

Resilient emotionality and molecular compensation in mice lacking the oligodendrocyte-specific gene *Cnp1*

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Altered oligodendrocyte structure and function is implicated in major psychiatric illnesses, including low cell number and reduced oligodendrocyte-specific gene expression in major depressive disorder (MDD). These features are also observed in the unpredictable chronic mild stress (UCMS) rodent model of the illness, suggesting that they are consequential to environmental precipitants; however, whether oligodendrocyte changes contribute causally to low emotionality is unknown. Focusing on 2'-3'-cyclic nucleotide 3'-phosphodiesterase (*Cnp1*), a crucial component of axoglial communication dysregulated in the amygdala of MDD subjects and UCMS-exposed mice, we show that altered oligodendrocyte integrity can have an unexpected functional role in affect regulation. Mice lacking *Cnp1* (knockout, KO) displayed decreased anxiety- and depressive-like symptoms (i.e., low emotionality) compared with wild-type animals, a phenotypic difference that increased with age (3–9 months). This phenotype was accompanied by increased motor activity, but was evident before neurodegenerative-associated motor coordination deficits (\leq 9–12 months). Notably, *Cnp1*^{KO} mice were less vulnerable to developing a depressive-like syndrome after either UCMS or chronic corticosterone exposure. *Cnp1*^{KO} mice also displayed reduced fear expression during extinction, despite normal amygdala c-Fos induction after acute stress, together implicating dysfunction of an amygdala-related neural network, and consistent with proposed mechanisms for stress resiliency. However, the *Cnp1*^{KO} behavioral phenotype was also accompanied by massive upregulation of oligodendrocyte- and immune-related genes in the basolateral amygdala, suggesting an attempt at functional compensation. Together, we demonstrate that the lack of oligodendrocyte-specific *Cnp1* leads to resilient emotionality. However, combined with substantial molecular changes and late-onset neurodegeneration, these results suggest the low *Cnp1* seen in MDD may cause unsustainable and maladaptive molecular compensations contributing to the disease pathophysiology.

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Introduction

Disruptions in neuronal signaling long have been the focus of research on major depressive disorder (MDD). More recently, however, glial disruptions have been postulated to contribute to the pathophysiology of MDD,^{1–3} and the roles of specialized glial subtypes are now being investigated. Although alterations in microglia^{4–6} and astrocyte-related components^{7–10} are observed in MDD, oligodendrocyte alterations in MDD are becoming a primary focus of research.² Oligodendrocytes, the main myelin-forming cells of the central nervous system, provide both structural and trophic support for neurons, and facilitate axonal conduction. In the amygdala and prefrontal cortex of MDD subjects, previously reported decreases in glial cell number were attributed to reduced oligodendrocyte number,^{11,12} consistent with patterns of downregulation of oligodendrocyte-related transcripts in amygdala¹³ and nearby temporal cortex.¹⁴ Changes in NG2 cells, a cell type sharing a common lineage with oligodendrocytes,^{15,16} have also been associated with depression-related characteristics in rodents. For instance, both chronic corticosterone exposure and

chronic stress in rodents decreased cortical and limbic oligodendrocyte¹⁷ and NG2 cell proliferation.^{18,19} Rats given electroconvulsive seizure therapy, an antidepressant treatment for otherwise non-responsive MDD subjects, showed increased proliferation of NG2 cells in the amygdala and hippocampus.^{18,20,21} Thus, evidence suggests that oligodendrocytes, a crucial element for maintaining optimal neuronal function, may be vulnerable to stress-related insults and may contribute to the pathophysiology of MDD.

We previously identified a set of dysregulated gene transcripts in the amygdala of male subjects with familial MDD, including robust downregulations of multiple oligodendrocyte-related genes.¹³ This pattern was also observed in mice following unpredictable chronic mild stress (UCMS)²² and appeared restricted to the amygdala (not anterior cingulate cortex or dentate gyrus). 2'-3'-Cyclic nucleotide 3'-phosphodiesterase (*Cnp1*) was one of several oligodendrocyte-specific genes significantly downregulated across species. In addition, *Cnp1* was previously implicated in MDD^{14,23} and schizophrenia,^{24–26} supporting the hypothesis

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that altered *Cnp1* function may participate in the pathophysiology of psychiatric disorders. Although *Cnp1* is expressed at low levels in the periphery, it is expressed at higher levels in NG2 cells and it comprises ~4% of myelin-associated proteins in mature oligodendrocytes.²⁷ *Cnp1* is localized to non-compact myelin in the inner mesaxon and paranodal loops,²⁸ the principle sites of contact with the axon. *Cnp1* binds microtubules and regulates mRNA expression and transport at the paranode,²⁹ although its exact role in axoglial communication is not known. The lack of *Cnp1* protein in mice results in disorganization of nodal sodium channels and paranodal adhesion proteins (e.g., *Caspr* and *Nav*).^{30,31} Adult *Cnp1*^{KO} mice do not initially show an overt behavioral phenotype, but develop progressive axonal degeneration and motor deficits after 6 months of age, leading to premature death.^{30,31}

Healthy oligodendrocytes are necessary to maintain optimal axon function;³² however, it remains to be determined whether oligodendrocyte-specific alterations are causal to MDD, as opposed to compensatory or neutral side effects in psychiatric disorders. Here, we investigated the effect of a lack of *Cnp1* on emotionality (i.e. anxiety and depressive-like behaviors) in mice, under baseline conditions (trait) and after chronic stress or corticosterone exposure, two validated paradigms for inducing high-emotionality states. To summarize the number of behavioral tests that were performed, and to obtain comprehensive and integrated measures in each group, emotionality- and locomotion-related data were normalized using a Z-score methodology previously described.³³ As the amygdala is a central region in affect regulation and as our initial findings showed low *Cnp1* in the amygdala of MDD patients,³⁴ we also assessed amygdala function in *Cnp1*^{KO} mice using fear conditioning (FC), c-Fos induction and gene expression profiling. The experiments show that lack of *Cnp1* results in low emotionality under baseline and induced states, along with reduced corticolimbic fear expression, a suggested mechanism for stress resilience.

Materials and methods

Animals. *Cnp1*^{KO} mice were obtained from KA Nave and C. Lappe-Siefke (Max Planck Institute, Göttingen, Germany). Adult male and female *Cnp1*^{KO} and wild-type (WT) littermate mice (C57BL/6 background)³¹ were obtained from heterozygous crossings and microchipped for identification. Five cohorts were used: Baseline Cohort ($N=11-17$ per group; 6 and 9 months baseline), UCMS Cohort ($N=8-12$ per group, 6 months of age), CORT Cohort ($N=10-14$ per group; 3-month baseline and CORT exposure), FC Cohort ($N=16-19$ per group, 3 and 6 months of age) and cFos Cohort ($N=6$ per group, 6 months of age). Mice were maintained under standard conditions (group housed, 12/12-hour light/dark cycle, 22 ± 1 °C, food and water *ad libitum*). All testing was conducted in compliance with the National Institutes of Health laboratory animal care guidelines and with protocols approved by the Institutional Animal Care and Use Committee.

