Disorders of the Nervous System

Chemogenetic Inhibition of Infrahlimbic Prefrontal Cortex GABAergic Parvalbumin Interneurons Attenuates the Impact of Chronic Stress in Male Mice

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Abstract

Hypofunction of the prefrontal cortex (PFC) contributes to stress-related neuropsychiatric illnesses. Mechanisms leading to prefrontal hypoactivity remain to be determined. Prior evidence suggests that chronic stress leads to an increase in activity of parvalbumin (PV) expressing GABAergic interneurons (INs) in the PFC. The purpose of the study was to determine whether reducing PV IN activity in the Infrahlimbic (IL) PFC would prevent stress-related phenotypes. We used a chemogenetic approach to inhibit IL PFC PV INs during stress. Mice were first tested in the tail suspension test (TST) to determine the impact of PV IN inhibition on behavioral responses to acute stress. The long-term impact of PV IN inhibition during a modified chronic variable stress (CVS) was tested in the forced swim test (FST). Acute PV IN inhibition reduced active (struggling) and increased passive coping behaviors (immobility) in the TST. In contrast, inhibition of PV INs during CVS increased active and reduced passive coping behaviors in the FST. Moreover, chronic inhibition of PV INs attenuated CVS-induced changes in Fos expression in the prelimbic cortex (PrL), basolateral amygdala (BLA), and ventrolateral periaqueductal gray (vPAG) and also attenuated adrenal hypertrophy and body weight loss associated with chronic stress. Our results suggest differential roles of PV INs in acute versus chronic stress, indicative of distinct biological mechanisms underlying acute versus chronic stress responses. Our results also indicate a role for PV INs in driving chronic stress adaptation and support literature evidence suggesting cortical GABAergic INs as a therapeutic target in stress-related illnesses.

Key words: DREADDs; GABA; interneurons; parvalbumin; prefrontal cortex; stress coping

Significance Statement

Stress-related diseases are associated with prefrontal hypoactivity, the mechanism of which is currently not known. In this study we showed that by inhibiting prefrontal GABAergic parvalbumin interneurons (PV INs), we can attenuate some of the chronic stress-related phenotypes. Additionally, we showed that modulation of PV IN activity during acute stress had opposing effects on stress coping strategies, suggesting different plasticity mechanisms in PV INs following acute versus chronic stress. Our findings indicate that GABAergic PV INs may be involved in driving stress-related phenotypes and may therefore be an important target for treatment of stress-related illnesses. Our data suggest that reducing PV IN activity to promote prefrontal output may be a potential treatment strategy for stress-related disorders.

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Introduction

Mood disorders [e.g., posttraumatic stress disorder (PTSD) and major depressive disorder (MDD)] are associated with alterations in ventromedial prefrontal cortex (PFC; Broadman area 25) structure, activity and connectivity (Rogers et al., 2004; Drevets et al., 2008; Hasler et al., 2008; Murray et al., 2011; Holmes et al., 2018). To date, no universally efficacious therapeutic strategy exists for these neuropsychiatric conditions, despite having a lifetime prevalence of over 20% (Kessler et al., 2005; Duman and Duman, 2015). Studies in both humans and animal models have shown that chronic stress impairs functioning of the PFC, potentially making prefrontal hypofunction an important factor in the etiology of mood disorders (Drevets et al., 1997; Radley et al., 2006; Duman and Duman, 2015; Li et al., 2011; McKlevene et al., 2016).

Various clinical and preclinical studies implicate altered GABAergic circuitry and prefrontal hypofunction in the generation of depression in humans as well as depression-related behaviors in rodent chronic stress models (Luscher et al., 2011; Duman, 2014; Veeraiah et al., 2014; Musaazi et al., 2015; McKlevene et al., 2016). Recent functional and electrophysiological studies indicate increased infralimbic (IL) PFC (rodent homolog of the human ventromedial PFC) GABAergic transmission (e.g., increased inhibitory synaptic drive and increased expression of GABAergic marker) following chronic variable stress (CVS). These findings suggest that enhanced interneuron (IN) activity may be involved in disruption of prefrontal cortical signaling, leading to over inhibition and prefrontal hypofunction (McKlevene et al., 2016; Shepard et al., 2016; Page et al., 2019).

GABAergic parvalbumin INs (PV INs) synapse onto cell bodies of PFC pyramidal neurons and exert strong control over medial PFC (mPFC) output, maintaining appropriate excitatory/inhibitory (E/I) balance (Cardin et al., 2009; Courtin et al., 2014a; Tremblay et al., 2016) and coordinating oscillatory activity (γ oscillation) required for efficient PFC signaling. Consequently PV INs are well positioned to play an important role in stress-mediated prefrontal dysfunction (Rymar and Sadikot, 2007; Sherwood et al., 2007; DeFelipe et al., 2013; Ferguson and Gao, 2018). Increases in expression and activity of PV INs and enhancement of glutamatergic transmission onto PV INs following chronic stress are associated with prefrontal hypofunction, anxiogenesis, and impaired coping behaviors in forced swim test (FST) (Shepard et al., 2016; Page et al., 2018, 2019; Shepard and Coutellier, 2019). Reduction in PV expression in PFC is also associated with antidepressant efficacy (Ohira et al., 2013; Zhou et al., 2015; Page and Coutellier, 2019). In contrast to chronic stress, acute inhibition of PV INs in the PFC has opposing effects, resulting in increase in passive coping behavior such as learned helplessness in mice (Perova et al., 2015). This suggests differential role of PV INs in response to acute versus chronic stress, indicative of distinct brain circuitry being involved in modulating acute versus chronic stress-mediated phenotypes.

This study was designed to specifically investigate the role of IL PV INs in driving somatic and behavioral manifestations of acute and chronic stress-related phenotypes. We employed a chemogenetic strategy using designer receptors exclusively activated by designer drugs (DREADDs) to specifically inhibit PV INs in the IL mPFC during exposure to acute stress and throughout exposure to a modified CVS paradigm. Our results indicate that acute inhibition of PV INs increases passive and decreases active coping behavior in tail suspension test (TST). In contrast, chronic inhibition of IL PV INs reduces passive and increases active coping behavior in FST. PV IN inhibition attenuates CVS-mediated reduction in Fos expression in stress-related brain regions downstream of the IL. Additionally, we show that inhibition of PV INs prevents CVS-induced somatic effects such as adrenal hypertrophy and body weight loss. These data suggest that IL PV INs play a role in driving behavioral, neuronal and physiological adaptations associated with chronic stress and also indicate differential role of these INs in the context of acute versus chronic stress-related phenotypes.

