bFGF expression is differentially regulated by cocaine seeking versus extinction in learning-related brain regions

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In cocaine use disorder, relapse can be elicited by drug-associated cues despite long periods of abstinence. The persistence of drug-associated cues in eliciting drug seeking suggests enduring changes in structural and functional plasticity, which may be mediated by basic fibroblast growth factor (bFGF, FGF2). Stimulant drug use increases bFGF expression in reward- and learning-related brain regions, such as the infralimbic medial prefrontal cortex (IL-mPFC), and we previously found that this increase was reversed by extinction. However, whether bFGF expression is similarly modified in other brain regions is unknown. Therefore, we used the conditioned place preference (CPP) paradigm to assess bFGF expression following cocaine-associated CPP or extinction of that CPP within the mPFC, nucleus accumbens (NAc), hippocampus (Hipp), and basolateral amygdala (BLA). bFGF expression was increased in IL-mPFC and NAc-Core and -Shell following a cocaine-associated CPP, an effect reversed by extinction. Conversely, bFGF expression was increased in BLA following extinction, but no significant changes were observed in PL-mPFC or either dorsal or ventral Hipp. These results demonstrate differential regulation of bFGF following cocaine-associated CPP or extinction of that CPP in discrete brain regions. Changes in bFGF expression may regulate long-lasting drug-induced plasticity that underlies persistent drug-associated memories, and therefore present potential prophylactic targets.

There are currently no FDA-approved treatments for cocaine use disorder, and individuals who abuse cocaine remain vulnerable to relapse despite treatment or long periods of abstinence (McEllan et al. 2000). This long-lasting vulnerability is maintained by exposure to cues associated with the drug, which can promote craving (Ehman et al. 1992) and lead to relapse (Kosten et al. 2006). Although the longevity of drug-associated memories suggests persistent plastic changes, characterizations of such changes are lacking. Drug use can induce increases in neurite outgrowth and spine density in brain regions such as the prefrontal cortex (PFC) and nucleus accumbens (NAc; Robinson and Kolb 1997, 1999). However, whether such changes directly subserve persistent drug-associated memories, or result from drug exposure generally, is currently unknown.

One probable regulator of drug-induced neural plasticity is basic fibroblast growth factor (bFGF, FGF2). Repeated stimulant administration increases bFGF expression in the infralimbic medial PFC (IL-mPFC), Hafenbreidel et al. (2015), striatum, hippocampus (Maggio et al. 1998; Roceri et al. 2001; Fumagalli et al. 2006), and ventral tegmental area (VTA; Flores et al. 1998). As a growth factor, the drug-induced increase in bFGF expression may promote plastic changes. For example, in VTA, bFGF is necessary for amphetamine-induced neurite outgrowth (Mueller et al. 2006), which is a persistent form of plasticity seen in other reward-related brain regions (Robinson and Kolb 1997, 1999). Besides drug-induced plasticity, bFGF is required for learning and memory. Increased bFGF mRNA in the dentate gyrus corresponds with learning (Gómez-Pinilla et al. 1998), and bFGF administration improves learning and alleviates memory-related cognitive impairments (Srivastava et al. 2008). Moreover, bFGF facilitates both the acquisition (Graham and Richardson 2009b) and extinction (Graham and Richardson 2009a) of contextual fear conditioning, and reduces reinstatement when administered immediately following extinction either systemically (Graham and Richardson 2010) or directly within the basolateral amygdala (BLA; Graham and Richardson 2011b). Thus, bFGF has an important role not only in drug-induced structural changes, but in learning and memory as well.

Although bFGF is important for learning and memory, stimulant drug use results in an overexpression of bFGF (Flores et al. 1998; Fumagalli et al. 2006; Hafenbreidel et al. 2015), which can have behavioral consequences. bFGF is required for amphetamine-induced behavioral sensitization (Flores et al. 2000) and neutralizing bFGF in IL-mPFC facilitates extinction of cocaine seeking following self-administration (Hafenbreidel et al. 2015). Interestingly, extinction itself can reverse drug-induced increases in bFGF expression in IL-mPFC (Hafenbreidel et al. 2015). However, whether other reward- and learning-related brain regions demonstrate similar changes in bFGF expression or whether extinction can ameliorate these changes is unknown. Therefore, we used the cocaine conditioned place preference (CPP) paradigm, wherein rats were conditioned to associate a distinct context with the rewarding effects of the drug, to examine changes in bFGF expression within reward- and learning-related brain circuitry following both conditioning and extinction.

