

TITLE

Overexpression of the schizophrenia risk gene C4 in PV cells drives sex-dependent behavioral deficits and circuit dysfunction.

AUTHORS

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ABSTRACT

Fast-spiking parvalbumin (PV)-positive cells are key players in orchestrating pyramidal neuron activity and thus serve an indispensable role in cognitive function and emotional regulation. Feed-forward excitatory inputs, essential for the function of PV cells, are disrupted in various neurological conditions, including schizophrenia (SCZ). However, it is not clear how disease-associated stressors such as immune dysregulation contribute to defects in particular cell types or neuronal circuits. We have developed a novel transgenic mouse line that permits conditional, cell-type specific overexpression (OE) of the immune complement component 4 (C4) gene, which is highly associated with SCZ. Using this genetic approach, we demonstrate that specific global OE of mouse C4 (*mC4*) in PV cells causes pathological anxiety-like behavior in male, but not female mice. In the male medial prefrontal cortex (mPFC), this sexually dimorphic behavioral alteration was accompanied by a reduction in excitatory inputs to fast-spiking cells and an enhancement of their inhibitory connections. Additionally, in PV cells, elevated levels of *mC4* led to contrasting effects on the excitability of cortical cells. In males, PV cells and pyramidal neurons exhibited reduced excitability, whereas in females, PV cells displayed heightened excitability. Contrary to the behavioral changes seen with elevated *mC4* levels in PV cells, pan-neuronal overexpression did not increase anxiety-like behaviors. This indicates that *mC4* dysfunction, particularly in fast-spiking cells, has a more significant negative impact on anxiety-like behavior than widespread alterations in the neuronal complement. Consequently, by employing a novel mouse model, we have demonstrated a causal relationship between the conditional overexpression of the schizophrenia risk gene *C4* in fast-spiking neurons and the susceptibility of cortical circuits in male mice, resulting in changes in behaviors associated with prefrontal cortex function.

INTRODUCTION

Cortical parvalbumin (PV)-positive fast-spiking cells are a distinct class of inhibitory neurons characterized by their expression of the Ca^{2+} -binding protein PV (1–3). Their unique biophysical properties allow them to drive potent, precise inhibition, effectively controlling the temporal dynamics of the excitatory and inhibitory inputs (4–6) that support critical brain functions such as sensory processing, motor control, and cognition (7–9). At a network level, PV cells are responsible for generating and regulating gamma oscillations (10), 30-80 Hz rhythmic fluctuations in brain activity that correlate with cognitive performance (11–13) and are impaired in anxiety disorders (14), schizophrenia (SCZ) (15,16), and Alzheimer's Disease (AD) (17,18). Besides controlling the temporal dynamics of excitation and inhibition and orchestrating oscillatory activity, PV neuron activity tightly regulates cortical maturation during critical developmental windows (19–21).

Despite significant progress in understanding the function of this inhibitory interneuron in the healthy and diseased brain, it remains to be determined how specific genetic alterations associated with neuropsychiatric disorders lead to the dysfunction of inhibitory microcircuits. It is also unclear whether particular circuitry or synaptic inputs underlying the function of PV cells are susceptible to disease-associated genetic alterations. For example, in post-mortem tissue of patients with SCZ, the density of excitatory synapses is decreased on PV cells with concomitant downregulation of PV and other inhibitory markers (22–25), suggesting that excitatory drive to fast-spiking cells is compromised in this brain disorder. Moreover, SCZ-associated genetic alterations have been found to disrupt the molecular machinery underlying feed-forward excitatory inputs to PV neurons (26–28), suggesting these particular excitatory synaptic connections are susceptible to genetic perturbations.

The association between the Major Histocompatibility Complex (MHC) and SCZ could provide a link between immune dysfunction and the disruption of molecular mechanisms that regulate the wiring of synaptic circuits (29–32). In support of this, Sekar et al. (33) showed that mice that lack *C4b* (mouse *C4*, *mC4*), which in humans is harbored in the MHC locus, exhibit deficits in the developmental refinement of retinogeniculate synapses. Furthermore, Comer et al. (34) demonstrated that increasing levels of the human (*C4A*) and mouse *C4* homologs in layer (L) 2/3 pyramidal neurons (PYRs) of the medial prefrontal cortex (mPFC) – a brain region associated with the pathology of SCZ (35–37), anxiety disorders (38), and other neuropsychiatric conditions (39,40) – led to pathological synaptic loss during early postnatal development and social behavioral deficits in mice. This research suggests a link between immune dysfunction and brain disorders, particularly SCZ, through the role of the MHC and its impact on synaptic development and plasticity.

As a consequence of their unique properties and role in controlling neuronal network function, PV cells exhibit high metabolic demands, which make them vulnerable to oxidative stress and chronic inflammation (41–45); these pathological processes are often linked to a range of brain disorders (46–48). Consequently, determining the molecular pathways through which immune imbalances can impact PV neuron functionality offers significant potential for unraveling the origins of neurodevelopmental diseases. Despite this, there is a notable gap in the availability of models to determine how neuroimmune dysfunction alters specific circuitry in the brain. In the same vein, alterations in the complement pathway have been linked to the pathology of several brain disorders (49). However, it remains an open question whether particular brain cell types are especially vulnerable to complement dysfunction.

Here, we developed and validated a novel mouse line that permits cell-type specific overexpression of *mC4* (mC4-OE). Utilizing this unique knock cell mouse line, we demonstrate that increased *mC4* levels in PV cells (PV-mC4-OE) drives pathological anxiety-like behaviors in male, but not female mice. In both sexes, PV-mC4-OE led to changes in a subclass of social behavior, indicating that elevated expression of this immune gene in fast-spiking cells disrupts the circuitry governing social behaviors. We used electrophysiological recordings to show that excitatory and inhibitory inputs to mPFC PV cells are altered with increased levels of mC4 in male, but not female mice, mirroring the sexually dimorphic anxiety-like behavior. Additionally, elevated *mC4* in PV neurons had differing impacts on cortical cell excitability: hypoexcitability in male fast-spiking cells and PYRs, as opposed to enhanced excitability in female fast-spiking cells. In contrast to PV-cell-specific effects of mC4-OE, increased *mC4* levels in all neurons had no effects on anxiety-like behavior. This implies that *C4* perturbations in PV cells are more harmful to brain function than pan-neuronal alterations. In summary, our results show that *mC4* dysregulation in PV neurons led to alterations in anxiety-like behavior in male mice, which were associated with mPFC circuit dysfunction. Our results provide crucial insights into the molecular interplay between cell-type-specific increased levels of *mC4*, defects in mPFC circuitry, and abnormal behavior associated with the mPFC.

METHODS

Ethics Statement and Animals

All experimental protocols were conducted according to the National Institutes of Health (NIH) guidelines for animal research and were approved by the Boston University Institutional Animal Care and Use Committee (IACUC; protocol #2018–539). All mice were group-housed (2–4 animals/cage). The light/dark cycle was adjusted depending on the behavioral task (see below). Unless otherwise stated, food and water were provided *ad libitum* to all mice. Experimental offspring were reared in the cage with the dam until weaning at postnatal day (P) 21. Stimulus CD-1 (Charles River Laboratories, strain code: 022, RRID:IMSR_CRL:022) mice 3–5 weeks of age were used in social assays. During experiments and analysis, the experimenter was blinded to mouse genotype and experimental conditions whenever possible.

Generation of the mC4-KI mouse: Generation of the mC4-KI mouse was accomplished using Cyagen/Taconic services (Santa Clara, CA). Briefly, the “adenovirus SA-Frt-CAG promoter-Frt-loxP-3*polyA-loxP-Kozak-mouse C4b CDS-polyA” cassette was cloned into intron 1 of ROSA26. The homology arms were generated by PCR using BAC clones as templates. C57BL/6 ES cells were used for gene targeting. In the targeting vector, the positive selection marker (Neo) is flanked by SDA (self-deletion anchor) sites. Diphtheria toxin A (DTA) was used for negative selection. Targeted ES cell were injected into C57BL/6J albino embryos, which were then re-implanted into CD-1 pseudo-pregnant females. Founder animals were identified by their coat color, and their germline transmission was confirmed by breeding with C57BL/6J females and subsequent genotyping of the offspring. Three male and two female heterozygous targeted mice were generated from clone 1F3 and were bred to establish a colony. Lastly, the constitutive knock cell allele was obtained after Cre-mediated recombination (see *Breeding* section).

Breeding: C57BL/6J (Jackson Laboratory, strain #: 000664, RRID:IMSR_JAX:000664) mice were paired with heterozygous mC4-KI transgenics of the same genetic background. Genotyping (Transnetyx) was used to plan breeding schemes and identify specific genotypes.

Heterozygous mC4-KI transgenics were paired with homozygous PV-Cre mice (Jackson Laboratory, B6.129P2-Pvalb^{tm1(Cre)Arbr}/J, strain #: 017320, RRID:IMSR_JAX:017320). Our breeding scheme generated mice that inherited the floxed mC4-KI allele, and thus overexpress (OE) *mC4* in PV cells (PV-mC4-KI, or simply KI), and wild type littermates, which were used as control mice (PV-mC4-WT, or simply WT).

To OE *mC4* in all neurons, heterozygous mC4-KI transgenic mice were paired with homozygous BAF53b Pan-neuronal-Cre (Jackson Laboratory, STOCK Tg(Actl6b-Cre)4092Jiwu/J, Strain #:027826, RRID:IMSR_JAX:027826, (50)) mice. Breeding these two mouse lines yielded mice that inherited the floxed mC4-KI allele, and thus OE *mC4* in all neurons (PanN-mC4-KI), and wild type control littermates (PanN-mC4-WT).

Our breeding scheme ensured that all experimental mice carried the cre recombinase gene to control for its effects. PV-mC4-KI or PanN-mC4-KI mice did not exhibit any gross brain abnormalities, indistinguishable from controls. Additionally, they had similar weights to their controls, suggesting that besides the described cellular and behavioral deficits, these mice were otherwise healthy and had no significant defects.

A strength of the mC4-KI transgenic mouse is that more moderate levels of mC4-OE can be driven in a cell-specific manner by crossing the mC4-KI mice to Flp transgenic mice, which substitutes the strong CAG promoter for the weaker ROSA26 promoter (51,52), allowing the effects of more moderate transgene expression to be studied.

Multiplex fluorescence in situ hybridization

Tissue preparation and staining: For multiplex fluorescence *in situ* hybridization (M-FISH) experiments, brains of PV-mC4-WT and PV-mC4-KI mice P21-22 or P58-65 were extracted and immediately embedded in O.C.T. (Tissue-Tek, 4583), flash-frozen on dry ice, and stored in -80°C until being cut. Prior to slicing, brains were moved to -20°C for 30 min. Slices were cut on a Leica CM 1800 cryostat at 16 µm at -16 to -19°C and adhered directly onto microscope slides (Fisher Brand Superfrost Plus, #1255015), which were then stored in -80°C until ready for M-FISH (<1 week). M-FISH experiments were then performed as directed by the commercially available kit (RNAScope, Advanced Cell Diagnostics), from which all probes and reagents were purchased. Fluorescent probes for *mC4* (Mm-C4b, 445161), parvalbumin (Mm-Pvalb-C2, 421931-C2), and somatostatin (Mm-Sst-C3, 404631-C3) were used. To confirm cell bodies in the slice, nuclei were stained with RNAScope DAPI (320858). After staining, the slices were mounted with ProLong Diamond Antifade Mountant (ThermoFisher, P36961). Each round of M-FISH performed contained tissue from both PV-mC4-WT and PV-mC4-KI mice.

Imaging and analysis: M-FISH images were acquired at 40x on a confocal laser scanning microscope (Nikon Instruments, Nikon Eclipse Ti with C2Si⁺ confocal), controlled by NisElements (Nikon Instruments, 4.51) including four laser lines (405, 488, 561, and 640 nm), at a step size of 0.4 µm for nearly the entire thickness of the tissue slice (16 µm). For each round of M-FISH, consistent imaging parameters were used. Tissue slices imaged and analyzed belonged to mPFC divisions (prelimbic (PrL), infralimbic (IL), and anterior cingulate (AC) cortices) of the mouse brain. Images predominantly included L2/3, though deeper cortical layers were also included in the region of interest (ROI). For analysis, a maximum intensity projection of each z-stack was made (ImageJ, National Institute of Health, Bethesda, Maryland) and transferred into CellProfiler (53,54) (Broad Institute). Cells were identified and segmented using DAPI, and the contour was expanded by 10 pixels (approximately 3.1 µm) to capture the majority of PV, SST, and *mC4* puncta surrounding the nucleus. Cells were classified as PV cells

or SST cells if their expanded contour contained an empirically-derived minimum of 13 or 10 PV or SST puncta, respectively. Once identified as a PV cell, SST cell, or non-PV/non-SST DAPI+ 'Other' cell, CellProfiler was used to quantify the number of *mC4* puncta within each contour (i.e., within each cell).

PV cell density

Perfusion and immunohistochemistry: Mice P55-74 were anesthetized with a 4% isoflurane-oxygen mixture (v/v) and perfused transcardially with phosphate-buffered saline (PBS, Gibco, Life Science Technologies, 70011044) followed by 4% paraformaldehyde (PFA) in PBS. Extracted brains were further fixed in PFA for 24 h before being transferred to a 30% (w/v) sucrose solution and stored at 4°C. Brain slices were cut at 40 µm on a freezing stage sliding microtome (Leica SM2000) and stored in PBS. From each mouse, two brain slices were selected for immunostaining: one at approximately Bregma +1.98 mm and another at Bregma +0.98 mm along the anterior-posterior (A-P) axis. Slices were first blocked and made permeable in a solution containing 10% donkey serum (Sigma-Aldrich, S30-100ML) and 1% TritonX100 (Sigma-Aldrich, X100-100ML) in PBS. Next, after applying the primary antibody (rabbit anti-PV, Abcam, ab11427) at a 1:250 dilution, slices were placed on a shaker at 4°C for 48 h. Slices were then washed 4 x 15 min with 0.025% TritonX100 in PBS. Next, the secondary antibody was applied (anti-rabbit STAR RED, Abberior, STRED) at a 1:500 dilution and returned to the shaker for 48 h at 4°C. Slices were then washed 4 x 15 min in PBS, and mounted onto 1 mm microscope slides (Globe Scientific, #1324) with DAPI Fluoromount (Thermo Fisher Scientific, Cat. #: 00-4959-52).

Imaging and Analysis: Cell density imaging was acquired (laser lines 405 and 640 nm) at a step size of 1 µm for nearly the entire thickness of the tissue slice (40 µm). For each animal, the mPFC was imaged at 20x in both the slice at Bregma +1.98 mm (6 ROIs total, 3/hemisphere, that include the PrL, IL, and AC cortices) and the slice more posterior at Bregma +0.98 mm (4 ROIs, 2/hemisphere, all AC). Consistent imaging parameters were maintained across all imaging sessions. Images were analyzed as TIFFs in ImageJ and compared to a brain atlas to identify brain regions (55,56). The intensity value in the PV channel for each ROI (in a brain slice) and the average background signal for each brain slice were quantified. To binarize cells as PV-positive, we calculated an intensity threshold and classified cells as PV-positive if their intensity value was higher than this threshold. These data were used to calculate the number of PV cells that were positive. To calculate density, we determined the area (excluding L1) of the ROI in which PV cells were counted and calculated the 3D volume (in mm³) by multiplying the 2D area of each slice by the depth of the tissue imaged (the Z-stack).

Behavior

General experimental conditions: P40-60 mice were used for all behavioral assays, group housed in sets of 2-3 mice per cage. Mice were used in either (1) a series of three anxiety-related assays or (2) a series of two sociability assays. The specific sequence of anxiety-related assays that mice were exposed to was consistent across all mice and proceeded in the following order: open field (OF) and elevated zero maze (EZM, performed on the same day), light-dark box (LDB), and novelty-suppressed feeding (NSF) (Novel arena and cage NSF, performed back-to-back).

Similarly, the specific sequence of sociability assays that mice were exposed to was also consistent across all mice: Object and juvenile interaction (performed immediately back-to-back

for each mouse, see *Object and social interaction*) was followed by the three-chamber sociability assay.

Seven days prior to the first day of handling (see *Handling* below), mice were genotyped (Transnetyx), transferred to a fresh cage, and placed with a *Do Not Handle* card to minimize human handling and stress. Mice were reliably identified using a set of ear hole-punches throughout behavioral experiments.

All behavioral assays were performed at a similar time of the day. Mice used in anxiety-related assays were reared on a 12 h light/dark cycle with lights on at 7 AM and lights off at 7 PM, and assays were performed under white light (Adorama, 13" Dimmable LED Ring Light). The intensity of light used for each assay was consistent each time the assay was performed, but varied dependent on the assay (see specific assays below). Each day, the lux was measured and adjusted to the appropriate level for the assay being performed (Dr.meter LX1330B). Mice used in sociability assays were reared on an inverted light/dark schedule, with lights on at 7 PM and lights off at 7 AM, and assays were performed under red light (Amazon Basics 60W Equivalent, Red) to minimize the stress-inducing effects of bright white light and to remain consistent with their inverted light/dark schedule. Behavioral assays in a given series were always separated by at least two days but never more than four days. On all days of behavioral experiments and handling, mice were retrieved from the facility and left in the behavior room to acclimate to the environment for at least 1 h. Once all mice in a cage completed the assay, all were returned to their original home cage.