Materials and Methods

Mice

Male breeders from B6 PV-Cre knock-in homozygous mouse line (B6;129P2-Pvalbtm1(cre)Arbr/J, JAX stock #017320, The Jackson Laboratory) were bred with WT C57BL/6J females (JAX stock #000664) to generate an in-house colony of heterozygous PV-Cre C57BL/6J at the University of Cincinnati animal housing facility. The PV-Cre mouse line has been characterized extensively in prior publications (Zhao et al., 2019; Cummings and Clem, 2020; Grosisman et al., 2020). Mice were maintained under standard conditions (12/12 h light/dark cycle, 22 ± 1°C, food and water ad libitum; four mice per cage on arrival) in accordance with the University of Cincinnati Institutional Animal Care and Use Committee, which specifically approved all acute and chronic stress regimens employed in this proposal. Mice were single housed following surgeries and continued to be housed singly throughout the duration of the experiment, to prevent aggression and injury to...
animals following surgery and during the stress paradigms (Lidster et al., 2019). Enrichment for housing cages included mouse hut and nestlets. All protocols conformed to the Society’s Policies on the Use of Animals in Neuroscience Research. All experiments were performed on adult male mice (~7.5 months of age at surgery).

**Stereotaxic viral vector injection with AAV vectors**

PV-Cre mice were anaesthetized with isoflurane, scalp shaved and placed in the stereotaxic frame. The incision site was disinfected using chlorohexidine and 70% ethanol. An incision at the midline was made using a single-edged blade. Cre-dependent adeno-associated virus 2 (AAV2) vectors AAV2-hsyn-DIO-hM4Di(Gi)-mCherry (Gift from Bryan Roth; Addgene viral prep #44362-AAV2) and AAV2-hsyn-DIO-mCherry (Gift from Bryan Roth; Addgene viral prep #50459-AAV2) were injected bilaterally at a volume of 300 nl (~10^-12 genome copies/ml) into the IL mPFC. A pilot study was conducted (data not shown) to optimize viral load, volume and stereotactic coordinates to ensure viral spread is restricted to the IL. We specifically chose AAV serotype 2, which has a limited volume of spread enabling us to primarily target the IL and prevent spread to other regions (Burger et al., 2004; Cearley and Wolfe, 2006). The coordinates used were as follows: anterior/posterior range defined as +1.78 mm anterior to bregma, medial/lateral range defined as ±0.2 mm lateral to the midsagittal suture; dorsal–ventral range defined as −2.9 mm ventral to skull (Franklin and Paxinos, 2007). Viruses were infused using a 2-μl Hamilton syringe at a rate of 60 nl/min for 5 min. To ensure that virus injection was restricted to bilateral spread in the IL, we took particular care at a few key points: (1) mouse skulls were always carefully vertically leveled based on bregma and lambda to ensure bilateral spread, (2) Hamilton syringes were always checked to run properly before injections and were carefully cleaned to remove excess virus particles after viral loading to prevent spread when the needle is lowered into the brain, (3) following infusion, the injector was kept at the site for 8 min to allow for the virus to diffuse and prevent spread to other regions, and (4) after injection, the needle was very slowly removed from the injection site to avoid spreading of the virus throughout the injector. The injection site on the skull was covered with gel foam and incision site sutured; 2.5 mg/kg meloxicam was administered for 3 d following surgery. Behavioral studies and stress protocols were initiated three weeks post injection to allow sufficient time for viral expression. Diagrammatic representation of experimental timeline is outlined in Figure 1A.

**Modified CVS procedure**

During the CVS procedure, mice were subjected to a series of randomly alternating stressors administered twice daily over a period of 14 d. The CVS procedure used in this study is a modification of prior published protocols from the literature (Ostrander et al., 2009; Ghosal et al., 2017; Wohleb et al., 2018; Cotella et al., 2019), such that each individual stress session did not last for >2 h to ensure that CNO was on-board throughout the stressor (Jendryka et al., 2019). CNO has been shown to reach maximal plasma levels 30 min after intraperitoneal injection and has been shown to rapidly clear from plasma in rodents (Baldessarini et al., 1993; Guettier et al., 2009; MacLaren et al., 2016; Jendryka et al., 2019). As a result, extended stress periods and overnight stressors typically used in CVS paradigms were excluded from the modified CVS paradigm. Additionally, swim stress was not included in the CVS paradigm since animals were tested in FST (a novel stressor) following completion of the CVS procedure. The unpredictable stressors used were as follows: restraint (30 min), cold room exposure (15 min, 4°C), shaker stress (1 h, 100 rpm), hypoxia (30 min, 8% oxygen and 92% nitrogen), Y maze (8 min), shallow water (30 min), wet bedding (2 h), and cage tilt (2 h, 45°). Body weights were measured on days 1, 4, 8, and 14 during the CVS procedure.

**Drug administration**

Clozapine N-oxide (CNO; NIMH Chemical synthesis and Drug Supply Program) was used as the DREADD activator to activate the inhibitory DREADD. CNO was dissolved in 5% dimethyl sulfoxide (Sigma) and then diluted with 0.9% saline and administered intraperitoneally at a dose of 1 mg/kg twice a day, 30 min before start of each stressor. We chose intraperitoneal injections for CNO delivery to effectively control the dose and timing of DREADDs during our stress paradigm. All animals received chronic injection of CNO for 14 d. Previous work has shown that chronic CNO for 14 d does not lead to desensitization of hM4Di DREADDs, confirming efficacy of chronic activation of DREADDs (Soumier and Sibille, 2014; Xia et al., 2017; Stedehouder et al., 2018; Phillips et al., 2019). A maximum time period of 6 h was given between stressors, to allow sufficient time for CNO to be cleared from the body (MacLaren et al., 2016; Jendryka et al., 2019). To ensure inhibiting PV INs had no effects on locomotor activity, total distance traveled and velocity of mice was measured during Y maze task on day 12th of the CVS regimen.

