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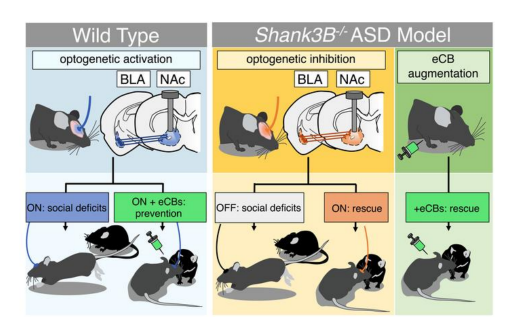
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An Endocannabinoid-Regulated Basolateral Amygdala-Nucleus Accumbens Circuit Modulates Sociability

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ABSTRACT

Deficits in social interaction (SI) are a core symptom of Autism Spectrum Disorders (ASD), however treatments for social deficits are notably lacking. Elucidating brain circuits and neuromodulatory signaling systems that regulate sociability could facilitate a deeper understanding of ASD pathophysiology and reveal novel treatments for ASD. Here we found that in vivo optogenetic activation of the basolateral amygdala-nucleus accumbens (BLA-NAc) glutamatergic circuit reduced SI and increased social avoidance in mice. Furthermore, we found that 2arachidonoylglycerol (2-AG) endocannabinoid (eCB) signaling reduced BLA-NAc glutamatergic activity, and that pharmacological 2-AG augmentation via administration of JZL184 blocked SI deficits associated with in vivo BLA-NAc stimulation. Additionally, optogenetic inhibition of the BLA-NAc circuit significantly increased SI in the Shank3B^{-/-}, an ASD model with substantial SI impairment, without affecting SI in wild-type mice. Finally, we demonstrated that JZL184 delivered systemically or directly to the NAc also normalized SI deficits in Shank3B^{-/-} mice, while ex vivo JZL184 application corrected aberrant NAc excitatory and inhibitory neurotransmission and reduced BLA-NAc-elicited feedforward inhibition of NAc neurons in Shank3B-/- mice. These data reveal circuit-level and neuromodulatory mechanisms regulating social function relevant to ASD and suggest 2-AG augmentation could reduce social deficits via modulation of excitatory and inhibitory neurotransmission in the NAc.

INTRODUCTION

Autism Spectrum Disorders (ASD) are neurodevelopmental disorders characterized by two core symptom domains, repetitive behaviors and abnormal social communication and social behaviors (1). Currently, there are no FDA-approved pharmacological treatments for the core symptoms of ASD. Therefore, elucidation of new therapeutic approaches is urgently needed. In particular, there is an unmet need for pharmacological treatment of social interaction and social communication deficits in ASD (2-6).

One such novel target could be the endocannabinoid (eCB) system. eCBs regulate several central nervous system (CNS) functions including motor control, repetitive behaviors, pain perception, anxiety, stress, learning and memory, and social behaviors (7, 8) all of which are implicated in ASD (1). Regarding social behaviors, the eCB 2-arachidonoylglycerol (2-AG) has a demonstrably significant role in regulating social behavior, as increasing levels of 2-AG increases social play in juvenile rats, while decreasing 2-AG results in decreased acquisition of social conditioned place preference (CPP) (9, 10) in mice. Despite these data, the neural circuit and synaptic mechanisms by which eCB signaling affect social function are not well understood.

In animal models, the nucleus accumbens (NAc) and amygdala are two central regions for eCB regulation of sociability (9, 11, 12). Canonically, the NAc enhances motivated behavior and social function via a dopamine-dependent mechanism (11, 13-16). Although disruptions in amygdala and NAc function have been demonstrated in ASD (17-19), how amygdala-NAc circuit activity relates to sociability deficits in ASD is not well understood. Existing literature on the amygdala, specifically the basolateral amygdala (BLA), indicates that the BLA-NAc circuit regulates learned behavioral choice including reward seeking, risk-based decision making, fear, and depressive-like behavior, all of which may contribute to changes in social approach and social

behaviors (20-27). Because the BLA-NAc circuit underlies decision making in response to rewarding cues and subsequent behavioral responses (21, 23), we hypothesize that activity in the BLA-NAc circuit may regulate multiple aspects of sociability under physiological conditions, and perhaps under conditions of impaired social function.

ASD is a highly heritable neuropsychiatric disorder, as evidenced by twin and familial studies, and one of the most common genetic findings in ASD patients are mutations in Srchomology domain 3 (SH3) and multiple ankyrin repeat domains 3 (SHANK3) encoding genes (28-32). SHANK3 is a postsynaptic density protein expressed in glutamatergic neurons that serves as a scaffold for key glutamatergic receptors (33, 34). The *Shank3B*-/- mouse model is characterized by an excision of exons 13-16 that causes removal of two major *Shank3* isoforms, and replicates the genetic findings commonly presented in patients (33, 35-38). This excision results in a loss-of-function of the SHANK3 protein. *Shank3B*-/- mice demonstrate excessive, repetitive grooming and deficits in social behavior, making the model a good candidate for exploring novel ASD therapeutic targets and circuit dysfunction (39, 40). Previous studies suggest that the significant changes in social behavior in the *Shank3* loss-of-function models may be mediated by signaling changes in the NAc (41), warranting further investigation of the NAc and NAc-associated circuitry, in the regulation of social function and pathophysiology of ASD.

Here we utilized optogenetic circuit mapping to demonstrate that activation of the BLA-NAc circuit disrupted physiological social function and that inhibition of this circuit reversed social deficits in *Shank3B*-/- mice. We further demonstrated that 2-AG signaling is a negative regulator of BLA-NAc glutamatergic transmission and that pharmacological augmentation of 2-AG levels mitigated social avoidance induced by BLA-NAc circuit activation and those observed in *Shank3B*-/- mice, possibly via decreasing BLA-NAc-elicited feed-forward inhibition.

RESULTS

Optogenetic activation of the BLA-NAc circuit disrupts social behavior

To examine the role of the BLA-NAc circuit in social function, we utilized an in vivo optogenetic approach in which we injected a ChR2 (AAV5-CaMKIIa-hChR2(H134)-EYFP) viral vector into the BLA and delivered blue-light stimulation into the NAc via fiber optic cannulas. First, we tested the effects of 20Hz blue light stimulation (5ms light pulses, 10-13mW, 5s on 5s off) on social function using the 3-chamber social interaction (SI) task in male and female C57BL/6J mice. We found that blue light delivery into the NAc impaired sociability in ChR2expressing compared to YFP-expressing (AAV5-CaMKIIa-EYFP) control animals (Figure 1 A-B and Figure S1 A-B for cannula placements). This light stimulation pattern did not affect distance traveled in this apparatus (Figure 1 C), but significantly decreased time ChR2-expressing mice spent in the social chamber relative to YFP-expressing mice (Figure 1 D). We also found that ChR2-expressing mice, relative to YFP controls, spent significantly less time in close proximity (5cm) to the target animals (Figure 1 E) and spent less time sniffing or investigating the cup in which the target mouse was contained (Figure 1 F). Importantly, BLA-NAc stimulation for 5 minutes prior to the 3-chamber SI task did not alter sociability (Figure S2 A-B), suggesting that the sociability-impairing effects of BLA-NAc activation required ongoing circuit stimulation. We next validated our optogenetic stimulation protocol using ex vivo whole-cell electrophysiological recordings. These studies revealed 20Hz blue light stimulation within the NAc triggered high-fidelity neuronal firing, confirming that this frequency of stimulation increased NAc neuronal activity and increased spontaneous glutamate release onto NAc medium spiny neurons (MSNs) (Figure S3 A-E). Optogenetic stimulation of BLA afferents to the NAc also resulted in optically-evoked excitatory postsynaptic currents (oEPSCs) onto D2 dopamine

receptor expressing MSNs (D2+ MSNs), and D2-(presumptive D1) MSNs, indicating BLA glutamatergic afferents to the NAc target both subtypes equally (Figure S3 F-H). Lastly, we sought to determine whether BLA-NAc activity could bi-directionally modulate sociability. Thus, we repeated the 3-chamber SI task in animals that bilaterally expressed either NpHR (AAV5-CaMKIIa-eNpHR3.0-EYFP) or YFP in order to inhibit BLA-NAc activity in vivo (Figure S2 E). However, we found no effects of constant orange-light (620 nm, 10-13mW) inhibition of the BLA-NAc circuit on sociability in the 3-chamber SI test (Figure S2 F-G), suggesting BLA-NAc activity is not necessary for physiological expression of SI.