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Results

Extinction reverses cocaine-induced increases in bFGF expression in IL-mPFC and NAc

The effects of extinction of cocaine CPP on bFGF expression were examined by measuring bFGF immunoreactivity in reward- and learning-related brain regions. Following an initial pretest, rats were matched into three groups with no overall chamber bias. Next, rats were conditioned to associate one chamber, but not another, with cocaine (10 mg/kg, i.p.), and a control group was conditioned to associate both chambers with saline. Following conditioning, some rats (Sal-Ext and Coc-Ext) underwent two 30-min extinction sessions, whereas others (Coc-NoExt) remained in their home cages (Fig. 1A). To determine if rats demonstrated a preference for the cocaine-paired chamber following conditioning, extinction day 1 was analyzed with a one-way ANOVA. As expected, following conditioning, Sal-Ext rats did not express a CPP during the first extinction session (Fig. 1B; $F_{(2,9)} = 1.38, P = ns$). Coc-NoExt rats did not undergo extinction testing (Fig. 1C; denoted in the figure as blank CPP extinction sessions). In contrast, Coc-Ext rats expressed a CPP for the previously cocaine-paired chamber during the first extinction session (Fig. 1D; $F_{(2,15)} = 9.92, P = 0.002$). Post hoc tests confirmed that rats spent more time in the previously cocaine-paired chamber than saline-paired chamber ($P = 0.007$), indicating that rats expressed a cocaine CPP. Next, to determine if rats demonstrated extinction, extinction day 2 was analyzed with a one-way ANOVA. During the second extinction session, Sal-Ext rats again did not express a significant CPP ($F_{(2,9)} = 4.11, P = ns$). Similarly, Coc-Ext rats did not express a CPP during the second extinction session ($F_{(2,15)} = 0.40, P = ns$), demonstrating that extinction had occurred.

The next day, all rats were tested for extinction during a 30-min session (Fig. 1E). Sal-Ext rats again did not express a significant CPP, as a one-way ANOVA revealed no significant effect of chamber ($F_{(2,9)} = 1.20, P = ns$). Similarly, Coc-Ext rats did not express a cocaine CPP, demonstrating extinction recall, as a one-way ANOVA revealed no significant effect of chamber ($F_{(2,15)} = 2.11, P = ns$). However, Coc-NoExt rats did express a cocaine CPP, as a one-way ANOVA revealed a significant effect of chamber ($F_{(2,15)} = 9.30, P = 0.002$). Post hoc tests confirmed that Coc-NoExt rats spent more time in the previously cocaine-paired chamber than in the saline-paired chamber during the extinction test ($P = 0.001$), indicating that rats expressed a cocaine CPP. Importantly, Coc-NoExt rats did not demonstrate within session extinction when the 30-min extinction test was analyzed in 3-min bins (Fig. 1F). A two-way repeated measures ANOVA revealed no significant effect of bin ($F_{(9,270)} = 0.01, P = ns$), or chamber by bin interaction ($F_{(18,540)} = 0.79, P = ns$). Following the extinction test, all rats were immediately euthanized and brain tissue was collected and prepared for bFGF immunohistochemical analysis.