**Behavioral assessments**

**TST**

The TST (Can et al., 2011) was used as the first stressor in the CVS group to observe acute effects of inhibiting PV INs on passive coping behavior. Mice were suspended 55 cm above ground using a 17 cm long tape that was attached to a suspension bar, for a total time period of 6 min. Sessions were video recorded from the side to allow full body visualization of mice behaviors-active coping (struggling) behavior, which comprised of strong shaking of the body and movement of all four limbs, and passive coping (immobility) behavior which comprised of not making any active limb movements. Latency to reach immobility was also measured. Behaviors were quantified by an experimenter blinded to the group assignments using behavioral scoring software Kinoscope 3.0.4. Behaviors during the 6-min block were reported.
Figure 1. Experimental design and targeting of PV INs in the PFC using DREADDs. A, Experimental design and timeline. C57BL/6J PV-Cre mice —7.5 months of age underwent surgery to inject AAV2-hM4Di-mCherry (inhibitory DREADD) or AAV2-mCherry (control virus). Mice were allowed three weeks to recover to enable sufficient time for DREADD expression. Animals were then subjected to CVS procedure twice a day for 14 d or served as controls. The first stressor was a TST to determine acute effects of PV IN inhibition in animals within the CVS group. Animals were dosed with 1 mg/kg CNO before each stressor to inhibit PV INs during the CVS procedure; 24 h after the end of CVS, animals were subjected to FST, following which mice were euthanized, body was perfused, and brains and organs were collected. No CNO was administered during the testing phase in FST. B, Design of AAV2-hSyn-hM4Di-mCherry (top) and AAV2-hSyn-mCherry (bottom) vectors employing the DIO strategy. Two pairs of heterotypic, antiparallel loxP recombination sites (blue triangles) achieve Cre-mediated transgenes inversion and expression under the control of hSyn promoter. ITR, left-inverted terminal repeat; hSyn, human synapsin; WPRE, woodchuck hepatitis DREADD posttranscriptional regulatory element. C, Successful Cre-mediated recombination of DREADDs demonstrated by presence of red mCherry (left); green fluorescence identifies PV INs (right); hM4Di receptors selectively expressed in PV INs as illustrated by mCherry (red) and PV (green) co-expression (merged, yellow, bottom middle image). Scale bar: 100 μm.
**Euthanasia and tissue collection**

Mice were euthanized with an overdose of sodium pentobarbital after FST, and transcardially perfused with 0.9% saline followed by 4% paraformaldehyde in 0.01M PBS, pH 7.4. Brains were removed and postfixed in 4% paraformaldehyde at 4°C for 24 h, then transferred to PBS, pH 7.4. Thymi and adrenal glands were collected, cleaned, and weighed.

**Immunohistochemistry**

Brains were sectioned into 30-μm coronal sections using a freezing microtome (−20°C). Sections were collected into 12 wells (1/12) containing cryoprotectant solution [30% sucrose, 1% polyvinyl-pyrolidone (PVP-40), and 30% ethylene glycol, in 0.01 M PBS]. Immunohistochemistry was performed at room temperature (RT) and 0.01 m PBS was used to rinse brain slices before each treatment described below.

**PV and DREADD co-localization**

Targeting of IL mPFC PV neurons and recombination of hM4Di DREADD was verified by co-localization of PV immunoreactivity with virally expressed mCherry fluorescence. Free floating sections were incubated in blocking solution [4% normal goat serum (NGS), 0.1% Triton X-100, 0.1% bovine serum albumin (BSA) in 0.01 m PBS] for 1 h at RT. After that, sections were incubated with rabbit anti-PV (1:1000, Abcam, ab11427) overnight, followed by visualization with donkey-anti-rabbit Alexa Fluor 488 conjugate (1:500, Invitrogen, A10343). Images were acquired using Nikon Confocal Microscope at 40× magnification.

**Injection site and viral spread**

To determine whether virus spread was restricted to the IL, sections were incubated with a rabbit anti-mCherry (1:500, Abcam, ab167453) for 2 h, followed by visualization with goat anti-rabbit Cy5 conjugate (1:500, Invitrogen, A10523). Images were acquired using Carl Zeiss Imager Z1 at 2.5× and 5× magnification. Viral spread was mapped onto its respective focal plane and bregma level by outlining the spread of the infection from confocal images onto corresponding brain atlas illustrations (Lein et al., 2007). Fluorescent cells were counted using semi-automated analysis macro in the ImageJ software package (National Institutes of Health). The threshold was adjusted to detect the fluorescence and cells counted using the Analyze Particle feature. The percentage of infected cells in IL was calculated by dividing the number of mCherry-positive cells in IL by the total number of mCherry-positive cells in a specific focal plane. Brain bregma coordinates used for IL cytoarchitecture were defined in the Franklin and Paxinos mouse brain atlas (third edition; Franklin and Paxinos, 2007).

**Fos immunoreactivity.** Neuronal activation was measured using Fos as a marker. Free floating sections were incubated in 1% sodium borohydride for 20 min and then in 3% hydrogen peroxide in PBS for 20 min. After that, slices were incubated in blocking solution (NGS, 0.3% Triton X-100, 0.2% BSA in 0.01 m PBS) for 1 h. Sections were then incubated with Fos rabbit polyclonal antibody (1:200, Santa Cruz, sc-52) in blocking solution overnight and was followed by incubation in secondary antibody (biotinylated goat anti-rabbit, 1:400; Vector Laboratories, BA1000) in blocking solution for 1 h the next day. Sections were then treated with avidin-biotin horseradish peroxidase complex (1:800 in 0.01 m PBS; Vector Laboratories, PK6100) for 1 h and then developed with an 8-min incubation in 3,3′-diaminobenzidine (DAB)-Nickel solution: 10 mg DAB tablet (Sigma, DF905), 0.5 ml of a 2% aqueous nickel sulfate solution, 20 μl of 30% hydrogen peroxide in 50 ml of 0.01 m PBS. Sections were mounted on superfrost slides (Fisherbrand, Fisher), allowed to dry, dehydrated with xylene, and then cover-slipped with DPX mounting medium (Sigma).