Behavioral measures. Major depression is defined as a syndrome (i.e., collection of symptoms) including low mood or anhedonia, accompanied by cognitive (e.g., attention,

concentration) and physiological symptoms (e.g., weight, locomotor and sleep pattern changes), and frequently comorbid with significant anxiety symptoms. Thus, the emotionality component is best characterized in mice by a comprehensive panel of behavioral tests for anxiety-like and depressive-like emotionality, and for antidepressant-like behavior. Hence, Baseline, UCMS and CORT Cohorts were tested in the elevated plus maze (EPM), open field (OF), novelty suppressed feeding (NSF), forced swim test (FST) and rotarod as described,³⁵ and in the following order: EPM, OF, NSF, FST, rotarod, separated by a minimum of 1–2 days. Detailed methods are provided in the Supplementary Material.

Behavioral Z-scoring. To address behavioral variability and obtain comprehensive and integrated measures in each group, emotionality- and locomotion-related data were normalized using a Z-score methodology previously described.³³ Briefly, for each behavioral measure, Z-scores for individual animals were calculated using the formula below, which indicates how many s.d.'s (σ) an observation (X) is above or below the mean of a control group (μ).

$$z = \frac{X - \mu}{\sigma}$$

Z-scores for behavioral measures were first averaged within test, and then across test to ensure equal weighting of the four tests comprising the final Z-score (Figure 1b). Separate Z-scores were calculated for the Baseline, UCMS and CORT Cohorts using the means and s.d.'s of the respective control groups (i.e., 'WT' for Baseline or 'WT control' for UCMS and CORT). Locomotion Z-scores were similarly obtained from EPM (total crosses) and OF (total distance traveled) data.

Estrous phase. Female mice in the UCMS and 3-month Baseline Cohorts were assessed for estrous phase on the day of behavioral testing after the EPM, OF and NSF tests to control for potential effects of hormonal fluctuations on behavior. Estrus, metestrus, diestrus and proestrus were determined via vaginal cytology³⁶ as previously described.³³

Unpredictable chronic mild stress. Single-housed mice were subjected to 4 weeks of a randomized schedule of 1–2 mild stressors per day, 7 days per week. Detailed methods are in the supplements and full table of stressors in Supplementary Table 1.

Chronic corticosterone treatment. As described previously,³³ mice were given corticosterone ($35 \mu\text{g ml}^{-1}$; Sigma-C2505, St Louis, MO, USA) dissolved in 20% cyclodextrin (Sigma H107) as their only water supply for 4 weeks. Owing to the reduced cohort size, behavior of all mice was tested both before (3-month baseline) and following CORT exposure.

Fear conditioning. The protocol was performed over 3 days using a computer-controlled system (Coulbourn Instruments, Allentown, PA, USA) (Figure 3a). On day 1, mice were trained to associate a conditioned stimulus (CS; 80 db, 2 kHz, 15 s tone) to an unconditioned stimulus (US; 0.5 mA, 0.5 s footshock) in context A (shock floor, silver/aluminum walls, 70% ethanol cleanser). On day 2, mice