To extend upon the observation that BLA-NAc activation disrupts sociability, we next analyzed the effects of BLA-NAc activation in a juvenile reciprocal SI test using a wireless in vivo optogenetic system (Figure 2). In this task, we found that unilateral 20 Hz blue light stimulation significantly increased time fleeing or withdrawing from the social target (Figure 2 A) and decreased time spent social sniffing (Figure 2 B) in mice that expressed ChR2 relative to YFP controls. In contrast, we found no effects of blue light stimulation in ChR2-expressing compared to YFP-expressing controls on the time the test animals spent following the social target (Figure 2 C) or engaged in passive social behavior (Figure 2 D). Lastly, immobility time was increased (Figure 2 E) while exploration time (Figure 2 F) and self-grooming (Figure 2 G) were decreased in ChR2 animals compared to YFP-expressing controls. These data are consistent with our 3-chamber SI data and indicate that BLA-NAc pathway activation reduces social behavior and induces social avoidance in a more naturalistic setting.

We next tested the effects of BLA-NAc activation on motivational aspects of SI seeking using a social CPP task (**Figure 2 H**). After conditioning, we found that bilateral (but not unilateral, **Figure S2 C-D**) blue light delivery during social CPP testing resulted in decreased time spent in

the social-paired chamber and lower CPP scores (post-test chamber time minus pre-test chamber time) (Figure 2 I-K), in ChR2-expressing animals compared to YFP controls. Of note, virus expression did not alter pre-test chamber preference (Figure 2 L). Taken together, data from these three assays indicate BLA-NAc activation impairs motivational aspects of SI seeking, reduces social preference, and induces social avoidance, in male and female mice.

Optogenetic activation of the BLA-NAc circuit is rewarding but not anxiogenic

The NAc is well-known to regulate reward-related processes (42, 43) suggesting BLA-NAc stimulation could reduce sociability via an occlusion of the rewarding aspects of SI. To examine this possibility, we first tested the reinforcing effects of BLA-NAc stimulation using a real-time place preference (RtPP) assay. Consistent with previous studies (24), we found animals expressing ChR2 spend more time in the ON chamber, in which blue light stimulation was delivered, versus the OFF, chamber in the RtPP task (Figure 3 A-B), without significantly affecting locomotor activity (Figure 3 C). Taken together with our previous data above, these results indicated BLA-NAc glutamatergic circuits can support RtPP and social avoidance in parallel. These data also suggest BLA-NAc stimulation could indeed impair social function via occlusion of the rewarding aspects of social behavior. If this were the case, one would also predict occlusion of natural reward seeking, such as food seeking, behavior by BLA-NAc stimulation. To test this hypothesis more explicitly, we measured time in a chamber containing a highly palatable food (vanilla Ensure) in lieu of a social target in a modified 3-chamber task design (Figure 3 D). In this task we observed BLA-NAc stimulation did not disrupt chamber preference (Figure 3 E), alter time spent in close proximity to the Ensure-containing cup (Figure 3 F), or affect time spent drinking Ensure (Figure 3 G). Overall these data suggest BLA-NAc activation does not reduce

sociability via occlusion of the rewarding aspects of SI. Additionally, because changes in anxiety and anxiety-like behavior can affect social behavior (1, 44, 45), we next tested the effects of BLA-NAc stimulation on anxiety-like behavior and found activation did not alter behavior in the light-dark box (**Figure S4 A-D**) and neither activation nor inhibition affected anxiety-like behavior in the elevated-plus maze (**Figure S4 E-L**). Taken together that these data indicate social avoidance observed during BLA-NAc stimulation was not secondary to increased anxiety-like behavior.

eCB signaling broadly regulates BLA-NAc glutamatergic transmission

Given the prominent role of eCB signaling in the regulation of central glutamatergic transmission (46), and the role of eCB signaling in social function (11, 12, 47), we next examined the possibility that eCB signaling regulates BLA-NAc glutamatergic activity. Indeed, the cannabinoid receptor agonist CP55940 significantly decreased oEPSC amplitude at BLA-NAc synapses, confirming the presence of functional cannabinoid receptors in this pathway (Figure 4 A). Because previous investigations have shown differential eCB signaling at glutamatergic synapses onto distinct striatal MSN subtypes and within distinct BLA-NAc subcircuits (24, 48-50), we further examined the effect of 2-AG signaling and CB1 function at BLA-NAc D1+ and D1- (presumptive D2) MSNs glutamatergic synapses. We first found that increasing 2-AG levels via inhibition of monoacylglycerol lipase (MAGL) with JZL184 significantly reduced the frequency, but not amplitude, of asynchronous EPSCs onto both D1+ and D1- NAc MSNs (Figure **4 B-C)** and that WIN55212-2, another direct cannabinoid receptor agonist, resulted in suppression of oEPSC amplitude and long-term depression onto both D1+ and D1- MSNs (Figure 4 D). We next investigated the expression of depolarization-induced suppression of excitation (DSE), a protocol used to measure endogenous retrograde synaptic 2-AG signaling, and found DSE was

present at both BLA to D1+ and D1- MSN synapses (Figure 4 E), although the suppression was more robust in D1+ relative to D1- MSNs (Figure 4 F). Lastly, we examined the role of JZL184 on spontaneous synaptic activity in the NAc. JZL184 did not alter spontaneous EPSC (sEPSC) frequency onto either D1+ or D1- MSNs (Figure 4 G) but significantly decreased spontaneous inhibitory post synaptic current (sIPSC) frequency, which was statistically significant, via post hoc analysis for D1+ MSNs (Figure 4 H). There was no effect of JZL184 on D1+ or D1- sIPSC amplitude (Figure 4 I). These data suggest that eCB regulation of BLA-NAc glutamatergic transmission is present on both primary MSN cell types in the NAc and that 2-AG signaling can also affect NAc local GABAergic neurotransmission.

2-AG augmentation prevents BLA-NAc activation-induced social impairment

Our data thus far suggest 2-AG signaling can reduce presynaptic glutamate release at BLA-NAc synapses, suggesting that pharmacological 2-AG augmentation could counteract the social impairment observed after BLA-NAc circuit activation. To test this hypothesis, we administered JZL184 (8mg/kg IP) 1 hour prior to BLA-NAc stimulation in the 3-chamber SI task. We found that JZL184 prevented blue light delivery-dependent suppression of sociability in the 3-chamber SI test in ChR2-expressing animals but did not affect SI in YFP-expressing controls (Figure 5 A); JZL184 did not affect distance travelled in this assay (Figure 5 B). ChR2-expressing JZL184-treated mice also spent more time in the close interaction zone of the 3-chamber apparatus (Figure 5 C) and more time investigating the mouse-containing cup relative to the empty cup (Figure 5 D) compared to vehicle-treated ChR2-expressing mice. Lastly, neither pharmacologically increasing (JZL184 8mg/kg IP) nor decreasing (DO34 50 mg/kg IP) 2-AG levels affected SI in naive, non-surgical animals (Figure S5 A-D). These data indicate that, while pharmacological

modulation of 2-AG levels has little effect on physiological expression of sociability, 2-AG augmentation is able to attenuate the social avoidance behavior induced by BLA-NAc circuit activation.