Figure 1. Extinction and extinction test prior to quantifying bFGF-immunoreactivity. (A) Schematic showing design of CPP conditioning and testing. (B) Sal-Ext rats had no significant preference for either saline-paired chambers during extinction day 1 and 2. (C) Coc-NoExt rats remained in their home cages while the other two groups underwent extinction, and therefore did not undergo extinction or receive exposure of the conditioning contexts. No behavioral data was collected for these rats during extinction day 1 and 2 (denoted as a blank space). (D) Coc-Ext rats had a significant preference for the previously cocaine-paired chamber during extinction day 1, but not extinction day 2. (E) Coc-NoExt rats had a significant preference for the previously cocaine-paired chamber during the extinction test, whereas Sal-Ext and Coc-Ext rats did not. (F) Coc-NoExt rats do not demonstrate within session extinction during the extinction test. Error bars are ±SEM. (**) $P < 0.01$. HC, Home Cage.
To determine whether bFGF expression was changed following extinction of cocaine CPP in mPFC, we assessed differences in the density of cells expressing bFGF between groups in the prelimbic mPFC (PL-mPFC) and IL-mPFC (both: Sal-Ext, n = 4; Coc-NoExt, n = 6; Coc-Ext, n = 6). Figure 2A shows representative tissue collection areas (bregma, +3.00 mm; Paxinos and Watson 2007) and Figure 2B shows representative images from each group. In PL-mPFC (Fig. 2C, left), a one-way ANOVA revealed no significant differences in bFGF expression between Sal-Ext, Coc-Ext, and Coc-NoExt rats ($F_{(2,13)} = 2.92, P = \text{ns}$). In IL-mPFC (Fig. 2C, right), a one-way ANOVA revealed a significant difference in bFGF expression between groups ($F_{(2,13)} = 4.28, P = 0.037$). Post hoc tests confirmed that Coc-NoExt rats had significantly higher bFGF expression than Sal-Ext rats ($P = 0.040$) and Coc-Ext rats ($P = 0.020$), but Sal-Ext rats were not different from Coc-Ext rats ($P = \text{ns}$). Thus, bFGF expression was increased following cocaine CPP conditioning in IL-mPFC, but not PL-mPFC, and this increase was reversed by extinction.

Next, to determine whether bFGF expression in NAc-Core or -Shell was changed following extinction, we analyzed expression between groups (both: Sal-Ext, n = 4; Coc-NoExt, n = 6; Coc-Ext, n = 6). Figure 2D shows a representative area of tissue collection for NAc-Core and -Shell (bregma, +2.52 mm; Paxinos and Watson 2007), and Figure 2E shows representative images from each group. In NAc-Core (Fig. 2F, left), a one-way ANOVA revealed a significant difference in bFGF expression between groups ($F_{(2,13)} = 5.77, P = 0.016$). Post hoc tests confirmed that Coc-NoExt rats had significantly higher bFGF expression than Sal-Ext rats ($P = 0.009$) and Coc-Ext rats ($P = 0.020$), but Sal-Ext rats were not different from Coc-Ext rats ($P = \text{ns}$). In NAc-Shell (Fig. 2F, right), a one-way ANOVA revealed a significant difference in bFGF expression between groups ($F_{(2,13)} = 4.32, P = 0.036$). Post hoc tests confirmed that Coc-NoExt rats had significantly higher bFGF expression than Sal-Ext rats ($P = 0.022$) and Coc-Ext rats ($P = 0.034$), but Sal-Ext rats were not different from Coc-Ext rats ($P = \text{ns}$). Thus, bFGF expression was increased following cocaine CPP conditioning in NAc-Core and -Shell, and this increase was reversed by extinction.

bFGF expression is increased in BLA following extinction, but is unchanged in Hipp

To determine whether bFGF expression was altered in BLA following extinction, we analyzed expression between groups (Sal-Ext, n = 4; Coc-NoExt, n = 6; Coc-Ext, n = 6). Figure 3A shows a representative area of tissue collection for BLA (bregma, −2.28 mm; Paxinos and Watson 2007), and Figure 3B shows representative images from each group. In BLA, a one-way ANOVA revealed a significant difference in bFGF expression between groups ($F_{(2,13)} = 9.78, P = 0.002$). Post hoc tests confirmed that Coc-NoExt rats had significantly higher bFGF expression than Sal-Ext rats ($P = 0.002$) and Coc-Ext rats ($P = 0.019$), but Sal-Ext rats were not different from Coc-Ext rats ($P = \text{ns}$). Thus, bFGF expression was increased following cocaine CPP conditioning in BLA, and this increase was reversed by extinction.

Figure 2. bFGF expression was increased in IL-mPFC and NAc-Core and -Shell following acquisition of cocaine CPP, an effect reversed by extinction. Representative areas of tissue collection containing (A) PL-mPFC and IL-mPFC (Sal-Ext, n = 4; Coc-NoExt, n = 6; Coc-Ext, n = 6). (B) Representative image and region of interest (ROI) analyzed from each group containing PL-mPFC and IL-mPFC. (C) bFGF expression was not significantly different between groups in PL-mPFC. However, in IL-mPFC, bFGF expression was increased in Coc-NoExt rats compared to Sal-Ext and Coc-Ext rats. (D) Representative areas of tissue collection containing (A) NAc-Core and -Shell (Sal-Ext, n = 4; Coc-NoExt, n = 6; Coc-Ext, n = 6). (E) Representative image and ROI analyzed from each group containing NAc-Core and -Shell. (F) bFGF expression was increased in Coc-NoExt rats compared to Sal-Ext and Coc-Ext rats, in NAc-Core and -Shell. Error bars are ±SEM. Data normalized to % Sal-Ext. (*) $P < 0.05$. 

