharmacol.* (2011) 14:1091–8. doi: 10.1016/s1474-7044(10)00182-P

98. Unsuld PG, Ising M, Roeske D, Erhardt A, Specht M, Kloiber S, et al. Gender-specific association of galanin polymorphisms with HPA-axis dysregulation, symptom severity, and antidepressant treatment response. *Neuropsychopharmacology.* (2010) 35:1583–92. doi: 10.1038/npp.2010.30

99. Rao U, Hammen CL, Poland RE. Mechanisms underlying the comorbidity between depressive and addictive disorders in adolescents: interactions between stress and HPA activity. *Am J Psychiatry.* (2009) 166:361–9. doi: 10.1176/appi.ajp.2008.08030412

100. Ulrich-Lai YM, Herman JP. Neural regulation of endocrine and autonomic stress responses. *Nat Rev Neurosci.* (2009) 10:397–409.

101. Mizoguchi K, Ishige A, Aburada M, Tabira T. Chronic stress attenuates glucocorticoid negative feedback: involvement of the prefrontal cortex and hippocampus. *Neuroscience.* (2003) 119:887–97.

102. Modell S, Yassouridis A, Huber J, Holshofer F. Corticosteroid receptor function is decreased in depressed patients. *Neuroendocrinology.* (1997) 65:216–22.

103. Young EA, Lopez JF, Murphy-Weinberg V, Watson SJ, Akil H. Mineralocorticoid receptor function in major depression. *Arch General Psychiatry.* (2003) 60:24–8.

104. Anacker C, Zunszain PA, Carvalho LA, Pariante CM. The glucocorticoid receptor: pivot of depression and of antidepressant treatment? *Psychoneuroendocrinology.* (2011) 36:415–25. doi: 10.1016/j.psyneuen.2010.03.000

105. Lee M-S, Park W-S, Kim YH, Kwon S-H, Jang Y-I, Han D, et al. Antidepressant-like effects of cortex mori radicis extract via bidirectional phosphorylation of glucocorticoid receptors in the hippocampus. *Behav Brain Res.* (2013) 236:56–61. doi: 10.1016/j.bbr.2012.08.028

106. Johnson SA, Fourrier NM, Kalynchuk LE. Effect of different doses of corticosterone on depression-like behavior and HPA axis responses to a novel stressor. *Behav Brain Res.* (2006) 168:280–8. doi: 10.1016/j.bbr.2005.11.019

107. Horstmann S, Dose T, Lucase S, Kloiber S, Menke A, Hennings J, et al. Suppressive effect of mirtazapine on the HPA system in acutely depressed women seems to be transient and not related to antidepressant action. *Psychoneuroendocrinology.* (2009) 34:238–48. doi: 10.1016/j.psyneuen.2008.09.004

108. Espallergues J, Givalois L, Temsamani J, Laruelle C, Maurice T. The 3β-hydroxysteroid dehydrogenase inhibitor trilostane shows antidepressant properties in mice. *Psychoneuroendocrinology.* (2009) 34:644–59. doi: 10.1016/j.psy neuuen.2008.11.003

109. Pace TW, Hu F, Miller AH. Cytokine-effects on glucocorticoid receptor function: relevance to glucocorticoid resistance and the pathophysiology and treatment of major depression. *Brain Behav Immun.* (2007) 21:9–19. doi: 10.1016/j.bbi.2006.08.009
110. Sapolsky R, Rivier C, Yamamoto G, Plotsky P, Vale W. Interleukin-1 stimulates the secretion of hypothalamic corticotropin-releasing factor. *Science*. (1987) 238:522–4.

111. Besedovsky H, Del Rey A, Klausman I, Furukawa H, Arditi GM, Kabiersch A. Cytokines as modulators of the hypothalamic-pituitary-adrenal axis. *J Steroid Biochem Mol Biol*. (1991) 40:613–8.

112. Antonioli M, Rybka J, Carvalho LA. Neuroimmune endocrine effects of antidepressants. *Neuropsychiatric Dis Treat*. (2012) 8:65.

113. Dantzer R, O’connor JC, Freund GG, Johnson RW, Kelley KW. From inflammation to sickness and depression: when the immune system subjugates the brain. *Nat Rev Neurosci*. (2008) 9:46–56. doi: 10.1038/nrm2297

114. Sulser F. New perspectives on the molecular pharmacology of affective disorders. *Eur Arch Psychiatry Neurol Sci*. (1989) 238:231–9.

115. Vetulani J, Sulser F. Action of various antidepressant treatments reduces reactivity of noradrenergic cyclic AMP-generating system in limbic forebrain. *Nature*. (1975) 257:495–6. doi: 10.1038/257495a0

116. Millon C, Flores-Burgess A, Narváez M, Borroto-Escuela DO, Santín L, Gago B, et al. Galanin (1–15) enhances the antidepressant effects of the 5-HT1A receptor agonist 8-OH-DPAT: involvement of the raphe-hippocampal 5-HT neuron system. *Brain Struct Funct*. (2016) 221:4491–504. doi: 10.1007/s00429-015-1180-y

117. Jackowski A, Crockard A, Burnstock G, Lincoln J. Alterations in serotonin and neuropeptide Y content of cerebrovascular sympathetic nerves following experimental subarachnoid hemorrhage. *J Cerebral Blood Flow Metab*. (1989) 9:271–9. doi: 10.1038/jcbfm.1989.45

118. Webling KE, Runesson J, Bartfai T, Langel U. Galanin receptors and ligands. *Front Endocrinol*. (2012) 3:146. doi: 10.3389/fendo.2012.00146

119. Šípková J, Kramáriková J, Hynie S, Klenerová V. The galanin and galanin receptor subtypes, its regulatory role in the biological and pathological functions. *Physiol Res*. (2017) 66:5.

120. Kaufman AS, Buentle J, Fraley GS, Rissman EF. Effects of galanin-like peptide (GALP) on locomotion, reproduction, and body weight in female and male mice. *Hormones Behav*. (2005) 48:141–51. doi: 10.1016/j.yhbeh.2005.01.010

121. Krasnow SM, Fraley GS, Schuh SM, Baumgartner JW, Clifton DK, Steiner RA. A role for galanin-like peptide in the integration of feeding, body weight regulation, and reproduction in the mouse. *Endocrinology*. (2003) 144:813–22. doi: 10.1210/en.2002-220982

122. Wang Q, Deng F, Zhu D. Superoxide anions modulate the effects of alarin in the paraventricular nucleus on sympathetic activity and blood pressure in spontaneously hypertensive rats. *Neuropeptides*. (2020) 80:102021. doi: 10.1016/j.npep.2020.102021

123. Zhang Y, Gao Y, Li C-Y, Dong W, Dong Y, Li M-N, et al. Galanin receptor 1 plays an antinociceptive effect via inhibiting PKA activation in the nucleus accumbens of rats with neuropathic pain. *Physiol Res*. (2019) 68:511–8. doi: 10.33549/physiolres.933941

124. Bhandari M, Kawamoto M, Thomas A, Barreto S, Schlothe A, Carati C, et al. Galanin receptor antagonist m35 but not m40 or c2 ameliorates cerebral-induced acute pancreatitis in mice. *Pancreatology*. (2010) 10:682–8.