Deficits in KCC2 and activation of the HPA axis lead to depression-like behavior following social defeat

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Abstract
Background: Chronic social stress triggers the development of major depression in humans and depression-like behavior in animal models. Hyperexcitability of the hypothalamic-pituitary-adrenal (HPA) axis has been implicated in the pathophysiology of depression. The HPA axis is tightly controlled by GABAergic inhibition at the level of corticotrophin-releasing hormone (CRH) neurons in the paraventricular nucleus (PVN) of the hypothalamus. Recently, our lab has demonstrated plasticity in GABAergic control of the HPA axis following stress, involving downregulation of the K⁺/Cl⁻ co-transporter 2 (KCC2) which is critical for the inhibitory actions of GABA. The purpose of this study was to determine if chronic social defeat stress activates the HPA axis using similar mechanisms which may contribute to the development of depression-like behavior.

Methods: The activation of the HPA axis following social defeat was quantified by measuring circulating levels of corticosterone (CORT). The role of KCC2 in activation of the HPA axis following social defeat was assessed by measuring KCC2 levels in the PVN by Western blot analysis. The impact of HPA axis activation on depression-like behavior was examined using the forced swim test. The therapeutic potential of blocking the activation of the HPA axis for depression-like behavior was determined by pharmacological blockade of the HPA axis activation, using Antalarmin.

Results: Here, we demonstrate that chronic social defeat stress increased plasma levels of CORT and depression-like behavior in submissive mice. The activation of the HPA axis in submissive mice following chronic social defeat stress was associated with a dephosphorylation and downregulation of KCC2 in the PVN, which has previously been demonstrated to play a key role in mounting the physiological response to stress. Elevations in corticosterone levels and the development of depression-like behavior were restricted to submissive mice and were not observed in dominant animals. Treatment of dominant mice with exogenous corticosterone induced submissive behaviors and depression-like behaviors in these animals. In addition, blocking CRH signaling with Antalarmin prevented the social stress-induced development of depression-like behavior in submissive mice.

Conclusions: Our study suggests that plasticity in the GABAergic regulation of the HPA axis may underlie elevations in corticosterone following chronic social defeat and that blocking the activation of the HPA axis may have therapeutic potential to combat social stress-induced depression.

Keywords: Stress, social defeat, KCC2, GABA, depression, corticotropin-releasing hormone (CRH), corticosterone

Introduction
Major depressive disorder is one of the most prevalent mood disorders, affecting approximately 6.7% of the population at any one time [1] and up to 20% of the population at some point in their lives [2]. Depression is associated with numerous long-term, adverse consequences on health, longevity, and quality of life [2-7]. Chronic social and psychological stress trigger the development of depression in both humans [8-9] and animal models [9-11]. For example, bullying in both children at school and in adults in the workplace is extremely stressful for the victims and causes many negative consequences, including depression [12-17]. However, the mechanism(s) through which stress leads to depression remains unclear.

Although the pathophysiological causes of depression are undoubtedly varied [2,18-21], it is clear that major depression is associated with stress and hyperactivity of the hypothalamic-pituitary-adrenal (HPA) axis [22,23]. Dysregulation of the HPA axis precedes the development of depression and is implicated in mediating the effects of stress on depression-like behavior. Consistent with the role of the HPA axis in depression, antidepressant treatment normalizes the activity of the HPA axis in depressed patients (for review see [26]), which precedes the therapeutic effects (for review see [26]).

Social, and other types of, stressors stimulate the release of corticotropin-releasing hormone (CRH) from the hypothalamus, which then acts in the pituitary gland to signal the release of adrenocorticotropic hormone (ACTH) to elicit the release of cortisol from the adrenal cortex in humans or corticosterone in rodents (CORT) [27-31]. Chronic stress leads to dysregulation of the HPA axis and increased baseline levels of CORT [28-32], which has been proposed to play a role in the pathophysiological mechanisms underlying depression. However, the mechanism(s) through which the HPA axis becomes dysregulated following chronic stress is unknown but understanding these mechanisms will likely provide insight into the pathophysiology of depression. The HPA axis is tightly regulated by GABAergic inhibition at
the level of the CRH neurons in the paraventricular nucleus (PVN) of the hypothalamus [28,33-35]. The inhibitory actions of GABA on CRH neurons requires the maintenance of the chloride gradient [36-42], a task primarily accomplished by the potassium chloride cotransporter 2 (KCC2) in the adult brain [36,37,41-43]. Recent studies demonstrated a collapse in the chloride gradient in CRH neurons following acute restraint stress due to dephosphorylation of KCC2 residue Ser940 and downregulation of KCC2 in the PVN [44], which led to decreased GABAergic control of the HPA axis [44,45]. KCC2 residue Ser940 controls the surface expression and function of KCC2 [46]. Thus, dephosphorylation of KCC2 residue Ser940 following acute stress likely has a significant impact on maintaining the chloride gradient in these neurons. Although these studies defined the role of KCC2 in GABAergic control of the HPA axis following acute stress [44], it remains unclear how these regulatory mechanisms controlling the HPA axis are altered following chronic stress and if dysregulation may contribute to the development of depression-like behavior.

