

# Absence of VGLUT3 expression leads to impaired fear memory in mice

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53	
54	Abstract
55	Fear is an emotional mechanism that helps to cope with potential hazards. However, when fear is generalized
56	it becomes maladaptive and represents a core symptom of Post-Traumatic Stress Disorder (PTSD). Converg-
57	ing lines of research show that dysfunction of glutamatergic neurotransmission is a cardinal feature of trau-
58	ma and stress related disorders such as PTSD. However, the involvement of glutamatergic co-transmission in
59	fear is less well understood. Glutamate is accumulated into synaptic vesicles by vesicular glutamate trans-
60	porters (VGLUTs). The atypical subtype, VGLUT3 is responsible for the co-transmission of glutamate with
61	acetylcholine (ACh), serotonin (5-HT) or GABA.
62	To understand the involvement of VGLUT3-dependent cotransmission in aversive memories, we used a Pav-
63	lovian fear conditioning paradigm in VGLUT3 <sup>-/-</sup> mice. Our results revealed a higher contextual fear memory
64	in these mice, despite a facilitation of extinction. In addition, the absence of VGLUT3 leads to fear generali-
65	zation, probably due to a pattern separation deficit. Our study suggests that the VGLUT3 network plays a
66	crucial role in regulating emotional memories. Hence, VGLUT3 is a key player in the processing of aversive
67	memories and therefore a potential therapeutic target in stress-related disorders.
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### Introduction

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Fear is an emotion in response to a threat that is essential for survival. However, generalization of fear is a core symptom of major psychiatric disorders such as anxiety disorders, phobia, panic disorder and posttraumatic stress disorder (PTSD) (Lissek and van Meurs, 2015; Besnard and Sahay, 2016). Major progress has been made thanks to animal studies of aversive memories through the Pavlovian fear conditioning paradigm (LeDoux, 2012). This test consists of associating an initially neutral stimulus (such as a tone, a light or a context) to an aversive event (such as a footshock) (Maren et al., 2013). This paradigm is widely used to dissect mechanisms underlying fear learning and memory, and to better understand fear-related disorders. Fear generalization is defined by the fact that a distinct, but perceived by the animal as similar, context elicits fear responses. The balance between contextual discrimination and generalization is a crucial aspect of the expression of fear. Fear generalization is currently considered a central feature of generalized anxiety and PTSD (Lissek, 2012; Mahan and Ressler, 2012). An effective treatment for fear generalization is based on extinction training to reduce acquired fear (Craske et al., 2008; Rothbaum and Davis, 2003). Fear extinction consists of new inhibitory learning after repeated presentations of fear-associated stimulus, in the absence of the aversive event, leading to a gradual decrease in the magnitude of the fear response (Myers and Davis, 2007). However, after extinction fear memory is not erased, but inhibited, as it can reappear in spontaneous recovery, external disinhibition, renewal, and reinstatement (Maren and Holmes, 2016). Therefore, it is important to better characterize neural circuits underlying the formation and maintenance of aversive memories if we want to understand and treat generalized fear more efficiently. The neuronal circuits and the neuromodulators regulating emotional memories are well characterized. Emotional memories rely on a complex network including the amygdala, the hippocampus and the prefrontal cortex (Tovote et al., 2015). The amygdala is necessary for fear processing from acquisition to expression, whereas the hippocampus is mainly involved in contextual memory processing (Myers and Davis, 2007; Sierra-Mercado et al., 2011; Fanselow, 2000; Marek et al., 2018). Finally, the infra-limbic (IL) and the prelimbic areas of the prefrontal cortex are essential for fear extinction (Sierra-Mercado et al., 2011; Marek et al., 2019).