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We found that bFGF is differentially expressed in a region-specific manner following cocaine seeking and extinction. In IL-mPFC, NAc-Core, and NAc-Shell, bFGF expression was significantly up-regulated following cocaine CPP conditioning, but expression decreased to baseline levels following extinction. Conversely, in BLA, bFGF expression was unaltered following cocaine CPP conditioning, but was increased following extinction. Finally, no significant changes in bFGF expression were observed in PL-mPFC, dHipp, or vHipp. Overall, these results indicate that bFGF expression is differentially regulated in reward- and learning-related brain regions following drug seeking or extinction, and suggests neural plasticity in circuits mediating different aspects of drug-cue associative learning and extinction.

Consistent with previous findings (Flores et al. 1998; Flores and Stewart 2000; Fumagalli et al. 2006; Hafenbreidel et al. 2015; Flagel et al. 2016), cocaine use and drug-associated learning specifically affected bFGF in IL-mPFC and NAc. These regions have reciprocal projections (Sesack et al. 1989; Groenewegen et al. 1999) and differentially mediate drug seeking. Specifically, projections between PL-mPFC and NAc-Core are thought to drive drug seeking, whereas the projections between IL-mPFC and NAc-Shell are thought to inhibit drug seeking following extinction (McFarland et al. 2003, 2004; Lalumiere and Kalivas 2008; Peters et al. 2009). Here we replicated our previous findings in IL-mPFC (Hafenbreidel et al. 2015), but did not see a significant change in PL-mPFC. It is possible that cocaine CPP increased bFGF expression transiently in PL-mPFC following conditioning, but expression was down-regulated by the time point selected to examine the effect of extinction. Others (Fumagalli et al. 2006) have reported increased bFGF expression in mPFC up to 72 h following five daily cocaine injections, but they did not differentiate between PL-mPFC and IL-mPFC. Similarly, we saw increased bFGF expression in NAc-Core and -Shell following drug seeking (Coc-NoExt), and a reversal of this increase following extinction. These results suggest, despite the putatively opposing roles of NAc subregions, that bFGF expression is regulated similarly in the NAc-Core and -Shell during drug seeking and extinction.

Interestingly, we found that bFGF expression was increased in BLA following extinction. The BLA is necessary for reconsolidation, or restabilization, of drug-associated memories following retrieval (Otis et al. 2013), drug-associated memory storage (Young et al. 2016), and stress- or cue-induced reinstatement (Erb et al. 2001; Kantak et al. 2002; Leri et al. 2002; McLaughlin and See 2003; McFarland et al. 2004; Hiranita et al. 2006), but not cocaine-induced reinstatement (McFarland and Kalivas 2001) or extinction in a self-administration paradigm (Peters et al. 2008). Thus, our results demonstrating the increased bFGF expression following extinction, but not following drug seeking, in BLA were surprising. However, BLA does have a role in extinction of conditioned fear (Quirk and Mueller 2008; Maren 2015), and infusions of bFGF into BLA can facilitate extinction (Graham and Richardson 2011a,b). These results, in combination with ours, suggest that bFGF has a unique role in BLA during extinction of behaviors driven by associative memories.

Our lack of significant findings in dHipp was also surprising, considering the important role of dHipp in contextual learning (Fanselow and Dong 2010) and reported increases in dHipp bFGF expression following psychostimulant administration (Fumagalli et al. 2006). These increases are transient, however, as bFGF is increased for only 2 h following five daily cocaine injections, and expression is not significantly different by 72 h after the final injection (Fumagalli et al. 2006). Our rats were euthanized ~72–96 h

Figure 3. bFGF expression was increased in BLA following extinction. (A) Representative areas of tissue collection containing BLA. (B) Representative image and ROI analyzed from each group containing BLA (Sal-Ext, n = 4; Coc-NoExt, n = 6; Coc-Ext, n = 6). (C) bFGF expression was significantly increased in Coc-Ext rats compared to Sal-Ext and Coc-NoExt rats, in BLA. Error bars are ±SEM. Data normalized to % Sal-Ext. (** P < 0.01.)
after the final cocaine injection (temporal variability arose due to the counterbalanced sequence of cocaine or saline administration on the final conditioning day, in addition to 2 d of extinction and 1 d for the extinction test). Therefore, we likely missed the transient increase in expression following cocaine injections during conditioning. Furthermore, our data suggest that neither cocaine-associated learning nor extinction in the CPP paradigm result in significant prolonged changes to bFGF expression in dHipp or vHipp.