Here we utilized the chronic social defeat paradigm to investigate the role of the HPA axis in the development of depression-like behavior. The chronic social defeat stress paradigm mimics bullying behavior in humans and leads to depression-like behavior in defeated animals [10,12,47]. We hypothesized that plasticity in the GABAergic regulation of the HPA axis may be the mechanism through which the HPA axis becomes activated following social defeat and may contribute to the development of depression-like behavior. Our data demonstrate that chronic social defeat caused dephosphorylation and downregulation of KCC2 in the PVN, which was associated with activation of the HPA axis and depression-like behavior in submissive mice. We demonstrate that exogenous administration of CORT was sufficient to induce depression-like behavior in previously dominant mice. Furthermore, blocking the activation of the HPA axis with the CRH antagonist, Antalarmin, prevented the development of depression-like behavior that was associated with chronic social defeat stress. These findings suggest that blocking the physiological response to stress may be a therapeutic target for depression.

Methods

Animal housing

Male 10- and 6-week-old C57BL/6 mice were purchased from Jackson Laboratory and housed at the Tufts University School of Medicine, Division of Laboratory Animal Medicine. Mice were housed in a temperature- and humidity-controlled environment with a 12 h light/dark cycle (lights on at 0700 h) with food and water available ad libitum. Mice were housed in our colony for at least 5 days before delivery for acclimation after transport prior to experimentation. Separate groups of animals were used for each experiment unless otherwise noted. Animals were handled according to protocols approved by the Tufts University Institutional Animal Care and Use Committee.

Chronic social defeat stress

Ten-week-old mice for dominant groups were separated and singly-housed 3-5 days before experimental procedures. Mice that were consistently aggressive when paired with 6-week-old mice were selected as dominants. Naïve mice (10 weeks of age) were then individually introduced into the home cage of a dominant mouse. After a 10 minute interaction, the mice were physically separated and housed in custom cages with a perforated divider (Ancare) so the naïve, now submissive mice, were protected from physical interaction but could still see, smell, and hear the dominant mice. If the animals became injured during the interaction, the test was immediately stopped, the animals separated, and excluded from further experimentation. Thereafter, every day for a total of 14 days, the divider was removed for a 10 minute social defeat interaction (see Figure 1 for paradigm schematic and timeline). Prior to social defeat each day, submissive mice were switched to a new dominant’s cage to avoid habituation to the same dominant animal. Social defeat interactions took place between 1300 and 1700 h daily. Interactions were either observed and scored live or videotaped and scored later for attack latency, number of attacks, and number of tail rattles to determine dominance relationships. Dominant mice were chosen which defeated 100% of the submissive mice. In contrast, submissive mice were chosen in which they never defeated a dominant mouse (0% defeats). Control animals were singly-housed in the social defeat cages and during the experimental groups' daily social defeat, controls had their divider removed for 10 minutes and then replaced. Dominant and control groups were not significantly different in either their forced swim behavior or plasma CORT levels (Figure 2).

![Figure 1. Schematic representation of the chronic social defeat stress paradigm.](image)

Dominant mouse singly-housed after training. Submissive introduced, 10 min interaction. Submissive housed with visual, auditory and olfactory cues. Social Defeat for 14 Days.

Dominant mice (10-weeks-old) were selected out based upon dominance when paired with 6-week-old mice. These dominant mice were singly-housed after training and individual naïve mice (10-weeks-old) were introduced into the dominants’ homecage. The mice were allowed to interact for 10 mins per day for 14 days and were housed together in custom cages with a perforated divider so that the mice could see, smell, and hear one another but were physically separated by a barrier. During the 10 min interaction, the dominance relationship was evaluated.
Figure 2. Chronic social defeat activated the HPA axis and induced depression-like behavior.
(A) The average plasma CORT levels after chronic social defeat were significantly elevated in submissive mice compared to dominant mice and controls. There was no significant difference in plasma CORT levels between dominant and control mice. (n=8-10 mice per experimental group). (B) The average total time spent immobile during the forced swim test was increased in submissive mice compared to dominant mice and controls. There was no significant difference in the total time spent immobile in the forced swim test between dominant and control mice. (n=7-10 mice per experimental group).