Acquisition of behavior data for all anxiety-related assays and the three-chamber assay was recorded using Logitech C922x Pro Stream Webcams at 30 frames per second (fps) via the open-source UCLA miniscope software (57). Acquisition of object and juvenile interaction data was recorded at 30 fps using a Teledyne Flea3 USB3 camera (Model: FL3-U3-13E4C-C: 1.3 MP, e2v EV76C560) via an in-house, python-based, open-source video acquisition software, REVEALS (<https://github.com/CruzMartinLab/REVEALS>) (58).

Throughout all components of behavioral assays and handling, gloves and a lab coat were worn. Gloves, behavioral arenas, and any relevant objects or cups used during the assay were sprayed with 70% ethanol in between handling mice or between each new mouse performing a given assay.

All behavioral assays were performed blind to condition, and analysis was performed via custom code written in MATLAB (MathWorks) (see *Behavior Analysis* below).

Handling: The first assay of each series was always preceded by three consecutive days of handling. Mice involved in anxiety-related assays were handled under standard, ambient room lighting (270 lux), and mice involved in sociability assays were handled under red light.

Anxiety-related assays

Anxiety-inducing arenas were custom-made as described in Johnson et al. (59).

Open field (OF): Mice were placed in the center of a custom-made OF, a (45 × 45 × 38 cm (length x width x height) black arena under 200 lux of white light, and were free to explore for 10 min. The OF was used to measure locomotion by measuring the distance traveled by the mice.

Elevated zero maze (EZM): The EZM is an elevated (63 cm) circular platform with a 5 cm track width and diameter of 60 cm. It is comprised of two closed arms with a wall height of 14 cm and two open arms that lack walls. The EZM was run under 200 lux of white light for 10 min.

Light-dark box (LDB): The LDB uses the frame of the OF, but inserted into the OF is a black divider (45 x 38 cm, length x height) that divides the OF into two distinct zones 1/3 and 2/3 the width of the OF, but features a small passage-way at the bottom (7.6 x 7.6 cm, width x height) to allow the mice to move freely between zones. Over the smaller zone (45 x 15 cm, length x width) is a black lid that blocks all light: this is the dark zone. Because there is no lid over the remaining 2/3 of the OF (45 x 30 cm, length x width), this is the light zone. The LDB was run under 300 lux of white light for 10 min.

Novelty-suppressed feeding (NSF): 24 h before the start of experimentation, mice were transferred to a clean cage that possessed no food. During the assay, mice were placed in a novel, open arena (50 x 35.5 x 15 cm, length x width x height) with a single, fresh food pellet strapped down in the center of the arena with a rubber band. The experimenter watched the live video feed to observe when the mouse traveled to the center of the maze and began feeding on the food pellet, which concluded the assay. Simple investigation or sniffing of the pellet was not considered feeding. The NSF was the only hand-scored assay, and was done so by a trained, blinded experimenter. To measure the latency to feed, the researcher watched the trial back and determined the exact frame that the mouse first bit the pellet. In the rare event that the mouse did not eat the pellet in the time allotted (10 min recording), that mouse was excluded from the NSF analysis. The NSF was run under 200 lux of white light.

Cage NSF: Immediately following the NSF, mice were placed in a fresh cage with approximately 10 food pellets placed in one corner of the cage. Again, the latency to feed was recorded by the experimenter. The Fresh Cage NSF served to determine and verify the anxiety-inducing nature of the arena relative to the more familiar environment of a standard mouse cage.

Z-Anxiety quantification: Z-Anxiety quantification was adapted from Guilloux et al. (60). For each individual assay, the mean (μ) and standard deviation (σ) of the relevant metric for that assay (e.g., EZM: time spent in open arms) for all WT animals was calculated. For any given mouse in any single assay, whose performance in that assay (e.g., EZM: time spent in open arms) is given by x , the z-score is the following:

$$z - score_{assay} = \frac{x - \mu}{\sigma}$$

To be consistent with a positive z-score being indicative of increased anxiety, the sign of the z-score in the EZM and LDB was multiplied by -1. In this way, less time spent in the open arms (EZM) or in the light zone (LDB), indicators of increased anxiety-like behavior, yielded positive z-score values. For the NSF, because an increased latency to feed was indicative of anxiety-like behavior, these z-scores were not multiplied by -1. Each animal's Z-Anxiety is simply an average across all three assays, given by the following:

$$Z - Anxiety = \frac{(z - score_{EZM} * -1) + (z - score_{LDB} * -1) + (z - score_{NSF})}{3}$$

For the sex-separated quantification of Z-Anxiety, the only modification made was that the z-score for each assay for all males was found in reference to the WT male average and standard deviation; likewise, z-scores for females were made in reference to the WT female average and standard deviation for each assay.

Sociability Assays

Object and social interaction: Mice were habituated to the empty, clear arena (46 x 23 cm). Once 120 s elapsed, a single novel object made from two glued 6-well plates was temporarily

secured at one end of the arena with a magnet. Mice were free to explore this object for another 120 s. After this time elapsed, the novel object was removed, and was immediately replaced by a novel, juvenile (3-5 weeks old), sex-matched CD-1 stimulus mouse. These mice were then free to interact with one another unencumbered for 120 s.

Three-chamber sociability assay: One day prior to the three-chamber assay, mice were habituated to the three-chamber apparatus (three 46 x 23 cm chambers connected by passageways (10 cm wide)). These mice were free to explore the entire three-chamber arena with an empty wire-mesh pencil cup in each of the two end chambers for 5 min. On that same day, stimulus CD-1 mice were habituated to being placed underneath these wire-mesh pencil cups for 5 min. The next day, the three-chamber assay was performed as follows. A novel, juvenile, sex-matched CD-1 stimulus mouse was placed underneath a cup in the chamber at one end of the arena, and an empty cup was placed in the chamber at the other end of the arena. Weights sat on top of the cups to ensure that the CD-1 stimulus mouse was secure in the cup and that the experimental mouse would not move either cup. To begin the assay, dividers were placed in the three-chamber passageways to block movement between chambers, and the experimental mouse was placed in the center chamber. After the mouse explored the center chamber for 120 s, the inserts were removed, and the mouse was free to explore the entire three-chamber arena, as well as the empty cup and mouse cup for 10 min. The chambers in which the mouse cup and empty cup were placed was alternated randomly across mice. The percent time spent in each of the three chambers was scored.

Behavior Analysis

In all behavioral assays (except for the NSF and Fresh Cage NSF), mice and relevant behavioral arena components were tracked using DeepLabCut (DLC) (61). To ensure the accuracy of tracking by DLC, a random sampling of videos from each day of experimentation were inspected. Next, a trained experimenter watched annotated videos to verify consistent tracking of fiducial points. Fiducial points included the snout, the nose bridge, the head, left and right ears, and five points that ran along the sagittal axis of the mouse body from the neck to the base of the tail. All corners of all arenas were labeled, as well as any relevant features of the arenas, including corners of objects and cups, and the thresholds separating the open and closed arms of the EZM. To calculate the interaction times, binary behavior matrices (vectorized behavior) indicating the location of the relevant key points of the mouse with respect to relevant key points of the arena (e.g., head of mouse and corner of object) were created using custom MATLAB scripts, available at (<https://github.com/CruzMartinLab/PV-mC4> [Project/tree/main/behavior_code](#)).

Neonatal viral injections

To genetically tag and identify PV cells in electrophysiology experiments, P1-3 pups were injected with 360 nL of AAV1-FLEX-tdTomato (titer: 2.5×10^{13} vg/mL, Addgene #28306) per cortical hemisphere. Borosilicate pipettes (BF150-117-10, Sutter Instrument Co., Novato, California) were pulled to a fine tip (approximately 3-15 μ m) and back-filled with mineral oil and inserted into the Nanoject Injector (Drummond, Nanoject II, 3-000-204/205A/206A). After cutting the tip of the pipette and emptying roughly half of the mineral oil, the pipette was filled with virus solution from the open tip. Prior to injection, pups were anesthetized via a cold-plate (approximately 15 min) and remained on the cold surface of an ice pack during injection to ensure continued anesthesia throughout the entire process. The mPFC was targeted along the A-P axis and hit consistently using empirically derived landmarks and the Allen Brain Atlas (55). From here, the tip of the pipette was moved medially into position immediately adjacent to the

midline. Injections at several depths were made to ensure effective labeling of the entire dorsal-ventral depth of mPFC target sub-regions. Fine spatial navigation of the tip was made using a stereotax (Kopf Instruments, Tujunga, California). Post-injection, pups recovered in a plastic chamber that was placed on top of a heated blanket. Pups were returned to the dam once fully recovered.

Electrophysiology

Acute slice preparation and recording: Mice (P40-62) were anesthetized with a 4% isoflurane-oxygen mixture (v/v) and perfused intracardially with ice-cold Perfusion/Slicing artificial cerebrospinal fluid (P/S-aCSF) bubbled with 95% O₂/ 5% CO₂ containing the following (in mM): 3 KCl, 26 NaHCO₃, 1.25 NaH₂PO₄, 212 sucrose, 10 glucose, 0.2 CaCl₂, and 7 MgCl₂ (300-310 mOsm). Thirty min before slicing, 200 mL of P/S-aCSF was transferred to -20°C until turned to a slushy consistency. Coronal slices 300-μm thick were cut in this slushy P/S-aCSF using a VT1000 S (Leica) vibratome and were then transferred to a Recording aCSF (R-aCSF) solution bubbled with 95% O₂/ 5% CO₂ containing the following (in mM): 125 NaCl, 2.5 KCl, 25 NaHCO₃, 1.4 NaH₂PO₄, 16 glucose, 0.4 Na-ascorbate, 2 Na-pyruvate, 2 CaCl₂, and 1 MgCl₂ (300-310 mOsm). Slices were incubated in this R-aCSF for 30 min at 35°C before being allowed to recover at room temperature for 1 h prior to recording.

Whole-cell voltage- and current-clamp recordings were performed in Layer (L) 2/3 of the PrL, IL, and AC cortex divisions of the mPFC (34). For all recordings, tdTomato-positive PV cells were identified using a Prior Lumen 200 Light Source (Prior Scientific) and a CMOS camera (ROLERA-BOLT-M-12; 1.3 MP, MONO, 12-BIT, UNCOOLED, QImagingBolt) mounted on an Olympus BX51WI microscope (Olympus America, Inc.). Pyramidal neurons (PYRs) were identified based on morphological and electrophysiological properties. All recordings were performed at 29-31°C. Signals were recorded with a 5X gain, low-pass filtered at 6 kHz, and digitized at 10 kHz using a patch-clamp amplifier (Multiclamp 700B, Molecular Devices). Nearly all recordings were made using 3-5 MΩ borosilicate pipettes (Sutter, BF-150-117-10). Series (R_s) and input resistance (R_{in}) were monitored throughout the experiment by measuring the capacitive transient and steady-state deflection in response to a -5 mV test pulse, respectively. Liquid junction potentials were calculated and left uncompensated.

Miniature post synaptic currents (mPSCs): For recording miniature excitatory post synaptic currents (mEPSCs), borosilicate pipettes were filled with an internal recording solution that contained the following (in mM): 120 Cs-methane sulfonate, 8 NaCl, 10 HEPES, 10 CsCl, 10 Na₂-phosphocreatine, 3 QX-314-Cl, 2 Mg²⁺-ATP, and 0.2 EGTA (292mOsm, adjusted to pH 7.3 with CsOH). PV cells and PYRs were voltage clamped at -70 mV in the presence of tetrodotoxin (TTX, 1 μM, Tocris) and picrotoxin (PTX, 100 μM, HelloBio). For each condition, 6-9 mice from 3-4 litters per sex per condition were used to collect all data for PV cells and PYRs.

For recording miniature inhibitory post synaptic currents (mIPSCs), borosilicate pipettes were filled with a high-chloride internal recording solution that contained the following (in mM): 60 Cs-methane sulfonate, 8 NaCl, 70 CsCl, 10 HEPES, 10 Na₂-phosphocreatine, 0.2 EGTA, and 2 Mg²⁺-ATP (290 mOsm, adjusted to pH 7.3 with CsOH). PV cells and PYRs were voltage clamped at -70mV in the presence of TTX (1 μM), CNQX (20 μM), and DL-APV (50 μM). Because this high-chloride internal solution has a chloride reversal potential of -13 mV, mIPSCs were inward. For each condition, 4-6 animals from 3-5 litters per sex per condition were used to collect all data for PV cells and PYRs.

mPSC analysis: mPSCs were identified and their amplitude, frequency, rise, and decay determined using custom scripts written in MATLAB. At least 120 s were analyzed for each cell. All mPSC raw traces were first lowpass filtered in Clampfit (Molecular Devices) using a boxcar filter. Next, local minima in the trace were recognized by identifying potential synaptic events using the native *islocalmin()* MATLAB function. After these events were filtered, a series of steps were taken to remove false-positives while simultaneously limiting the number of false negatives. More specifically, we calculated a threshold based on the standard deviation of the noise of the raw trace within a 1 s temporal window to differentiate the background noise from mPSCs, thus setting an amplitude threshold. Next, a series of thresholds based on the rise and decay times were used to filter subsequent postsynaptic events. For all remaining mPSCs, amplitude is given as the difference between the baseline current value (determined using a highly smoothed line of the raw data that effectively serves as a moving baseline of the trace) at the time when the peak reaches a minimum and the average of the 10 points around the absolute minimum of that mPSC peak.

Frequency (in Hz) of postsynaptic events is given by the number of mPSCs per min. Rise₁₀₋₉₀ is defined as the time (ms) it takes for the mPSC to progress from 10 to 90% of the peak of that mPSC. To find the Decay_{tau}, for each event, the trace from the peak of the mPSC to its return to baseline is isolated and fit to a single-term exponential. Between groups, R_s values for each cell type were not statistically significant.

R_s for PV cells (mEPSCs) were as follows (in M Ω): WT males: 16.85 ± 1.17 ; WT females: 17.35 ± 1.26 ; KI males: 16.27 ± 1.33 ; KI females: 15.56 ± 1.21 . R_s for PV cells (mIPSCs) were as follows (in M Ω): WT males: 15.79 ± 1.06 ; WT females: 13.93 ± 1.09 ; KI males: 15.40 ± 1.46 ; KI females: 14.07 ± 0.79 . R_s for PYRs (mEPSCs) were as follows (in M Ω): WT males: 15.94 ± 0.92 ; WT females: 14.81 ± 0.75 ; KI males: 15.07 ± 0.73 ; KI females: 15.00 ± 0.80 . R_s for PYRs (mIPSCs) were as follows (in M Ω): WT males: 14.43 ± 1.05 ; WT females: 13.81 ± 0.92 ; KI males: 16.13 ± 1.34 ; KI females: 14.88 ± 1.26 .

Active and passive properties: To determine the active and passive properties of PV cells and PYRs, borosilicate pipettes were filled with an internal solution that contained the following (in mM): 119 K-gluconate, 6 KCl, 10 HEPES, 0.5 EGTA, 10 Na₂-phosphocreatine, 4 Mg²⁺-ATP, and 0.4 Na-GTP (292 mOsm, adjusted to pH 7.3 with KOH). Cells were held at -65 mV during recording, and the bath was perfused with CNQX (20 μ M), DL-APV (50 μ M), and PTX (100 μ M). Excitability was assessed by measuring membrane voltage changes (i.e., current-evoked Action Potentials (APs)) to a spiking protocol that applied 500 ms square current pulses to the patched cell, beginning at -250 pA and rising in 30 pA steps to a max current injection of 470 pA. Passive properties of the patched cells were determined via a 500 ms, -20 pA square pulse that preceded the square pulse of increasing current amplitude. This protocol was run and recorded 2-3 times per cell, and final values were averaged across recordings for each cell.

Active properties analysis: To quantify spike frequency, the number of spikes (temporally defined as when the rising phase of the spike crossed 0 mV) was divided by the length of the current pulse (0.5 s). Rheobase was defined as the minimum current injection that evoked at least a single AP. The inter-spike interval (ISI) was determined by finding the difference (in ms) between the crossing of 0 mV of one spike to the crossing of 0 mV by the next spike. To capture time-dependent changes in the frequency of APs, ISI 1/9 and 4/9 were determined by dividing the first ISI by the fourth and ninth ISI, respectively. ISI ratios were taken from the first sweep with at least 10 spikes.

Passive properties analysis: To obtain the R_{in} , the difference between the baseline voltage (holding membrane voltage of approximately -65 mV) and the average voltage response to a -20 pA injection was divided by that current injection value of 20 pA. The membrane time

constant, τ , was the fitted response to the -20 pA injection. Membrane Capacitance (C_m) was determined by dividing τ by the R_{in} . The resting membrane potential (V_m) was measured as the potential before any current was injected. Finally, Voltage Sag Ratio was determined by dividing the difference between the minimum voltage at the peak deflection to a -500 pA current injection and the voltage of the steady state response by the difference between the minimum voltage at the peak deflection and the baseline voltage.

Electrophysiological data were analyzed using custom routines written in MATLAB, available at (https://github.com/CruzMartinLab/PV-mC4_Project/tree/main/ephys_code).

RESULTS:

A novel transgenic mouse line permits PV cell specific overexpression of complement component 4.

We generated a tunable conditional transgenic mouse based on a design by Dolatshad et al. (51) to reliably drive mC4-OE in specific cell types, and to incorporate genetic recombination switches that allow the conversion of different OE alleles for tunable transgene expression (**Fig. 1A**). Comer et al. (34) previously showed that mPFC neurons in postnatal day (P) 30 control mice express low levels of *C4b* transcript, which were not present in tissue from *C4b* knock-out mice (62).