Optogenetic BLA-NAc circuit inhibition reduces social deficits in Shank3B-/- mice

That BLA-NAc inhibition did not affect social behavior in wild-type (WT) mice (see Figure S2) suggests that the BLA-NAc circuit does not physiologically regulate sociability under basal conditions. However, it is possible that BLA-NAc activity could contribute to social impairment under pathological conditions, such as those observed in the Shank3B-/- model of syndromic ASD. Indeed, we found, using a counterbalanced cross-over design, that bilateral orange-light stimulation of NpHR, and thus inhibition of the BLA-NAc circuit, normalized time in the mouse chamber during orange light-ON relative to light-OFF testing in Shank3B^{-/-} mice (Figure 6 A and see Figure S1 C-D for cannula placements). Consistent with previous data, YFPexpressing Shank3B^{-/-} controls showed a lack of sociability (39, 51, 52) under both light-ON and light-OFF conditions in the 3-chamber test, while NpHR-expressing Shank3B-/- mice exhibited social-chamber preference only under light-ON conditions (Figure 6 B, F). There were no changes in the distance travelled between groups (Figure 6 C, G). Orange light inhibition of the BLA-NAc circuit in Shank3B^{-/-} mice also resulted in a significant preference for time investigating the mousecontaining cup in NpHR-expressing mice under light ON, but not light OFF, conditions, while YFP-expressing mice show no preference under either condition (Figure 6 D, H). These data support the hypothesis that reducing BLA-NAc activity can enhance sociability under pathological conditions associated with reduced SI, such as those observed in the *Shank3B*-/- model.

2-AG augmentation reduces social deficits and feed-forward GABAergic signaling in Shank3B-/- mice

Finally, because acute treatment with JZL184 was able to reverse the sociability deficits resulting from overactivation of the BLA-NAc circuit, and inhibiting the BLA-NAc circuit reverses social deficits in *Shank3B*-/- mice, we hypothesized that acute treatment with JZL184 (8mg/kg IP) would increase social behavior in *Shank3B*-/- mice (Figure 7 A). While vehicle treatment did not increase SI in *Shank3B*-/- animals, JZL184 pretreatment resulted in significant social preference in *Shank3B*-/- animals in the 3-chamber SI task (Figure 7 B-E) but had no effect on distance travelled (Figure 7 C). JZL184 treatment also resulted in a preference for investigating the target mouse relative to the empty cup in *Shank3B*-/- mice (Figure 7 D). In addition to social deficits, *Shank3B*-/- animals display aberrant repetitive grooming behavior (39, 51, 52), modeling the second core feature of ASD: restricted and repetitive behaviors. We subsequently demonstrated that the amount of time grooming is significantly reduced by JZL184 (8mg/kg IP) treatment in *Shank3B*-/- animals but did not affect grooming behavior in WT mice (Figure 7 F).

While our data clearly show systemic pharmacological augmentation of 2-AG reduced social impairment upon BLA-NAc stimulation, and in *Shank3B*-/- mice, these data do not conclusively link the two findings in a manner that supports the site of action of JZL184 as being within the BLA-NAc circuit. To directly test the hypothesis that systemic JZL184 administration effectively reduces social impairment via NAc-mediated mechanisms, we administered JZL184 (5μg/μL) or vehicle directly into the NAc using a microinfusion approach prior to conducting the 3-chamber SI task in *Shank3B*-/- mice (**Figure 8 A**, and **Figure S1 E** for cannula placements). JZL184 microinfusion did not affect distance traveled in this task (**Figure 8 B**); however, it increased time *Shank3B*-/- mice spent in the mouse chamber and time investigating the target mouse

under the cup (**Figure 8 C-E**). Vehicle microinfusion did not result in increased time spent in the mouse containing chamber or increase time in social investigation of mouse relative to empty cup. These data indicate local actions of JZL184 within the NAc are sufficient to normalize SI deficits in *Shank3B*^{-/-} mice

Given that elevating 2-AG selectively within the NAc is sufficient to increase social behavior in the Shank3B^{-/-} mouse, we investigated the effects of JZL184 on the synaptic signaling in the NAc as well as specifically after BLA-NAc optogenetic stimulation to gain insight into the potential mechanisms by which JZL184 may affect sociability in Shank3B^{-/-} mice. Firstly, we found no changes in the intrinsic excitability of NAc MSNs as measured by the number of action potentials generated per current injection, action potential threshold, resting potential, or input resistance between WT and Shank3B-/- mice (Figure S6 A-D). However, we found that Shank3B-¹⁻ animals display a robust increase in sIPSC frequency (Figure 9 A) but not amplitude (Figure 9 **B)**, as well as increased sEPSC frequency (Figure 9 C), but not amplitude (Figure 9D) in the NAc. Moreover, we observed that JZL184 significantly reduced sIPSC and sEPSC frequency in the NAc of Shank3B^{-/-} mice (Figure 9 A, C). In response to ex vivo BLA-NAc optogenetic stimulation, we observed no genotype-specific effects on either BLA-NAc oEPSCs or di-synaptic feed-forward (FF) oIPSCs (Figure 9 E-H). However, JZL184 robustly decreased BLA-NAc feed-forward oIPSC amplitude in *Shank3B*^{-/-} but not WT mice (**Figure 9 E-F**). JZL184 did not affect BLA-NAc oEPSC amplitude in either Shank3B-/- or WT mice (Figure 9 G-H). These data suggest 2-AG augmentation could normalize social function in Shank3B-/- mice via modulation of BLA-NAcelicited FF inhibition (possibly mediated via local GABAergic interneurons), which could result ultimately in disinhibition of NAc MSNs (Figure 10). This model could also explain increased sociability after BLA-NAc circuit inhibition in Shank3B-/- mice, as reduced BLA glutamatergic

drive to local NAc GABAergic interneurons would also disinhibit MSNs (Figure 10). How the balance of direct BLA excitation of NAc MSNs and local GABAergic interneurons are sculpted to ultimately direct optimal sociability remains to be determined.

DISCUSSION

A main finding of the present study is that activation of the BLA-NAc glutamatergic circuit is sufficient to decrease sociability in the 3-chamber SI task, increase social avoidance in a naturalistic juvenile reciprocal interaction task, and reduce SI-seeking in a social CPP assay. Previous studies have shown that glutamatergic projections from the BLA, including those to the ventral hippocampus (vHPC) and medial prefrontal cortex (mPFC) also decrease SI time when activated (53, 54). However, activation of BLA-vHPC and BLA-mPFC circuits also increase anxiety-like behavior (53, 55). Because social behavior deficits are often associated with increases in anxiety (44, 45), we measured the effects of BLA-NAc activation in anxiety-like behavior. However, we did not observe anxiogenic effects of BLA-NAc stimulation in the elevated-plus maze or light-dark box assays. Interestingly however, we did observe increased immobility, a potential indicator of a fear response, during BLA-NAc stimulation in the juvenile reciprocal SI task, suggesting reduced SI in this task may be secondary to fear or anxiety-like states. Alternatively, it is possible mice are remaining immobile to avoid social interaction due to the target mouse freely interacting with the test mouse, rather than being contained within a cup, as is the case in the 3-Chamber SI task. Lastly, the BLA-NAc circuit has been strongly implicated in reward-seeking behavior, and stimulation of the BLA-NAc circuit is reinforcing (present data and (24, 43)). These data suggest BLA-NAc stimulation could reduce SI via an occlusion of the rewarding aspect of social behavior. However, BLA-NAc simulation did not reduce interaction with, or avoidance of, palatable non-social reward (Ensure) in our modified 3-Chamber assay, arguing against this possibility. Taken together, these data suggest BLA-NAc glutamatergic circuit activation reduces sociability and social reward seeking, which are not likely secondary to increases in anxiety or mediated via occlusion of the rewarding effects of SI.