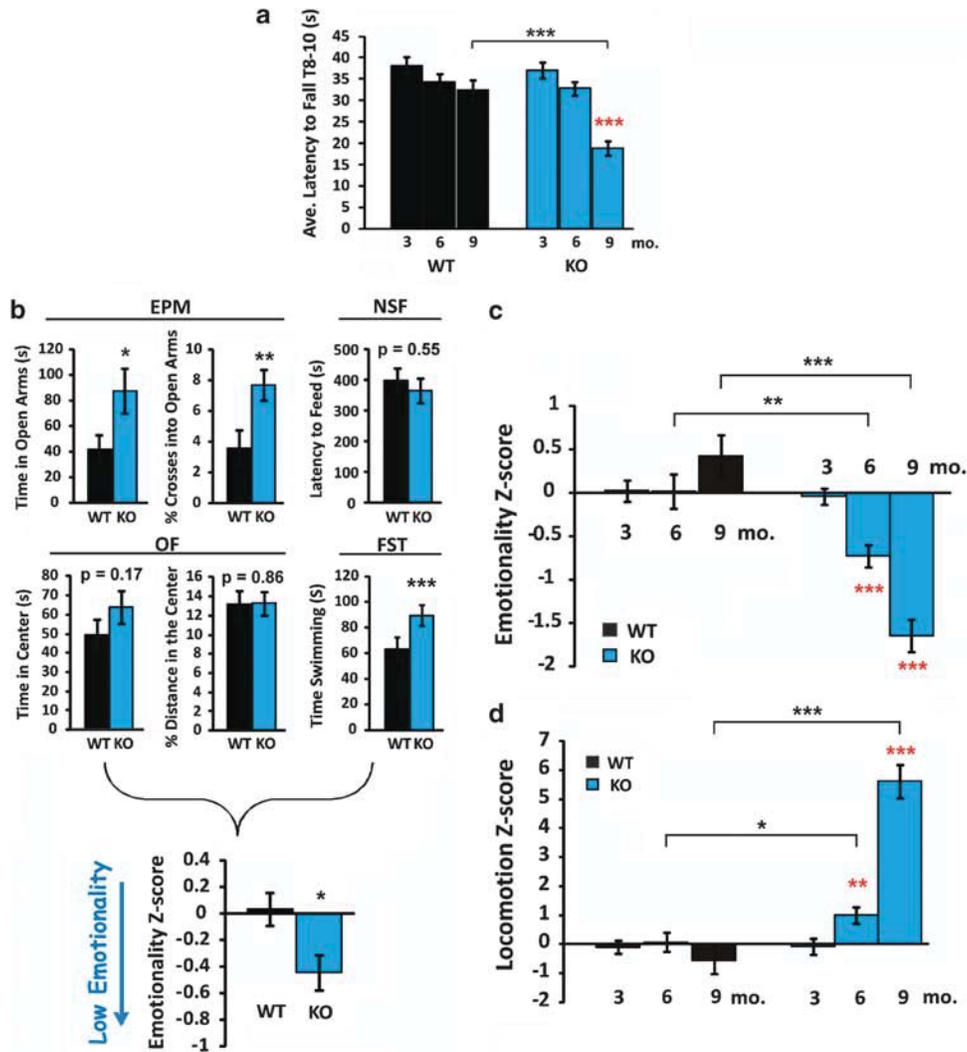


Figure 1 *Cnp1*^{KO} baseline behavior at 3, 6 and 9 months of age. (a) Latency to fall on the rotarod. *Cnp1*^{KO} mice show significant motor coordination deficits at 9 months. (b) Example of individual behavioral measures combined in the emotionality Z-scores (6-month time-point). See also Materials and methods, and Guilloux *et al.*³³ for Z-scoring methodology. *Cnp1*^{KO} mice showed significant changes in elevated plus maze (EPM) and forced swim test (FST), and nonsignificant changes in novelty suppressed feeding (NSF) and open field (OF). In NSF, no differences in weight loss or post-test food consumption were noted (data not shown). EPM, OF, NSF and FST measures were normalized using WT means and s.d.'s, and averaged per group.³³ Breakdown of Z-scores for other age groups are presented in Supplementary Figures 1–3. (c) High emotionality Z-scores indicate elevated anxiety-related and depressive-like behaviors. (d) Similarly derived locomotion Z-scores indicate elevated locomotor activity. Z-scores are normalized to the 3-month WT group for both emotionality and locomotion. Red asterisks represent within-genotype age comparisons. Black asterisks represent across genotype comparisons. Data represent mean ± s.e.m. (N = 9–18 per group). *P < 0.05, **P < 0.01, ***P < 0.001.

received 30 trials of the CS only in context B (non-shock grid floor, black walls, Windex cleanser). On day 3, mice were placed back in context B (extinction recall) followed by context A (fear renewal) 1 h later; and freezing was recorded during five CS exposures in each context. On each day, a 2-min acclimation period preceded and followed testing. Trials were presented with a variable inter-trial interval (25–35 s) and percent freezing was measured during the 15-s CS.

Immunohistochemistry. Stress-induced c-Fos immunoreactivity was performed as described.³⁷ Brains were collected by perfusion at 120 min following a 15-min restraint stress. Sections (40 μm, six per mouse) were incubated with c-Fos antibody (polyclonal rabbit anti-c-Fos; 1:5000 dilution; Calbiochem, San Diego, CA, USA), followed

by secondary biotinylated goat-anti-rabbit IgG (1:500 dilution; Vector Laboratories, Burlingame, CA, USA), and visualized with diaminobenzidine solution (DAB Peroxidase Substrate Kit; Vector Laboratories). Total numbers of c-Fos-positive cells were counted in the basolateral amygdala (BLA).

Gene array. Brains from the UCMS Cohort (N = 11–13 mice per group) were selected, based on emotionality Z-scores that were closest to the means of their group. Following dissection, brains were immediately flash frozen on dry ice. As the other hemisphere was allotted for other experiments, the left BLA was dissected directly on the cryostat using a 0.5-mm micropunch, tissue samples were frozen at –80 °C until extraction and total RNA was extracted using an RNeasy Mini Kit (Qiagen, Germantown, MD, USA).

Concentration and purity were tested using a Bioanalyzer and Nanodrop Spectrophotometer ($260/280 \geq 1.8$). One half of each RNA sample (~150 ng) was processed on Illumina Mouse WG-6v2 Expression BeadChips (San Diego, CA, USA) at the Keck Microarray facility (Yale University, New Haven, CT, USA) and expression levels were determined using the Illumina BeadArray Reader. The other half was used for independent verification by real-time quantitative PCR. Two internal controls verified the validity of the arrays (see Results): (1) *Cnp1* is downregulated/absent in *Cnp1*^{KO} and (2) *Cnp1* is downregulated in WT mice following UCMS.²²

Real-time quantitative PCR. Total RNA was converted into cDNA using the qScript cDNA synthesis kit (Quanta Biosciences, Gaithersburg, MD, USA). As previously described, real-time quantitative polymerase chain reaction reactions were assessed by SYBR green fluorescence signal (Invitrogen, Carlsbad, CA, USA) using the Opticon Monitor DNA Engine (Bio-Rad, Berkeley, CA, USA).¹³ Briefly, samples were run in quadruplicates and ΔC_t values were determined by comparison with the geometric mean of three reference genes (housekeeping genes: *actin*, *GAPDH* and *cyclophilin*). Signal intensities ($SI = 100 \times 2^{-\Delta C(t)}$) were used for comparison with microarray expression values.

Statistical analysis. There were no significant effects of sex or estrous phase in all examined cohorts, hence, male and female groups were combined, where appropriate. Genotype differences in Baseline and CORT Cohorts were assessed using repeated measures analysis of variance (ANOVA) for age and treatment, respectively. Genotype comparisons with the 3-month Baseline group were made using one-way ANOVA. The UCMS Z-scores were assessed using a two-way ANOVA for genotype and treatment. Weekly measures in UCMS (fur rating, body weight and corticosterone levels) were assessed using two-way repeated measures ANOVA for week, with genotype and treatment as cofactors. Genotype differences in the FC Cohort were assessed using repeated measures analysis of covariance with age as a covariate for days 1 (conditioning) and 2 (extinction) and one-way analysis of covariance for day 3 (recall and renewal). Genotype differences in the c-Fos Cohort were determined using two-way ANOVA for genotype and stress. Gene expression changes across genotype were assessed in four subgroups: WT-Control, WT-UCMS, KO-Control and KO-UCMS, using ANOVA followed by two-group *post hoc* tests for gene selection (Supplementary Table 2). Selection criteria were set at $P < 0.01$ and effect size greater than 30%. Genotype differences in real-time quantitative PCR were assessed with one-way ANOVA.

Results

***Cnp1*^{KO} mice display low baseline/trait emotionality and high locomotor activity.** Consistent with prior reports, *Cnp1*^{KO} mice showed normal motor coordination in the rotarod test at 3 and 6 months of age, and showed deficits in mice aged 9 months (Figure 1a). Mice were grossly impaired at 12 months of age and could not perform the rotarod test

(data not shown). Therefore, we tested the impact of reduced *Cnp1* on emotionality in male and female mice at 3, 6 and 9 months of age (i.e., before and during mild neurodegenerative-related stages), using four behavioral paradigms (EPM, OF, NSF and FST). To extract stable patterns underlying behavioral variability and summarize results across tests, emotionality-related measures were normalized to the 3-month WT group, and averaged per mouse across tests to obtain integrated 'emotionality Z-scores'³³ (See Materials and methods, and example of 6-month time-point in Figure 1b). *Cnp1*^{KO} mice were comparable to WT mice at 3 months of age and displayed progressively decreasing emotionality Z-scores at 6 and 9 months of age (Figure 1c and Supplementary Figures 1–3). It is unlikely that changes were due to repeated testing (at 6 and 9 months) as a pattern of decreased emotionality in WT mice would be expected, although *Cnp1*^{KO}-specific memory-related events cannot be ruled out (See Fear conditioning tests). Although EPM and OF emotionality measures were controlled for locomotor activity, FST results could reflect differences in activity. Removing the FST from the emotionality Z-score decreased power, but the same progressive pattern of low emotionality was observed in *Cnp1*^{KO} mice (6 months $P = 0.07$; 9 months $P < 0.0004$; Supplementary Figure 4).