Several studies have highlighted the involvement of neurotransmitters including glutamate, GABA, acetyl
choline and serotonin signaling in fear processing (Ballinger et al., 2016; Baratta et al., 2016; Christianson et al., 2016; Chri
al., 2010; Craske et al., 2008; Jiang et al., 2016; Johnson et al., 2015; Knox, 2016; Krabbe et al., 2018; Wil
son and Fadel, 2017). Interestingly several subpopulations of neurons and fibers of the amygdala, the hippo
campus or the prefrontal cortex release more than one neurotransmitter (for review, see El Mestikawy et al.
2011; Trudeau and El Mestikawy, 2018). Most of these bilingual neurons in the fear circuit express the atype
ical vesicular glutamate transporter type 3 (VGLUT3) (Rovira-Esteban et al., 2017; Sengupta and Holmes
2019; Amilhon et al., 2010; Fasano et al., 2017; Herzog et al., 2004; Omiya et al., 2015). Studies have illus
trated the involvement of VGLUT3 neurons in psychiatric disorders (Sakae et al., 2015; Favier et al., 2020)
Several studies have demonstrated that the absence of VGLUT3 in VGLUT3 neurons led to the abolishmen
of glutamatergic currents mediated by mGlu receptors in the striatum or the hippocampus (Sakae et al, 2005
Fasano et al, 2017, Favier et al, 2020) whereas others showed the abolition of a glutamatergic ionotropic currents.
rents (Higley et al., 2011; Varga et al., 2009). Interestingly, VGLUT3-/- mice show a persistent hyper-
reactivity to stress (Amilhon et al., 2010) and a dysregulation of their HPA-axis (Balázsfi et al., 2018), but
only a few studies focused on the role of VGLUT3 in the regulation of emotion and fear. A couple of studies
previously showed that VGLUT3 deficient mice have a higher contextual fear memory and tend to general
ize their fear to unrelated situations (Balázsfi et al., 2018) with no major other memory deficits (Fazekas e
al., 2019).
In this context, our aim was to confirm the role of VGLUT3 in aversive memories and to deepen our under
standing of it by using a combination of behavioural paradigms. Using a Pavlovian fear conditioning paradigm
we report that VGLUT3-/- mice express more stable and generalized contextual memories associated with a
deficit of pattern separation. Interestingly, VGLUT3-/- mice have no deficit in non-aversive learning or in
working memory, spatial reference memory, or in recognition memory. These results highlight the specific
role of the VGLUT3-positive network in the establishment and maintenance of aversive memories and mos
notably in the generalization of fear. They also provide evidence that VGLUT3 could be considered as a po-
tential target for the treatment of stress-related disorders.

## Materials and methods

### Animals

Animal care and experiments were conducted in accordance with the European Communities Council Directive for the Care and the Use of Laboratory Animals (86/809/EEC) and in compliance with the French Ministère de l'Agriculture et de la Forêt, Service Vétérinaire de la Santé et de la Protection Animale. All efforts were made to minimize the number of animals and to ensure their well-being. Animals were group caged and housed in a temperature-controlled room (20±2°C) with free access to water and food under a light/dark cycle of 12h (light 7:30am – 7:30pm).

VGLUT3<sup>-/-</sup> mice (Gras et al., 2008) were on a C57BL6/J background. Heterozygous mice were bred to obtain VGLUT3<sup>-/-</sup> mice and wildtype (VGLUT3<sup>+/+</sup>) littermates. Pups were weaned around 22 days old, marked by ear punch and genotyped using the ear sample. Experiments were performed with 2–4-month-old mice (159 males and 50 females). Animals were randomly allocated to experimental groups and investigators were blinded for experimental procedures. Total animal number used in each paradigm is presented in Table 1.

### Behavioural paradigms

#### The Watermaze task

The WM test was performed as described previously (Daumas et al., 2008). The mice were monitored with a video tracking system (AnyMaze). First mice went through a 4 days cuetask protocol where the 1.8m diameter pool is surrounded with curtains, and a cue placed on the platform (60sec trials, 4 trials a day, ITI=20min). For the spatial reference memory (SRM) task, the platform was centered in one of the four quadrants and kept stable throughout the task (without any cue on it). The protocol lasted 5 days (90sec trials, 4 trials a day, ITI=10min). Ten minutes after the last trial on day 5, a 60-sec probe test (SRM-10 min) was conducted during which the platform was removed. In order to avoid extinction, an additional trial with the platform was done immediately after each probe test. A second probe test was performed 72 hrs after assessing the long-term memory of the mice (SRM-72 hrs). For the SRM Reversal (SRM-R) task, which was

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conducted immediately after the second probe test, learning flexibility was assessed by moving the platform to the opposite quadrant used for the SRM task. The animals were trained for 3 days (90sec trials, 4 trials a day, ITI=10min) and spatial memory was assessed at 10 min (SRM-R-10 min) and 72 hrs (SRM-R-72 hrs) after the last SRM-R trial. Data for the following parameters were collected: latency to reach the platform location, path length, swim speed, thigmotactic behavior and the percentage of time spent in the quadrant zones.