One possible explanation for how BLA-NAc activation results in the social avoidance may relate to underlying BLA-NAc sub-circuits. Specifically, previous studies have shown that activation of D2-MSNs directly in the NAc, and in vivo optogenetic activation of the cholecystokinin (CCK)-expressing BLA neurons that project preferentially to NAc D2-MSNs, significantly decreases time in the SI zone after social defeat stress (24). In contrast, the reinforcing effects of BLA-NAc stimulation can be blocked by a D1 receptor antagonist (22), indicating BLA-NAc D1-MSN activation could underlie the rewarding effects of BLA-NAc circuit activation. Context-dependent differential engagement of these distinct subcircuits could explain parallel roles in reward and social avoidance observed after BLA-NAc circuit manipulations. Future studies directly examining and manipulating BLA-NAc subcircuit activity in social and non-social contexts may help clarify the neural mechanisms underlying BLA-NAc circuit effects on sociability and reward.

Given that BLA-NAc circuit activation reduces sociability, we next sought to explore molecular mechanisms that reduce BLA-NAc circuit activity in an attempt to reveal therapeutic approaches to increase social behavior. Previous work has demonstrated that activation of the CB1 receptor by eCBs, including 2-AG, dampens glutamatergic activity (46) and has a functional role in social behavior via modulation of both NAc signaling and endogenous BLA-NAc activity (9, 10, 12, 24, 56). Therefore, we investigated whether 2-AG signaling modulated BLA-NAc glutamatergic transmission. Indeed, we show that the CB1 receptor activation reduced glutamate release at BLA-NAc synapses and that enhancement of 2-AG signaling via MAGL inhibition by JZL184 reduced aEPSC frequency at BLA-NAc D1 and D2-MSNs. It is noteworthy that these results are in contrast to a recent study indicating cannabinoid signaling selectively modulates BLA-NAc D2-MSN synapses (24). Consistent with the ability of CB1 activation and 2-AG

augmentation to reduce optogenetically-elicited glutamatergic signaling efficacy at BLA-NAc synapses ex vivo, we showed that systemic JZL184 administration prevented the decreased sociability elicited by BLA-NAc optogenetic stimulation in vivo. Our data suggest pharmacological 2-AG augmentation could serve as a therapeutic approach to increase sociability under pathological conditions associated with amygdala-NAc circuit hyperactivity.

While we show that activation of the BLA-NAc circuit is sufficient to reduce SI behavior, we also showed that activity is not necessary for the physiological expression of social behavior, as optogenetic inhibition of the circuit did not alter sociability under basal conditions. This finding differentiates the BLA-NAc circuit from other BLA circuits, such as the BLA-vHPC, which bidirectionally alter social behavior (54). Although, our data suggest BLA-NAc activity is not physiologically recruited to regulate sociability, it is possible that BLA-NAc circuit inhibition could increase or normalize sociability in mice with pre-existing social impairment. Indeed, we found that inhibition of the BLA-NAc circuit robustly increased social behavior in the Shank3B^{-/-} mouse, a model of ASD which has a well-established social deficit (39, 51, 52). Consistent with our data, clinical reports have demonstrated that the amygdala is hyper-functional in ASD patients during anticipation of social cues, and that the NAc of ASD patients has increased functional connectivity relative to typically developing peers (17, 18, 57). Continued research into the functional connectivity of the BLA-NAc circuit in ASD patients and additional preclinical mouse models of ASD is needed to fully understand the role the BLA-NAc circuit in ASD-associated social deficits.

We also showed that 2-AG augmentation via systemic JZL184 administration reverses both social and restricted, repetitive behavior deficits in *Shank3B*-/- mice. Additionally, direct administration of JZL184 into the NAc is sufficient to rescue social behavioral deficits in

Shank3B^{-/-} mice. Taken together, these data add to the growing body of evidence suggesting pharmacological eCB augmentation could serve as a therapeutic approach for treating ASD and ASD-associated syndromes (47, 58-62). Currently there are no FDA-approved pharmacological treatments for ASD, and current medications to improve social deficits are limited. Given the recent advancement of MAGL inhibitors to early clinical trials (63, 64), our data could support examination of MAGL inhibitors for the treatment of social deficits in ASD, and other disorders characterized by social domain deficits.

In the present study we showed that, relative to WT littermates, *Shank3B*-/- animals have increased sIPSC and sEPSC frequency, but not amplitude, in the NAc, revealing increased GABA and glutamate release probability, but no changes in the intrinsic excitability of NAc MSNs. Interestingly, JZL184 restored both sIPSC and sEPSC frequency to that of WT levels. These data provide a potential mechanism for the restoration of social function elicited via systemic and intra-NAc JZL184 infusions in *Shank3B*-/-. Our findings of increased IPSC frequency in the NAc of the *Shank3B*-/- animals are consistent with previous findings showing hypoactivity of the NAc during SI in a complete *Shank3* knockout model (41). Taken together, these data may reveal a convergent mechanism of heightened GABAergic activity in the NAc of *Shank3*-/- models. Future studies of NAc activity in additional ASD models may provide insight to the role of NAc synaptic physiology in social behavior more broadly.

Because we found SI deficits were normalized by the BLA-NAc circuit inhibition in *Shank3B*-/- mice, we also analyzed the BLA-NAc circuit physiology in *Shank3B*-/- animals. Unexpectedly, we found no measurable differences in BLA input into the NAc in *Shank3B*-/- mice compared to WT animals as oEPSC and feed-forward IPSC amplitudes were not different between genotypes; however, JZL184 was able to significantly reduce FF IPSC amplitude in *Shank3B*-/-,

but not WT, mice. This data seemingly contrasts with our finding showing inhibition of the BLA-NAc restores sociability behavior in *Shank3B*-/- animals as glutamatergic drive onto NAc neurons is not altered. However, it is possible that inhibition of GABAergic interneurons in the NAc of Shank3B^{-/-} animals, by either NpHR-induced reductions in FF-inhibition or JZL184 application, is sufficient to restore social function. One possible way that JZL184 could improve SI is via inhibition of glutamatergic BLA synapses and GABAergic fast-spiking interneurons (FSIs) onto MSNs, both of which are regulated by CB1 receptors at the presynaptic level (Figure 10) (26, 65, Increases in 2-AG may therefore inhibit activation of NAc-FSIs in response to BLA stimulation as well as inhibit GABAergic release from FSIs. Because FSIs in the NAc have been shown to gate impulsive behavior in mice (67), we hypothesize that JZL184 inhibition of FSIs signaling could promote SI via increasing directed social behavior output in a similar manner. Additionally, reductions in FSI inhibition of NAc MSNs could allow for restoration of proper signaling from other NAc inputs such as the prefrontal cortex, which could be dysregulated in Shank3B^{-/-} mice. Further investigation into the role of FSIs, GABAergic circuitry, and additional glutamatergic inputs to the NAc in the Shank3B^{-/-} model, as they relate to social behavior, may uncover additional synaptic mechanism contributing to social deficits in ASD.

In summary, we show that BLA-NAc circuit activation reduces sociability and is highly regulated by 2-AG-mediated eCB signaling. We further show that systemic and NAc-specific 2-AG augmentation and optogenetic BLA-NAc circuit inhibition normalize social deficits in the *Shank3B*-/- model of ASD. We hypothesize that 2-AG augmentation reduces social deficits in *Shank3B*-/- mice via normalization of hyperactive GABAergic and glutamatergic signaling in the NAc and via reductions in BLA-elicited FF-inhibition onto NAc MSNs. These data reveal the BLA-NAc circuit as a critical regulator of social function and suggest pharmacological 2-

AG augmentation could represent a promising approach to the treatment of social domain symptoms in ASD.