*indicates statistical significance of p<0.05 using a one-way ANOVA with Tukey’s post hoc test.

Corticosterone measurements
Pairs of dominant and submissive mice were briefly anesthetized with isoflurane, decapitated using a guillotine, and trunk blood was collected 30 minutes following subjection to the final social defeat interaction. For the exogenous corticosterone experiments, corticosterone levels were measured 24 hours after the forced swim test. The plasma was isolated from the trunk blood by centrifugation at 14,000 rpm for 5 minutes and then stored at -20°C until use. CORT levels were measured by enzyme immunoassay according to manufacturer’s specifications (Enzo Life Sciences). Briefly, plasma samples were assayed in duplicate and absorbance was measured at 415 nM and compared to a standard curve. Samples from different experimental groups were run in parallel.

Forced swim test
Thirty minutes following the final social defeat interaction, pairs of dominant and submissive mice were placed into separate, adjacent clear plastic containers 22 cm in diameter, which were filled with approximately 14 cm of 24°C water, for 6 minutes. Containers were cleaned and refilled with fresh water between subjects. After the 6 min forced swim test, animals were removed from the containers, gently patted dry, and then returned to their cages. Sessions were videotaped and then later blindly scored for duration of time spent immobile. A subject was considered to be immobile when it was floating in the water without struggling and using only the small movements necessary to keep its head above water. Testing took place between 1000 h and 1700 h, with groups being counterbalanced to control for time of day.

Corticosterone pellet implantation
Mice that were trained as dominants were briefly anesthetized using 2-3% isoflurane then anesthesia was maintained via a nose cone with 1.5-2% isoflurane during the implantation surgery. The back of each subject’s head and neck was shaved, cleaned, and sterilized using ethanol and betadine. A small, approximately 1 cm incision was made in the skin on the back of the neck, and a CORT pellet (10 mg/pellet, 21-day release, Innovative Research of America, Sarasota, FL, Catalogue No. G-111,) was implanted (CORT dominant group). An equal number of mice were sham implanted as controls (sham dominant group). Subjects were allowed to recover for 5 days before further experimentation. Following recovery, subjects underwent the chronic social defeat paradigm as dominants.

Note that not all naïve animals introduced into CORT dominants’ cages were consistently defeated, in contrast to the sham implanted mice, but will be referred to as submissives for consistency and clarity. Thirty minutes after the last social defeat, mice were tested in the forced swim test. Whole blood was collected for analysis of plasma corticosterone levels 24 hours after the forced swim test.

Antalarmin injections
Submissive mice received daily i.p. injections of either 10 mg/kg Antalarmin (Sigma, A8727) or vehicle (5% ethanol and 5% cremophor, in saline) 30 minutes prior to social defeat throughout the 14 day social defeat paradigm. Antalarmin- and vehicle-treated submissive mice were tested along with dominant animals in the forced swim test 30 min after their final social defeat on Day 14.