Images were acquired using microscope Carl Zeiss Imager Z1 at a 5× objective. For analysis, we counted minimum of three bilateral sections per brain region/animal covering the prelimbic cortex (PrL; bregma 2.80 to 1.98 mm), basolateral amygdala (BLA; bregma −1.06 to −1.58), and ventrolateral periaqueductal gray (vIPAG; bregma −4.16 to −4.36) as defined in the Franklin and Paxinos mouse brain atlas (Franklin and Paxinos, 2007). The number of Fos-positive nuclei was counted using a semi-automated analysis macro in the ImageJ software package (National Institutes of Health). The macro was generated using the Analyze Particle tool, with a defined common level of background intensity, nuclei circularity, and size (previously validated manually). The relative density of the population of immunopositive cells was calculated by dividing the number of Fos-positive cells by the respective brain area.

**Statistical analysis**

The experiment was setup as a 2 × 2 study design, with stress (CVS or No CVS) and DREADD (hM4Di or control) as factors with a sample size of n = 10 per group (for experimental design and timeline, see Fig. 1A). Statistical analyses for FST and Fos protein quantification were performed using a two-way ANOVA with stress (No CVS, research article: New Research 7(5) ENEURO 0423-19.2020 eNeuro.org
CVS) and DREADD (Control, hM4Di) as main factors. TST data were analyzed using Student’s t test. FST measurements over time were done using two-way repeated measure ANOVA with stress (No CVS, CVS) and DREADD (Control, hM4Di) as main factors analyzed over time. Tukey’s post hoc test was performed in cases with significant interaction between factors. Because specific hypotheses were formed a priori on the effects of CVS within groups, planned comparisons using Fisher’s least significant difference (LSD) were performed in cases with no significant interaction effect. Data were analyzed by STATISTICA 7.0 (Statsoft) and GraphPad Prism 8.1.2 (GraphPad Software). Outliers were detected using the Grubbs’ test (GraphPad Software) and removed from analysis. After exclusion of outliers, data were assessed for normal distribution (Shapiro–Wilk) and appropriate parametric and/or non-parametric tests used. Data are presented as mean ± SEM with statistical significance set at p < 0.05. See Table 1 for details regarding data structure and type of test used. Superscript letters listed with p values correspond to the statistical tests shown in Table 1.

### Results

**Selective targeting of PV INs achieved using DREADDs**

The AAV constructs used in this experiment are shown in Figure 1B. Following Cre recombination, the viral

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**Table 1: Data structure, type of test to analyze the data, and observed power of key results**

| Data structure | Type of test | Power |
|----------------|--------------|-------|
| a Normal distribution | Unpaired sample t test (struggling) | Main effect hM4Di DREADD: Cohen’s d = 1.2 |
| b Normal distribution | Unpaired sample t test (immobility) | Main effect hM4Di DREADD: Cohen’s d = 1.3 |
| c Normal distribution | Unpaired sample t test (latency to immobility) | Main effect hM4Di DREADD: Cohen’s d = 1.3 |
| d Normal distribution | Two-way ANOVA (swimming duration) | Main effect of stress: 0.91 Main effect of hM4Di DREADD: 0.55 |
| e Normal distribution | Two-way repeated measures ANOVA (swimming duration over time) | Main effect stress: Power = 0.91 Main effect hM4Di DREADD: Power = 0.55 Main effect of time: Power = 1 |
| f Normal distribution | Two-way ANOVA (immobility duration) | Main effect stress: Power = 0.85 |
| g Normal distribution | Two-way repeated measures ANOVA (immobility duration over time) | Main effect stress: Power = 0.85 Main effect time: Power = 1 |
| h Normal distribution | Two-way ANOVA (PrL) | Main effect stress: Power = 0.98 Interaction (stress × hM4Di DREADD): Power = 0.76 |
| i Normal distribution | Two-way ANOVA (IL) | Main effect stress: Power = 0.99 |
| j Normal distribution | Two-way ANOVA (BLA) | Main effect stress: Power = 0.89 |
| k Normal distribution | Two-way ANOVA (vIPAG) | Main effect stress: Power = 0.94 |
| l Normal distribution | Two-way ANOVA (adrenal weight) | Main effect stress: Power = 0.90 Interaction (stress × hM4Di DREADD): Power = 0.57 |
| m Normal distribution | Two-way ANOVA (thymus weight) | Main effect stress: Power = 1 |
| n Normal distribution | Two-way ANOVA (final body weight) | Main effect stress: Power = 0.70 |
| o Normal distribution | Two-way repeated measures ANOVA (body weight over time) | Main effect stress: Power = 0.6 Main effect time: Power = 0.9 |
| p Normal distribution | Unpaired sample t test (viral spread in IL) | Main effect hM4Di DREADD in IL: Cohen’s d = 1.2 |

Table depicts the data structure, type of statistical test used, and power of key results for each of the statistical tests used in the manuscript. Each analysis includes a letter indicator linking the test in the table to the analysis in the text.
construct expresses inhibitory DREADD sequence hM4Di along with a fluorescent reporter (mCherry), allowing visualization of cells undergoing recombination. Control virus was a Cre-inducible mCherry lacking the DREADD hM4Di construct. Cre-mediated recombination of hM4Di-mCherry and cell type specificity were conferred by immunostaining. Expression of red hM4Di-mCherry demonstrates successful recombination (Fig. 1C, left). hM4Di-mCherry expression was restricted to PV INs only (Fig. 1C, right), confirmed by colocalization of red mCherry with green (Alexa Fluor 488) PV immunostaining, resulting in yellow cells in the merged image (Fig. 1C, bottom).

Targeting of IL and viral spread

Figure 2A depicts viral injection site and spread of virus determined by the presence of mCherry and far-red (Cy5) fluorescence in animals injected with hM4Di DREADD (n = 9). It shows the stereotaxic injection sites and that AAV DREADD expression was primarily restricted to the IL, with some minor spread into the PrL and dorsal peduncular (DP) cortex located above and below IL, respectively. Injection sites and viral spread were located primarily within the bregma range 1.78–1.54 (Franklin and Paxinos, 2007). Figure 2B is a representative image from an animal demonstrating red mCherry fluorescence primarily in the IL delineated by white dashed lines. Quantification of percentage of transfected neurons from animals injected with hM4Di DREADD, revealed expression was significantly contained, with ~72% of transfected cells localized in the IL (t = 4.6, df = 16, p = 0.0003; Fig. 2C).