To achieve specific OE of *mC4* in parvalbumin (PV)-positive cells under the strong CAG promoter (63,64), we crossed PV-Cre mice with a novel conditional mC4-OE KI mouse line, mC4-KI mice (**Fig. 1A, B**). Crossing the mC4-KI mouse to the PV-cre line generated litters that consisted of a mixture of pups that either inherited the floxed mC4-KI allele and thus overexpressed *mC4* specifically in PV cells (PV-mC4-KI, or KI) or did not inherit the floxed mC4-KI allele and were used as a wild type littermate control (PV-mC4-WT, or WT) (**Fig. 1B**). To control for effects of Cre-recombinase expression (65,66), we crossed homozygous PV-Cre mice to heterozygous mC4-KIs to obtain Cre recombinase expression in all offspring (**Fig. 1B**). We focused on a P40-60 temporal window, roughly equivalent to the young-adult life stage that immediately precedes SCZ symptom onset (67). PV-mC4-OE did not lead to observable health or locomotor deficits, as measured via weight (**Fig. 1C**) and distance traveled in an open field (OF) (**Fig. 1D**) in postnatal day P40-60 mice. Moreover, the density of mPFC PV cells – including the prelimbic (PrL), infralimbic (IL), and anterior cingulate (AC) cortices – was not significantly changed in PV-mC4-KI mice, compared to WT controls (**Fig. 1E-G**).

To validate our genetic approach, we used multiplex fluorescence *in situ* hybridization (M-FISH) in the mPFC of WT and KI mice (**Fig. 1H-J**). As early as P21, we observed a significant increase in the number of *mC4* mRNA puncta in PV cells in PV-mC4-KI mice, indicating reliable mC4-OE in this interneuron type (**Fig. 1I**). In contrast, we did not observe an increase in *mC4* puncta in neighboring somatostatin (SST)-positive cells and non-PV- non-SST-expressing cells (“Other”), suggesting that our genetic approach led to specific *mC4* overexpression in PV cells (**Fig. 1I**). In PV-mC4-KI mice, the number of *mC4* puncta in PV cells was not different between P21 and 65 (Two-way ANOVA, Age x Cell-type: $F_{(2,18)}=2.454$, $p=0.1142$. Age: $F_{(1,18)}=1.253$, $p=0.2777$. Cell-type: $F_{(2,18)}=42.19$, **** $p<0.0001$. Post-test: P21 vs. P65 PV, $p=0.1897$; SST, $p=0.9998$; Other, $p=0.9998$) (**Fig. 1I, J**), indicating that we achieved stable overexpression across development. Additionally, SST cells and Other cells of both P21 and P65 KI mice did not show differences in *mC4* expression compared to WT mice, indicating cell-type-specific maintenance of the OE into adulthood (**Fig. 1I, J**). Finally, quantification of PV mRNA puncta revealed that PV-mC4-OE did not drive any significant changes in the average number of PV

puncta per PV cell at either P21 (**Fig. 1K**) or P65 (**Fig. 1L**), suggesting that PV expression was not altered by increased levels of *mC4* in this cell type.

In summary, we have developed and validated a novel conditional mouse model that, when combined with various cre-driver lines, facilitates the study of distinct cellular sources of *mC4* overexpression. These results also indicate that we can reliably and specifically overexpress *mC4* in PV cells and that these transgenic mice are devoid of gross motor or health defects.

PV-specific mC4-OE causes an increase in anxiety-like behavior in male mice.

Anxiety and mood disorders are highly prevalent in SCZ patients, manifesting during the early stages of the illness and prior to episodes of psychosis (68–71). Moreover, PV cells have been shown to directly modulate stress and depression-like behavior in mice (72,73). To determine if PV-mC4-OE drives changes in anxiety-like behavior in P40-60 young adult mice, we used a battery of three assays, yielding a robust and reliable measure of behavior (60) (**Fig. 2A**). To assess arousal levels in these experiments, we measured the time spent by mice in a field's anxiogenic regions, namely the open or lighted areas (**Fig. 2B**, for example, Light-Dark Box assay (LDB)).

In the Elevated Zero Maze (EZM), we observed no differences in the amount of time spent in the open arms between WT and KI mice, suggesting that mice with increased *mC4* expression in PV neurons did not lead to an overall change in anxiety-like behavior, relative to WT controls (**Fig. 2C**). However, when we separated these data by sex, we observed a 46% reduction in the amount of time that KI males spent in the open arms, suggestive of a sexually dimorphic anxiety-like deficit (**Fig. 2D**). In contrast, we did not see such change in the behavior of KI female mice, relative to their WT controls (**Fig. 2D**). In the LDB, we observed a significant reduction in the overall time the PV-mC4-KIs spent in the light zone, again indicative of an increased anxiety-like response (**Fig. 2E**). Separation of the data by sex did not reveal the same sexual dimorphism as observed in the EZM (**Fig. 2F**). Lastly, in the novelty-suppressed feeding assay (NSF), we observed no change in the latency to feed between groups (**Fig. 2G**). Despite this, much like the EZM, there was a significant increase in anxiety-like behavior in the KI male mice, observed via a 91% increase in the latency to feed, but no such changes were observed in the PV-mC4-KI females (**Fig. 2H**). As a control, we performed the NSF in a safer environment, a mouse cage, where mice are exposed to bedding and can navigate close to the cage wall. In the Cage NSF, we did not observe differences in the latency to feed between groups (**Fig. S1**), suggesting that the NSF in the novel environment is anxiety-inducing and that the male KI mice, indeed, have increased anxiety-like behavior levels. These results suggest that increased levels of *mC4* in PV cells lead to sexually dimorphic alterations in anxiety-like behavior.

To describe the anxiety-like behavior of mice in response to PV-mC4-OE, we used a z-scoring approach that effectively tracks the performance of each mouse across all three behavioral assays, yielding a single score for each mouse, termed its Z-Anxiety score (60). We observed that the Z-Anxiety score of KI mice was increased in comparison to the control group (**Fig. 2I**). However, when we divided the data by sex, we discovered that overall heightened anxiety-like behavior is solely driven by a significant increase in the Z-Anxiety of PV-mC4-KI male mice (**Fig. 2J**). In support of this, female KI mice did not exhibit an increased Z-Anxiety score compared to the WT controls (**Fig. 2J**). In summary, PV-mC4-OE causes sex-dependent behavioral changes, with male KI mice exhibiting increased anxiety-like behavior relative to WT controls.

Increased levels of *mC4* in PV cells disrupts active but not passive social behaviors.

Next, we determined whether increased levels of *mC4* in in fast-spiking cells led to social behavioral changes. We employed a naturalistic freely-moving interaction assay between experimental mice and a novel, juvenile, sex-matched CD-1 stimulus mouse (**Fig. 3**). To disentangle deficits of general motivation in mice from alterations specifically in social interactions, social assays were preceded by an object interaction task (**Fig. 3A**). Both male and female KI mice spent a similar time interacting with a novel object as their WT counterparts (**Fig. 3B**). This suggests that PV-*mC4*-OE does not alter novelty-seeking behaviors or general motivation.

Immediately following the object interaction assay, the object was replaced with a juvenile stimulus CD-1 mouse, leaving it and the experimental mouse free to interact unencumbered (**Fig. 3C**). Interactions between WT or KI mice and the stimulus mouse were divided into one of five sub-classes that best described the interaction bout (**Fig. 3C**). Of all interaction sub-classes, experimental mice engaged in the active (active: experimental mouse initiating the interaction vs. passive: stimulus mouse initiating the interaction) snout-rear interaction (snout-ano-genital interactions) the most frequently, comprising half of all interaction time across all groups (**Fig. 3C, percentages**). Notably, PV-*mC4*-KI mice engaged less in active interactions than controls (**Fig. S2A**); this reduction was driven by a 24% decrease in active snout-rear interaction (**Fig. 3D**). These results indicate that increased levels of *mC4* in PV cells lead to a reduction of specific types of active behavior, snout-ano-genital interactions. Unlike the dimorphic nature of the anxiety-like behavior, deficits in sociability affected KI mice of both sexes (**Fig. 3D**).

In general, we observed more active than passive interactions by the experimental mice, which suggests that the experimental mice initiated more social interactions than the stimulus mice. (**Fig. 3C, Fig. S2A**). Furthermore, we did not notice any changes in passive interactions, indicating that stimulus mice behaved similarly when interacting with both KI and WT mice (**Fig. S2**). In the four remaining interaction sub-classes, which were less frequent than the active snout-rear interaction, WT and KI mice interacted similarly with their CD-1 stimulus mice (**Fig. S2B-I**).

Computing a Z-Sociability score that accounts for all five interaction sub-classes (the active and passive interactions from the freely-moving interaction assay) did not reveal a broad deficit in sociability (**Fig. 3E**). Separately, in a three-chamber assay of sociability, a more restricted social behavior assay, PV-*mC4*-OE did not alter the social preference of PV-*mC4*-KIs, relative to WT mice (**Fig. S3**), suggesting all experimental animals preferred to interact with a mouse rather than an object. Comparing the results from the two social tasks suggests that a naturalistic, freely-moving interaction assay allowed us to capture the complex behavioral phenotype that overexpressing mice exhibit.

Although female PV-*mC4*-KI mice did not exhibit changes in anxiety-like behavior, they had decreased social interactions as part of the KI group (**Fig. 3D**). This result indicates that the mechanisms driving the pathology in the anxiety-like and social behaviors in response to PV-*mC4*-OE likely function independently. Lastly, by deconstructing social behavior into sub-classes, we have also identified a deficit in the initiation of active snout-rear interactions in response to PV-*mC4*-OE that similarly impacts mice of both sexes. Overall, these results suggest that increased levels of *mC4* in PV cells disrupts the circuits that underlie emotional and social behavior in mice.

Sex-related differences in excitatory-inhibitory dynamics in mPFC PV cells with increased levels of *mC4* in PV cells.

To determine whether PV-mC4-OE changed the connectivity of circuits in the mPFC, we performed whole-cell voltage-clamp recordings in acute brain slices. Specifically, we first recorded miniature excitatory post-synaptic currents (mEPSCs) in PV neurons in layer (L) 2/3 of the mPFC in P40-60 mice (**Fig. 4A-G**), the same temporal window within which we identified anxiety-like deficits and social dysfunction in response to PV-mC4-OE. Though we observed no change in mEPSC amplitude in PV-mC4-KI mice (**Fig. 4A-C**), we observed a 39% reduction in PV cell mEPSC frequency specifically in KI male mice (**Fig. 4D**). In contrast, there were no changes in PV cell mEPSC frequency in the KI females (**Fig. 4D**). In support of this, there was a rightward shift in the cumulative frequency distribution of inter-event-intervals (IEIs) in PV neurons in PV-mC4-KI males, but not females (**Fig. 4E**), suggesting that PV-mC4-OE leads to a decrease in excitatory drive to this fast-spiking neuron. Finally, PV-mC4-OE did not alter the Rise₁₀₋₉₀ or Decay_{tau} of mEPSCs recorded in PV cells (**Fig. 4F, G**), suggesting that increased levels of *mC4* did not alter the kinetics of the excitatory postsynaptic response in PV cells.

To determine if PV-mC4-OE impacted the inhibitory drive to PV neurons, we recorded miniature inhibitory post-synaptic currents (mIPSCs) in this inhibitory cell type (**Fig. 4H-N**). Using a high-chloride internal recording solution with a chloride reversal potential of -13 mV yielded inward mIPSCs when recording at -70 mV (**Fig. 4H**). The voltage-clamp recordings revealed that PV-mC4-OE drove a 16% increase in the amplitude of PV cell mIPSCs (**Fig. 4I**) and a concomitant rightward shift in the cumulative distribution of mIPSC amplitudes (**Fig. 4J**) specifically in male, but not female mice, suggesting that inhibitory inputs are increased in PV cells in male PV-mC4-KI mice. Additionally, we observed no changes in mIPSC frequency (**Fig. 4K, L**), Rise₁₀₋₉₀ (**Fig. 4M**), or Decay_{tau} (**Fig. 4N**) between groups.

Taken together, these results suggest that PV-mC4-OE drives sex-dependent alterations in the excitatory and inhibitory connections to this cell type, mirroring the sexually dimorphic changes in anxiety-like behavior. The combined effects of reduced excitation and increased inhibition to PV cells in KI male mice suggests hypoactivity of mPFC inhibitory circuits in response to increased levels of *mC4* in fast-spiking cells.

PV-specific mC4-OE leads to opposing changes in the excitability of PV cells in male and female mice.

We next evaluated the passive and active properties of both mPFC L2/3 PV cells (**Fig. 5A-E, Table S1**) and PYRs (**Fig. 5F-J, Table S2**). To accomplish this, we injected steps of hyperpolarizing and depolarizing current pulses and recorded the membrane voltage changes. Changes in excitability of PV neurons in male and female mice in response to PV-mC4-OE diverged: while there was a significant decrease in PV cell spike frequency in KI male mice relative to WT males (**Fig. 5A, B**), we observed a significant increase in the spike frequency of PV cells in female overexpressing mice, compared to WT females (**Fig. 5A, C**). This increase in excitability in PV cells in KI females was also accompanied by a 26% reduction in their rheobase, another indication of increased excitability (**Fig. 5D**). Finally, we observed a significant shift towards a more depolarized resting membrane voltage in KI mice overall (**Fig. 5E, Table S1**). These results suggest that there is a sex-dependent divergence in the fast-spiking cell's excitability with higher *mC4* levels in PV cells.

Though mC4-OE is limited to PV cells in this mouse model, it is possible that disruption in the activity of PV cells may elicit compensatory changes in PYRs. To this end, we used the same

hyperpolarizing and depolarizing current steps and recorded the membrane voltage response as before, now in mPFC L2/3 PYRs (**Fig. 5F-J**). Similar to PV neurons, in male mice PV-mC4-OE drove a reduction in the spike frequency of PYRs (**Fig. 5F, G**). Unlike PV cells, we observed no changes in spike frequency in the PYRs of KI female mice (**Fig. 5F, H**). Moreover, PV-mC4-OE did not alter the rheobase (**Fig. 5I**) or resting membrane voltage (**Fig. 5J**) of PYRs. These results indicate that PV-mC4-OE induced changes in PYR excitability in male mice.

Overall, increased *mC4* levels in PV cells caused sexually dimorphic effects on the excitability of cortical cells. Fast-spiking cells and PYRs in males showed a decrease in excitability, while fast-spiking cells in females exhibited hyperexcitability. This divergent outcome suggests that the male and female mouse brain respond to complement dysfunction in opposing ways.

PV-specific mC4-OE alters the kinetics of mEPSCs in PYRs of female mice.

To further investigate whether PV-mC4-OE drives responsive changes in PYRs, we recorded mEPSCs (**Fig. S4A-E**) and mIPSCs (**Fig. S4F-J**) in L2/3 mPFC PYRs. PV-mC4-OE did not lead to changes in PYR mEPSC amplitude (**Fig. S4B**) or frequency (**Fig. S4C**) compared to controls. While the Rise₁₀₋₉₀ of the mEPSCs in PYRs was also not altered in PV-mC4-KI mice (**Fig. S4D**), PV-mC4-OE caused a 14% reduction in Decay_{tau} of the mEPSCs in KI female, but not KI male mice (**Fig. S4E**). This is perhaps suggestive of a change in receptor subunit composition in PYRs (74) or a change in the location of PYR excitatory synapses along its somatodendritic axis (75,76). Finally, PV-mC4-OE did not induce any changes in PYR mIPSC amplitude (**Fig. S4G**), frequency (**Fig. S4H**), or kinetics (**Fig. S4I, J**) relative to controls. These results suggest that PV-mC4-OE largely does not alter the synaptic drive of mPFC PYRs.

No changes in anxiety-like behavior with pan-neuronal overexpression of *mC4*.

After observing that conditional overexpression of *mC4* in PV cells resulted in consequential cellular and behavioral abnormalities in male mice, we investigated whether increasing the levels of *mC4* in all neurons would lead to similar pathology. To this end, we crossed the mC4-KI mouse to the BAF53b-Cre transgenic mouse line (50) that express Cre recombinase under the control of the mouse *Actl6b* gene promoter to drive mC4-OE in all neurons (PanN-mC4-OE). The expression of the BAF53b gene in neurons can first be detected during embryonic day 12.5 in the brain and spinal cord (77). Using a similar breeding strategy as with the PV-mC4-KI mice, litters consisted of a mix of mice that inherited the floxed mC4-KI allele and thus overexpressed *mC4* in all neurons (PanN-mC4-KI), or littermates that did not inherit the floxed mC4-KI allele, and thus were effectively WT (PanN-mC4-WT) (**Fig. 6A, top**).

We employed the same battery of three assays – the EZM, LDB, and NSF – to test anxiety-like behavior in P40-60 PanN-mC4-WT and PanN-mC4-KI mice (**Fig. 6A, bottom**). First, overexpression of *mC4* in all neurons did not alter the distance traveled compared to controls (**Fig. 6B, C**), suggesting that PanN-mC4-KI mice exhibited intact locomotor activity. Moreover, compared to controls, we did not observe any deficit in anxiety-like behavior in PanN-mC4-KI mice in the EZM (**Fig. 6D, E**), LDB (**Fig. 6F, G**), or NSF (**Fig. 6H, I**). Also, we did not observe any increase in Z-Anxiety in PanN-mC4-KI mice overall (**Fig. 6J**), nor in either sex specifically (**Fig. 6K**). Taken together, PanN-mC4-OE does not drive anxiety-like behavior in mice.