Western blot
Western blot analysis was conducted as previously described
Chronic social defeat stress activated the HPA axis
To determine if chronic social defeat alters HPA axis responsiveness, we measured circulating CORT 30 min following the last social defeat interaction. Submissive mice showed significantly increased plasma CORT levels (66.50±10.15 ng/ml) compared to either dominant mice (29.41±4.96 ng/ml) or control mice (21.07±4.83 ng/ml) (Figure 2A). Interestingly, there was no significant difference in the CORT levels between dominant (29.41±4.96 ng/ml) and control mice (21.07±4.83 ng/ml) (Figure 2A), suggesting that HPA axis reactivity is only altered in submissive mice (n=8-10 mice per group; F(2,25)=3.39; p<0.05). The mechanism(s) underlying activation of the HPA axis in submissive mice is unclear, but likely directly related to the stress of the social defeat interaction. Previous studies in our lab demonstrated a role for dephosphorylation and downregulation of KCC2 in activation of the HPA axis following acute stress [44]. To determine if this same regulatory mechanism plays a role in dysregulation of the HPA axis following chronic social defeat stress, we examined the total levels of KCC2 and the phosphorylation of KCC2 residue Ser940 in the PVN of dominant and submissive mice using Western blot analysis. We did not observe a difference in the expression of KCC2 or phosphorylation of KCC2 Ser940 between control and dominant mice. Therefore, dominant mice were used for comparison. Submissive mice had significantly decreased expression of total KCC2 in the PVN (93.59±2.28 O.D. units/50µg total protein) compared to dominant mice (104.41±4.25 O.D. units/50µg total protein) (Figure 3, Left) (n=10 mice per group; t(18)=2.24; p<0.05). In addition, the phosphorylation of KCC2 residue Ser940, which regulates the expression and function of KCC2 [46], was significantly decreased in the PVN of submissive mice (38.77±6.58 O.D. units/50µg total protein) compared to dominant mice (58.69±5.47 O.D. units/50µg total protein) (Figure 3, Right), which likely compromises KCC2 function [46] (n=10 mice per group; t(18)=2.33; p<0.05). However, we did not observe any difference in the expression of β-tubulin in the PVN, used as a loading control, between dominant (84.7±3.1 O.D. units/50µg total protein) compared to submissive mice (80.8±2.0 O.D. units/50µg total protein) (n=10 mice per group; t(18)=1.06; p=0.29). These data demonstrate increased activation of the HPA axis only in submissive mice following chronic social defeat stress and may utilize regulatory mechanisms similar to those following acute restraint stress [44].

Chronic social defeat stress induced depression-like behavior
To determine the impact of social defeat stress on depression-like behavior, we assessed the total time spent immobile during the forced swim test in dominant, submissive, and control mice following subjection to 14 days of chronic social defeat. Submissive mice spent significantly more time immobile in the forced swim test (198.00±10.77 s) compared to dominant mice (152.60±9.53 s) and control mice (156.27±9.81 s) (Figure 2B), suggesting increased depression-like behavior in submissive mice. Interestingly, dominant mice (152.60±9.53 s) did not differ from control mice (156.27±9.81 s), similar to the results we observed with circulating CORT levels (Figure 2B) (n=7-10 mice per group; F(2,28)=3.34; p<0.05). These data suggest that depression-like behavior is altered in the submissive mice, not the dominant mice. Furthermore, increased depression-like behavior in submissive mice is associated with elevated levels of CORT.

Exogenous CORT was sufficient to induce submissive and depression-like behavior
As we found significantly elevated levels of plasma CORT in submissive mice following chronic social defeat stress, we wanted to determine whether elevated CORT itself contributes...
to submissive and depression-like behavior. To evaluate the effect of CORT on submissive behavior, we compared the social defeat behavior of dominant mice that received subcutaneous CORT pellets (CORT dominants) to dominant mice receiving a sham surgery (sham dominants) and submissive mice. To ensure that the CORT pellets successfully elevated circulating levels of CORT, we compared basal plasma corticosterone levels of CORT dominants to sham dominants and submissives 24 hours after the forced swim test. CORT levels were significantly elevated in CORT dominants (57.78±6.40 ng/ml) compared to sham dominants (17.57±1.27 ng/ml) (data not shown) (n=10 mice per group; p<0.05). Importantly, the levels of CORT in CORT dominants (57.78±6.40 ng/ml) are similar to what we observed in submissive mice 30 minutes after their final social defeat (66.50±10.15 ng/ml). To evaluate the effects of exogenous CORT on behavior, we assessed both the degree of dominance and depression-like behavior. We found that out of the 14 days tested, CORT dominants defeated submissive mice significantly less often (4.30±1.41 days dominant) than sham dominants (10.00±1.44 days dominant) (Figure 4A) (n=10 mice per group; p<0.05). In fact, CORT dominants (4.30±1.41 days dominant) were not significantly more dominant than submissive mice (0.80±0.29 days dominant) (Figure 4A). In contrast, sham dominants were significantly more often dominant (10.00±1.44 days dominant) than submissives (0.80±0.29 days dominant) (Figure 4A) (n=10 mice per group; F(2,27)=3.35; p<0.05). These data suggest that exogenous CORT is sufficient to abolish dominant behavior. In order to determine whether exogenous CORT affected depression-like behavior in dominant mice, we examined the total time spent immobile in the forced swim test in CORT dominants, sham dominants, and submissive mice. CORT dominants spent significantly more time immobile in the forced swim test (192.80 sec±6.99) compared to sham dominants (159.60±8.49 s) (Figure 4B), suggesting that CORT treatment induced depression-like behavior in dominant mice. Interestingly, the total time spent immobile was not significantly different between CORT dominants (192.80±6.99 s) and submissives (201.10±5.74 s) (Figure 4B). In contrast, sham dominants
spent significantly less time immobile (159.60±8.49 s) than submissive mice (201.10±5.74 s) (Figure 4B) (n=10 mice per group; F(2,27)=3.35; p<0.05). These data suggest that exogenous CORT is sufficient to increase depression-like behavior in dominant mice.