Growth factors, such as bFGF, are associated with stimulant-induced structural and functional neural plasticity in reward- and learning-related brain regions. These plastic changes may sustain long-lasting drug-associated memories and therefore promote persistent drug-seeking behavior. For example, bFGF can enhance neurite outgrowth following stimulant administration (Mueller et al. 2006), increase expression of GluR1-containing AMPA receptors (Cheng et al. 1995; Chew et al. 1997), and promote long-term potentiation (Ishiyama et al. 1991; Zhao et al. 2007). bFGF can conversely also decrease NMDA receptor expression (Mattson et al. 1993), inhibit voltage-gated Na+ (Hilborn et al. 1998) and K+ currents, and decrease evoked action potentials (Cuppini et al. 2009). Together, these findings indicate that the effects of bFGF can be varied and the collective influence of bFGF on neural plasticity can be nuanced. Functionally, systemic bFGF administration can facilitate later acquisition of cocaine self-administration (Turner et al. 2009), and bFGF in VTA is necessary for amphetamine-induced increases in locomotion and sensitization (Flores et al. 2000). Moreover, neutralizing bFGF in IL-mPFC can facilitate extinction following cocaine self-administration (Hafenbreidel et al. 2015), and neutralizing bFGF in the dorsal striatum can reduce alcohol seeking and reinstatement (Even-Chen et al. 2017). However, intra-BLA bFGF infusions can facilitate extinction of conditioned fear (Graham and Richardson 2011a,b). Whether bFGF induces neural plasticity that promotes (e.g., facilitated extinction of conditioned fear) or disrupts new learning (e.g., perseverative drug seeking) appears to be region specific (e.g., BLA versus IL-mPFC).

As suggested, certain reward- and learning-related brain regions may be specifically vulnerable to changes in bFGF elevations (Fig. 5), which could drive long-term plasticity and persistent drug seeking. In agreement with our previous findings (Hafenbreidel et al. 2015), we found increased bFGF expression in IL-mPFC and NAc-Core and -Shell following cocaine CPP conditioning (Fig. 5, left, black), which suggests that bFGF-associated adaptations within these brain regions promotes the expression of cocaine CPP. Similarly, we also found that bFGF expression was decreased in these brain regions following extinction (Fig. 5, right, white).

**Figure 4.** bFGF expression was unchanged in dHipp and vHipp. (A) Representative areas of tissue collection containing dHipp (Sal-Ext, n = 4; Coc-NoExt, n = 5; Coc-Ext, n = 6). (B) Representative image and ROI analyzed from each group containing dHipp. (C) bFGF expression was not significantly different between groups in dHipp. (D) Representative areas of tissue collection containing vHipp (Sal-Ext, n = 4; Coc-NoExt, n = 5; Coc-Ext, n = 6). (E) Representative image and ROI analyzed from each group containing vHipp. (F) bFGF expression was not significantly different between groups in vHipp. Error bars are ±SEM. Data normalized to % Sal-Ext.
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Materials and Methods

Subjects

Male Long-Evans rats (Harlan) weighing 175–200 g at study onset were housed and handled as previously described (Otis and Mueller 2011). Protocols were approved by the Institutional Animal Care and Use Committee at the University of Wisconsin-Milwaukee, in compliance with National Institute of Health guidelines.

Drugs

Cocaine HCl (NIDA) was dissolved in sterile 0.9% saline at a concentration of 10 mg/mL, and administered at a dose of 10 mg/kg (i.p.).