Integrated Z-score measures of locomotor activity revealed a concomitant progressive increase from 3 to 9 months of age in *Cnp1*^{KO} mice (Figure 1d and Supplementary Figures 1–3). *Cnp1*^{KO} mice also displayed lower body weight at 6 and 9 months compared with WT, but had normal levels of stress hormones (Supplementary Figures 5a and b). Together, these results are supportive of a time-dependent decrease in emotionality in *Cnp1*^{KO} mice, which is paralleled by a progressive elevated locomotor phenotype.

***Cnp1*^{KO} mice are resistant to developing high emotionality states in two distinct rodent models of depression.** To test whether low baseline emotionality also conferred reduced vulnerability to develop high emotionality states, we exposed two independent cohorts of *Cnp1*^{KO} mice to environmental (UCMS) or neuroendocrine (CORT) stressors for a period of 4 weeks. In both experiments, WT groups responded with robust and characteristic increases in emotionality (Figures 2a and b). *Cnp1*^{KO} mice responded with increased emotionality after UCMS compared with non-stressed *Cnp1*^{KO} mice (Figure 2a; Supplementary Figure 6), but displayed a trend toward lower emotionality following CORT exposure (Figure 2b; Supplementary Figure 7). Notably, following both paradigms, *Cnp1*^{KO} mice remained at emotionality levels that were lower than WT non-stressed groups (Figures 2a and b). These differences remained when FST was excluded from Z-score analyses (Supplementary Figures 8a and b) and were confirmed in an independent cohort (Supplementary Figure 9). Note that the UCMS Cohort was 6 months of age at testing, hence, the low emotionality and high locomotion of *Cnp1*^{KO} mice match prior results (Figure 1), thus, independently confirming the Baseline phenotype.

UCMS increased locomotion in WT (trend level) but not in *Cnp1*^{KO} mice (Figure 2c; Supplementary Figure 6). CORT exposure reduced locomotion in WT and increased locomotion in *Cnp1*^{KO} mice (Figure 2d; Supplementary Figure 7).

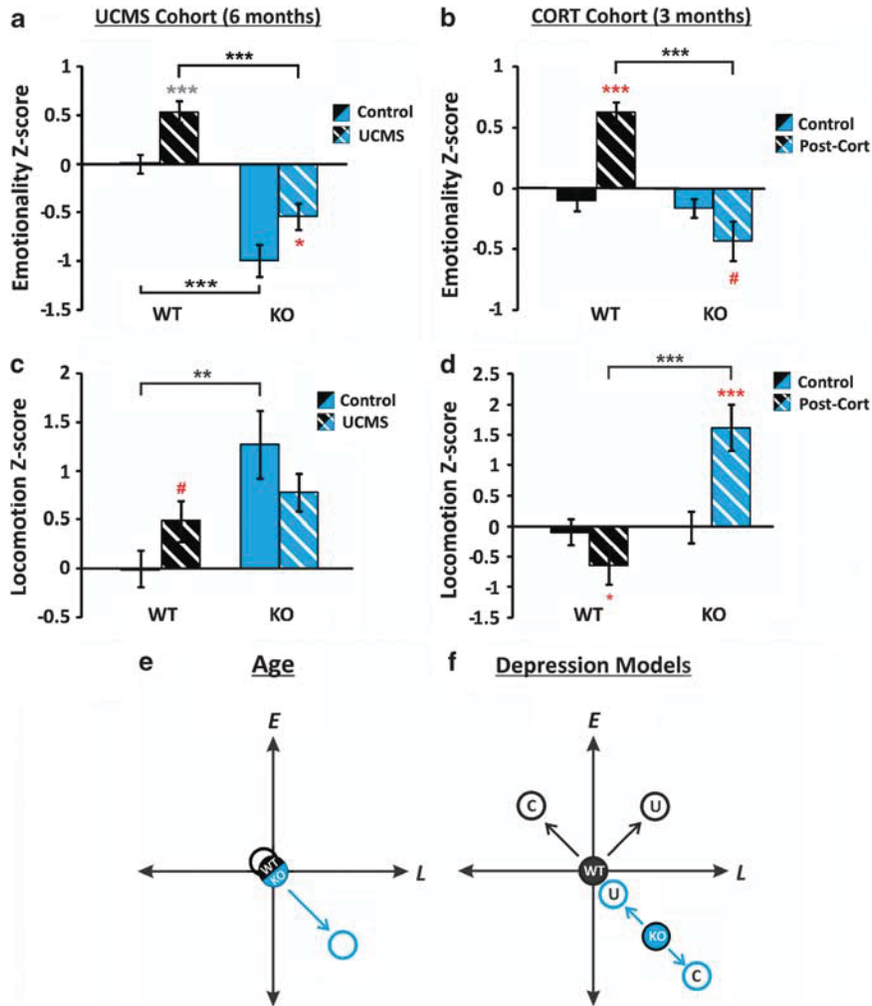


Figure 2 Effects of two rodent models of depression in *Cnp1^{KO}* mice. Emotionality Z-scores in the unpredictable chronic mild stress (UCMS) (a) and chronic corticosterone treatment (CORT) (b) Cohorts. Locomotion Z-scores in the UCMS (c) and CORT (d) Cohorts. Z-scores were normalized to wild-type (WT) control groups within each experimental cohort. Differences in *Cnp1^{KO}* control groups between the cohorts reflect age differences (UCMS, 6 months; CORT, 3 months). Schematic diagrams of emotionality and locomotion interactions in WT and *Cnp1^{KO}* mice across age (e) and after UCMS or CORT exposure. (f) WT mice showed dissociation in emotionality/locomotion effects between the two depression models, whereas changes in *Cnp1^{KO}* remained along the same axis. Red asterisks represent within genotype age comparisons. Black asterisks represent across genotype comparisons. Data represent mean \pm s.e.m. ($N = 10-14$ per group). * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$ and # represent statistical trends ($P < 0.1$).

No notable genotype differences were seen in weight gain, fur rating or in corticosterone levels in either paradigm (Supplementary Figures 10a–c and 11a–c). In summary, *Cnp1^{KO}* mice displayed reduced vulnerability to develop high emotionality behaviors after chronic environmental and neuroendocrine challenges, as emotionality Z-scores never reached WT levels. Notably, dissociations between changes in emotionality and locomotor activity were observed between genotype groups (Figures 2e and f).