Overall, we have demonstrated that increased levels of *mC4* in PV neurons resulted in a strong, sex-specific anxiety-like phenotype not observed in pan-neuronal *mC4* overexpressors. This

suggests that specific complement changes in PV cells leads to developmental dysfunction of inhibitory circuits that is more detrimental to brain function than pan-neuronal alterations.

DISCUSSION

Using a new model to conditionally overexpress *mC4*, we have discovered that mPFC PV cells in male mice are susceptible to complement dysfunction. Additionally, we have established a connection between the *mC4*-driven alterations in the circuitry of the mPFC and pathological anxiety-like behavior in male mice. Increased levels of *mC4* in PV neurons disrupted both excitatory and inhibitory inputs to fast-spiking cells in male but not female mice. Furthermore, we have also demonstrated that specific overexpression of *mC4* in PV cells led to opposing effects on the excitability of cortical cells. While *mC4*-OE in PV cells drove a decrease in the excitability of both male fast-spiking cells and PYRs, it led to hyperexcitability of female PV cells. By utilizing a Cre-driver line to induce *mC4* overexpression in all neurons, we also observed that specific *mC4* dysfunction in PV cells has a greater adverse effect on anxiety-like behavior than widespread neuronal complement alterations. Overall, these results establish a causative link between the schizophrenia-associated gene *C4* and the vulnerability of fast-spiking cells, which are crucial for the function of the mPFC.

Synaptic alterations in fast-spiking cells with PV-specific *mC4* overexpression

Here, we demonstrate that in male mice, conditionally targeting *mC4* overexpression to PV cells leads to a drastic loss of excitatory drive on this inhibitory cell type that is accompanied by increased inhibition. Several lines of evidence point to synaptic dysfunction and pathological excitatory synaptic loss as prominent features of SCZ (22,78–80). In support of this, our group previously demonstrated that increased levels of *C4* in L2/3 mPFC PYRs is sufficient to cause a significant loss of excitatory synapses during early postsynaptic development, leading to mPFC circuit dysfunction (34). Our electrophysiological recordings reveal a significant decrease in the frequency of mEPSCs on PV cells without alterations in their amplitude. Although the underlying mechanism is not clear, our results suggest that the decline in excitatory drive to fast-spiking cells is either a reduction in the probability of presynaptic release or synapse number. In conjunction with our previous results (34), this suggests that synapse loss is the most likely mechanism of hypoconnectivity in fast-spiking neurons.

While it is yet to be established whether pathological dysfunction in SCZ is confined to specific microcircuitry, a study conducted in post-mortem tissue of patients with SCZ demonstrated that there is a decrease of excitatory synapses on PFC PV cells relative to control subjects (22). In support of this, dysregulated ErbB4, a receptor of the SCZ-linked protein neuregulin-1, may contribute to lower activity of PV cells by reducing their excitatory inputs (81). A decrease in the excitatory drive to fast-spiking interneurons has also been observed in mouse models of AD (82,83) and other neurodevelopmental disorders (72,84,85), suggesting that dysfunction in feed-forward excitatory synapses to fast-spiking cells is a common denominator in brain pathology.

A long-standing hypothesis is that defects in the GABAergic inhibitory system can contribute to SCZ (86). Additionally, cognitive dysfunction in SCZ could be the result of dysfunction in the convergence of glutamatergic and GABAergic systems (87). One possible outcome of decreased feed-forward excitation on PV cells in the male PV-*mC4*-KI mouse is a disruption in

the dynamics of excitation and inhibition, tipping the scales towards the side of unchecked excitation and excess glutamatergic release. This also aligns well with the NMDA-hypofunction SCZ model (88–90), where the loss of NMDA receptors, specifically on interneurons, results in hypoactivity of PV neurons. Separate from the NMDA centric-hypothesis, alterations in inhibitory circuitry could alter the timing of excitation and inhibition (91) that controls oscillatory activity and information flow (10).

In addition to a reduction in the excitatory drive to fast-spiking cells in response to mC4-OE in PV cells, we also observed a significant increase in the amplitude of mIPSCs in PV cells in PV-mC4-KI male, but not female mice, potentially suggesting an enhancement of inhibitory inputs to fast-spiking interneurons. Naturally, this effect would amplify the putative decrease in PV cell activity in the male PV-mC4-KI mice, already caused by the reduction in the excitatory drive to this interneuron. As increased inhibition of PV cells is counterintuitive to the effects that increased complement activity would have on inhibitory synapses or a compensatory change to enhance the drive of fast-spiking cells, we can only conclude that these are mC4-driven maladaptive changes in the male brain. Whether this increase in mIPSC amplitude is driven by presynaptic changes in quantal size or postsynaptic changes in GABA receptor subunit composition or sensitivity is unknown and will require deeper investigation.

Microglia-dependent synaptic engulfment is an established mechanism for complement-driven synaptic loss in the normal and diseased brain (49,92–94). Studies using mice that lack specific complement pathway members have shown that these immune molecules contribute to synaptic plasticity (95–97). In fact, complement upregulation has been observed in several neurodegenerative diseases where synaptic loss is a prominent feature (98–100).

A recent study showed that C1q, the initiating member of the classic complement pathway, binds neuronal activity-regulated pentraxin (Nptx2) (101), an immediate early gene highly enriched at excitatory synapses on PV cells, where we observed the most drastic phenotype. Furthermore, deletion of Nptx2 caused increased activation of the classical complement pathway and microglia-mediated elimination of excitatory synapses on PV cells (101), supporting this established mechanism of synaptic loss in excitatory inputs on PV neurons. Still, other non-glia mechanisms could underlie excitatory synaptic loss in interneurons. In support of this, our group previously (102) used super-resolution imaging in mPFC slices (103) to demonstrate that increased levels of *mC4* accelerate the accumulation of the postsynaptic receptor GluR1 in neuronal LAMP1-positive lysosomes, leading to pathological synaptic loss.

The mPFC and neuropsychiatric disorders

The lifetime prevalence of anxiety disorders is close to 30% (104) and it is highly comorbid with other neuropsychiatric disorders, including SCZ (68,105,106). Grasping the mechanisms behind the comorbidity of neuropsychiatric disorders is challenging, given the complex interplay of genetic, neurobiological, psychological, and environmental influences. Nonetheless, our approach of conditionally targeting *mC4* in fast-spiking cells provides a unique example that establishes a causal relation between elevated levels of the SCZ-associated gene *C4* in these cells and enhanced anxiety-like behavior and mPFC circuit dysfunction in male mice, shedding light on the intricate dynamics of neuropsychiatric disorders.

We have focused on the mPFC to establish a connection between altered circuitry and disrupted emotional behavior. The mPFC is responsible for learning and predicting the potential outcomes of different actions (107). Genetic insults and chronic stress have lasting effects on the PFC that lead to alterations in cognitive and social function (108–111). In the mouse mPFC, inhibitory neurons respond to a variety of social and emotional stimuli (59,112). Additionally, PV cells coordinate and enhance the neuronal activity of PFC projection neurons to drive fear expression in the mouse (113). Consistent with its function in regulating emotional behavior, we observed that increased levels of *mC4* in PV neurons lead to synaptic alterations in fast-spiking cells and opposing effects in the excitability of cortical cells in male and female mice.

The PFC is a pivotal brain region in the regulation of various dimensions and aspects of social behavior (108,114,115). It plays a crucial role in social cognition, enabling us to understand and interpret the actions of others, which is fundamental for effective social interaction (114,115). Here we show that while increased levels of *mC4* in PV cells did not cause a drastic deficit in social behavior, overexpressing mice exhibited deficits in subclasses of exploratory social behavior – namely snout-ano-genital interactions – linking defects in inhibitory circuits to the initiation of social behaviors. In support of the role of mPFC PV neurons in the regulation of social behavior, Bicks et al. (116) demonstrated that PV cell activity in the mPFC preceded an active social episode, or an episode initiated by the experimental mouse, but not during a passive interaction when a stimulus mouse explored the experimental mouse. Similarly, we showed that increased levels of *mC4* in PV cells lead to deficits in active but not passive social interactions. Finally, in contrast to the deficits in anxiety-like behavior, *mC4*-driven social deficits were not sexually dimorphic, suggesting that in mice social deficits might have a distinct etiology from pathological anxiety-like behavior.

The combination of high metabolic demand, unique properties, and sensitivity to environmental stressors means that fast-spiking PV cells are particularly vulnerable to damage in conditions where oxidative stress and neuroinflammation are present (117). In support of this, gamma oscillations in the hippocampus require high complex I gene expression and strong functional performance of mitochondria (118). Despite the lack of understanding surrounding the mechanisms underlying *mC4* dysfunction in PV cells, the resulting impairments in synaptic connectivity and excitability are in line with disturbances in the metabolic processes of fast-spiking cells.

Complement dysfunction and the mPFC

Although there is a strong link between immune dysfunction and neuropsychiatric disorders (49,119–124), more research is needed to establish a connection between complement dysfunction and specific circuitry underlying emotional and social behavior. Disruption of *Csmd1*, which is a C4 inhibitor, induces behaviors reminiscent of blunted emotional responses, anxiety, and depression (125). Additionally, Crider et al. (126) found a significant increase in C3 expression, a downstream effector of *mC4*, in the PFC of depressed suicide subjects. In this study conducted by Crider et al.(126), mice lacking the complement C3a Receptor (*C3aR KO*) exhibited a specific rise in innate, unconditioned anxiety, whereas *C3 KO* mice displayed a particular elevation in learned, conditioned fear, suggesting a possible link between complement dysfunction and emotional dysregulation. Crider et al. (126) also showed that expression of *Cacna1d*, which encodes the Cav1.3 subunit of L-type voltage-gated calcium channels and is

a SCZ susceptibility gene, was increased in the mPFC of *C3aR KO* mice, providing a link between altered complement levels and mPFC dysfunction. Together with previous results from our group (34), we showed that *mC4* alterations in specific cell-types are linked to mPFC-related pathologies. In summary, these results suggest that the prefrontal cortex is a brain region susceptible to pathological complement activity.

Increased levels of *mC4* in PV neurons, cortical function, and sex-related pathologies

Using a new mouse model, we show that targeted overexpression of mouse *C4* (*mC4*), specifically in fast-spiking cells, induces pathological anxiety-like behavior in male mice while sparing females. In the male mPFC, this sex-specific behavioral change correlates with a decrease in excitatory synaptic inputs to fast-spiking neurons, coupled with an increase in their inhibitory synapses, potentially resulting in reduced activity of this interneuron. During development, the maturation of fast-spiking cells contributes to the wiring of the neural networks, controlling the critical window of plasticity (127–130). Therefore, alterations in the developmental plasticity windows driven by increased levels of *mC4* in PV cells may cause the synaptic and excitability deficits we observed in the mPFC. There are also sex-dependent differences in the developmental cortical mechanisms of plasticity (131), which are regulated, in part, by PV cell activity, including their feed-forward circuits (128,129). Therefore, increased levels of *mC4* in fast-spiking cells could alter the function of the mPFC through distinct mechanisms in males and females, explaining the sex-divergent outcomes.

Since female mice with increased levels of *mC4* in PV neurons did not exhibit enhanced anxiety-like behavior, our data suggests that the hyperexcitability of female PV cells is a compensatory adaptation that protects them from *mC4* dysfunction. However, as the hyperexcitability of PV cells was one of the only phenotypes we observed in female mice, it could contribute to deficits in social behavior in females. More work will be needed to determine how specific cellular deficiencies contribute to the behavioral abnormalities of overexpressing *mC4*, or whether some of the *mC4*-driven cellular deficits are compensatory mechanisms.

In humans, there is a marked divergence in the incidence, cast of symptoms, and age of onset of SCZ in men compared to women. This sex-dependent imbalance is most obvious in the prevalence of SCZ between sexes, impacting men vs. women at a rate of roughly 1.4:1 (132,133) but is further supported by an earlier onset of symptoms and poorer functional outcome in males (134–137). The SCZ odds ratio associated with the four most common *C4* alleles are all higher in men than in women (138) and protein levels of *C4* and downstream effector, *C3*, are also higher in the cerebrospinal fluid of men compared to women (138). In our mouse model, overexpression of *mC4* is comparable in male and female PV-*mC4*-KI mice, suggesting that differences in the levels of this immune gene cannot explain circuit and behavioral deficits, and it is most likely a consequence of its impact on downstream *mC4* effectors in male and female mice.

Microglia activity, which is triggered by complement activation, is sexually dimorphic (139) and could potentially explain the stronger phenotype in male mice. Additionally, other factors not related to neuroimmune function could contribute to *mC4*-driven pathology. For example, SCZ symptom severity in women tends to fluctuate in accordance with circulating estrogen levels, improving when the levels of this sex hormone are highest during the earlier follicular menstrual stage and during pregnancy, and worsening when estrogen levels drop *postpartum* and during

menopause (140–142). Further research is needed to understand the contribution of sex to molecular and cellular pathologies associated with neuropsychiatric disorders.

Weaknesses of this study

Overexpression of *mC4* in fast-spiking interneurons caused a sexually dimorphic cellular phenotype in the mPFC that accompanied changes in anxiety-like behavior in male mice. However, overexpression of *mC4* in PV cells is global, and we did not include recordings in other brain regions related to emotional regulation. Alterations in the inhibitory microcircuitry of other anxiety-related areas may underlie the behavioral effects that we have captured (143,144). For example, Wolff et al. (145) demonstrated that in the mouse basolateral amygdala, interneurons expressing PV- and SST cells bidirectionally control the acquisition of fear conditioning through two distinct disinhibitory mechanisms. Continuous monitoring of activity in the BNST and insula also enables the detection of changes in environmental threat levels and supports hypervigilant threat-monitoring processes in highly anxious individuals (146), suggesting a connection between BNST activity and anxiety. However, our findings demonstrate that conditional overexpression of *mC4* in fast-spiking cells results in synaptic and excitability deficits that are consistent with the role of mPFC PV cells in regulating emotional behavior.

SCZ is a complex disorder and it's likely that multiple genetic and non-genetic factors contribute to its pathogenesis, each potentially impacting synaptic function and the excitability of cortical cells in different but converging ways. Nevertheless, we provide a new mouse model where complement dysfunction in PV cells causes cellular and behavioral dysfunction reminiscent of PFC-associated neurological conditions.

Concluding Remarks

Here, we have generated a unique mouse model to overexpress *C4* conditionally. Diseases linked to increased *C4* levels often have autoimmune or inflammatory aspects. Therefore, this mouse can be used to target specific cell types and tissues to determine the role of this important gene in various diseases outside of the nervous system or test the efficacy of pharmacology to target complement-related diseases. Together with previous studies, we have established *C4* as an important regulator of pathological synaptic loss in the prefrontal cortex, a region associated with several neuropsychiatric disorders. Furthermore, by conditionally overexpressing *C4* in fast-spiking PV cells, we have identified a connection between dysfunction of inhibitory circuits in the prefrontal cortex and pathological anxiety-like behavior in male mice.

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AUTHOR CONTRIBUTIONS

L.A.F. and A.C-M. conceptualized experiments including formulating composition, goals, and scope of the paper and approaches for analyses. L.A.F., R.A.P., M.S., A.B., S.B., J.R.G and N.M.P.L. collected the data and performed experiments. L.A.F. and R.A.P. performed data curation. L.A.F., R.A.P., A.B., M.S., and A.C.M analyzed data. L.A.F. and R.A.P. contributed code for data analysis. L.A.F. and A.C-M. contributed to parts of the original draft, including figure design and generation. All authors contributed to revision and editing of the draft. A.C-M. obtained funding and supervised the project providing mentorship, oversight, and project administration.

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COMPETING INTERESTS

The authors declare that they have no competing interests.