METHODS

Animals:

Mice were housed on a 12:12 light-dark cycle with lights on at 06:00. All experiments were conducted during the light phase. Food and water were available *ad libitum*. Mice on a C57/BL6J background were used for all experiments. Wild-type mice were ordered from JAX (stock #00064) at 5 weeks of age. Male and female *Shank3B*^{+/-} mice were ordered from JAX laboratories (Stock number: 017688) and bred in house. male and female transgenic bacterial artificial chromosome (BAC) *Drd1a-tdTomato* and BAC Drd2-EGFP mice were obtained from JAX laboratories and bred to C57BL/6J wild type females.

Viruses:

For in vivo optogenetic studies we used AAV5-CaMKIIa-hChR2(H134)-EYFP for optical excitation or AAV5-CaMKIIa-eNpHR3.0-EYFP for optical inhibition and AAV5-CaMKIIa-EYFP (250nL, UNC Vector Core, Chapel Hill, NC, USA) as a control in all studies.

Stereotaxic surgery:

Male and female mice at 5–8 weeks old underwent unilateral or bilateral stereotaxic surgery, as indicated in the figures, under isoflurane anesthesia. Viruses were infused into the BLA (AP: -1.25, ML: ±3.25, DV: 5.05) at a rate of 100 nl min⁻¹. At least 4 weeks following viral injection, mice underwent unilateral or bilateral fiber optic implantation. Implants were lowered to the ipsilateral NAc of the viral surgery (AP: 1.25, ML: ±0.55, DV: 4.10, +/- 10° tilt for bilateral implants) (0.20mms⁻¹). For microinfusion studies, bilateral, stainless-steel infusion guide cannulas (26 gauge, cut to 3mm length, 2mm center to center distance, C235GS-5-2.0/SPC- Plastics One,

Roanoke VA) were placed above the NAc (AP: 1.35, ML: ±1.00, DV: 3.00) and fitted with a dummy cannula (C235DCS-5/SPC, Plastics One, Roanoke VA) and dust cap (303DC/1 Plastics One, Roanoke VA). Implants were secured with dental cement. Animals recovered for at least one week prior to any experimental manipulation.

Electrophysiology:

Acute slice preparation and whole cell patch clamp recordings were performed similar to those previously described (68-70). MSNs in the NAc were identified based on resting membrane potential and visual appearance (size, morphology) at 40X magnification with an immersion objective with differential interference contrast microscopy (DIC). Putative D1 receptor or D2 receptor neurons were identified by the presence or absence of fluorophore expression in BAC transgenic mice expressing td-Tomato under the D1 receptor promoter or GFP under the D2 receptor promoter. For more detailed electrophysiological methods see supplementary materials.

Behavior testing:

All mice were single housed with the exception of animals tested in repetitive grooming assay. All experiments were conducted at least 48 hours apart. *Shank3B*-/- animals were run in parallel with WT littermate controls. All animals were handled at least 3 days prior to testing.

1) 3-chamber social interaction test: After a 10-minute habituation phase, the test mouse was permitted to freely explore the arena containing a sex-matched, juvenile (4-6 week-old) mouse inside one inverted pencil cup, and an empty inverted pencil cup in the opposite chamber for the test phase (5 minutes). When *Shank3B*-/- animals were used, animals were paired with agematched, sex-matched targets and given a 5-minute habituation while restricted to the center

- chamber, as previously described (39). For in vivo optogenetic studies, stimulation was delivered during the test phase unless otherwise noted (**Figure S2 A-B**). Videos were coded post-hoc for time investigating each cup by two blinded coders. Time investigating represents time in which the test mouse is sniffing or interacting with the cup.
- Ensure- 3-Chamber Task: Animals were trained for 3 consecutive days in home cage to drink commercially available Ensure. On test day, animals were placed in the 3-chamber apparatus (as described above). An ensure-filled sipper bottle was affixed to an inverted pencil cup so the animals could easily reach and drink from the sipper. Stimulation was turned on and animals were permitted to freely explore the apparatus for 5-minutes. Videos were analyzed post-hoc for time drinking Ensure or time investigating (sniffing or interacting) the empty cup by two blinded coders.
- 3) Grooming behavior assay: Animals were placed into an empty, clean, novel cage for 15 minutes. Videos were recorded during the interaction and analyzed post-hoc by two independent, blinded coders beginning after a 5-minute habituation period.
- 4) <u>Juvenile Reciprocal Social Interaction Test</u>: Animals were permitted to freely interact with a sex-matched, juvenile mouse (~4 weeks old) for 10 minutes in a clean, novel cage. Videos were analyzed post-hoc by two independent, blinded coders.
- 5) Real-time place preference: When animals entered a randomly assigned chamber, light stimulation was delivered; stimulation ended when animals left the stimulation chamber.

 Animals were tested for 5 minutes.
- 6) <u>Social Conditioned Place Preference:</u> We used a 3-chamber apparatus with two distinct chambers (dots and stripes) and a connecting corridor. After a pre-test trial (10-minutes), mice were assigned in a counter balanced, unbiased manner in either the dots or stripe chamber with

a novel, juvenile sex-matched mouse for 10 minutes, immediately followed by 10 minutes in in the opposite chamber alone. This repeated for 3 consecutive days total. Finally, on test day, light stimulation was turned on and animals were able to freely move in the apparatus for 10 minutes.

For complete behavioral methods see Supplemental Materials.

In vivo optogenetic stimulation:

Stimulation patterns were delivered from the PlexBright 4 Channel Optogenetic Controller and controlled by Radiant v2 Software (Plexon, Dallas TX). The optogenetics controller box was attached to the PlexBright Dual LED, rotatable commutator in which a blue (465 nm) or orange (620 nm) LED lights were affixed. PlexBright Optical Patch Cables (.5NA) were then attached to the commutator. In blue-light stimulation studies, animals received 20Hz blue light stimulation (Plexon, Dallas, TX, USA) in a 5s on 5s off pattern at 10-13mW. For inhibition studies, animals received a constant orange light (~10mW). One week following implantation surgery, mice were habituated to false patch cables in 20-minute bins for 3 consecutive days. On test day, the patch cable was connected to fiber optic implants and animals were habituated for at least 5 minutes before the start of any behavior test.

Microinfusion Studies:

After 7 days of recovery from stereotaxic surgery, animals were habituated to handling for 3 consecutive days in which animals were restrained in increasing amounts (30s, 60s, 120s) and dummy cannula were replaced daily to prevent blocking and habituate animals to restraint. JZL184

was bilaterally infused into the NAc using a bilateral, 4mm cut length internal, infusion cannula (C235IS-5/SPC, Plastics One, Roanoke VA) at a dose of 0 or 5μg/L and at a volume of 0.2μL per hemisphere over 1 min as previously described (71). 30 minutes following infusion, animals were placed into the 3-chamber SI task.

Histology and Imaging:

After animals completed behavior experiments, brain tissue was collected in order to validate implant and viral placement. For validation experiments, brain tissue slices were mounted in VECTASHIELD + DAPI (Vector laboratories, Burlingame, CA, USA) onto charged slides and sealed with a clear nail polish. Images were collected using an upright Axio Imager M2 epifluorescent microscope at a 5x and 20x objective.

Exclusion Criteria:

For all viral and implant studies, animals were excluded based on a priori standards. The injection site of all viral injections was identified by the presence of GFP or eYFP fluorescent marker. For all fiber optic implantations, location was determined by implantation track. Animals were excluded from all data sets if the viral expression or implantation was not in the targeted regions. For all in vivo studies the location of the viral expression and implantation is displayed in figure **S1**. *Shank3B*^{-/-} and WT controls were genotyped following sacrifice for confirmation. Cells were excluded from all analyses for four reasons. 1: if the holding current dropped below - 200 pA at any time during the recording. 2: if the access resistance was > 20 M Ω . 3: if the access resistance fluctuated by more than 20% throughout the recording. 4: There was no optogenetically-evoked response. In all data sets, data were analyzed using Grubbs' outlier test

(alpha = 0.05) and removed accordingly. If animals were excluded from a behavioral test for outlying data points, all data collected from the animal in the experiment were removed.