#### Novel place recognition (NPR) / Novel object recognition (NOR)

The NPR/NOR task was performed in a square open-field (25cm) with sawdust on the floor and cues on the walls. Habituation consisted of 1) a 10-min exploration period of the open-field with cagemates (Day1), 2) two 5-min periods during which each mouse was placed individually in the empty open-field on two consecutive days (Day2-3), and 3) a 5-min period during which the mice were placed in the open-field with two identical objects (Day4). On the training day (Day5), mice were allowed to explore two new identical objects until they had accumulated 15 sec then 10 sec of total inspection time during the first and second training session, respectively. Since VGLUT3-- mice are more anxious, the protocol was adapted in this way rather than a fixed 10-min training session, to ensure that all animals explore the objects sufficiently to establish memory formation. Therefore, the length of the session was different between animals, but the exploration of the objects was identical. On Day6, the mice were tested for the NOR paradigm (10 min) during which one of the original objects was replaced with a new object. On Day7 we started the NPR paradigm during which two new objects were placed in the open-field. As for NOR, 2 sessions of training were run and consisted in accumulating 15 sec and 10 sec of total exploration time. Twenty- four hours later (Day8), the mice were tested in the NPR paradigm (10 min): the same pair of familiar objects was used but one of the objects was displaced in another corner of the open-field. The percentage of time exploring the new object was calculated as a discrimination index: [novel /(novel + familiar)].

## Y Maze

Working memory was assessed with a Y maze apparatus (Imetronic, Pessac, France). Mice freely explored the maze for 10 min. The total number of entries was counted as well as the spontaneous alternation. Spontaneous alternation occurs when a mouse enters a different arm of the maze 3 consecutive times. The percent-

1/9	age of spontaneous afternation was calculated by dividing the number of spontaneous afternations by the to-
180	tal number of arm entries minus 2 and multiplied by 100.
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182	Fear Conditioning Experiments
183	The Fear Conditioning Apparatus (BIOSEB) is made of black methacrylate walls, a grid floor and transpar-
184	ent ceiling and front door. Panlab software (BIOSEB) was used to carry out the experiments and record
185	freezing behaviour. A video recording system (Multimedia Video Record) allowed manual scoring of freez-
186	ing levels to validate the automatic counts.
187	Shock sensitivity paradigm. Because VGLUT3 is present in peripheral sensory neurons and contributes to
188	mechanical pain (Seal et al, 2009), we assessed the sensitivity to electric footshocks in VGLUT3 <sup>-/-</sup> mice. A
189	train of electric footshocks (ES, 1-sec duration) was delivered starting from 0.1 mA and gradually increasing
190	by 0.05 mA every 30 sec. Shock delivery was stopped when all expected behavioural responses were ob-
191	served: increased locomotor activity (movement), vocalization, running and jumping. The intensity of the
192	electric shock that first triggered each of these behaviours was recorded.
193	The fear conditioning paradigm was used to study learning and memory of aversive stimuli as previously
194	reported (LeDoux, 2003; Daumas, 2005). Since VGLUT3-/- mice are deaf (Ruel et al., 2008; Seal et al.,
195	2008), a flashing light was used as the conditioned stimulus (CS: 20 sec, 2 sec ON / 2 sec OFF, 80 lux) and a
196	0.25 mA electric footshock as the unconditioned stimulus (US, 2 sec).
197	After 3 days of habituation (6 min per day), the conditioning session took place on day 4. After 2 min in the
198	chamber, the CS was triggered and its final 2 sec coincided with the US. After a 30 sec interval, a second
199	CS-US pairing was presented. Memory tests were done on day 5. Contextual memory was assessed with the
200	contextual test, and cue memory was assessed by the cue test 2 hours later. For the contextual test, mice were
201	placed in the conditioning context for 6 min without CS (light) or US. The cue test consisted of 3 min of ex-
202	ploration of a modified context (color, shape, light and odor), followed by 4 CS presentations with an
203	intertrial interval of 30 sec.
204	A Pattern Separation protocol was conducted for 11 days in two highly similar contexts: the shock associ-
205	ated context A and the safe context B as described by (Sahay et al., 2011). On day 0 mice were introduced

tistics are presented in Extended Tables.

206	into context A and after 185 sec received a 0.75 mA US for 2 sec. During the following ten days, mice were
207	exposed to the US-associated context A (183 sec exploration - 2 sec US - 15 sec exploration, before being
208	removed to home cage) and one hour later to the safe context B (180 sec exploration) in a defined order.
209	Freezing behaviour was assessed during the first 180 sec for each context.
210	Immediate shock procedure. Mice were submitted to a no-shock (NS) or an immediate shock (IS) proce-
211	dure. For the NS, mice were free to explore the conditioning cage for 30sec. In the IS procedure, mice re-
212	ceived an immediate shock (0.25mA, 2sec) immediately after their placement in the conditioning chamber
213	and were removed after 30sec. Generalized fear was evaluated 24h later by placing the animals in the condi-
214	tioning chamber (same context; SC) or in a novel box (novel context; NC) for 5 min.
215	Fear extinction learning and memory were studied for 15 days. Mice were habituated to the conditioning
216	chamber for 2 min before ten CS-US were delivered at 75 sec intervals. From day 2 to day 8, extinction took
217	place in the modified context. Mice were exposed to 10 presentations of CS with an interval of 85 sec under
218	red light illumination. A learning index (LI) was calculated daily. This index is used to ascertain the daily
219	extinction rate by calculating the difference between the first and last CS-induced freezing. On day 15, mice
220	were re-exposed to the conditioning context with ten CS presentations to assess fear recall. On day 18, they
221	were placed in a new context and ten CS were once again presented in order to evaluate renewal in a new
222	context.
223	
224	Statistics
225	Statistical comparisons were performed with Prism 9 (GraphPad software Inc. USA for Mac OS, La Jolla,
226	CA). Each statistical test was appropriately chosen for the relevant experimental design. Sidak's multiple
227	comparisons test was performed for post-hoc analysis when required unless otherwise indicated. All data are
228	presented as the mean $\pm$ SEM, with differences considered significant at p<0.05. Complete analysis and sta-