Place preference apparatus

Behavioral testing and conditioning were conducted as previously described (Mueller and Stewart 2000; Otis and Mueller 2011). Briefly, a three-chamber apparatus was used, in which two contextually distinct larger conditioning chambers were separated by a smaller center chamber. Rats had access to all three chambers during pre- and post-conditioning sessions, but were isolated to one of the larger chambers during conditioning. Time was recorded during CPP trials by four infrared photobeams located in the larger chambers, and total photobeam breaks made in all three chambers were used to record locomotor activity. During all phases of the experiments, the room and chambers were kept in semidarkness.

Behavioral testing

Baseline preferences were determined during a pretest as previously described (Otis and Mueller 2011). Briefly, during a pretest, rats were placed into the center chamber with free access to all three chambers for 15-min, and time spent in each chamber was recorded. A one-way ANOVA revealed a significant effect of chamber ($F_{2,45} = 99.046, P < 0.0001$), and post hoc analysis confirmed that an equivalent amount of time was spent in the conditioning chambers ($P > 0.05$), but less time was spent within the center chamber than in either conditioning chamber ($P < 0.0001$). As rats spent equivalent time in the conditioning chambers, but less in the center chamber, an unbiased procedure was used. Additionally, all three groups displayed similar locomotor behavior during the pretest before conditioning, as a one-way ANOVA did not reveal a significant difference between groups in total photobeam breaks made in all three chambers ($F_{2,11} = 2.51, P = 0.120, P = ns$). Rats were matched into two groups, in which rats were conditioned to associate one chamber, but not the other, with cocaine in a counterbalanced manner over 8 d. An additional control group was included, which was conditioned to associate both chambers with saline. Cocaine or saline injections were given immediately prior to each 20-min conditioning session, during which the rats were confined to their respective chambers.

To determine the effect of extinction of drug seeking on bFGF protein expression, following conditioning, saline-treated controls and half of the cocaine-treated group were given two daily CPP sessions (Sal-Ext, Coc-Ext, respectively) during which they were placed in the center chamber and given free access to the entire apparatus for 30-min to induce extinction. The other half of the experimental group was home-cage confined (Coc-NoExt). A cocaine CPP was defined when significantly more time was spent in the previously cocaine-paired chamber. Following the last extinction day, all rats were tested for extinction during a 30-min session. Immediately following the extinction test, rats were overdosed with sodium pentobarbital (120 mg/kg, i.p.) and perfused transcardially with 150 mL of phosphate-buffered saline (PBS) followed by 150 mL of 4% paraformaldehyde in 0.1 M PB solution. Brains were extracted and stored at 4°C in 4% paraformaldehyde overnight, then submerged in 30% sucrose for subsequent immunohistochemistry (IHC) processing and analysis.

Immunohistochemistry and imaging

Coronal sections (50 µm) from each group that contained mPFC, NAc, BLA, dHipp, and vHipp were collected and processed with IHC to measure bFGF cellular expression, as previously described (Mueller et al. 2006). Briefly, sections were quenched in 30% hydrogen peroxide and phosphate buffer (PB), preblocked in normal goat serum and triton TBS for 30-min, and incubated in a 1:500 concentration of primary anti-bFGF (Millipore) overnight. Next,
sections were incubated in anti-mouse secondary for 1-h, followed by incubation in Vectastain standard ELITE ABC kit for 2-h, before undergoing staining with DAB and NiCl2. Processed sections were imaged under a Nikon Eclipse 55i microscope using Nikon NIS Elements software. Next, regions of interest were defined (Paxinos and Watson 2007), and the total number of bFGF-immunoreactive cells in each region was quantified bilaterally in a semiautomated manner across three sections to calculate average cell densities per region. Researchers were blind to conditions throughout the analysis. Data are expressed as percent immunoreactivity relative to Sal-Ext.

Data analysis

Drug-seeking behavior during each CPP session was analyzed by comparing time spent in the previously cocaine-paired, saline-paired, and center chambers using a one-way ANOVA. To examine within session behavior, a two-way repeated measures ANOVA was used to examine differences in time spent in each chamber during 3-min bins. For IHC analysis, average cell densities (number of bFGF expressing cells per mm²) were analyzed using a one-way ANOVA. Due to tissue damage, some samples were discarded. Final group sizes for each brain region are reported in their respective results section and figure captions. Locomotor activity (total photobeam breaks made in all three chambers during the pretest session) was analyzed using a one-way ANOVA. All post hoc tests were conducted, when appropriate, using Fisher’s least significant difference (LSD) test.

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