Statistics:

Data represented as means ± SEM, and individual plot points overlaid on a mean bar graph. Statistical analysis conducted using Prism 8 (Graphpad, La Jolla, CA). Statistical tests and parameters are indicated in figure legends. Significance set at alpha=0.05. A P value less than 0.05 was considered significant. Student's t-tests were two-tailed.

Study Approval:

All studies were carried out in accordance with the National Institute of Health Guide for the Care and Use of Laboratory Animals and approved by the Vanderbilt University Institutional Animal Care and Use Committee.

For full methods please see Supplemental Materials.

AUTHOR CONTRIBUTIONS:

SP and OF conceived the study, designed experiments, and co-wrote the paper. OF performed experiments, acquired, and analyzed data in laboratory of SP. RB, DM, and NH, completed, assisted in the design, and analyzed ex vivo electrophysiological experiments and data in laboratory of SP, while BT completed, designed and analyzed electrophysiological data in the lab of BG. CG and OF designed and completed microinfusion experiments in the laboratory of BG. VK, JB, and MA assisted in the completion of experiments. VK, JA, and MM coded behavioral data.

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FIGURES

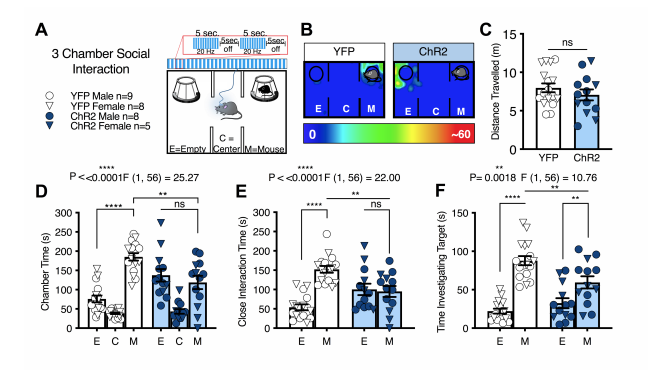


Figure 1: Activation of BLA terminals in the NAc decreases sociability A) 3-chamber SI task and optogenetic stimulation protocol. Animals were continuously stimulated during the 5-minute test using a 20Hz pattern (10mW, 5s on 5s off). B) Representative heat maps of chamber time in a ChR2 and YFP control mouse. C) Optogenetic stimulation of the BLA-NAc circuit did not affect distance traveled (ns p = 0.3224). D) Animals that express ChR2 showed reduced social preference (**p = 0.0010, ****p < 0.0001, ns p = 0.5565), E) decreased close interaction time (**p = 0.0011, ****p < 0.0001, ns p = 0.9353) and F) reduced time investigating, or sniffing, the target mouse compared to animals that express YFP (M-M **p = 0.0026, E-M **** p < 0.001, **p = 0.0071). YFP n= 17, ChR2 n=13 (C-F). Data analyzed via Two-Way Mixed-Effects ANOVA followed by Sidak's multiple comparisons test (D-F) and unpaired two-tailed t-test (C). P and F values for Chamber x Virus interaction shown in D-F.

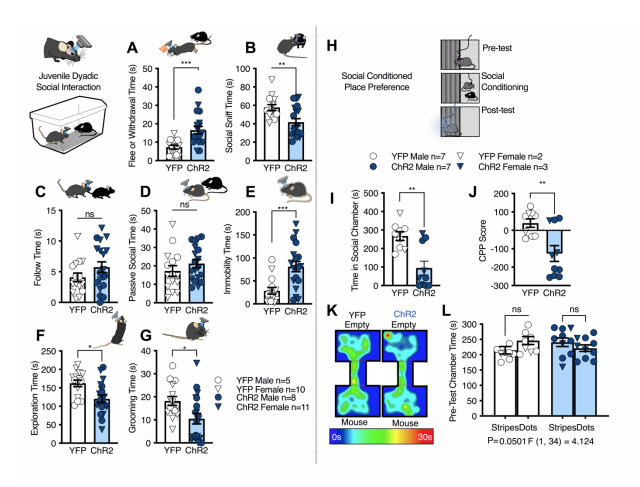


Figure 2: Activation of BLA terminals in the NAc increases social avoidance and reduces social interaction seeking A) Optogenetic stimulation of the BLA-NAc circuit increased fleeing and withdrawal behaviors (***p = 0.0007) and (B) decreased social sniffing behaviors (**p = 0.0041) compared to YFP expressing controls. There was no effect of BLA-NAc activation on C) following (p = 0.1573) or D) passive social behavior (p = 0.2604). Animals expressing ChR2 were significantly more E) immobile (***p = 0.0007) and F) explored less (*p = 0.0049) than YFP controls. G) Animals that expressed ChR2 self-groomed significantly less than YFP-expressing controls (*p = 0.0184) YFP n=15, ChR2 n=19 (A-G). H) Social CPP paradigm. I) Bilateral activation of BLA-NAc circuit significantly decreased time in social-paired chamber during the post-test (**p = 0.0013) J) and reduced CPP score relative to YFP controls (**p = 0.0041). K) Representative heat maps for social CPP experiment. L) No pre-test preference to either chamber was detected (YFP dots vs stripes, p = 0.1777, ChR2 dots vs stripes, p = 0.4619. YFP n=9, ChR2 n=10 (I, J, L). Data analyzed via unpaired, two-tailed t-test (A-G and I-J) and Two-Way Mixed Effects ANOVA with Sidak's multiple comparisons post-hoc test (L), with P and F values for Chamber x Virus interaction shown in L.

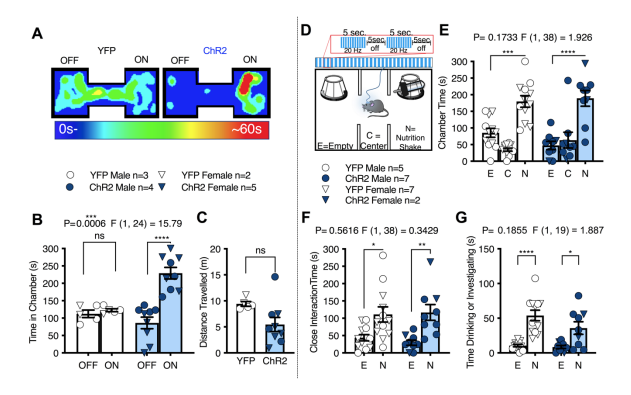


Figure 3: Activation of BLA terminals in the NAc is rewarding but does not reduce palatable food-seeking. A-C) Effects of BLA-NAc stimulation in the RtPP assay A) Representative heat maps of RtPP results. B-C) Animals expressing ChR2 spent significantly more time in the stimulation-paired (ON) compared to non-paired (OFF) side in RtPP assay (ns p = 0.9083, ****p < 0.0001) without any effect on total distance travelled (unpaired t-test, p = 0.0568). YFP n=5, ChR2 n=9 (B, C). **D-G**) Effects of BLA-NAc stimulation on Ensure seeking behavior. **D)** In a modified 3-chamber task a sipper bottle of Ensure was added to one chamber while stimulation was delivered to animals. **E)** Activation of the BLA-NAc circuit did not alter time spent in in the chamber with (***p = 0.0003, ****p = 0.0001) (F) close to (*p = 0.0102, **p = 0.0037), or G) time spent drinking (****p < 0.0001, *p = 0.0128) the nutrition shake in ChR2-expressing animals compared to YFP-expressing animals. YFP n=12, ChR2 n=9 (E-G). Data analyzed via unpaired, two-tailed t-test (C), and Two-Way Mixed-Effects ANOVA with Sidak's multiple comparison post-hoc tests (B, E-G), with P and F values for Chamber x Virus interaction shown in (B, E-G).