## 230 Results

Fear conditioning is based on learning/memory and on the propensity of mice to feel and react to electric footshock. VGLUT3 is expressed in the hippocampus where it contributes to hippocampal plasticity and network properties (Fasano et al., 2017). On the other hand, VGLUT3 is also found in subsets of neurons in pain circuits (Sakai et al., 2020; Larsson and Broman, 2019; Peirs et al., 2015; Draxler et al., 2014; Seal et al., 2009; Landry et al., 2004). Therefore, prior to using the fear conditioning paradigm, we assessed learning, spatial memory and pain threshold (i.e., response to foot shock) in VGLUT3<sup>-/-</sup> mice.

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## The absence of VGLUT3 does not impair learning and memory in mice

To explore the consequences of VGLUT3 deletion on spatial and non-spatial memories, we first used the watermaze task (WM) (Fig.1; statistics details can be found in extended figure 1-1). Relative to wildtype littermates, VGLUT3-/- mice displayed no impairment of learning in either the non-spatial (Fig.1D) or the spatial (Fig.1E) task. We observed a main effect of time but no main effect of genotypes or interaction between time and genotype. Therefore, both genotypes improved their learning during the training days (Fig.1D-E p<0.0001). To challenge them and assess their learning flexibility, a 3-day reversal task was performed immediately after the spatial reference learning task (Fig.1E: R1-R3). On day 1 of reversal learning (R1 in Fig.1E) both groups increased their latency to reach the new platform location, and then similarly improved their performance (Fig.1E; time: p<0.0001, genotype: p=0.882). Spatial memory was assessed 10 min and 72h after training completion for SRM and SRM-R. In all tested conditions, control littermates as well as VGLUT3-- mice spent significantly more than 25% of probe trial time in the targeted quadrant, indicating intact spatial reference memory (group performance vs. 25% p<0.05; Fig. 1F). However, during the long-term memory test, VGLUT3-- mice showed better performances (SRM-PT2) than controls (Fig.1F). Since VGLUT3-/- mice are more vulnerable to anxiety than WT mice (Amilhon et al, 2010), we explored the contribution of anxiety to memory formation and learning in VGLUT3--- mice in a more stressful condition, when the water temperature was lowered to 19°C (Sandi et al., 1997). At 19°C, we observed no main effect of genotype or interaction between genotype and time, but a main effect of time for both cuetask and SRM/SRM-R (Fig.1G-H). A 3 way ANOVA revealed no main effect of genotype, tests or water temperature and no interactions between these parameters except for the temperature x genotype (p=0.04; see Extended

258	Figure 1-1 for statistical details). Moreover, in all tested conditions, VGLUT3-/- and control mice show simi-
259	lar performances and spent more than 25% of their time in the correct quadrant (Fig.11). These data show no
260	deficit of learning and memory in VGLUT3 <sup>-/-</sup> mice in the WM paradigm.
261	
262	Figure 1 AROUND HERE
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264	We then studied spontaneous learning and memory using the object recognition paradigm. We observed for
265	both genotypes a significant difference from chance level (score 0.5) revealing long-term recognition
266	memory for objects (Fig.2A) and position (Fig.2B) in all animals. VGLUT3 mice show higher perfor-
267	mances than control littermates in the object recognition task (Fig.2A). Since VGLUT3 mice spent signifi-
268	cantly more time in the open field, we wondered whether the time spent during training was correlated with
269	the memory score obtained in the object recognition test. The correlation curve (Fig.2C) and the linear re-
270	gression revealed no correlation between memory score and the length of the session in VGLUT3-/- mice
271	$(R^2=0.03, F(1,11)=0.31, P>0.05;$ equation: Y=-0.001889*X+0.6951). These experiments do not reveal major (R=-0.001889*X+0.6951).
272	learning or memory impairment in VGLUT3 <sup>-/-</sup> mice.
273	
274	Figure 2 AROUND HERE
275	
276	Footshock sensitivity is not altered by VGLUT3 deletion
277	Deletion of VGLUT3 did not affect the behavioral responses (movement, vocalization, running, jump) elicit-
278	ed by footshock stimuli of varying intensity (Fig. 3A and statistics in in extended figure 3-1). This result
279	shows that pain sensitivity to electric footshocks is unaffected in VGLUT3-/- mice.
280	
281	Visual and Contextual Fear Conditioning are altered in the absence of VGLUT3
282	During conditioning we observed no main effect of genotype, no interaction between genotype and time,
283	only a main effect of time (Fig.3B /Table4). We then assessed contextual memory 24h after conditioning