Acute inhibition of PV INs: TST

The behavioral consequences of acute inhibition of PV INs in the IL were tested by using the TST as the first stressor in the CVS paradigm (Fig. 3). Animals were dosed with 1 mg/kg CNO 30 min before the start of TST. We observed significant changes in coping behavior following acute inhibition of PV INs in the IL. Compared with control mice, mice expressing hM4Di showed significant reduction in struggling duration (t = 2.7, df = 18, p = 0.02a; Fig. 3A), significant increase in immobility duration (t = 2.9, df = 18, p = 0.009b; Fig. 3B) and decreased latency to immobility (t = 2.5, df = 17, p = 0.02c; Fig. 3C), respectively.

Figure 2. Injection site, viral spread, and targeting of IL. A. Injection site and viral spread mapped onto respective mouse bregma coordinates, following bilateral injection of hM4Di DREADD into the IL of PV-Cre mice (n = 9). Light green rectangular box depicts the IL region with the injection sites restricted to the region. Each animal’s injection site and viral spread has been represented by a unique color. B. Representative image from one animal demonstrating the viral spread in PFC detected by red mCherry fluorescence. Image shows the spread was restricted to the IL with the white dashed lines outlining the IL cortex region of the PFC. Scale bars: 1000 and 500 μm. C. Percentage of DREADD transfected PV INs in the IL of PV-Cre mice. The percentage of infected cells in IL was calculated by dividing the number of mCherry-positive cells in IL by the total number of mCherry-positive cells in a specific focal plane and bregma coordinate. Brain bregma coordinates used for IL cytoarchitecture were taken from Franklin and Paxinos (2007). *indicates significant effect p < 0.05 compared to outside IL group. Values represent mean ± SEM, n = 9 per group.
Animals were tested for coping behaviors in the FST 24 h after cessation of CVS. Our purpose was to test whether inhibition of PV INs during the chronic stress regimen could block the aggregate effect of repeated stress on subsequent coping behavior (FST used as a novel stressor), brain activation patterns (Fos expression) and somatic endpoints (organ and body weights). All subjects received viral (Control or hM4Di) and CNO treatments, and because CNO was only administered during CVS, any phenotypes observed during FST were interpreted as reflecting an impact of PV IN manipulation during CVS on subsequent stress coping behavior. We observed significant differences in FST coping behaviors following chronic inhibition of PV INs during CVS. Specifically, chronic PV IN inhibition during stress increased active coping (swimming) and reduced passive coping (immobility) behaviors in the FST. Two-way ANOVA of total swimming duration showed a significant main effect of stress ($F_{(1,35)} = 11.7; \ p = 0.002$; Fig. 4A), DREADD ($F_{(1,35)} = 4.7; \ p = 0.037$) and time ($F_{(9,315)} = 68.2; \ p < 0.0001$) but no interaction effects were observed among the three groups time $\times$ stress $\times$ DREADD ($F_{(9,315)} = 0.6; \ p = 0.78$). Planned comparisons revealed significant increase in swimming duration in the CVS hM4Di group compared with No CVS hM4Di group ($p = 0.004$, $\ p = 0.009$, $\ p = 0.01$, and $\ p = 0.009$, respectively; Fig. 4B).

Figure 3. Impact of acute chemogenetic inhibition of PV IN in the IL mPFC on coping behavior in the TST. Acute inhibition of PV INs in the IL mPFC during TST, reduced total time spent struggling (A), increased total time spent immobile (B), and reduced latency to immobility (C). All mice were treated with CNO (1 mg/kg, i.p.) 30 min before TST. Behaviors were analyzed for a total time of 6 min. Values represent mean $\pm$ SEM, $n = 9$–10 per group; * indicates significant effect $p < 0.05$ versus corresponding control group.

Two-way ANOVA of total immobility duration showed a significant main effect of stress ($F_{(1,35)} = 9.5; \ p = 0.004$; Fig. 4C), no main effect of DREADD ($F_{(1,35)} = 1.7; \ p = 0.2$) and no stress $\times$ DREADD interaction ($F_{(1,35)} = 3.9; \ p = 0.057$). Planned comparisons revealed a significant reduction in immobility duration in the CVS hM4Di group compared with No CVS hM4Di group ($p < 0.0009$). There was a significant main effect of stress ($F_{(1,35)} = 9.5; \ p = 0.004$; Fig. 4C) and time ($F_{(9,315)} =
156.4; \( p < 0.0001 \)) on immobility duration but no interaction effects were observed among the three groups time \( \times \) stress \( \times \) DREADD \( (F_{(0,315)} = 1.1; p = 0.4) \). Planned comparisons revealed a significant decrease in immobility in the CVS hM4Di group at 2-, 3-, 4-, and 8-min time points compared with No CVS hM4Di group; \* indicates planned comparison significant effect \( p < 0.05 \) versus corresponding No CVS hM4Di group; \# indicates planned comparison significant effect \( p < 0.05 \) versus corresponding CVS Control group. Extended Data Figure 4-1 demonstrates chronic inhibition of PV INs did not affect locomotor activity. Extended Data Figure 4-2 demonstrates chronic CNO administration did not lead to any changes in FST behavior following CVS or control. Extended Data Figure 4-3 demonstrates CVS effects on FST did not depend on the age of the animal and also shows chronic injection stress and housing conditions had no effect on FST behavior in control animals.

Chronic inhibition of PV INs during CVS: Impact on Fos induction by FST
To test for Fos activation, animals were perfused after FST and brains were collected to analyze neuronal