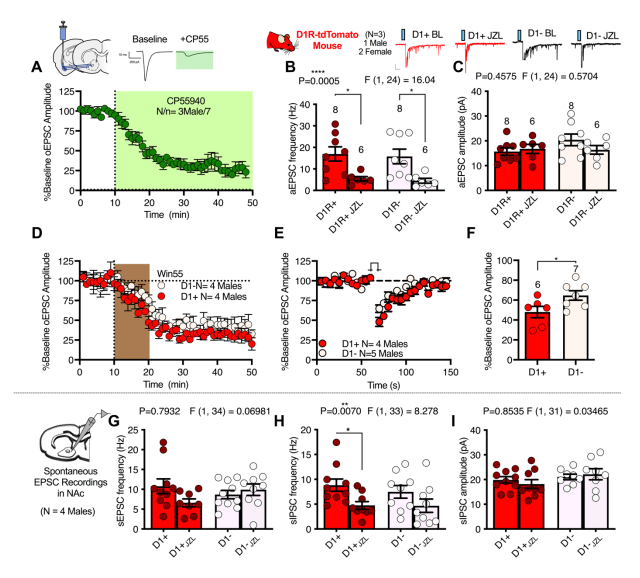


Figure 4: CB1 receptors and 2-AG augmentation regulate BLA-NAc synapses. A) The cannabinoid receptor agonist CP55940 decreased oEPSC amplitude in NAc; N= 3. **B)** JZL184 reduced aEPSC frequency in D1R+ (*p = 0.0171) and D1R- cells (*p = 0.0197), **(C)** but did not affect aEPSC amplitude after BLA-NAc optogenetic stimulation; D1+ N=3, D1- N=3 (B, C). **(D)** WIN55212-2 (Win55), a cannabinoid receptor agonist, uniformly decreased oEPSC amplitude in D1+ and D1- NAc cell types; D1+ N=4, D1- N= 4. **E)** DSE was present in both D1+ and D1- NAc cells, **F)** although DSE magnitude was increased in D1+ compared to D1- cells (*p = 0.0450); D1+ N=4, D1- N= 5 (E, F). **G)** sEPSC frequency in NAc recordings were unaffected by JZL184, **H)** but there was a significant effect of JZL184 on sIPSC frequency (*p = 0.0400). **I)** There are no effects of JZL184 on sIPSC amplitude; D1+ N=4, D1- N= 4. Data analyzed via Two-Way Mixed-Effects ANOVA followed by Sidak's multiple comparisons test (B-C, G-I) and unpaired two-tailed t-test (F). P and F values for Drug Effect shown in (B-C and G-I).

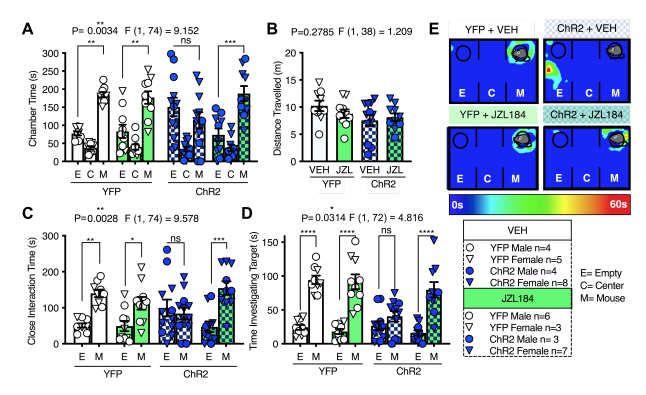


Figure 5: JZL184 pretreatment blocks BLA-NAc activation-induced decreases in sociability A) YFP-expressing animals that received JZL184 or VEH treatment showed a social preference (Veh: **p = 0.0015, JZL: **p = 0.0041), (C) spent more time in close interaction zone (Veh: **p = 0.0032, JZL:*p = 0.0429), and (D) and increased time investigating mouse, relative to empty, target (Veh: **** p <0.0001, JZL **** p <0.0001). ChR2 animals pre-treated with VEH had no A) social chamber preference (ns p = 0.4699), C) preference for time spent near the target (ns p = 0.9302), or D) increased time investigating mouse, relative to empty, target (ns p = 0.5315). ChR2 animals treated with JZL184 have a significant preference for the A) mouse chamber (***p = 0.0003), C) and spend increased time in close interaction (***p = 0.0001) and D) investigating the mouse, relative to empty, target (****p < 0.0001). B) There was no effect of virus or drug treatment on distance traveled. E) Representative heat maps of each treatment condition. YFP + VEH n=9, YFP + JZL184 n=9, ChR2 + VEH n=12, ChR2 + JZL184 n=10 (A-D). Data analyzed via Three-Way Mixed-Effects ANOVA followed by Sidak's multiple comparisons test (A, C-D) or Two-Way ANOVA with Sidak's multiple comparisons test (B). P and F values for Chamber x Virus x Drug Interaction shown in (A, C-D) or Virus x Drug (B).

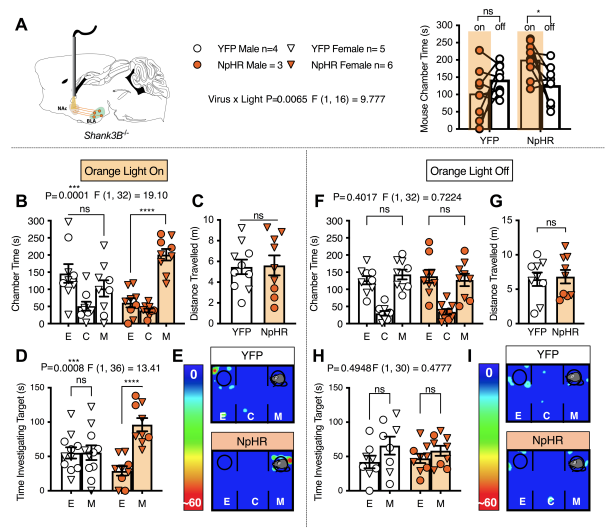


Figure 6: Inhibition of the BLA-NAc circuit normalized SI deficits in Shank3B^{-/-} mice. A) Shank3B^{-/-} mice expressing NpHR in the BLA with bilateral orange light stimulation delivered to the NAc spend significantly more time in the mouse chamber during orange light delivery (ON) compared to light-OFF conditions (*p = 0.0230). YFP animals do not show a preference for the mouse chamber (ns p = 0.2529). **B)** Shank $3B^{-1}$ mice expressing YFP under light ON conditions do not exhibit social preference (ns p = 0.2831), while animals that express NpHR have a preference for the mouse chamber (****p < 0.0001). **D)** Mice that express NpHR have a significant increase in time investigating the mouse cup under light ON conditions (****p < 0.0001). C) No effect on distance traveled was observed (ns p = 0.9057) in light ON paradigm. E) Representative heat maps under light ON conditions. F) Neither YFP (ns p = 0.7166) nor NpHR (ns p = 0.8731)-expressing Shank3B^{-/-} showed mouse-chamber preference H) or preference for time investigating mouse target (YFP ns p = 0.1617, NpHR ns p = 0.6193) under light OFF conditions. G) There are no effects on distance travelled under light OFF conditions (p = 0.7959). I) Representative heat maps under light OFF conditions. YFP n=9, NpHR n=9 (B-D, F-H). Data analyzed via Two-Way Mixed effects ANOVA with Sidak's multiple comparisons test (A, B, D, F, H) or unpaired, two-tailed t-test (C, G). P and F values for Light x Virus interaction shown in (A) and Chamber x Virus interaction shown in (B, D, F, H).