(Fig.3C-D). The time-course analyses of the freezing rate during the 6 min test shows an interaction and a time effect, but no main effect of genotype (Fig.3C/Table4). When we analyzed the test by 3 min bins (Fig.3D), a clear genotype difference arises. Post-hoc analysis revealed higher freezing rate in VGLUT3<sup>-/-</sup> than in VGLUT3<sup>+/+</sup> mice in the last 3-min of the test (Fig.3D: 0-3 min: t(44)=0.4722, p=0.87; 3-6 min: t(44)=2.464, p=0.03; Sidak's multiple comparisons test). The cue test was then done by exposing mice to the flashing light in a novel environment. Mice were free to explore the new context for three minutes before the light (CS) was triggered (Fig.3E-F). The global analysis reveals only a main effect of time but no main effect of genotype or interaction between genotype and time (Fig.3E/Table4). Remarkably, the freezing rate significantly increased in VGLUT3<sup>+/+</sup> mice but not in VGLUT3<sup>-/-</sup> mice after CS presentation in the new context (Fig.3F: respectively t(22)=2.541, p=0.03; and t(22)=1.395, p=0.32; Sidak's multiple comparisons test). One possible explanation of the higher fear expression observed in VGLUT3-/- mice in the new context could be that once conditioned, they show a higher fear response to a new context with either no specific freezing responses associated with the US or too low to be observed.

### Figure 3 AROUND HERE

### The absence of VGLUT3 leads to a deficit in pattern separation

Cued memory alterations in VGLUT3<sup>-/-</sup> mice (Fig.3E-F) might be caused by a deficit to discriminate between the two contexts, that associated with an US versus the safe one, a process governed by pattern separation. To examine this possibility, we submitted a group of mice to a pattern separation protocol (Fig.4A) where context A is always associated with an electric shock (ES), whereas context B is safe and free of ES. In VGLUT3<sup>+/+</sup> mice, we observed no main effect of context, but a main effect of time and an interaction between context and time (Fig.4B and extended figure 4-1) Over time VGLUT3<sup>+/+</sup> mice learn to dissociate the 2 contexts since they significantly freeze less from Day 7 to Day 10 (Fig.4B; Day7, t(11)=3.031, p=0.02; Day8, t(11)=2.933, p=0.03; Day9, t(11)=2.917, p=0.03; Day10, t(11)=5.038, p<0.0001; Sidak's multiple comparisons test). Strikingly, in VGLUT3<sup>-/-</sup> mice we observed no main effect of context, or interaction between context and time but a main effect of time (Fig.4C and extended figure 4-1). VGLUT3<sup>-/-</sup> mice did not learn to discriminate the two contexts as high freezing levels were maintained over the 10 days of the test

312	(Fig. 4C). Furthermore, VOLO13 indee showed comparable revers of spontaneous freezing on Day 0 below
313	the occurrence of the first ES (Fig.4D).
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315	Figure 4 AROUND HERE
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317	However, on Day1, after conditioning, we observed a main effect of genotype, but no main effect of contex
318	or interaction between context and genotype (Fig.4E and extended figure 4-1). On Day7 and 10, we observed
319	no main effect of genotype, but a main effect of context and an interaction between context and genotype
320	(Fig.4F-G and extended figure 4-1). VGLUT3 <sup>+/+</sup> mice clearly dissociated context A from B (Fig.4F; Day7
321	t(11)=4.04, p=0.001; Fig.4G, Day10, t(11)=7.934, , p<0.0001; Sidak's multiple comparisons test). This was
322	not the case with VGLUT3 mice. Altogether, these results illustrate a deficit in pattern separation in
323	VGLUT3 <sup>-/-</sup> mice.
324	
325	The absence of VGLUT3 leads to generalized fear after aversive experiences
326	The observed deficit in pattern separation could also represent generalized fear in VGLUT3-/- mice. To in
327	vestigate this point, we submitted a group of VGLUT3+++ mice and VGLUT3 mice to an immediate shock
328	paradigm (Fig.5). On day 1 mice were introduced to a context and either immediately received a footshool
329	(immediate shock (IS) condition) or nothing (no shock (NS) condition). The next day, they were tested in the
330	same context (SC) or in a new context (NC). As expected the immediate shock (IS) did not elicit freezing
331	behaviour on day 2 in VGLUT3*/+ mice, in either context (Fig.5 IS-SC or IS-NC). VGLUT3*/- mice showed
332	no freezing when they were not shocked (Fig 5, NS), however significant higher freezing levels were ob
333	served after the IS procedure in both contexts (Fig.5, IS-SC & IS-NC). These results revealed increased
334	freezing levels in VGLUT3 mice after experiencing an aversive stimulus.
335	
336	Figure 5 AROUND HERE
337	