DATA AVAILABILITY

Data are available at <https://osf.io/je38k/>

CODE AVAILABILITY

Custom-written routines are available at <https://github.com/CruzMartinLab>

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FIGURE LEGENDS

Figure 1. A novel transgenic mouse line permits PV cell specific overexpression of complement component 4. (A) Genetic targeting strategy that permits Cre- and Flp-dependent OE of *mC4*. Adenovirus SA-Frt-CAG promoter-Frt-loxP-3*-polyA-loxP-Kozak-mouse C4b CDS-polyA cassette was cloned into intro 1 of the ROSA26 locus, located on mouse chromosome 6. In the targeting vector, the positive selection marker (Neo) is flanked by self-deletion anchor (SDA) sites. Diphtheria toxin A (DTA) sites were used for negative selection. The constitutive KI allele can be obtained after Cre- or Flp-mediated recombination, yielding relatively strong or weak OE of *mC4*, respectively. (B) Schematic of mouse breeding scheme used to generate litters to overexpress *mC4* in PV cells under the strong CAG promoter. (C) In both sexes, increased levels of *mC4* in PV cells did not alter mouse weights compared to controls (Condition x Sex: $F_{(1,44)}=3.204$, $p=0.0803$. Condition: $F_{(1,44)}=0.6311$, $p=0.4312$. Sex: $F_{(1,44)}=103.2$, $****p<0.0001$. Post-test: WT vs. KI males, $p=0.9793$, WT, $N=13$, KI, $N=12$; females, $p=0.3947$, WT, $N=12$, KI, $N=11$). For both WT and KI mice, males weighed more than females (WT males vs. females, $****p<0.0001$, KI males vs. females, $****p<0.0001$). (D) OE of *mC4* in PV cells did not impact the distance traveled by mice in an open field (Condition x Sex: $F_{(1,34)}=0.04903$, $p=0.8261$. Condition: $F_{(1,34)}=0.004672$, $p=0.9459$. Sex: $F_{(1,34)}=2.657$, $p=0.1123$. Males: WT, $N=12$, KI, $N=7$. Females: WT, $N=7$, KI, $N=12$). (E) Representative confocal images of PV cells (magenta) in mouse mPFC in P55-74 WT (left) and KI (right) animals. scale bar = 100 μ m. L, layer. (F) PV cell density was analyzed in the mPFC, indicated by the yellow box. Coordinates relative to Bregma. AP, Anterior-posterior axis. (G) No difference in PV cell density in the mPFC between WT and KI (Mann-Whitney test, $p=0.2319$, WT, $N=8$, KI $N=7$). (H, top two rows) Representative confocal images (40x) of a PV cell (first row) and somatostatin (SST) cell (second row) in a WT mouse. (H, bottom two rows) Representative confocal images (40x) of a PV cell (first row) and SST cell (second row) in a KI mouse. For all rows in H, left to right: DAPI (gray), *mC4* (red), PV (green), SST (cyan), merged image without DAPI. (I) In P21 KI mice, *mC4* was significantly overexpressed in PV cells, but not in SST cells or all other DAPI-labeled cells (Condition x Cell-type: $F_{(2,18)}=41.62$, $****p<0.0001$. Condition: $F_{(1,18)}=113.7$, $****p<0.0001$. Cell-type: $F_{(2,18)}=38.22$, $****p<0.0001$. Post-test: WT vs. KI: PV, $****p<0.0001$, SST, $p=0.1793$, neighboring non-PV cell or non-SST cell (Other), $p=0.2166$). In P21 KI mice, *mC4* expression was significantly greater in PV cells than in SST cells (PV vs. SST, $****p<0.0001$) and all other DAPI-labeled cells (PV vs. Other, $****p<0.0001$). (J) In P65 mice, *mC4* was overexpressed in PV cells, but not in SST cells or all other DAPI-labeled cells (Condition x Cell-type: $F_{(2,24)}=33.42$, $****p<0.0001$. Condition: $F_{(1,24)}=66.63$, $****p<0.0001$. Cell-type: $F_{(2,24)}=32.59$, $****p<0.0001$. Post-test: WT vs. KI: PV, $****p<0.0001$, SST, $p=0.6948$, Other, $p=0.7839$). In KI mice, *mC4* expression was significantly greater in PV cells than in SST cells (PV vs. SST, $****p<0.0001$) and all other DAPI-labeled cells (PV vs. Other, $****p<0.0001$). (K, L) No differences in PV expression between groups (P21: WT vs. KI: Mann-Whitney test, $p=0.7857$; P65: WT vs. KI: Mann-Whitney test, $p=0.0952$). WT: $N=5$ at P21, $N=5$ at P65. KI: $N=3$ at P21, $N=5$ at P65. WT: blue squares, KI: red circles. For all statistics, $*p<0.05$, $**p<0.01$, $***p<0.001$, $****p<0.0001$, Two-way ANOVA, unless otherwise stated. Mean \pm SEM shown.

Figure 2. PV-specific *mC4*-OE causes an increase in anxiety-like behavior in male mice. (A) Schematic timeline of assays used to measure anxiety-like behavior. Mice were tracked in the light zone, represented, for example, by a green outline in the LDB cartoon (B). (B) Representative

trajectories of WT (*blue*) and KI (*red*) mice in the light zone of the LDB (colored traces, via DLC). Yellow triangle, opening of the light zone. **(C)** Percent time spent in the open arms of the EZM did not differ between WT and KI mice (Mann-Whitney test, $p=0.1912$). **(D)** Decreased time in the EZM open arms in male but not female KIs relative to WT. (Condition x Sex: $F_{(1,34)}=5.324$, $*p=0.0273$. Condition: $F_{(1,34)}=3.579$, $p=0.0671$. Sex: $F_{(1,34)}=0.06062$, $p=0.8070$. Post-test: WT vs. KI males, $*p=0.0109$, females, $p=0.9474$). **(E)** KI mice spent significantly less time in the light zone of the LDB compared to WT mice (t-test with Welch's correction, $*p=0.0130$). **(F)** No sex-dependent differences in LDB light zone time between groups (Condition x Sex: $F_{(1,34)}=0.3253$, $p=0.5722$. Condition: $F_{(1,34)}=4.137$, $*p=0.0498$. Sex: $F_{(1,34)}=6.147$, $*p=0.0183$. Post-test: WT vs. KI males, $p=0.5211$, females, $p=0.1430$). **(G)** Latency to feed in the NSF did not differ between groups (unpaired t-test, $p=0.1964$). **(H)** Significant increase in the latency to feed for KI males compared to WT controls (Condition x Sex: $F_{(1,33)}=6.718$, $*p=0.0141$. Condition: $F_{(1,33)}=2.061$, $p=0.1606$. Sex: $F_{(1,33)}=0.4862$, $p=0.4905$. Post-test: WT vs. KI males, $*p=0.0159$, females, $p=0.6584$). **(I)** Significant increase in the Z-Anxiety of KI mice relative to WT (unpaired t-test, $*p=0.0114$). **(J)** Compared to WT controls, there was a significant increase in the Z-Anxiety of KI male but not KI female mice. (Condition x Sex: $F_{(1,34)}=5.209$, $*p=0.0289$. Condition: $F_{(1,34)}=5.325$, $*p=0.0272$. Sex: $F_{(1,34)}=5.756$, $*p=0.0221$. Post-test: WT vs. KI males, $**p=0.0053$, females, $p=0.9998$). WT: blue squares, $N=12$ males, $N=7$ females. KI: red circles, $N=7$ males, $N=12$ females. For all statistics, $*p<0.05$, $**p<0.01$, Two-way ANOVA, unless otherwise stated. Mean \pm SEM shown.

Figure 3. Increased levels of *mC4* in PV cells disrupts active but not passive social behaviors. **(A)** Schematic (*left*) of object interaction task used to assess motivation to seek novelty. Representative trajectories of WT (*middle, blue*) and KI (*right, red*) exploring a novel object (*black rectangle*). **(B)** KI mice explored the novel object as much as WT controls (Condition x Sex: $F_{(1,44)}=1.068$, $p=0.3072$. Condition: $F_{(1,44)}=0.008471$, $p=0.9271$. Sex: $F_{(1,44)}=0.02598$, $p=0.8727$). **(C)** Representative images of interaction sub-classes in a social interaction task identified using DLC and custom-written algorithms (see *Methods*). *Active*, experimental mouse engaging the stimulus CD-1; *Passive*, stimulus CD-1 mouse engaging the experimental mouse. Percentage of each sub-class of behavior is indicated below the image. **(D)** *Left*: Relative to WT controls, KI mice spent significantly less time engaged in the active snout-rear interaction type, the most prominent type of social behavior (t-test with Welch's correction, $**p=0.0099$). *Right*: There were no sex-related differences in active-snout-rear interaction between groups. (Condition x Sex: $F_{(1,44)}=0.007274$, $p=0.9324$. Condition: $F_{(1,44)}=7.127$, $*p=0.0106$. Sex: $F_{(1,44)}=0.1121$, $p=0.7394$. Post-test: WT vs. KI males, $p=0.1325$, females, $p=0.1218$). **(E)** There was no significant change in Z-Sociability in either KI male or female mice (Condition x Sex: $F_{(1,44)}=1.808$, $p=0.1857$. Condition: $F_{(1,44)}=0.09738$, $p=0.7565$. Sex: $F_{(1,44)}=1.507$, $p=0.2261$). WT: blue squares, $N=13$ males, $N=12$ females. KI: red circles, $N=12$ males, $N=11$ females. For all statistics, $*p<0.05$, $**p<0.01$, Two-way ANOVA, unless otherwise stated. Mean \pm SEM shown.

Figure 4. Sex-related difference in excitatory-inhibitory dynamics in mPFC PV cells with increased levels of *mC4* in PV cells. **(A)** Representative 1 s voltage-clamp traces of mEPSCs recorded in PV cells of P40-60 WT (*blue*) and KI (*red*) mice, scale bar = 250 ms, 10 pA. **(B)** No significant change in mEPSC amplitude in KI mice, compared to controls (Condition x Sex:

$F_{(1,66)}=1.111$, $p=0.2956$. Condition: $F_{(1,66)}=0.7845$, $p=0.3790$. Sex: $F_{(1,66)}=5.552$, $*p=0.0214$. Post-test: WT vs. KI males, $p=0.3225$, females, $p=0.9910$). (C) No shift in the cumulative frequency distribution of mEPSC amplitudes in KI mice (Kolmogorov-Smirnov test, WT vs. KI males, $p=0.5758$, females, $p=0.7810$). (D) *mC4* overexpression led to a significant decrease in mEPSC frequency in KI male mice, but not KI female mice (Condition x Sex: $F_{(1,66)}=2.138$, $p=0.1485$. Condition: $F_{(1,66)}=8.979$, $**p=0.0038$. Sex: $F_{(1,66)}=0.3501$, $p=0.5561$. Post-test: WT vs. KI males, $**p=0.0051$, females, $p=0.4809$). (E) Increased *mC4* expression caused a rightward shift in the distribution of mEPSC inter-event-intervals (IEIs) of PV cells in KI male mice, but not KI female mice (Kolmogorov-Smirnov test, males $**p=0.0059$, females, $p=0.3079$). (F) mEPSC Rise₁₀₋₉₀ was not impacted in KI mice, relative to WT controls (Condition x Sex: $F_{(1,66)}=0.8620$, $p=0.3566$. Condition: $F_{(1,66)}=0.02164$, $p=0.8835$. Sex: $F_{(1,66)}=1.121$, $p=0.2935$). (G) mEPSC Decay_{tau} was not changed in KI mice (Condition x Sex: $F_{(1,66)}=1.426$, $p=0.2367$. Condition: $F_{(1,66)}=0.9701$, $p=0.3282$. Sex: $F_{(1,66)}=2.700$, $p=0.1051$). (H) Representative 1s voltage clamp traces of mIPSCs recorded in PV cells of P40-60 WT (blue) and KI (red) mice, scale bar = 250 ms, 10 pA. (I) mIPSC amplitude was significantly increased in KI male, but not KI female mice (Condition x Sex: $F_{(1,53)}=1.557$, $p=0.2176$. Condition: $F_{(1,53)}=5.831$, $*p=0.0192$. Sex: $F_{(1,53)}=0.2637$, $p=0.6097$. Post-test: WT vs. KI males, $*p=0.0232$, females, $p=0.6599$). (J) Increased expression of *mC4* caused a rightward shift in the distribution of mIPSC amplitudes in KI male mice (Kolmogorov-Smirnov test, WT vs. KI males, $*p=0.0404$, females, $p=0.9048$). (K) mIPSC frequency was not changed in KI mice (Condition x Sex: $F_{(1,53)}=1.117$, $p=0.2954$. Condition: $F_{(1,53)}=0.04095$, $p=0.8404$. Sex: $F_{(1,53)}=0.5669$, $p=0.4548$). (L) No shift in the distribution of mIPSC IEIs of KI mice (Kolmogorov-Smirnov test, WT vs. KI, males $p=0.4149$, females, $p=0.9988$). (M) Relative to controls, there were no changes in mIPSC Rise₁₀₋₉₀ with increased levels of *mC4* in PV cells (Condition x Sex: $F_{(1,53)}=7.666$, $**p=0.0077$. Condition: $F_{(1,53)}=0.2031$, $p=0.6541$. Sex: $F_{(1,53)}=0.4950$, $p=0.4848$. Post-test: WT vs. KI males, $p=0.0507$, females $p=0.2079$). (N) mIPSC Decay_{tau} was not changed in KI mice (Condition x Sex: $F_{(1,53)}=1.708$, $p=0.1969$. Condition: $F_{(1,53)}=0.01747$, $p=0.8953$. Sex: $F_{(1,53)}=0.1166$, $p=0.7341$). WT: blue squares. KI: red circles. *N* represents cells. mEPSC WT *N*=20 males, *N*=17 females; mEPSC KI: *N*=15 males, *N*=18 females. mIPSC WT *N*=14 males, *N*=14 females; mIPSC KI: *N*=15 males, *N*=14 females. For all statistics, $*p<0.05$, $**p<0.01$, Two-way ANOVA used, unless otherwise stated. Mean \pm SEM shown.

Figure 5. PV-specific *mC4*-OE leads to opposing changes in excitability of PV cells in male and female mice. (A) Representative current-clamp recordings of PV cells injected with -100 (black), 140 (darker colored shade), and 230 pA (lighter colored shade) of current from WT (blue) and KI (red) mice, scale bar = 200 ms, 200 pA/20 mV. Baseline membrane voltage was maintained at approximately -65 mV for all cells. (B) PV cells in KI male mice spike significantly less than PV cells in WT male mice (Repeated measure (RM) Two-way ANOVA, Condition x Current: $F_{(24,840)}=1.941$, $**p=0.0045$. Condition: $F_{(1,35)}=1.989$, $p=0.1672$. Current: $F_{(1,656,57,95)}=416.4$, $****p<0.0001$). (C) *mC4* overexpression led to a significant increase in the excitability of PV cells in KI female mice, relative to controls (RM Two-way ANOVA, Condition x Current: $F_{(24,816)}=3.428$, $****p<0.0001$. Condition: $F_{(1,34)}=4.178$, $*p=0.0488$. Current: $F_{(1,719,58,46)}=568.3$, $****p<0.0001$). (D) The Rheobase of PV cells was significantly decreased in KI female mice, relative to controls (Condition x Sex: $F_{(1,69)}=10.23$, $**p=0.0021$. Condition: $F_{(1,69)}=0.008666$, $p=0.9261$. Sex: $F_{(1,69)}=2.932$, $p=0.0913$. Post-test: WT vs. KI males, $p=0.0596$, females, $*p=0.0472$). (E) Overall, PV-*mC4*-OE drove a shift in PV cell resting membrane voltage towards a more depolarized potential (t-test with Welch's correction, $*p=0.0356$). (F)

Representative current-clamp recordings of PYRs injected with -100 (*black*), 140 (*darker colored shade*), and 230 pA (*lighter colored shade*) from WT (*blue*) and KI (*red*) mice, *scale bar* = 200 ms, 200 pA/20 mV. (G) Increased expression of *mC4* significantly decreased the spike frequency of PYRs in KI male mice, relative to controls (RM Two-way ANOVA, Condition x Current: $F_{(24,696)}=1.975$, $**p=0.0038$. Condition: $F_{(1,29)}=2.646$, $p=0.1147$. Current: $F_{(1,350,39.15)}=619.8$, $****p<0.0001$). (H) No differences in the excitability of PYRs in females between groups (RM Two-way ANOVA, Condition x Current: $F_{(24,816)}=0.2813$, $p=0.9998$. Condition: $F_{(1,34)}=0.1154$, $p=0.7361$. Current: $F_{(1,467,49.89)}=810.0$, $****p<0.0001$). (I) No changes in PYR Rheobase between groups (Condition x Sex: $F_{(1,63)}=5.148$, $*p=0.0267$. Condition: $F_{(1,63)}=0.2332$, $p=0.6308$. Sex: $F_{(1,63)}=0.8369$, $p=0.3638$. Post-test: WT vs. KI males, $p=0.1262$, females, $p=0.3502$). (J) No change overall in PYR resting membrane potential in KI mice, compared to controls (Mann-Whitney test, $p=0.6464$). WT: blue squares. KI: red circles. *N* represents cells. PV cell WT $N=19$ males, $N=17$ females; PV cell KI: $N=18$ males, $N=19$ females. PYR WT $N=15$ males, $N=18$ females; PYR KI: $N=16$ males, $N=18$ females. For all statistics, $*p<0.05$, $**p<0.01$, Two-way ANOVA, unless otherwise stated. Mean \pm SEM shown.

Figure 6. No changes in anxiety-like behavior with pan-neuronal overexpression of *mC4*. (A) Mouse breeding scheme (*top*) and same anxiety behavioral battery (*bottom*) as in Fig. 2. (B, C) Pan-neuronal *mC4* overexpression does not alter locomotion (B: t-test, $p=0.9092$. C: Condition x Sex: $F_{(1,36)}=1.039$, $p=0.3148$. Condition: $F_{(1,36)}=0.00415$, $p=0.9490$. Sex: $F_{(1,36)}=0.4105$, $p=0.5258$). (D, E) No change in time spent in the open arms of the EZM with pan-neuronal *mC4* overexpression (D: Mann-Whitney test, $p=0.6487$. E: Condition x Sex: $F_{(1,36)}=1.225$, $p=0.2757$. Condition: $F_{(1,36)}=0.1021$, $p=0.7512$. Sex: $F_{(1,36)}=0.00008$, $p=0.9928$). (F, G) Time spent in the light zone of the LDB was not altered by pan-neuronal *mC4* overexpression (F: Mann-Whitney test, $p=0.9360$. G: Condition x Sex: $F_{(1,36)}=4.620$, $*p=0.0384$. Condition: $F_{(1,36)}=0.07082$, $p=0.7917$. Sex: $F_{(1,36)}=2.552$, $p=0.1189$. Post-test: WT vs. KI males, $p=0.1977$, females, $p=0.3281$). (H, I) Overexpression of *mC4* in neurons did not alter the latency to feed in the NSF (H: t-test, $p=0.1761$. I: Condition x Sex: $F_{(1,29)}=4.093$, $p=0.0524$. Condition: $F_{(1,29)}=1.328$, $p=0.2586$. Sex: $F_{(1,29)}=0.4055$, $p=0.5293$). (J, K) Increased levels of *mC4* in neurons did not lead to drastic changes in overall anxiety-like behavior, as measured with Z-anxiety scores. (J: t-test with Welch's correction, $p=0.5358$. K: Condition x Sex: $F_{(1,36)}=0.3797$, $p=0.5416$. Condition: $F_{(1,36)}=0.03077$, $p=0.8617$. Sex: $F_{(1,36)}=1.008$, $p=0.3221$). WT: blue squares. KI: red circles. WT $N=9$ males, $N=10$ females; KI: $N=10$ males, $N=11$ females. For all statistics, $*p<0.05$, Two-way ANOVA, unless otherwise stated. Mean \pm SEM shown.