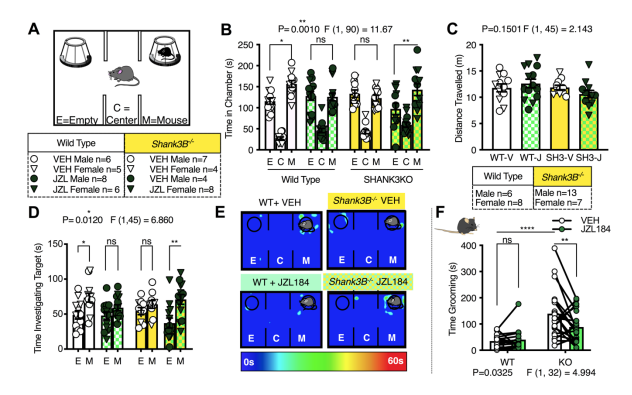


Figure 7: Systemic JZL184 treatment reverses the core behavioral abnormalities in Shank3B^{-/-} animals: A) Diagram of 3-chamber SI task. B) VEH treatment did not affect SI in WT mice (**p = 0.0242), and did not induce a preference in Shank3B^{-/-} (ns p = 0.9360), while JZL184 eliminated social preference in WT mice (ns p = 0.9998) but resulted in significant social preference in Shank3B^{-/-} animals (**p = 0.0094). C) There is no difference in distance travelled between groups. D) JZL184 treated Shank3B^{-/-} animals (***p = 0.0002) and VEH treated WT mice (*p = 0.0155) had preference for investigating the mouse, over empty, target while JZL184 WT (ns p = 0.7103) and VEH Shank3B / mice (ns p = 0.0.9529) did not; WT + VEH n=11, WT + JZL n=14, KO + VEH n=11, KO + JZL n=12 (B-D). E) Representative heat maps of 3-chamber SI task. F) Shank3B^{-/-} mice treated with VEH spend significantly more time grooming compared to WT VEH animals (****p < 0.0001). Shank3B^{-/-} mice treated with JZL184 spent significantly less time grooming compared to VEH treatment (**p = 0.0069) and WT VEH treatment, while JZL184 had no effect in WT mice (ns p = 0.9548); WT n=14, KO n=20. Data analyzed via Three-Way Mixed effects ANOVA with Sidak's multiple comparisons test (B, D), Two-Way ANOVA with Sidak's multiple comparisons test (C), or Two-way RM ANOVA with Sidak's Multiple Comparisons (F). P and F values for Chamber x Genotype x Drug interaction shown in (B, D) or Genotype x Drug interaction (C, F).

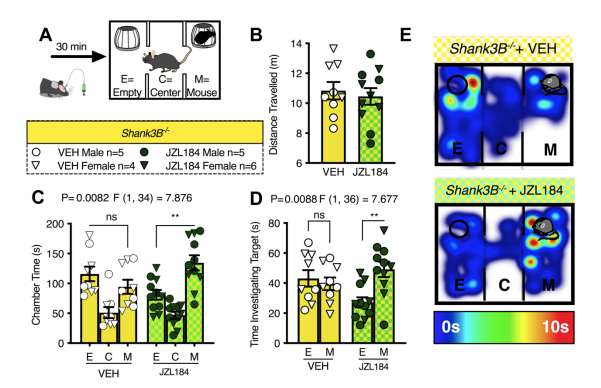


Figure 8: JZL184 microinfused into the NAc reverses the sociability deficit in *Shank3B*-/- **animals: A)** Diagram of microinfusion experimental design. **B)** JZL184 administered to the NAc does not alter distance traveled. **C)** VEH-treated *Shank3B*-/- animals did not exhibit a preference for the mouse chamber (ns p = 0.8566), while JZL184 increased social chamber preference (**p = 0.0014). **D)** Additionally, VEH -treated *Shank3B*-/- did not have a preference for investigating the mouse cup while JZL184 increased investigation time (**p = 0.0022); VEH n=9, JZL184 n=11 (B-D). **E)** Representative heat maps of 3-chamber SI task. Data analyzed via unpaired, two-tailed t-test (B) or Two-Way ANOVA with Sidak's multiple comparisons test (C, D). P and F values for Chamber x Drug interaction shown in (C, D).

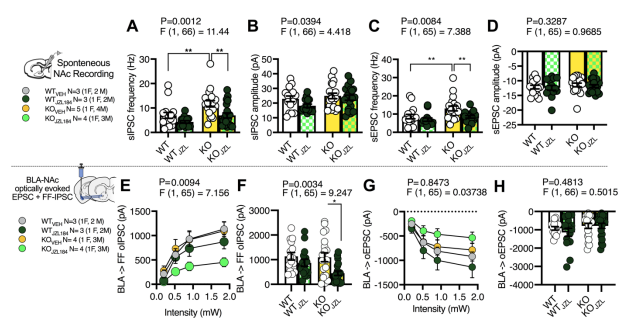


Figure 9: JZL184 significantly reduces sIPSC frequencies in the NAc and BLA-NAc feedforward inhibition in *Shank3B*-/ **mice**-: **A)** sIPSC frequency on the NAc MSNs is significantly increased in *Shank3B*-/- compared to WT mice (**p = 0.0060) and is restored by JZL184 (**p = 0.0033). **B)** There is no effect of *Shank3B*-/- on sIPSC amplitude. **C)** sEPSC frequency is significantly increased in *Shank3B*-/- mice relative to WT animals (**p = 0.0084) and is restored by JZL184 (** p = 0.0060). **D)** There was no effect of genotype or drug on sEPSC amplitude; WT + VEH N=3, WT + JZL184 N=3, KO + VEH N=5, KO + JZL184 N=4 (A-D). **E)** There was no difference between WT and *Shank3B*-/- BLA feedforward (FF) oIPSCs onto NAc cells, However **F)** at maximal stimulation (1.85mW) JZL184 significantly reduces BLA FF oIPSCs onto NAc MSNs (*p=0.0101). **G-H)** There is no effect of JZL184 or genotype on BLA oEPSC in the NAc across intensities; WT + VEH N=3, WT + JZL184 N=3, KO + VEH N=4, KO + JZL184 N=4 (E-H). Data analyzed via Two-Way ANOVA with Sidak's multiple comparisons test (A-H), P and F values for drug effect shown in (A-H). M= males, F= females.

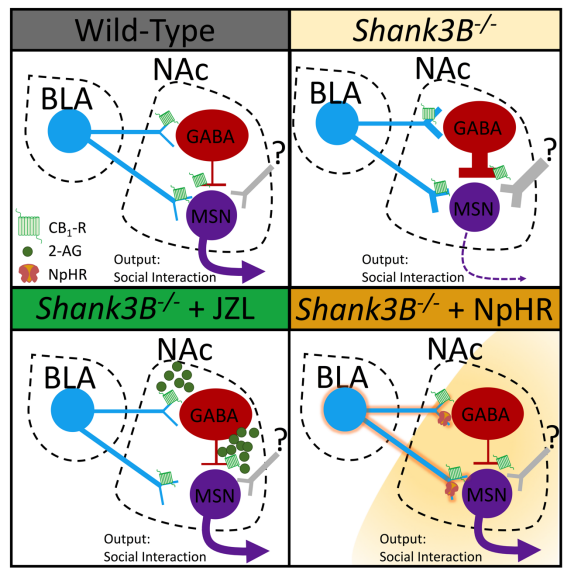


Figure 10: Working model of eCB-BLA-NAc interactions in *Shank3B*^{-/-} **mice.** *Shank3B*^{-/-} mice exhibit increased sEPSC and sIPSC frequency onto MSNs in the NAc relative to WT mice and exhibit social impairment. In *Shank3B*^{-/-} mice treated with systemic or intra-NAc JZL184 to increase 2-AG levels, sEPSC and sIPSC frequency is normalized as is BLA-NAc FF-inhibition, resulting in a reduction in social deficits. *Shank3B*^{-/-} mice with optogenetic inhibition of BLA-NAc circuit function are hypothesized to have similarly reduced FF-inhibition and reduced overall glutamatergic drive to GABA interneurons and MSNs, which also reduces social deficits.