363	Figure 6 AROUND HERE
362	
361	L1-5, t(12)=4.076, p=0.0004; L6-10, t(44)=1.292, p=0.36; Sidak's multiple comparisons test).
360	fect was confirmed when the first 5 recall sessions were analyzed separately from the last 5 sessions (Fig.6E;
359	p=0.03; L3 t(12)=3.773, p=0.002 and L4 t(12)=2.859, p=0.04; Sidak's multiple comparisons test). This ef-
358	level of VGLUT3 <sup>-/-</sup> mice and VGLUT3 <sup>+/+</sup> mice for the first CS presentations (Fig.6D, L2 t(12)=2.971,
357	no main effect of genotype (Table7). Post-hoc analysis revealed a significant difference between the freezing
356	(Fig.6D-E, Recall 1). We observed a main effect of time and an interaction between time and genotype but
355	On day 15, mice were re-exposed to the original square context and their fear memory was examined
354	higher performance than VGLUT3 <sup>+/+</sup> mice.
353	(Fig.6C). These findings suggest that VGLUT3 <sup>-/-</sup> mice properly learn to extinguish their fear, with an initial
352	VGLUT3 <sup>-/-</sup> mice have a higher LI than VGLUT3 <sup>+/+</sup> mice but that both groups show significant positive LI
351	Day3, t(12)=2.761, p=0.04; Sidak's multiple comparisons test). Cumulative analysis showed that overall,
350	followed by a similar pattern for the two genotypes during day 4-8 (Fig.6B: Day2: t(12)=2.922, p=0.02;
349	ble7). VGLUT3 <sup>-/-</sup> mice demonstrated a higher LI than VGLUT3 <sup>+/+</sup> mice during the first 2 days of the test,
348	We observed a main effect of genotype with no effect of time or interaction between time and genotype (Ta-
347	To determine the extinction of learning performances of mice, we calculated a learning index (LI, Fig.6B-C).
346	on the 10 <sup>th</sup> CS presentation (Fig.6A, t(12)=3.77, p=0.01; Sidak's multiple comparisons test).
345	started the test with an equivalent high level of freezing that progressively decreased, reaching significance
344	a clear main effect of time and an interaction between time and gentotype (Table7). On Day 3, both groups
343	assess cue extinction (Fig.6A). The overall analysis suggested a tendency for a main effect of genotype with
342	context, followed from Day 2 to Day 8 to a daily session of 10 CS-only presentations in a round context, to
341	tion protocol was performed (Fig.6). On Day 1, mice were exposed to 10 CS-US presentations in a square
340	fully conditioned with a discrete CS such as a light. To answer this question, a cue fear conditioning extinc-
339	Because of the impairment described in the cue-test (Fig.3.E-F), we wondered if VGLUT3 <sup>-/-</sup> mice were not
220	Visual Feat Extinction is affected in the absence of Vollo 15

To establish that the freezing behaviour observed during recall 1 was specific and was due to the occurrence of the light in the conditioning context, half of the animals were tested on day 18 in a completely new environment (Fig.6F; Recall 2). As can be seen from Fig.6F, we observed no main effect of genotype or time and no interaction between time and genotype (Table7). Freezing levels were similar (≈20-25%) for both groups, showing no evidence of generalized freezing behaviour after extinction. These data suggest that after an extinction procedure, VGLUT3<sup>-/-</sup> mice may have stronger original memory recall, with no generalized freezing responses to a new context.

## Working memory is intact in the absence of VGLUT3

The accelerated extinction observed in VGLUT3<sup>-/-</sup> mice during the first days of extinction (Fig.6A-B) could reflect altered working memory (WM). Hence, we compared WM of WT mice and VGLUT3<sup>-/-</sup> mice using the Y-Maze paradigm. Mice were free to explore the Y-Maze for 10 min and spontaneous alternation was quantified. In line with their anxiety phenotype, VGLUT3<sup>-/-</sup> mice made significantly fewer arm entries than controls (Fig.7A/Table8). However, both groups showed similar levels of spontaneous alternation, both above chance level (Fig.7B). Overall, VGLUT3<sup>-/-</sup> mice show normal working memory despite a lower exploration activity.