Normal fear conditioning and cellular reactivity to stress, but reduced fear expression, suggest low encoding of emotional salience in *Cnp1^{KO}* mice. Our motivating study indicated reduced *Cnp1* levels in the amygdala of MDD subjects and UCMS-exposed mice. To test whether the observed behavioral phenotype included disrupted amygdala function, we assessed *Cnp1^{KO}* mice in

the FC paradigm (Figure 3a), a test relying on amygdala processing. Similar to WT, *Cnp1^{KO}* mice learned to associate the CS with the US on day 1 (Figure 3b; left panel). On day 2, both groups showed intact extinction learning, but *Cnp1^{KO}* mice displayed lower CS-induced freezing from the onset, resulting in significantly lower freezing throughout the extinction paradigm (Figure 3b; right panel), an indication of low fear expression.³⁸ No significant differences in pain sensitivity were observed in the hot plate test (Supplementary Figure 12). These results were confirmed in a separate cohort ($N = 8-12$ per genotype; $P < 0.01$ for extinction; Supplementary Figure 13). This robust genotype difference was consistently observed during extinction recall and fear renewal (day 3), and was not due to lack of consolidation of fear memory (Supplementary Figure 14).

To assess whether the observed differences could reflect baseline changes in stress-induced activation of

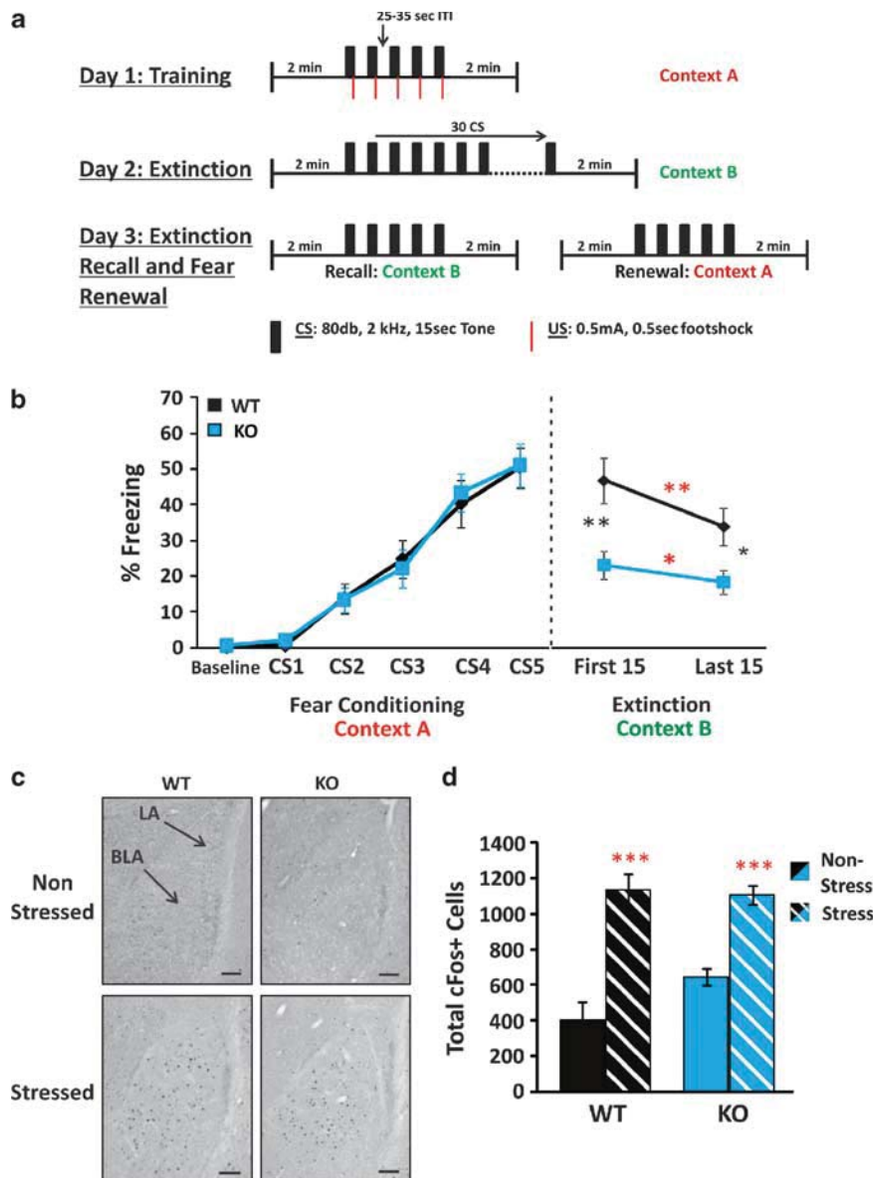


Figure 3 Fear conditioning (FC) and c-Fos analysis. **(a)** FC protocol. **(b)** Percent freezing during FC and extinction. **(c,d)** Analysis of stress-induced cFos expression in basolateral amygdala (BLA) of wild-type (WT) and *Cnp1*^{KO} mice. **(c)** c-Fos expression in the amygdala under 10 × magnification, scale bar = 0.1 mm. **(d)** Total c-Fos-positive cell counts. Although there was a trend ($P < 0.1$) toward a genotype difference in the non-stressed groups, no significant differences were found between WT and *Cnp1*^{KO}. In all the panels, red asterisks represent within-genotype comparisons. Black asterisks represent across-genotype comparisons. Data represent mean \pm s.e.m. ($N = 16$ – 19 per group and $N = 9$ – 18 per group). * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$.

the amygdala, we measured the expression of the immediate early gene c-Fos in the BLA of a separate cohort of animals following a 15-min restraint stress. WT and *Cnp1*^{KO} mice displayed similar baseline (although at trend level for increased reactivity in *Cnp1*^{KO} mice) and similar stress-induced number of c-Fos-positive cells (Figures 3c and d), indicating intact BLA response to acute stress in *Cnp1*^{KO} mice.

Upregulated oligodendrocyte- and immune-related gene transcripts in the BLA of *Cnp1*^{KO} mice. To investigate putative underlying amygdala-related mechanisms, large-scale gene expression was assessed by microarray in the BLA of

WT and *Cnp1*^{KO} mice from the UCMS Cohort (Figure 4a). Internal verification of the array data confirmed that *Cnp1* levels were undetectable in *Cnp1*^{KO} mice (Figure 4b) and downregulated in UCMS-exposed WT mice for two out of three probes (Figure 4c).^{13,22} UCMS-exposed *Cnp1*^{KO} (KO-UCMS) mice showed a similar pattern of transcript changes as UCMS-exposed WT mice ($R = 0.72$; Supplementary Figure 15), indicating that the broad biological response to stress is intact. However, owing to the phenotypic differences, these UCMS-related genes are unlikely to be related to the behavioral phenotype in *Cnp1*^{KO} mice. Instead, on the basis of consistent low emotionality in control and UCMS-exposed *Cnp1*^{KO} mice, we focused on genes

displaying similar changes under control and UCMS conditions, as potential ‘mediators’ of the behavioral phenotype. In total, 114 ‘*Cnp1*^{KO}-associated probesets’ were identified by the following criteria: (1) significant difference in *Cnp1*^{KO} under control (KO-Control vs WT-Control) and UCMS-exposed (KO-UCMS vs WT-UCMS) conditions ($P < 0.01$; effect size $> 30\%$), (2) consistent directionality of effect in both groups and (3) no change in WT after UCMS ($P > 0.1$).