SUPPLEMENTAL FIGURE LEGENDS

Supplemental Figure 1. No change in latency to feed in the Cage NSF in PV-*mC4*-KI mice relative to controls. (A) Schematic of the Cage NSF. Latency to feed (s) was measured when WT or KI mice were placed in a more familiar environment, a standard cage. (B) No change in the latency to feed between WT and KI mice (Mann-Whitney test, $p=0.5881$). (C) No change in the latency to feed between groups, separated by sex (Two-way ANOVA, Condition x Sex: $F_{(1,34)}=0.2861$, $p=0.5962$. Condition: $F_{(1,34)}=0.4357$, $p=0.5137$. Sex: $F_{(1,34)}=0.00207$, $p=0.9640$). WT: blue squares, $N=12$ males, $N=7$ females. KI: red circles, $N=7$ males, $N=12$ females. For all statistics, $*p<0.05$. Mean \pm SEM shown.

Supplemental Figure 2. No changes in less-frequent sub-classes of social behavior with increased levels of *mC4* in PV cells. (A) Both WT and KI mice initiate most interactions during the juvenile interaction task (Condition x Interaction class: $F_{(1,92)}=4.484$, $*p=0.0369$. Condition: $F_{(1,92)}=2.881$, $p=0.0930$. Interaction class: $F_{(1,92)}=236.1$, $****p<0.0001$. Post-test: Active vs. Passive WT, $****p<0.0001$, KI, $****p<0.0001$) with KI mice engaging less in active interactions compared to controls (WT vs. KI, $*p=0.0488$). (B, C) No change in reciprocal snout-snout interactions (13% of total social interactions) in KI mice, compared to controls (B: Mann-Whitney test, $p=0.4993$. C: Condition x Sex: $F_{(1,44)}=0.1482$, $p=0.7021$. Condition: $F_{(1,44)}=0.3293$, $p=0.5690$. Sex: $F_{(1,44)}=0.2833$, $p=0.5972$). (D, E) No change in the active snout-body interaction (23% of total social interactions) in KI mice (D: t-test with Welch's correction, $p=0.6501$. E: Condition x Sex: $F_{(1,44)}=0.0133$, $p=0.9088$. Condition: $F_{(1,44)}=0.2210$, $p=0.6406$. Sex: $F_{(1,44)}=3.028$, $p=0.0888$). (F, G) No differences in the passive snout-rear interaction (9% of total social interactions) between groups (F: Mann-Whitney test, $p=0.9186$. G: Condition x Sex: $F_{(1,44)}=2.601$, $p=0.1139$. Condition: $F_{(1,44)}=0.2085$, $p=0.6502$. Sex: $F_{(1,44)}=1.930$, $p=0.1718$). (H, I) No differences in the passive snout-body interaction (5% of total social interactions) between groups (H: Mann-Whitney test, $p=0.5392$. I: Condition x Sex: $F_{(1,44)}=3.745$, $p=0.0594$. Condition: $F_{(1,44)}=0.4934$, $p=0.4861$. Sex: $F_{(1,44)}=0.2645$, $p=0.6096$). WT: blue squares, $N=13$ males, $N=12$ females. KI: red circles, $N=12$ males, $N=11$ females. For all statistics, $*p<0.05$, $**p<0.01$, $***p<0.001$, $****p<0.0001$. Two-way ANOVA, unless otherwise stated. Mean \pm SEM shown.

Supplemental Figure 3. Overexpression of *mC4* in PV cells did not alter social interactions in the three-chamber assay. (A) (Top) Schematic representation of three-chamber assay. (Bottom) Representative trajectories (tracked with DLC) of WT (blue) and KI (red) mice in the three-chamber assay. E, empty cup; M, mouse cup. (B) WT and KI mice spent more time in the chamber containing the mouse cup compared to the empty-cup chamber, suggesting that they prefer social interactions (Condition x Chamber: $F_{(1,84)}=0.1733$, $p=0.6782$. Condition: $F_{(1,84)}=1.911$, $p=0.1706$. Chamber: $F_{(1,84)}=34.19$, $****p<0.0001$. Post-test: Mouse vs. Empty, WT $**p=0.0013$, KI, $***p=0.0002$). (C, D) No change in the social discrimination index (SI) between WT and KI mice (C: t-test with Welch's correction, $p=0.9463$. D: Condition x Sex: $F_{(1,40)}=1.455$, $p=0.2349$. Condition: $F_{(1,40)}=0.03420$, $p=0.8542$. Sex: $F_{(1,40)}=1.643$, $p=0.2074$). WT: blue squares, $N=10$ males, $N=12$ females. KI: red circles, $N=12$ males, $N=10$ females. For all statistics, $*p<0.05$, $**p<0.01$, $***p<0.001$, Two-way ANOVA, unless otherwise stated. Mean \pm SEM shown.

Supplemental Figure 4. PV-specific *mC4*-OE alters the kinetics of mEPSCs in PYRs of female mice. (A) Representative 1 s traces of mEPSCs recorded in PYRs in P40-60 young adult WT (blue) and KI (red) mice, scale bar = 250 ms, 10 pA. (B) No change in mEPSC amplitude in KI mice, relative to controls (Condition x Sex: $F_{(1,61)}=1.945$, $p=0.1681$. Condition: $F_{(1,61)}=0.4158$, $p=0.5214$. Sex: $F_{(1,61)}=1.397$, $p=0.2418$). (C) No change in mEPSC frequency in KI mice (Condition x Sex: $F_{(1,61)}=2.291$, $p=0.1353$. Condition: $F_{(1,61)}=0.1903$, $p=0.6642$. Sex: $F_{(1,61)}=3.341$, $p=0.0725$). (D) *mC4* overexpression did not alter mEPSC rise Rise₁₀₋₉₀ (Condition x Sex: $F_{(1,61)}=3.916$, $p=0.0524$. Condition: $F_{(1,61)}=0.3104$, $p=0.5795$. Sex: $F_{(1,61)}=0.3120$, $p=0.5785$). (E) Increased expression of *mC4* led to a significant decrease in mEPSC Decay_{tau} in

KI females (Condition x Sex: $F_{(1,61)}=12.74$, $***p=0.0007$. Condition: $F_{(1,61)}=0.3684$, $p=0.5461$. Sex: $F_{(1,61)}=2.183$, $p=0.1447$. Post-test: WT vs. KI males, $p=0.0822$, females, $**p=0.0083$). (F) Representative 1 s traces of mIPSCs recorded in PYRs in young adult WT (*blue*) and KI (*red*) mice, *scale bar* = 250 ms, 10 pA. (G) No change in mIPSC amplitude in KI mice, relative to controls (Condition x Sex: $F_{(1,60)}=0.5354$, $p=0.4672$. Condition: $F_{(1,60)}=0.00013$, $p=0.9909$. Sex: $F_{(1,60)}=0.00367$, $p=0.9519$). (H) mIPSC frequency was not different between groups (Condition x Sex: $F_{(1,60)}=0.03569$, $p=0.8508$. Condition: $F_{(1,60)}=0.2141$, $p=0.6452$. Sex: $F_{(1,60)}=2.690$, $p=0.1062$). (I) Increased expression of *mC4* did not impact mIPSC Rise₁₀₋₉₀ (Condition x Sex: $F_{(1,60)}=0.3837$, $p=0.5380$. Condition: $F_{(1,60)}=0.7667$, $p=0.3847$. Sex: $F_{(1,60)}=5.047$, $*p=0.0284$. Post-test: WT vs. KI males, $p=0.9795$, females, $p=0.5025$). (J) Decay_{tau} was not changed in KI mice (Condition x Sex: $F_{(1,60)}=1.153$, $p=0.2871$. Condition: $F_{(1,60)}=0.7059$, $p=0.4042$. Sex: $F_{(1,60)}=0.2624$, $p=0.6104$). WT: blue squares. KI: red circles. *N* represents cells. mEPSC WT *N*=17 males, *N*=16 females; mEPSC KI: *N*=15 males, *N*=17 females. mIPSC WT *N*=16 males, *N*=16 females; mIPSC KI: *N*=16 males, *N*=16 females. For all statistics, $*p<0.05$, $**p<0.01$, Two-way ANOVA, unless otherwise stated. Mean \pm SEM shown.

Supplemental Table 1: Active and passive electrophysiological properties of PV cells in PV-mC4-KI and WT mice. (*Top*) Table displays the main effects results of each Two-way ANOVAs for PV cells. Post-test analyses were only performed below if at least one of the main effects of the Two-way ANOVA were significantly different between groups. (*Bottom*) Table displays WT and KI means \pm SEM, and the associated *p*-value for the comparison between conditions if applicable. Data are separated by sex, male data on the left and female data on the right. For all statistics $*p<0.05$, Two-way ANOVA. In all KI mice, resting V_m was significantly more depolarized in response to PV-mC4-OE, relative to WT (WT: -67.25 ± 0.8696 mV vs. KI: -64.05 ± 1.209 mV, t-test with Welch's correction, $*p=0.0356$), data not shown in table.

Supplemental Table 2: Active and passive electrophysiological properties of PYRs in PV-mC4-KI and WT mice. (*Top*) Table displays the main effects results of each Two-way ANOVAs for PYRs. Post-test analyses were only performed below if at least one of the main effects of the Two-way ANOVA were significantly different between groups. (*Bottom*) Table displays WT and KI means \pm SEM, and the associated *p*-value for the comparison between conditions if applicable. Data are separated by sex, male data on the left and female data on the right. For all statistics $*p<0.05$, Two-way ANOVA.

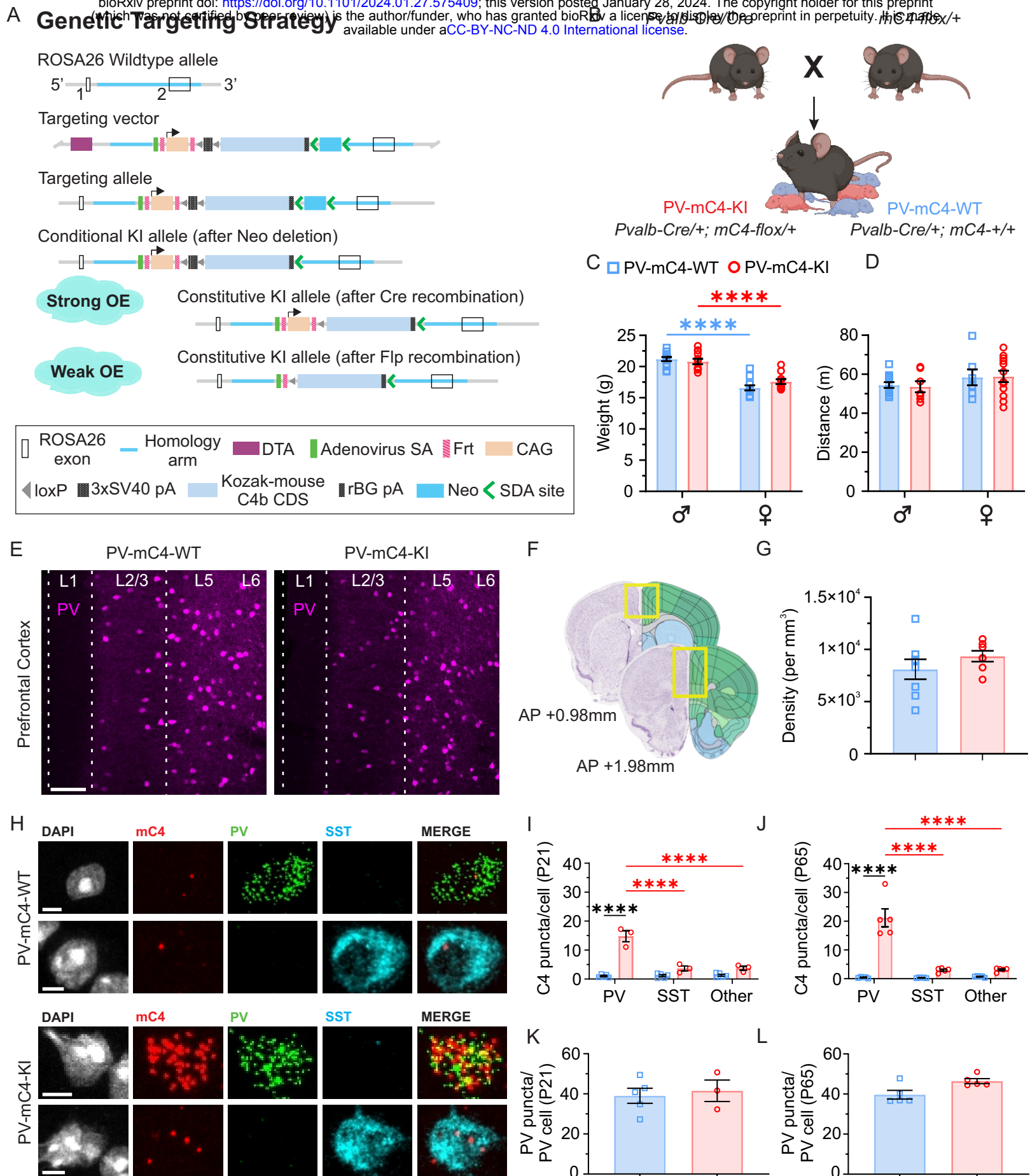


Figure 1. A novel transgenic mouse line permits PV cell specific overexpression of complement component 4.

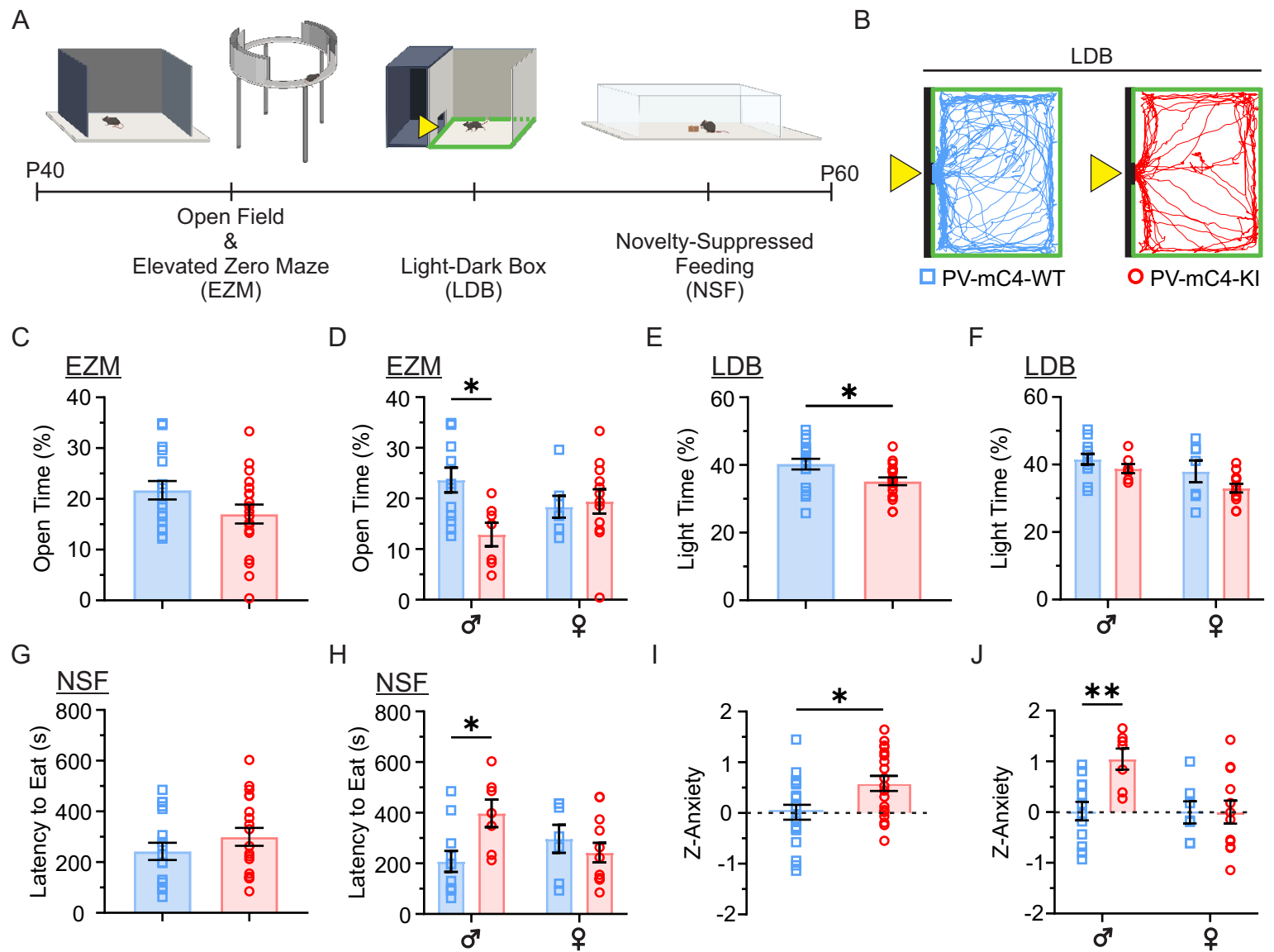


Figure 2. PV-specific mC4-OE causes an increase in anxiety-like behavior in male mice.