Figure 7 AROUND HERE

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## Discussion

The involvement of the VGLUT3 network in cognition and related psychiatric pathologies has been investigated in a few studies (Amilhon et al., 2010; Balázsfi et al., 2018; Favier et al., 2020; Fazekas et al., 2019; Sakae et al., 2015). For example, Balázsfi (2018) and Fazekas (2019) who focused on the study of learning and memory, concluded that the deficits in VGLUT3-deficient mice were very limited and mainly related to stress. Understanding how aversive memories are processed in the brain will help to decipher its dysfunction in trauma- and stress-related disorders. In the present study we explored the establishment and maintenance of fear-related memories in mice lacking VGLUT3. Using a Pavlovian fear conditioning paradigm, we report

that VGLUT3<sup>-/-</sup> mice express more stable and generalized contextual memories associated with a deficit of pattern separation. Interestingly, VGLUT3<sup>-/-</sup> mice have no deficit in non-aversive learning and memory, including working memory, spatial reference memory, and cue-based extinction learning. Our results partly confirm previous findings (Balázsfi et al., 2018; Fazekas et al., 2019) while deepening our understanding of the involvement of VGLUT3-dependent cotransmission in aversive memories.

Before studying aversive memories, we first assessed the consequences of the lack of VGLUT3 in learning,

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memory processing and cognitive flexibility in spatial and non-spatial tasks. No deficit was found in VGLUT3-deficient mice. Our results are in agreement with data obtained by Fazekas et al., (2019), who also found comparable spatial learning capacities in VGLUT3/- mice (although they trained only male mice, in a pool that was half the size of ours), supporting the robustness of the observed phenotypes. However, our approach of systematically testing spatial memories has revealed improved long-term spatial memory performance in VGLUT3<sup>-/-</sup> mice compared to control mice at 22°C in the watermaze task. Since memory performances of VGLUT3-/- mice were comparable to controls when the water temperature was dropped to 19°C, we hypothesized that the improved memory performance of VGLUT3<sup>-/-</sup> mice could be related to their anxiety trait (Amilhon et al. 2010) as well as to their hypothalamic-pituitary-adrenal axis dysfunctions (Balázsfi et al. 2018) in less-stressful watermaze conditions (i.e. at 22°C). This is in agreement with the literature in both humans and animals, highlighting that mild stress could have facilitating effects on memory consolidation (Sandi, Loscertales, et Guaza 1997; Sandi et Pinelo-Nava 2007; Cahill et McGaugh 1998). Nevertheless, depending on the behavioral paradigm used, this anxiety trait could interfere with appropriate data interpretation. In order to overcome this and accurately assess recognition memory (object and spatial) using an open-field, we had to adapt the protocol to ensure sufficient exploration of objects for recognition memory to occur. By using a fixed exploration time per session rather than a fixed session duration, we were able to circumvent the confounding effect of anxiety and ensure an unbiased assessment of recognition memory in VGLUT3<sup>-/-</sup> mice. We observed no deficit of recognition or spatial memories in VGLUT3<sup>-/-</sup>

mice. In conclusion, using different protocols or paradigms, we confirmed that the absence of VGLUT3 does

not impair spatial reference, non-spatial memory or associative-learning processes.

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We next explored fear-related memories in VGLUT3<sup>-/-</sup> mice using a Pavlovian fear conditioning paradigm. Because VGLUT3 is present in peripheral sensory neurons and contributes to mechanical pain detection (Seal et al, 2009), we assessed sensitivity to electric footshocks in VGLUT3--- mice and found unchanged sensitivity to electric foot shock in VGLUT3<sup>+/+</sup> mice. This result confirmed previous findings by Balazsfi et al (2018) using flinch and jump threshold as the readout. In the Pavlovian fear conditioning protocols used, the deletion of VGLUT3 led to normal fear learning but a higher and persistent contextual memory, which is consistent with the observations of Balazsfi et al (2018). However, our study highlighted an absence of cue memory, which could be explained by a contextual generalization deficit. To express fear when it is relevant, present and past associations have to be compared. This is adaptive, since it allows individuals to anticipate a threat by discerning pertinent cues in the environment. Increased interference between past and new memories could promote reactivation of traumatic memories and lead to overgeneralization of fear. Considerable evidence from the literature suggests the involvement of the hippocampal CA3-Dentate Gyrus (DG) circuit in contextual discrimination (McHugh et al. 2007; Besnard et Sahay 2016; Cravens et al. 2006). CA3 plays a major role in a process called pattern completion, which allows retrieval of a stored representation based on sparse cues in the environment. In contrast, the DG is also involved in pattern separation, to minimize the overlap between two similar representations. Precise memory requires remembering details with high specificity, so that memories can be discriminated from other similar memories to avoid interference. Pattern separation facilitates this discrimination by reducing the degree of similarities between overlapping experiences. The pattern separation paradigm (Sahay et al. 2011) was used to study the propensity of mice to discriminate among similar experiences (Yassa & Stark, 2011). At day 1, freezing levels were comparable between contexts A and B for both control and VGLUT3-/- mice, suggesting the degree of similarity between the two contexts was high enough to evoke generalization of contextual fear in both groups. However, control mice learned to discriminate the shocked context from the safe context as early as day 7, whereas VGLUT3deficient mice were unable to discriminate between the two contexts within the 10-day protocol used. These results highlight a significant deficit in pattern separation in VGLUT3-- mice (Fig.4 A2-3). Considerable evidence supports a role for the hippocampus in pattern separation to constrain the overgeneralization of fear. Previous work studied the hippocampal plasticity in VGLUT3 deficient mice (Fasano et al, 2017) and found that the absence of glutamate released by VGLUT3 hippocampal interneurons led to increased GA-BAergic transmission, altering the oscillatory activity of synchronized networks and inducing a metaplastic