**Blocking activation of the HPA axis prevented the development of stress-induced depression-like behavior**

To test the therapeutic potential of blocking CRH signaling in treating or preventing social stress-induced depression, we treated submissive mice with i.p. injections of either the CRH receptor antagonist Antalarmin (Antalarmin submissives) or vehicle (vehicle submissives) 30 minutes prior to social defeat every day. Antalarmin submissives had decreased CORT levels (39.9±3.3 ng/ml) compared to vehicle submissives (86.50±14.8 ng/ml). Depression-like behavior was then assessed in dominants, Antalarmin submissives, and vehicle submissives using the forced swim test. Vehicle-treated submissive mice spent significantly more time immobile (220.00±2.95 s) than dominants (186.90±3.90 s) (Figure 5), similar to untreated submissive mice (Figure 2B). In contrast, Antalarmin submissives spent significantly less time immobile (199.13±8.18 s) than vehicle submissives (220.00±2.95 s) (Figure 5) (n=14–15 mice per group; F(2,55)=10.18; p<0.05), suggesting that Antalarmin treatment decreased depression-like behavior in submissive mice. In fact, the amount of time Antalarmin submissives spent immobile (199.13±8.18 s) was not significantly different from dominants (186.90±3.90 s). These data indicate that Antalarmin significantly blocked the development of depression-like behavior in submissive mice.

**Discussion**

In the present study, we demonstrated that chronic social defeat stress (Figure 1) increased plasma levels of corticosterone and induced depression-like behavior (Figure 2). Activation of the HPA axis following chronic social defeat stress likely utilizes regulatory mechanisms similar to activation of the HPA axis following acute stress, namely dephosphorylation and downregulation of KCC2 (Figure 3). Exogenous CORT was sufficient to induce submissive behavior and to reproduce the depression-like behavior observed following chronic social defeat stress (Figure 4). Further, blocking the HPA axis prevented the social stress-induced development of depression-like behavior (Figure 5), suggesting that targeting the HPA axis may have therapeutic potential.

Our data demonstrate that chronic social defeat stress (Figure 1), an animal model of bullying and social stress in humans [12,47,48], increased corticosterone levels and induced depression-like behavior (Figure 2). These findings are consistent with previous studies using this paradigm [32,49-54], and confirm this as a useful model to examine how chronic social stress leads to depression. It is widely accepted that depression generally results from an interaction between genetic or developmental predispositions and environmental stress [2,20,22,24,31,55,56]. Thus, we chose a social stress paradigm for this study due to its translational relevance, as many of the stresses humans encounter are social in nature [8,17,49,57,58], and other types of stressors, for example physical stressors, are qualitatively different and have different consequences than social stressors [49,58-61]. While we know that chronic social stress leads to depression [9,11,31,56,62,63], we still don’t fully understand the mechanisms underlying this process [2,23,25,59]. Therefore, to fully combat social stress-induced depression, we need to gain a greater understanding of the role stress itself plays in the development of depression.

Our findings emphasize the important role that CORT plays in causing social stress-induced depression. As mentioned above, we found that submissive mice experienced increased depression-like behavior (Figure 2B) and concurrently increased levels of circulating CORT (Figure 2A), implicating increased CORT in the depression-like behavior. Furthermore, we demonstrate that dominant mice administered exogenous CORT were unable to maintain their status as dominants (Figure 4A) and exhibited increased depression-like behavior (Figure 4B), indicating that CORT itself is sufficient to cause submissive and depression-like behavior. These data are consistent with previous studies demonstrating CORT’s involvement in depression-like behavior in animal models [59,64-69] and humans [70,71]. The consistent finding across social defeat studies that submissive animals have increased...
levels of CORT implicate it as a potential causal step leading to subordinate behavior. However, to our knowledge, this is the first study to demonstrate that exogenous CORT can actually interfere with a rodent’s ability to be dominant in the social defeat paradigm, and indicates that CORT can directly cause submissiveness. This finding supports the theory that individual differences in regulation of the HPA axis could underlie vulnerability to stress-induced depression [23-25,57,72-74]. Our work confirms previous findings that exogenous CORT leads to depression-like behavior, but expands upon these past results by demonstrating that CORT itself is capable of causing submissive behavior in the social defeat paradigm and is also likely a direct part of the mechanism of how chronic social defeat stress leads to depression-like behavior.