Figure 4. Effects of chronic chemogenetic inhibition of PV INs during CVS on coping behavior in the FST following CVS. Chronic inhibition of PV INs in the IL mPFC during CVS, resulted in increased total time spent swimming (A) and decreased total time spent immobile (C) in the FST. B, D, Changes in swimming and immobility behavior, respectively, over the 10 min of FST. Values represent mean ± SEM, \( n = 9-10 \) per group; \* indicates planned comparison significant effect \( p < 0.05 \) versus corresponding No CVS hM4Di group; \# indicates planned comparison significant effect \( p < 0.05 \) versus corresponding CVS Control group. Extended Data Figure 4-1 demonstrates chronic inhibition of PV INs did not affect locomotor activity. Extended Data Figure 4-2 demonstrates chronic CNO administration did not lead to any changes in FST behavior following CVS or control. Extended Data Figure 4-3 demonstrates CVS effects on FST did not depend on the age of the animal and also shows chronic injection stress and housing conditions had no effect on FST behavior in control animals.
activation in brain regions typically activated by stress. We observed significant reduction in Fos induction in the CVS Control group compared with No CVS Control group, in the PrL, IL, BLA, and vlPAG. Inhibition of PV INs during CVS attenuated the reduction in Fos expression caused by CVS in the PrL, BLA, and vlPAG but not in the IL. Analysis of the PrL revealed a significant main effect of stress ($F_{(1,30)} = 17.5; p = 0.0002$; Fig. 5A) and a significant stress × DREADD interaction ($F_{(1,30)} = 7.7; p = 0.009$). Post hoc analysis using Tukey’s test revealed a significant reduction in Fos expression in the CVS Control group ($p = 0.0003$), which was attenuated by chronic PV IN inhibition in the CVS hM4Di group ($p = 0.75$). There was a significant main effect of stress only ($F_{(1,27)} = 21.2; p < 0.0001$; Fig. 5B) on Fos expression in the IL. There was a significant main effect of stress in the BLA ($F_{(1,20)} = 12.4; p = 0.004$; Fig. 5C) and vlPAG ($F_{(1,17)} = 20.5; p < 0.0001$; Fig. 5D) as well, with planned comparisons revealing significant reduction in Fos expression in the CVS Control group ($p = 0.008$ and $p = 0.0009$ in BLA and vlPAG, respectively) that was attenuated by chronic PV IN inhibition in the CVS hM4Di group ($p = 0.75$ and $p = 0.08$ in BLA and vlPAG, respectively). Analysis of Fos protein expression in the lateral septum (LS), anterior and ventral bed nucleus of the stria terminalis (BNST) and dorsolateral PAG (dlPAG) showed no significant treatment effects of PV IN modulation (Extended Data Fig. 5-1).

Chronic inhibition of PV INs during CVS: impact on somatic measurements

Organs and body weights were used to assess somatic effects of CVS. Adrenal gland hypertrophy and/or thymic

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**Figure 5.** Fos immunoreactivity in the PrL, IL, BLA, and vlPAG. Chronic inhibition of PV INs in the IL mPFC during CVS, attenuated CVS-mediated reduction in Fos expression in the PrL, BLA, and vlPAG, respectively (A, C, D) but did not prevent CVS-mediated reduction in Fos expression in the IL cortex (B) following FST. Values are presented as mean ± SEM; $n = 7–10$ per group; * indicates significant result $p < 0.05$ post hoc (A) and planned comparisons (C, D) compared with respective No CVS Control groups. PrL, IL, BLA, and vlPAG stands for prelaminic, infralaminic, basolateral amygdala and ventrolateral periaqueductal grey, respectively. Extended Data Figure 5-1 demonstrates Fos expression in various other brain regions where no significant effect of PV IN modulation was observed.
atrophy are often observed following chronic stress and are used as indicators of repeated/chronic hypothalamic pituitary adrenal (HPA) axis activation. In our experiments, there was a main effect of stress ($F_{(1,35)} = 11.2; p = 0.002^c$; Fig. 6A) and a significant stress $\times$ DREADD interaction ($F_{(1,35)} = 4.8; p = 0.035$) on adrenal weights. Post hoc analysis using Tukey’s test revealed that CVS Control group had significantly increased adrenal weight compared with No CVS Control group ($p = 0.002$), which was attenuated by CVS hM4Di when compared with No CVS hM4Di group ($p = 0.84$). There was a main effect of stress on thymus weight ($F_{(1,35)} = 161.4; p < 0.0001^m$; Fig. 6B), with no effect of DREADD ($F_{(1,35)} = 0.2; p = 0.7$) or stress $\times$ DREADD interaction ($F_{(1,35)} = 57; p = 0.1$).

Decreased body weight gain is observed following chronic mild stress exposure in rodents (Ghosal et al., 2017). We observed a significant main effect of stress on final body weight ($F_{(1,35)} = 6.6; p = 0.01^c$; Fig. 6C) with no effect of DREADD ($F_{(1,35)} = 0.2; p = 0.6$) or stress $\times$ DREADD interaction ($F_{(1,35)} = 0.006; p = 0.9$), consistent with known effects of CVS on body weight gain. Planned comparisons revealed final body weight in CVS group to be significantly lower than No CVS Control group ($p = 0.03$), which was attenuated by CVS hM4Di when compared with No CVS hM4Di group ($p = 0.08$; Fig. 6C). Two-way repeated measures ANOVA of body weight over time during the 14 d CVS paradigm showed a main effect of stress ($F_{(1,35)} = 5.3; p = 0.03^c$) and time ($F_{(3,108)} = 4.5; p = 0.005^c$; Fig. 6D). Planned comparisons revealed body weight in CVS Control group to be significantly lower than No CVS Control only on day 14 ($p = 0.03$; Fig. 6D). Control experiments to determine effects of chronic CNO on body weight showed no significant difference. Two-way ANOVA of total body weight showed a significant main effect of stress ($F_{(1,28)} = 18.6; p = 0.002$; Fig. 4-2C), no main effect of treatment ($F_{(1,28)} = 0.41; p = 0.52$) and no stress $\times$ treatment interaction ($F_{(1,28)} = 0.01$).

Figure 6. Impact of chronic stress on organ and body weights. Chronic inhibition of PV INs in the IL mPFC during CVS, attenuated CVS mediated increases in adrenal gland weight (A) and attenuated CVS mediated decreases in body weight (C, D). No change in CVS-induced decreases in thymus weight was observed following PV IN inhibition (B). Data are presented as absolute organ and body weights. Values represent mean $\pm$ SEM; $n$ = 9–10 per group; * indicates planned comparisons significant effect $p < 0.05$ versus corresponding No CVS Control groups.
Discussion

Our studies support a role for IL GABAergic PV INs in stress-mediated behavioral and somatic phenotypes. Using inhibitory DREADDs to inhibit the activity of PV INs during a modified CVS paradigm, we have established a causal role of IL PV INs in initiating and coordinating coping strategies and somatic outcomes in response to stress. Inhibition of PV IN during the CVS resulted in an increase in active coping strategy during FST, suggestive of dynamic behavioral remodeling during an aversive challenge. Chronic stress-induced behavioral alterations were accompanied by changes in neuronal activation patterns quantified by Fos expression following FST. Chronic PV IN inhibition attenuated CVS-induced reductions in Fos expression in PrL, BLA, and vPAG, indicating that inhibition of PV INs mitigates the impact of chronic stress on stress regulatory brain regions. PV IN inhibition during CVS also attenuated CVS-induced adrenal hypertrophy and body weight loss, further suggesting that PV IN signaling during CVS might be playing a role in somatic effects of chronic stress. Interestingly, PV IN inhibition during the TST, which was the first stressor in the modified CVS paradigm, resulted in an increase in passive coping and a decrease in active coping in stress naive animals. This finding suggests unique mechanisms of PV IN plasticity following acute versus chronic stress paradigms and also implies that an optimum level of prefrontal PV IN activity is important to maintain prefrontal E/I balance so that PFC can respond appropriately to stress. Overall, the data indicate that PV INs play a role in inhibiting IL output during chronic stress, suggesting a potential role in driving ventromedial PFC hypofunction. Exploring the role of PV INs in stress-related disorders is an important research direction for future experiments.