On the basis of prior identification of relative glial/neuronal enrichment of transcript origin,³⁹ 97% of the 114 identified transcripts were enriched in glial or mixed glial/neuronal origin. Specifically, a systematic upregulation of oligodendrocyte-related transcripts was observed (24% of *Cnp1*^{KO}-associated genes), including genes involved in structure, function and production of the myelin sheath (Table 1). In parallel, we observed a significant upregulation of

immune-related transcripts (32% of *Cnp1*^{KO}-associated genes), including genes associated with the immune complement system and major histocompatibility complex (Table 1; see also Supplementary Table 2). Ingenuity pathway analysis of *Cnp1*^{KO}-associated genes identified a gene network linking oligodendrocyte with immune genes, suggesting that the upregulation of immune-related genes may be synchronized with the dysregulation of oligodendrocyte-related genes in *Cnp1*^{KO} mice. Only two neuronal-enriched genes were identified (*Cortistatin* and *Serpina3n*), suggesting minimal structural and/or functional adjustment in neurons.

Discussion

Focusing on an oligodendrocyte gene (*Cnp1*) that is critical for neuronal support and that is dysregulated in MDD, we tested

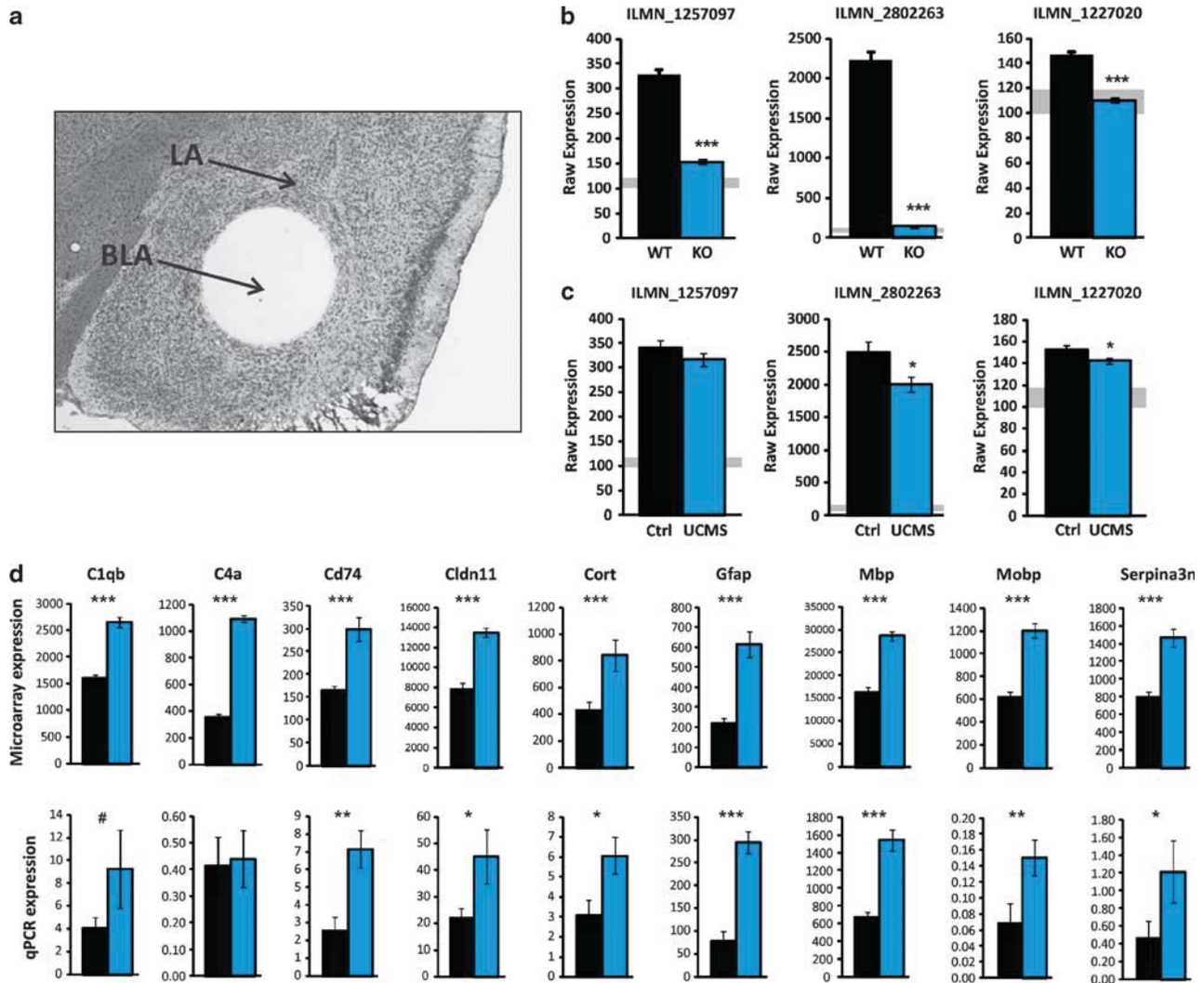


Figure 4 Basolateral amygdala (BLA) gene array analysis and real-time quantitative PCR validation (a) BLA tissue micropunch. (b) Absence of expression of the three *Cnp1* probes in *Cnp1*^{KO} mice. Expression levels were at the threshold of detection (~100–120 U; gray shading). (c) Downregulation of two out of three *Cnp1* probes in wild-type (WT) mice exposed to unpredictable chronic mild stress (UCMS). (d) Real-time quantitative PCR confirms significant upregulation of eight out of nine genes. Data represent mean \pm s.e.m. (Microarray $N = 11$ –13 per group; real-time quantitative PCR $N = 6$ per group). * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$ and # represent statistical trends ($P < 0.1$).