Our data also demonstrate that blocking HPA axis output via antagonizing CRH receptors with Antalarmin prevented the development of depression-like behavior in defeated mice (Figure 5), indicating this may be a potentially viable treatment for the prevention of social stress-induced depression. Previous studies also implicate the HPA axis in the development of stress-induced depression. In rodents, social defeat stress activates the HPA axis and this activation is mediated by CRH [31,58,75]. Previous work has shown that CRH receptor antagonism blocks depression-like behavior that is induced by other stressors [22,25,76], however, to our knowledge, ours is the first demonstration that CRH receptor antagonism prevents the development of depression caused by chronic social defeat stress (Figure 5). One study has attempted to treat depression in humans by blocking the HPA axis and found that when R121919, a CRH receptor antagonist, was administered, it reduced symptoms of depression and was well tolerated with little or no side effects and without impairing the normal responsiveness of the HPA axis [77-79]. Unfortunately, a parallel trial of this drug found elevated liver enzymes in patients, which halted further development of R121919 [80]. In addition, more recently, another CRH receptor antagonist, CP-316,311, also underwent clinical trial, and although it was safe and well tolerated, it failed to decrease depression symptoms [81]. However, other CRH receptor antagonists are currently undergoing Phase II/III trials and it is possible that these new potentials could be safer and more effective [80]. Interestingly, many antidepressants normalize the HPA axis dysregulation that is associated with depression [22,24,25,66,82-86]. Taken together, these studies conducted in animal models and the clinical trials in depressed patients indicate that, as our results suggest, blocking the HPA axis via the CRH receptors has the potential to be a powerful therapeutic intervention to treat or prevent depression, but we must continue to search for drugs that are both safe and effective.

Our lab recently discovered a novel mechanism controlling the acute stress-induced activation of the HPA axis, involving dephosphorylation and downregulation of KCC2 [44]. The HPA axis is tightly constrained by the GABAergic inhibition of CRH neurons in the PVN [28,33-35,44,45]. Following acute restraint stress, the GABAergic constraint of the HPA axis is lifted enabling the physiological response to stress to be mounted. Decreased GABAergic regulation of CRH neurons has also been implicated in HPA axis hyperexcitability following chronic stress [28,35,87,88], but it is unclear how GABAergic inhibition becomes altered. GABAergic inhibition requires the maintenance of the chloride gradient, which is accomplished by KCC2 [36-38,41-43], Hewitt and colleagues [45] showed that blocking KCC2 activity in the PVN increases HPA axis activity in the form of increased corticosteroid output, indicating that the actions of KCC2 in the PVN are crucial for maintaining inhibitory control over the HPA axis. These alterations in KCC2 may play a critical role in HPA axis dysregulation following chronic social stress. Here we demonstrate dephosphorylation and downregulation of KCC2 following chronic social defeat stress (Figure 3), which may play a role in HPA axis dysfunction. However, additional studies are required to fully investigate the role of KCC2 in HPA axis reactivity following chronic social defeat stress. To date, no studies have examined KCC2 expression after chronic social stress, making ours the first to implicate alterations in KCC2 in HPA axis hyperexcitability in this model. Future studies will determine whether restoration of the chloride gradient and of GABAergic inhibition is a viable therapeutic target for depression.

Conclusions
We have demonstrated that chronic social defeat caused elevated levels of plasma corticosterone (Figure 2A) and increased depression-like behavior (Figure 2B) in defeated mice. In addition, we have shown that CORT itself caused submissiveness (Figure 4A) and depression-like behavior (Figure 4B) and that blocking the HPA axis with a CRH antagonist blocked the development of
depression in submissive mice (Figure 5). We have also identified downregulation of the KCC2 transporter in the PVN as a likely mechanism of how chronic stress alters HPA axis function (see Figures 3 and 6). These data directly implicate stress hormones in the pathophysiological mechanisms underlying the development of depression-like behavior.

Competing interests
The authors declare that they have no competing interests.

Authors' contributions

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References


These references are from various journals and studies that explore the effects of stress on the hypothalamus-pituitary-adrenal (HPA) axis, the role of GABA in brain function, and the neurochemical mechanisms underlying stress-related behaviors. The studies indicate the complex interactions between stress, corticosteroids, and neurotransmitter systems, highlighting the importance of understanding these mechanisms for developing effective therapeutic strategies.

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