GABAergic PV INs are well positioned to provide strong, fast-spiking inhibitory signals to pyramidal projection neurons in the PFC and reduce network excitability, and therefore could be contributing to chronic stress-mediated hypoactivity (Winkelmann et al., 2014; Tremblay et al., 2016; Safari et al., 2017). Our data suggest that PV INs play an important role in chronic stress-mediated inhibition of the IL. Chronic inhibition of IL PV INs during CVS resulted in increased active and decreased passive coping behaviors in FST. A switch to active coping can be interpreted as an adaptive strategy to deal with chronic stress, and drugs that are effective antidepressants in humans typically promote active coping styles and reduce passive coping in the FST in mice (Porsolt et al., 1977; Marti and Armario, 1993). GABA receptor antagonists have been shown to have antidepressant and anxiolytic properties (Bhutada et al., 2010; Mehta et al., 2017; Zanos et al., 2017; Samad et al., 2018). It is known that antidepressants such as fluoxetine and ketamine reduce PV expression in the PFC (Ohira et al., 2013; Zhou et al., 2015; Page and Coutellier, 2019). Moreover, preventing the reduction in PV IN activity leads to loss of antidepressant efficacy, further suggesting that reduced activity of PV INs might be playing a role in therapeutic efficacy of antidepressants (Zhou et al., 2015; Page and Coutellier, 2019). Therefore, based on prior studies and our findings, inhibition of PV INs during chronic stress may lead to more adaptive stress coping strategies and reverse some of the behavioral deficits associated with chronic stress. It is important to note that the effects observed in FST are because of PV IN inhibition and not because of any changes in locomotor activity (Extended Data Fig. 4–1). Additionally, we did not detect any effects on FST because of chronic dosing of CNO alone, suggesting that repeated CNO dosing did not alter stress coping behavior (Extended Data Fig. 4–2).

Our experiments revealed that inhibition of PV INs in the IL during stress can attenuate chronic stress-induced decreases in Fos expression in key stress regulatory regions such as the PrL, BLA, and vPAG following FST (Keedwell et al., 2005; Berton et al., 2007; Vialou et al., 2014). Since we cannot verify direct PV IN modulation of IL projections to these regions, we cannot exclude the possibility that reversal of CVS-related inhibition of Fos induction is because of actions of the IL through other projection systems. Nonetheless, the data suggest that PV IN inhibition reduces inhibitory effects of CVS on IL outflow, permitting drive of downstream structures known to participate in physiological reactivity and stress coping behavior (Maier and Watkins, 2010). Notably, this includes the neighboring PrL, which is not targeted by our DREADD injections and thus has Fos excitability modulated by cortico-cortical connections. Involvement of the PrL is consistent with its prominent role in mediation of coping behavior (Fiore et al., 2015; Johnson et al., 2019; Molendijk and de Kloet, 2019).

PV IN modulation did not prevent CVS-induced changes in Fos in the IL following FST. The Fos data in IL represents an overall change in neuronal activity regardless of where the neurons project to. Thus, it is possible that PV IN modulation might be altering neural activity only in specific circuits, such as the IL-PrL or IL-BLA connections. Indeed, prior studies indicate that neurons projecting from IL to BLA receive strong innervation via PV INs (Marek et al., 2018). Additionally, activity of cortical PV INs is essential for microcircuit operations that correlate with behavioral events (Isomura et al., 2009; Kvitsiani et al., 2013; Kim et al., 2016). Thus PV INs might be affecting PFC microcircuits, thereby acting as a functional unit to synchronize the flow of information between IL-PrL (Courtin et al., 2014b; Kepecs and Fishell, 2014). Therefore, it will be important in the future to explore PV IN mediated plasticity in the specific circuits showing reduced Fos expression following PV IN modulation. It has been shown that chronic inhibition of neurons may also lead to plastic changes, resulting in rebound firing and enhanced excitability/hyperactivity when inhibition is removed (Stemmler and Koch, 1999; Sokolova and Mody, 2008; Wiegert et al., 2017; Purohit et al., 2018). Thus, it is also possible that removal of chronic inhibition of PV INs following CVS might have resulted in hyperactivity of PV INs, leading to reduced Fos expression in IL. CVS also causes a reduction in Fos on exposure to acute stress.
Repeated inactivation of PV INs during stress attenuated the CVS-induced increase of adrenal weight. The adrenals are highly sensitive to repeated stress, and it is believed that increased adrenal size is linked to cumulative increases in adrenocorticotropic hormone (ACTH) secretion (Ulrich-Lai et al., 2006). Blockade of adrenal hypertrophy suggests that PV INs participate in control of the central limb of HPA axis activation and provides additional confirmation of cumulative efficacy of chronic PV IN inhibition in control of stress endpoints. In contrast to the adrenals, CVS caused equivalent decreases in thymus weight, suggesting either sensitization of glucocorticoid sensitivity or enhanced autonomic activation by CVS, presumably mediated by mechanisms independent of PV INs. Chronic inhibition of PV INs also attenuates CVS-induced reduction in final body weight, suggesting PV IN signaling might also be playing a role in CVS mediated body weight effects.

As part of our design, we assessed the impact of IL PV IN inhibition acutely following the first stressor in our CVS regimen, the TST, which allows for behavioral readouts (duration of struggling, immobility and latency to immobility). Acute inhibition of IL PV INs resulted in decreased active coping (struggling) and increased passive coping (immobility) in the TST. Our data are consistent with a prior study indicating that reduced excitatory synaptic drive onto PV INs is linked to increased stress susceptibility and enhanced helplessness behavior (Perova et al., 2015). These data indicate that PV INs may play a role in driving active coping responses, when an animal with no history of prior stress is exposed to a novel acute stressor such as the TST. Together, these studies suggest that activation of PV INs is required for coping responses to acute stress.