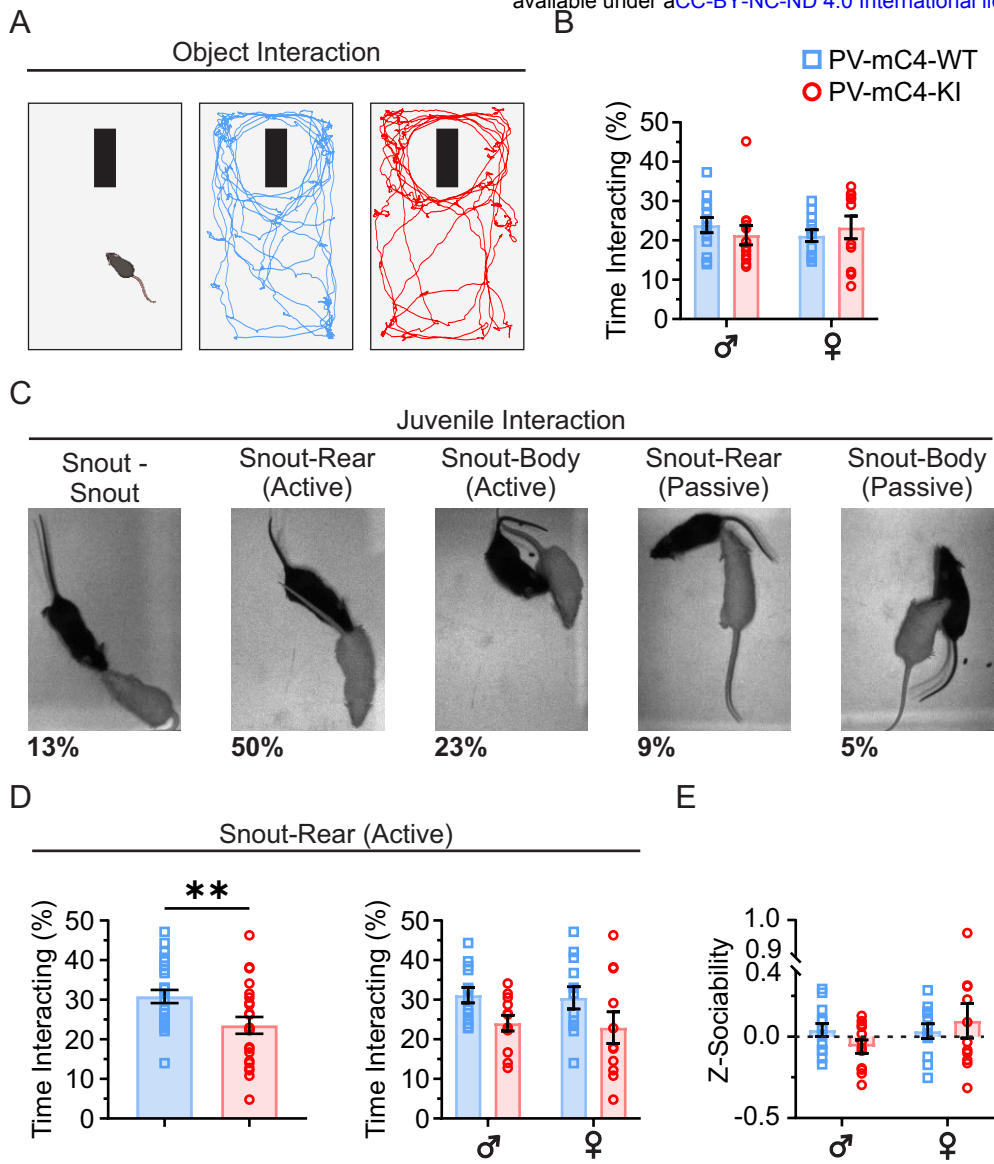


Figure 3. Increased levels of *mC4* in PV cells disrupts active but not passive social behaviors.

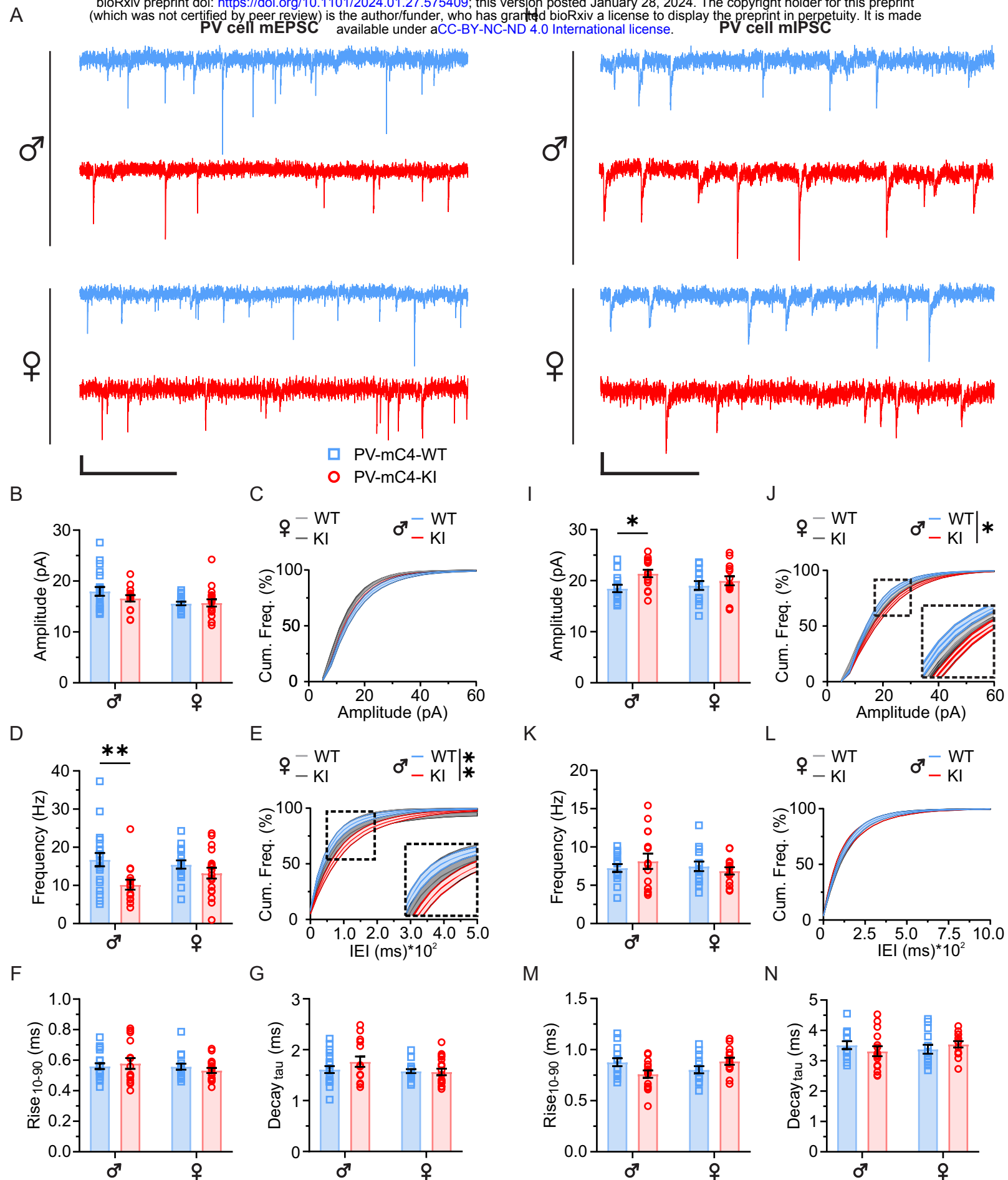


Figure 4. Sex-related difference in excitatory-inhibitory dynamics in mPFC PV cells with increased levels of *mC4* in PV cells.

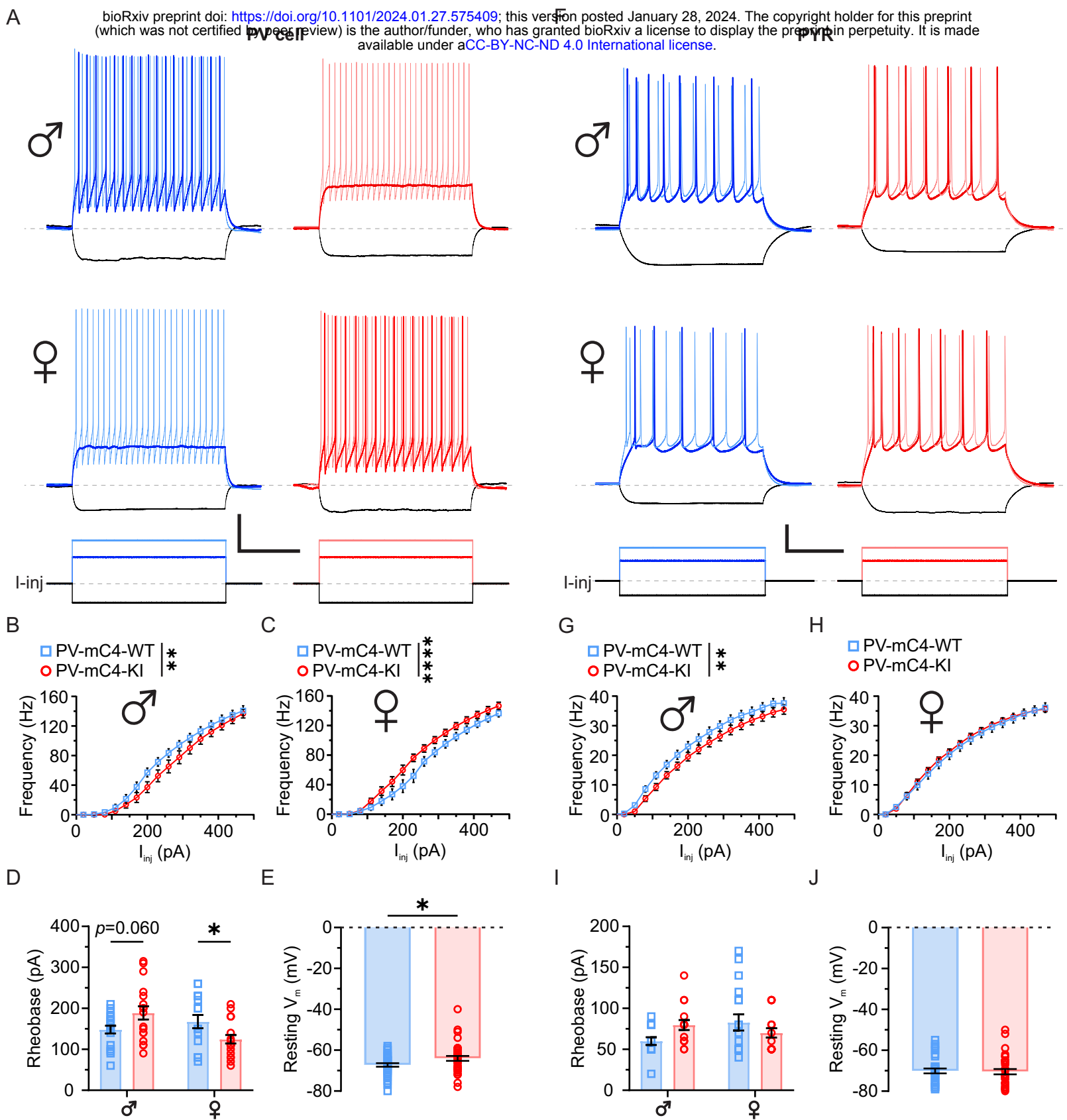


Figure 5. PV-specific mC4-OE leads to opposing changes in the excitability of PV cells in male and female mice.

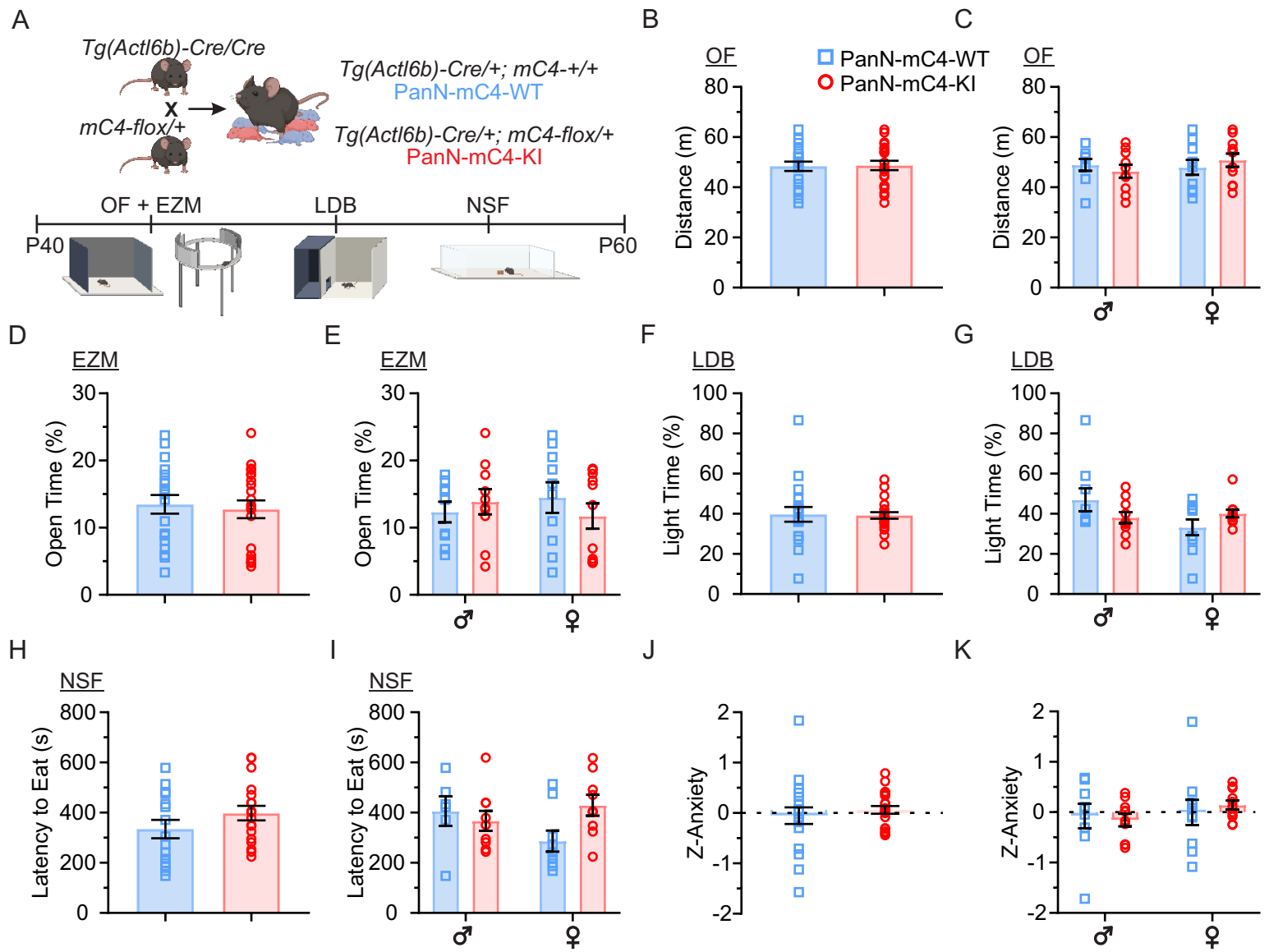
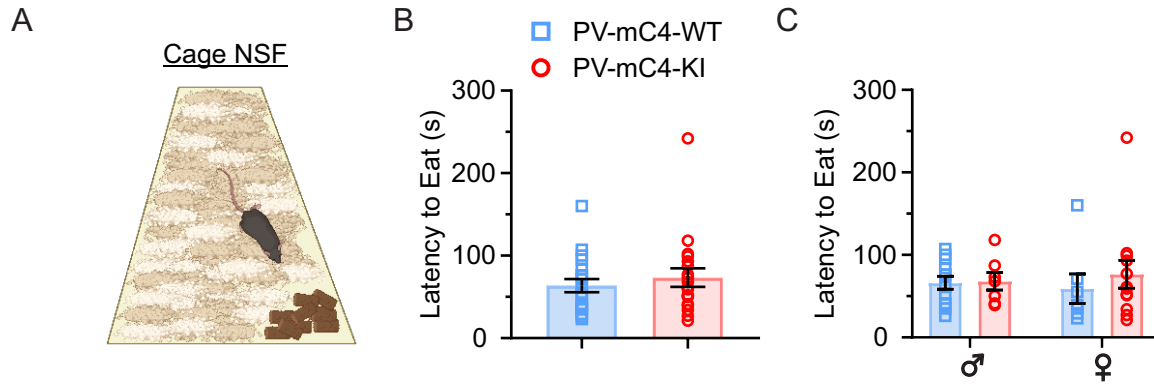
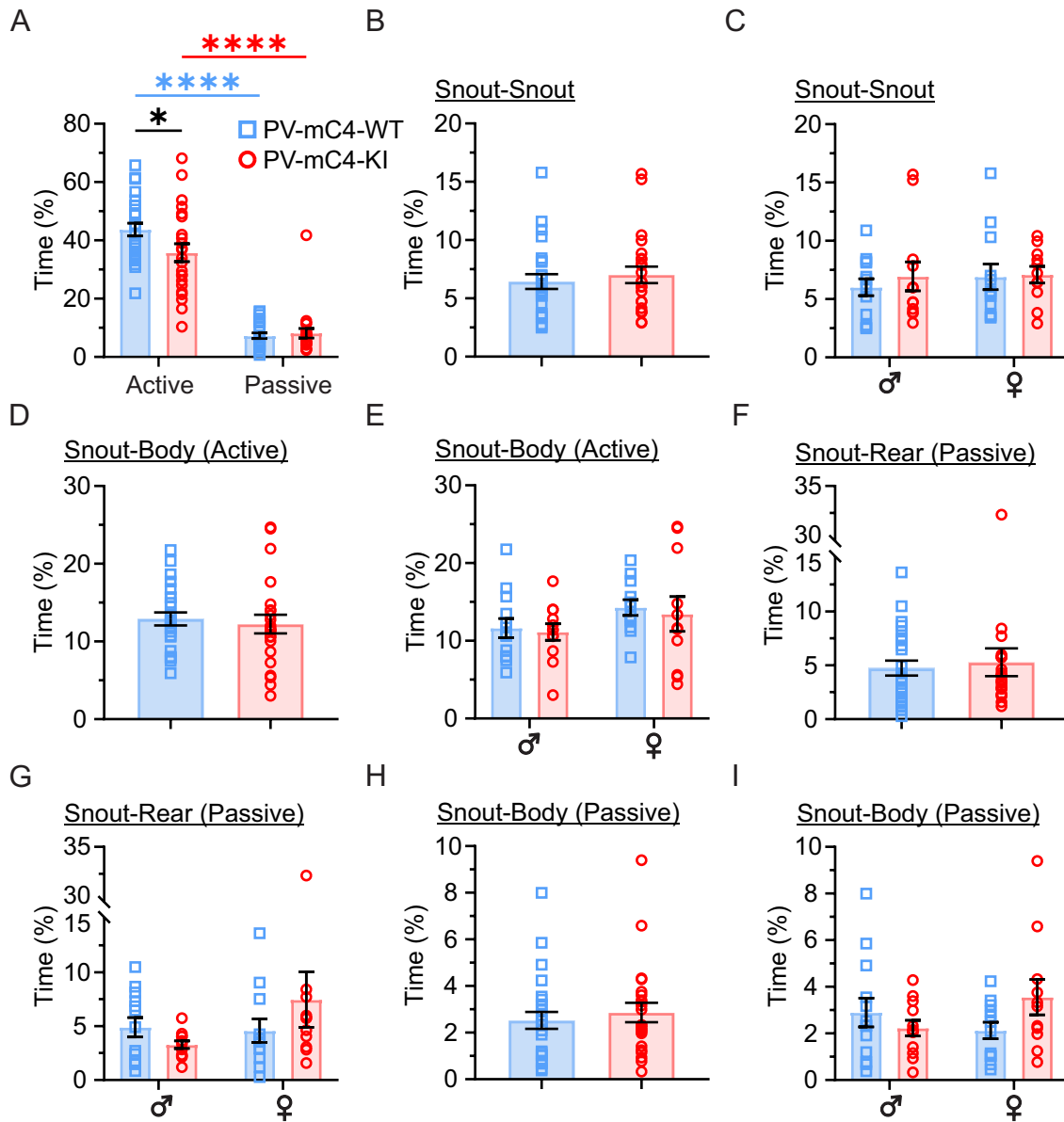