Table 1 Selection of significantly affected neuronal, oligodendrocyte and immune-related genes between *Cnp1*^{KO} and WT control mice

Gene Symbol	Gene Title	alr	p value (KO vs. WT)
<u>Serpina3n</u>	<u>serine (or cysteine) peptidase inhibitor clade A member 3N (Serpina3n)</u>	0.81	2.13E-10
<u>Cort (2)</u>	<u>cortistatin</u>	0.73	5.14E-06
<u>Maq</u>	<u>myelin-associated glycoprotein</u>	1.16	6.64E-18
<u>Tspan2</u>	tetraspanin 2	0.95	1.51E-16
<u>Tmem10</u>	transmembrane protein 10	1.10	6.76E-16
<u>Cldn11</u>	claudin 11	0.98	2.59E-15
<u>Adamts4</u>	<u>a disintegrin-like and metallopeptidase (reprolysin type) with thrombospondin type 1 motif, 4</u>	0.73	2.66E-15
<u>Mbp</u>	<u>myelin basic protein transcript variant 7</u>	0.83	3.68E-15
<u>Tmem125</u>	transmembrane protein 125	0.80	9.05E-14
<u>Mal</u>	<u>myelin and lymphocyte protein T-cell differentiation protein</u>	0.83	1.48E-13
<u>Slc44a1</u>	solute carrier family 44 member 1	0.73	3.90E-13
<u>Plip</u>	plasma membrane proteolipid	0.67	1.34E-12
<u>Plp1</u>	proteolipid protein (myelin) 1	0.48	2.99E-12
<u>Gltp</u>	glycolipid transfer protein	0.56	1.80E-11
<u>Mcam</u>	melanoma cell adhesion molecule	0.64	2.66E-11
<u>Gjc2</u>	gap junction protein gamma 2 (Gjc2) transcript variant 2	0.69	4.09E-11
<u>Fgfr2</u>	fibroblast growth factor receptor 2 transcript variant 2	0.48	5.44E-11
<u>Cd9</u>	<u>CD9 antigen</u>	0.75	9.35E-11
<u>Rhog</u>	ras homolog gene family member G	0.63	1.62E-10
<u>Elov11</u>	elongation of very long chain fatty acids-like 1 transcript variant 2	0.46	5.27E-10
<u>Mobp (4)</u>	myelin-associated oligodendrocytic basic protein	0.95	8.59E-09
<u>Ddr1</u>	discoidin domain receptor family member 1 transcript variant 1	0.50	1.09E-07
<u>Fa2h</u>	fatty acid 2-hydroxylase	0.44	5.64E-07
<u>Pmp22</u>	peripheral myelin protein	0.42	8.64E-07
<u>Moq (2)</u>	<u>myelin oligodendrocyte glycoprotein</u>	0.77	9.17E-07
<u>Nkx6-2</u>	NK6 transcription factor related locus 2 (<i>Drosophila</i>)	0.45	1.14E-05
<u>Lyz (2)</u>	lysozyme	1.78	8.29E-21
<u>Ly86</u>	lymphocyte antigen 86	0.77	3.06E-20
<u>Cd52</u>	CD52 antigen	1.39	4.74E-20
<u>Fcer1g</u>	Fc receptor, IgE, high affinity I, gamma polypeptide	0.74	5.96E-18
<u>C4a (2)</u>	complement component 4A (Rodgers blood group)	1.50	4.58E-16
<u>Tyrobp</u>	TYRO protein tyrosine kinase binding protein	0.73	5.52E-16
<u>Cd63</u>	CD63 antigen	0.59	7.72E-16
<u>Cyba</u>	cytochrome b-245 alpha polypeptide	0.65	1.49E-15
<u>C1qa</u>	complement component 1 q subcomponent alpha polypeptide	0.70	2.67E-15
<u>C1qb</u>	complement component 1 q subcomponent beta polypeptide	0.76	2.89E-15
<u>Trem2</u>	triggering receptor expressed on myeloid cells 2	0.70	8.75E-15
<u>C4b</u>	complement component 4B (Childo blood group)	1.24	9.36E-15
<u>C1qc</u>	complement component 1 q subcomponent C chain	0.69	3.47E-14
<u>Fcgr3</u>	Fc receptor IgG low affinity III	0.50	3.35E-13
<u>Lgals3</u>	lectin galactose binding soluble 3	0.74	5.33E-13
<u>Ctsc</u>	cathepsin C	0.54	7.47E-13
<u>Osmr</u>	oncostatin M receptor	0.45	2.12E-12
<u>Hvcn1</u>	hydrogen voltage-gated channel 1 transcript variant 1	0.41	2.77E-12
<u>B2m (2)</u>	beta-2 microglobulin	0.56	7.65E-12
<u>Kih16</u>	kelch-like 6 (<i>Drosophila</i>)	0.40	3.93E-10
<u>Litaf</u>	LPS-induced TN factor	0.54	6.06E-10
<u>Ndrq1</u>	N-myc downstream regulated gene 1	0.56	8.96E-10
<u>Lyz2</u>	lysozyme 2	0.73	1.39E-09
<u>Cd68</u>	CD68 antigen	0.56	1.98E-09
<u>Lag3</u>	lymphocyte-activation gene 3	0.68	2.27E-09
<u>Cd82</u>	CD82 antigen	0.55	4.69E-09
<u>Adss1</u>	adenylosuccinate synthetase like 1	0.48	7.37E-08
<u>Cd74 (2)</u>	CD74 antigen (major histocompatibility complex class II antigen-associated) transcript variant 2	0.66	1.33E-06

Abbreviations: Alr, average log ratio (*Cnp1*^{KO}/WT); KO, knockout; WT, wild type.

Number of redundant probes is shown following the gene symbol. Red, neuronal-enriched genes; blue, oligodendrocyte-enriched genes; underline, immune-related genes. The listed *P*-values represent the main genotype effect from the ANOVA analysis (see Supplementary Table 2).

the potential mechanistic link between altered oligodendrocyte function and emotionality in mice. We show that removal of *Cnp1* in mice leads to unexpected reduced baseline emotionality, reduced fear expression (during extinction) and lower vulnerability to develop high emotionality states using two rodent paradigms to induce depressive-like states.

This behavioral profile suggests the presence of a dysfunctional amygdala-related network that is consistent with proposed mechanisms for stress resiliency. Gene array analysis in the BLA of *Cnp1*^{KO} mice revealed a robust upregulation of oligodendrocyte- and immune-related transcripts, potentially representing functional compensations.

Cnp1^{KO} mice show a low baseline emotionality phenotype (trait) that appears progressively over time, but that is observed before the onset of motor coordination deficits (9–12 months), indicating that it is not due to the late-onset widespread axonal degeneration observed in *Cnp1*^{KO} mice.³¹ Rather, more discrete molecular changes in myelin structure have been observed at earlier ages, which may contribute to the observed phenotype. For instance, the direct functional consequence of *Cnp1* ablation is not known, but disorganization of critical proteins (e.g., Caspr, Nav) at the paranode region at 3 months and degeneration in some small diameter axons as early as post-natal day 15 were reported in *Cnp1*^{KO} mice.^{30,40} Hence, impaired paranode function may translate into suboptimal support for neuronal axons, hence priming the system for dysregulated physiological responses to stress/fear.