Our results with acute PV IN inhibition are in contrast to the results seen in the FST after chronic PV IN inhibition during a two-week CVS exposure. These data indicate different roles for these neurons in acute versus chronic stress adaptations. Our finding of divergent effects of IN function in PFC is in line with previous studies showing opposing effects on emotionality in acute versus chronic somatostatin (SST) IN inhibition in the PFC (Soumier and Sibille, 2014) and on auditory information processing in acute versus chronic IN inhibition in the auditory cortex (Seybold et al., 2012). Our data suggest that distinct neuronal ensembles and brain circuitry may be involved in modulating acute versus chronic stress-mediated behavioral outcomes. It is also possible that chronic stress may result in plastic changes in the same neuronal ensemble recruited by acute stress, leading to differences in stress response. However, it is not known what specific plasticity in the neural network underlies the emergence of opposing phenotypes following chronic stress and therefore further studies are needed to investigate the mechanisms. Moreover, to fully understand the differences observed here with acute versus chronic stress PV IN modulation, it would be beneficial to determine whether chronic stress is able to modify the potentiating effect of acute inactivation on active coping responses in the TST.

Our acute and chronic PV IN inactivation data also suggest that an optimum level of E/I balance needs to be maintained in the PFC for it to function appropriately and PV INs play a crucial role in maintaining that balance (Ferguson and Gao, 2018). Disruption of the balance can lead to abnormal behavioral endpoints and psychiatric illnesses (Selten et al., 2018; Berg et al., 2019; Page and Coutellier, 2019). Our data on TST agree with prior reports suggesting that reducing activity of PV INs acutely under baseline conditions can lead to a more stress vulnerable passive coping response which might be because of an over excitation of the PFC (Perova et al., 2015). On the other hand, chronic stress can disrupt the E/I balance leading to over inhibition by increasing the activity of PV INs, resulting in maladaptive behavioral outcomes (Page et al., 2019; Page and Coutellier, 2019). The current approach is designed to inhibit the over activation of PV INs in the context of chronic stress exposure. Indeed, our results show that inhibiting the activity of PV INs during stress leads to a more active coping response and attenuates CVS mediated alterations in Fos activation in brain regions controlling emotionality as well as attenuating the effects on body and organ weights in response to CVS.

There are a few caveats to the present study that must be considered in the interpretation of the data. First, the modified CVS paradigm used in this study did not produce the typical behavioral effects seen in FST when using alternative CVS procedures (Ghosal et al., 2017; Wohleb et al., 2018). Initially, we thought that older age and chronic injection stress (the later described previously by Moghaddam and Bolinao, 1994), might have resulted in high rates of immobility in control animals leading to a ceiling effect, hence reducing the window to detect an increase in immobility typically observed after CVS exposure. However, additional control experiments showed CVS did not affect FST behavior in younger mice (Extended Data Fig. 4–3). Additional experiments in control mice showed chronic injection stress and single housing also did not affect behavior in FST (Extended Data Fig. 4-3). These results further lead us to conclude that the lack of effect on FST is because of the exclusion of overnight and extended periods of stress in the modified CVS paradigm in our study. Nevertheless, the paradigm resulted in somatic effects leading to reduced body weight and adrenal hypertrophy, indicating that sustained HPA axis drive only occurred in the CVS group. Moreover, our CVS procedure also reduced Fos activation in stress regulatory brain regions, demonstrating alterations in neuronal activation typically observed following CVS (Ostrander et al., 2009; Moench et al., 2019). Second, although our injection site was primarily in the IL, there was some spread into the PrL and DP area of the cortex (~28% of cells). Third, this study was conducted only in male mice because prior research showed CVS mediated alteration in inhibitory synaptic drive in the IL of males (McKlveen et al., 2016, 2019). Because PV IN modulation may have sex specific effects (Shepard et al., 2016; Page et al., 2019), it would be important to examine the effects of PV IN modulation...
during stress in females. Fourth, in this study PV INs were inhibited during the 14 d of CVS. The degree to which the efficacy of DREADD-mediated inhibition might vary over the course of the 14-d treatment paradigm remains unclear and will require additional study. Finally, in this study we euthanized animals after exposure to one behavioral paradigm (FST) to obtain the anatomic Fos expression dataset to a novel stressor following CVS. Chronic stress can be characterized by cellular and behavioral changes spanning multiple interconnected neural network adaptations which were not explored in our current study. In order to get a clear representation of how PV IN modulation during stress is affecting emotionality, additional behaviors may be worth exploring in follow-up experiments.

Taked together, our data are consistent with a causal role of IL PV INs in initiating and coordinating coping strategies and physiological outcomes in response to stress. Chronic stress-mediated hypoactivity and aberrant behavioral responses may be mediated partly via plastic changes in PV IN function and may play a role in stress-related pathologies (e.g., depression and PTSD). Our data indicate that chemogenetic inhibition of PV INs during chronic stress, which reduces PV-initiated inhibition in the context of each individual stressor experience, may block or attenuate inhibition of glutamatergic neurons. In this case, maintenance of inhibitory synaptic inputs onto glutamatergic IL projection neurons is sufficient to attenuate some but not all behavioral and physiological consequences of chronic stress exposure, including decreased passive (immobility) and increased active coping behaviors (swimming) in the FST, attenuating CVS effects on reduction in neuronal Fos activity, adrenal hypertrophy, and body weight loss. Our findings suggest that reducing the activity of PV INs in the PFC during chronic stress may facilitate output of prefrontal neurons and could provide therapeutic benefits for stress-related disorders.

In conclusion, this study provides support that PV INs play a role in chronic stress-mediated coping behaviors and physiological phenotypes. Furthermore, the study adds to the current knowledge regarding possible mechanisms of hypoactivity of the PFC and how PV INs may be involved in driving chronic stress-related pathologies. The study also highlights opposing effects of acute and chronic PV IN inhibition, indicating different underlying mechanisms involved in acute versus chronic stress paradigms. Overall, this study shows that reducing PV IN activity to promote prefrontal output may be an effective treatment strategy for stress-related illnesses.

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