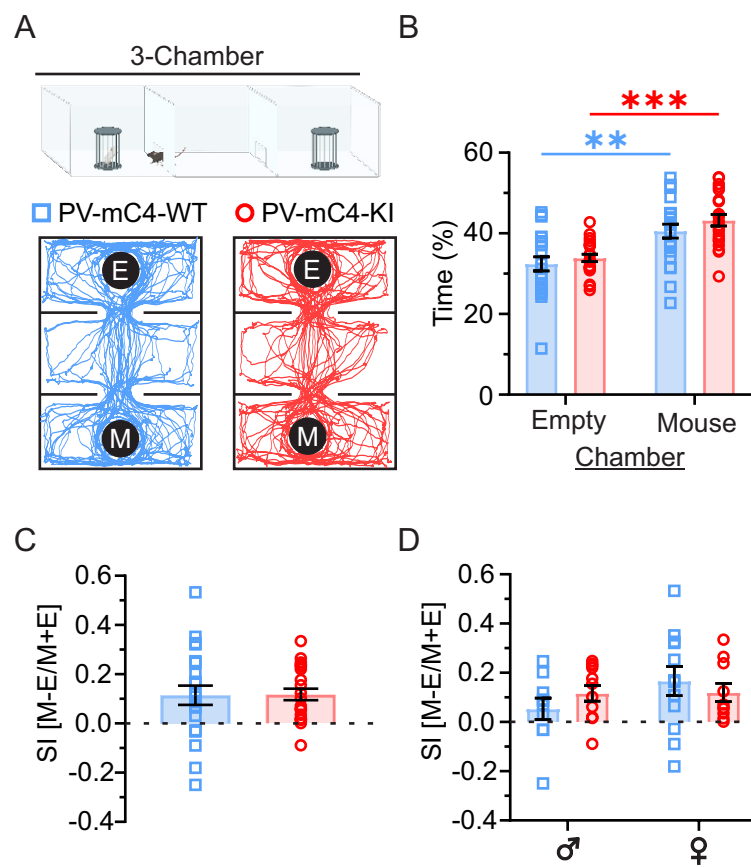
Figure 6. No changes in anxiety-like behavior with pan-neuronal overexpression of *mC4*.



Supplemental Figure 1. No change in latency to feed in the Cage NSF in PV-mC4-KI mice relative to controls.



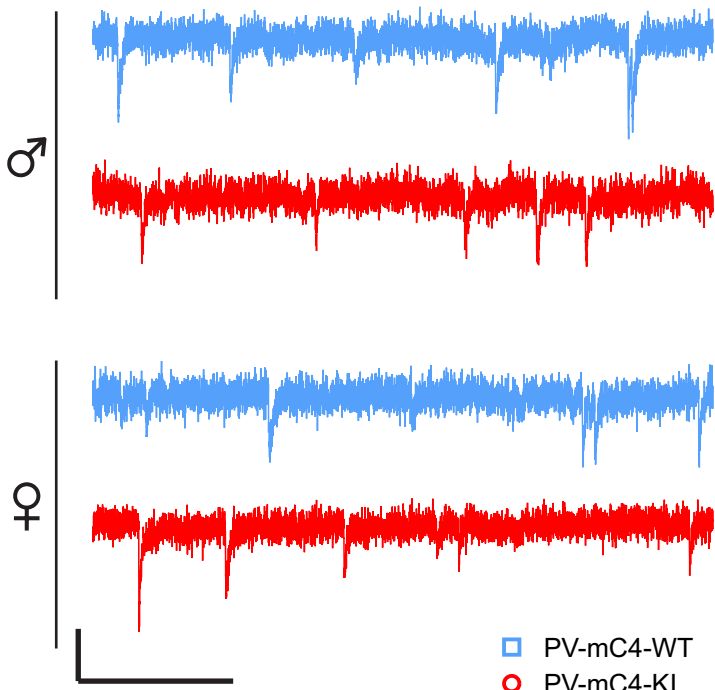
Supplemental Figure 2. No changes in less-frequent sub-classes of social behavior with increased levels of *mC4* in PV cells



Supplemental Figure 3. Overexpression of *mC4* in PV cells did not alter social interactions in the three-chamber assay.

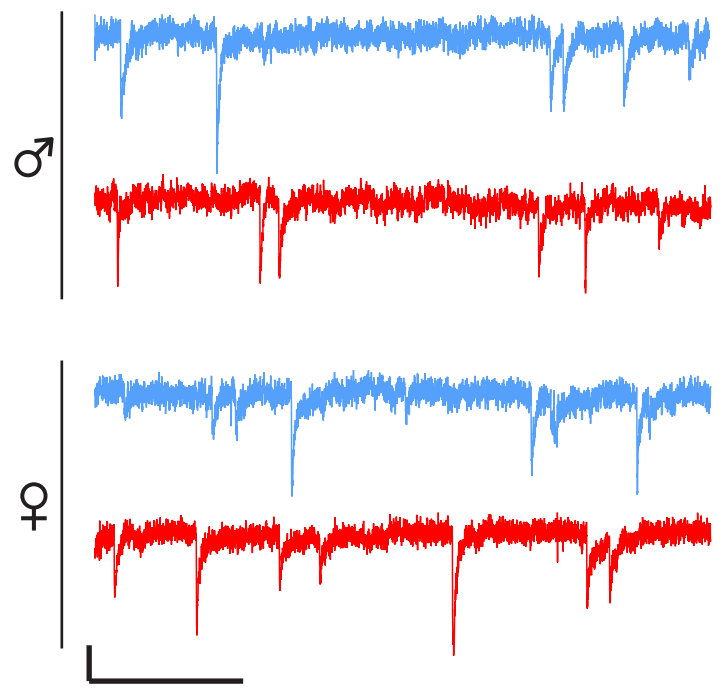
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PYR mEPSC

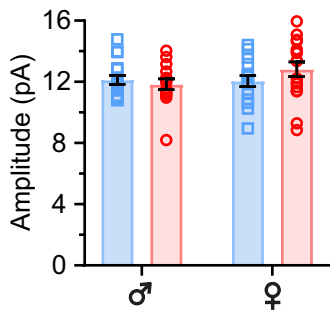


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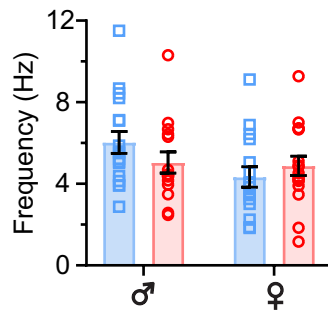
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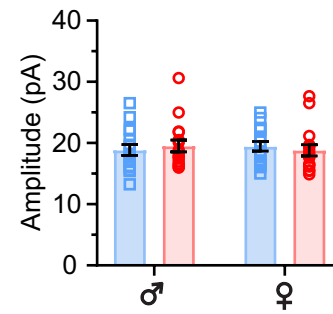
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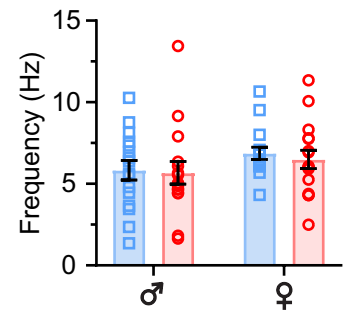
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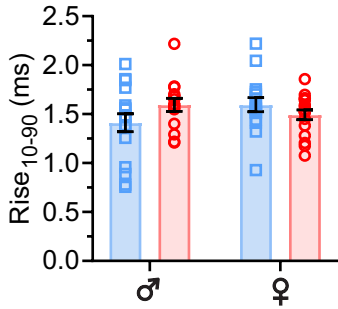
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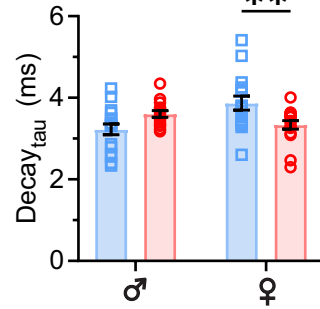
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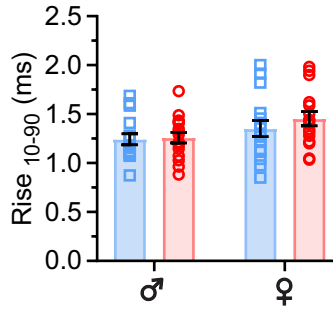
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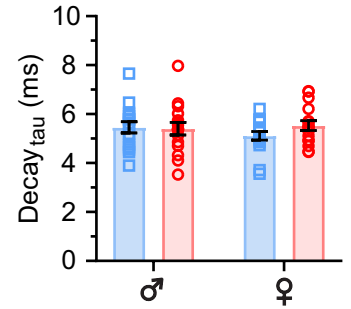
E



I



J



Supplemental Figure 4. PV-specific mC4-OE alters the kinetics of mEPSCs in PYRs of female mice.

PV cell	Two-way ANOVA. $F_{(1,69)}=$, $p=$		
	Sex x Condition	Sex	Condition
Resting V_m (mV)	0.0975, 0.7558	0.7875, 0.3779	4.309, *0.0416
Input Resistance, R_{in} ($M\Omega$)	2.925, 0.0917	2.057, 0.1560	0.2246, 0.6370
Membrane Capacitance, C_m (pF)	0.5158, 0.4750	0.0619, 0.8042	2.926, 0.0917
Decay constant, tau (ms)	2.100, 0.1518	4.507, *0.0373	1.615, 0.2080
ISI_1 / ISI_9	0.0082, 0.9281	2.703, 0.1084	0.0327, 0.8570
ISI_4 / ISI_9	0.0826, 0.7748	0.9215, 0.3405	0.2781, 0.5997
Sag Ratio, $F_{(1,60)}$	1.064, 0.3064	1.686, 0.1991	1.161, 0.2857

PV cell	♂			♀		
	Mean \pm SEM		<i>post-test</i> , <i>p</i> -value	Mean \pm SEM		<i>post-test</i> , <i>p</i> -value
	WT	KI		WT	KI	
Resting V_m (mV)	-68.11 \pm 5.37	-64.5 \pm 7.23	0.1781	-66.29 \pm 5.02	-63.63 \pm 7.65	0.3916
Input Resistance, R_{in} ($M\Omega$)	161.27 \pm 11.18	147.15 \pm 9.36	————	158.11 \pm 12.41	183.07 \pm 12.38	————
Membrane Capacitance, C_m (pF)	63.74 \pm 2.58	67.52 \pm 1.85	————	61.96 \pm 3.64	71.20 \pm 5.75	————
Decay constant, tau (ms)	9.78 \pm 0.52	9.64 \pm 0.58	0.9899	10.31 \pm 0.93	12.44 \pm 1.00	0.1171
ISI_1 / ISI_9	1.44 \pm 0.42	1.52 \pm 0.30	————	1.03 \pm 0.01	1.05 \pm 0.04	————
ISI_4 / ISI_9	0.98 \pm 0.01	0.98 \pm 0.02	————	1.00 \pm 0.01	0.99 \pm 0.01	————
Sag Ratio	0.040 \pm 0.003	0.040 \pm 0.004	————	0.049 \pm 0.004	0.041 \pm 0.004	————

Supplemental Table 1. Active and passive electrophysiological properties of PV cells in PV-mC4-KI and WT mice

PYR	Two-way ANOVA. $F_{(1,63)}=$, $p=$		
	Sex x Condition	Sex	Condition
Resting V_m (mV)	0.0104, 0.9191	0.1306, 0.7190	.0310, 0.8608
Input Resistance, R_{in} (M Ω)	3.661, 0.0602	0.2326, 0.6313	0.1358, 0.7138
Membrane Capacitance, C_m (pF)	0.1365, 0.7130	4.155, *0.0457	1.705, 0.1964
Decay constant, tau (ms)	2.826, 0.0977	0.6693, 0.4164	0.0719, 0.7894
ISI ₁ / ISI ₉	0.4917, 0.4857	1.259, 0.2660	0.3778, 0.5410
ISI ₄ / ISI ₉	1.396, 0.2419	4.856, *0.0312	0.0810, 0.7769
Sag Ratio, $F_{(1,59)}$	2.894, 0.0942	0.1244, 0.7256	0.3072, 0.5815

PYR	♂			♀		
	Mean \pm SEM		<i>post-test</i> , <i>p</i> -value	Mean \pm SEM		<i>post-test</i> , <i>p</i> -value
	WT	KI		WT	KI	
Resting V_m (mV)	-69.87 \pm 1.75	-70.0 \pm 2.0	————	-70.33 \pm 1.66	-70.83 \pm 1.75	————
Input Resistance, R_{in} (M Ω)	244.88 \pm 15.80	204.61 \pm 13.58	————	219.62 \pm 19.20	246.89 \pm 19.56	————
Membrane Capacitance, C_m (pF)	145.91 \pm 7.88	159.09 \pm 11.02	0.4488	132.77 \pm 5.35	140.14 \pm 6.76	0.7437
Decay constant, tau (ms)	34.32 \pm 1.92	31.22 \pm 1.99	————	28.84 \pm 2.50	33.16 \pm 2.15	————
ISI ₁ / ISI ₉	0.32 \pm 0.04	0.33 \pm 0.02	————	0.37 \pm 0.03	0.34 \pm 0.03	————
ISI ₄ / ISI ₉	0.78 \pm 0.02	0.81 \pm 0.02	0.5393	0.84 \pm 0.01	0.83 \pm 0.02	0.7618
Sag Ratio	0.065 \pm 0.005	0.075 \pm 0.011	————	0.083 \pm 0.009	0.063 \pm 0.008	————

Supplemental Table 2. Active and passive electrophysiological properties of PYRs in PV-mC4-KI and WT mice