Cnp1^{KO} mice also display a concomitant increase in locomotor activity. Tests of emotionality control for activity, but locomotor and emotionality phenotypes are often difficult to dissociate in rodent models. In fact, quantitative trait loci mapping studies have identified regions on chromosomes 2, 7 and 8 that encode for both anxiety-related behavior and locomotion in rats,⁴¹ and the well-characterized high emotionality and low locomotion phenotypes of mice⁴² or rats⁴³ lacking the serotonin transporter associate with common genetic loci.⁴³ Hence, instead of a confounding factor, altered locomotor activity may represent an epistatic outcome of genetic disruptions of emotionality regulators, which are nevertheless dissociable under certain genetic and/or stress-induced conditions (Figures 2e and f).

Cnp1^{KO} mice have normal amygdala cellular reactivity (c-Fos measures) and fear acquisition, but display low fear expression during extinction learning (Figure 3), which was not likely due to poor consolidation of the fear memory (Supplementary Figure 14). Instead, we speculate that reduced fear expression in *Cnp1*^{KO} mice may result from low encoding of emotionality salient stimuli (e.g., the original fear association). Accordingly, *Cnp1*^{KO} mice respond to chronic behavioral (UCMS) or physiological (CORT) stress (Figure 2a), but their induced emotionality states remained significantly below the levels associated with depressive-like states in WT mice (Figures 2a and b). Together, this behavioral pattern is consistent with stress resilient phenotype. Indeed, in humans, resilience is associated with the ability to adapt to chronic stress⁴⁴ and to perceive stressful events in a less threatening way,^{45–47} both characteristics observed here in *Cnp1*^{KO} mice. Fear expression is dependent on BLA function,^{48,49} but is also modulated by regions of the prefrontal cortex.^{38,50} Similarly, in humans, the ventromedial prefrontal cortex modulates amygdala function during fear extinction⁵¹, and potent ventromedial prefrontal cortex inhibition of the amygdala is postulated to occur in resilient individuals.^{44,52} Together, this suggests the presence of a dysfunctional amygdala-related neural network in *Cnp1*^{KO} mice, consistent with proposed mechanisms for stress resilience. As disruption in related networks including prefrontal and/or hippocampal regions cannot be excluded, future site-specific and time-dependent alterations could further refine the role of *Cnp1*.

Microarray evaluation of transcript levels within the BLA of *Cnp1*^{KO} mice indicated a robust pattern of upregulated

oligodendrocyte-related transcripts. This suggests an attempted compensation for the lack of structure/function due to missing *Cnp1*, and potentially reflects prior reports of enlargement of the myelin inner tongue in small diameter axons and whirls of excess redundant myelin sheaths seen early in *Cnp1*^{KO} mice.³⁰ Microarray assessment also revealed a pattern of upregulated immune-related transcripts (major histocompatibility complex class II, complement system, inflammatory mediators; Table 1), consistent with reports of reactive gliosis and microglial activation in *Cnp1*^{KO} mice.³¹ Interestingly, components of the complement system can be either deleterious or beneficial.⁵³ A recent report found upregulation of certain immune-related genes (e.g., *c1qc*, *cd74*, *Serpina3*, *tyrobp*) was associated with protected cognition in old subjects,⁵⁴ suggesting neuroprotective potential. Here, upregulated immune-related genes in *Cnp1*^{KO} mice show ~40% overlap with the gene set implicated in that study, suggesting potential mixed deleterious/neuroprotective effects of immune-related gene changes in *Cnp1*^{KO} mice. Surprisingly, only two neuronal-associated transcripts were found changed (upregulated) in *Cnp1*^{KO} mice, and both are associated with immune function. Cortistatin, a neuropeptide, has anti-inflammatory and neuroprotective properties in mice⁵⁵ and humans,^{56,57} whereas *Serpina3*, a serine peptidase inhibitor, is upregulated in response to inflammation in the rodent brain,^{58,59} suggesting a protective role for these genes in *Cnp1*^{KO} mice.

Whether increased immune- and oligodendrocyte-related transcripts may be an attempt of the system to 'repair' damage and/or protect against further damage owing to the lack of *Cnp1*-related function, is not known. However, ingenuity-based functional analyses suggest that the two processes are inter-related, a finding consistent with reports in most mental illnesses,^{60–62} and highlighted by the comorbidity of emotion regulation disorders (i.e., anxiety, depression) in patients with multiple sclerosis, an inflammatory demyelinating central nervous system disease.^{63,64} Consequently, *Cnp1* could be a critical component of the mechanistic link between oligodendrocytes and immune function underlying psychiatric and other central nervous system disorders.^{32,65}

Although the molecular phenotype of the *Cnp1*^{KO} appears confined to non-neuronal compartments, one of the primary functions of myelinating oligodendrocytes is to support electrical signal conduction along the axon.⁶⁶ The paranode region, where *Cnp1* is localized, is critical for maintaining axonal integrity.^{67,68} *Cnp1* and other gene products in this region are thought to maintain optimal functioning of axonal mitochondria and it is suggested that perturbations at the paranode could disrupt this fragile metabolic coupling.³² Electrophysiological changes in oligodendrocytes can modulate axonal conduction velocity,⁶⁹ and the firing of action potentials in NG2 cells is dependent on axonal synapses,⁷⁰ highlighting the functional connectivity between neurons and oligodendrocytes.⁷¹ In addition, structural changes in oligodendrocytes, specifically at the paranode region, are suggested to be a potential mechanism for subtle alterations in axonal conduction and associated loss of signal integrity.^{72,73} Accordingly, improper signal conduction in amygdala-related circuitry in *Cnp1*^{KO} mice (due to paranode disruption) could account for the low encoding of emotionally salient information, and related phenotypic abnormalities.

Previous studies reported low *Cnp1* in MDD and in response to chronic stress^{13,22} (Figure 4). We now demonstrate that disruption of oligodendrocyte function (via *Cnp1* ablation) can impact circuits mediating emotionality in mice, leading to a resilient emotionality phenotype. Thus, although these studies indicate that *Cnp1* disruption is not likely a causal factor in MDD; the current results suggest that low *Cnp1* and/or disruption of a critical axoglial junction may contribute to clinical symptoms of mood disorders in an unexpected and potentially maladaptive way (due to massive molecular changes), and should therefore be further examined in relation to psychiatric illnesses and potential treatments.

Conflict of interest

The authors declare no conflict of interest.

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