shift of synaptic plasticity in the ventral hippocampus. As hippocampal long-term plasticity is currently thought to underlie the cellular basis of such learning and memory processes, we cannot exclude that they might cause the observed contextual overgeneralization in VGLUT3-/- mice.

To better understand this discrimination deficit, we performed an immediate shock (IS) test. According to Fanselow (2000), in the IS test, animals do not have enough time to form an integrated memory representation of context features in order to associate it to the electric shock. In line with this hypothesis, wildtype mice do not form a contextual fear memory and show no freezing behaviour during retrieval tests. In contrast, when they were immediately shocked, VGLUT3<sup>-/-</sup> mice increased their level of freezing whatever the context used in the retrieval test (Fig.5). This observation suggests that in VGLUT3<sup>-/-</sup> mice, the mere occurrence of the traumatic event (i.e., the foot shock) elicited impaired fear expression. In our view, this increased fear expression reflects more than innate fear impairment in VGLUT3<sup>-/-</sup> mice, since innate fear experiences to natural threats need to be harmless (Silva, Gross, et Gräff 2016). When the animal experiences pain such as a foot shock as in our experiment, it is a conditioned response and a learned experience.

One major treatment of fear-related disorders, called exposure therapy in clinics or extinction fear learning in laboratory, involves repeatedly re-exposure of animals to the CS (the flashing light) previously associated with the aversive US (the foot shock) in a different context. With time, the animals learn that the CS is no longer associated with the US in this new context and thus the mice form a new « safer » memory (Myers et Davis 2007; Perusini et Fanselow 2015). Surprisingly, during the initial steps of this extinction learning, VGLUT3<sup>-/-</sup> mice show improved performance (Fig.6 A-C). This is particularly surprising since the processes governing extinction and generalization are thought to be similar (see Lopresto et al., 2016). The brain structures mainly involved are the prefrontal cortex (especially its infra limbic (IL-PFC) part) and the hippocampus as previously discussed regarding pattern separation. However, extinction mostly relies on the interaction of the IL-PFC with the basal-lateral amygdala. Those projections do not express VGLUT3 and might effectively control the amygdala activity as observed. One hypothesis is that context generalization (or lack of pattern separation) could be due to the dysfunction of the hippocampal network due to the absence of VGLUT3, whereas the cue-based extinction may depend on the IL-PFC projections to the amygdala.

Original memory was assessed at day 15 in the conditioning context (Fig.6D-E). Results confirm that the extinction procedure did not alter the original memory since both groups still displayed a high level of freezing (significantly higher in VGLUT3<sup>-/-</sup> than in control mice) to the context where they were originally shocked. Surprisingly, when tested in a third context on D18 VGLUT3-/- mice did not show fear generalization, indicating that the animals might have associated the aversive value of the CS only to the original context. This observation suggests that VGLUT3-/- mice could show an associative cue learning that can properly be recalled and specific to a context.

In regards to the initial facilitation of the extinction, we cannot exclude that this could be due to increased attention related to the anxiety trait in VGLUT3<sup>-/-</sup> mice, or in their working memory. Attentional processes are difficult to test in VGLUT3<sup>-/-</sup> mice, since those experiments classically require the use of sound (e.g., prepulse inhibition, fear startle tests...) and these mutants are deaf (Ruel et al, 2008). To rule out any working memory modification that could explain this initial extinction improvement, we subjected our mice to a Y-maze alternation protocol. Unlike Fazekas et al. (2019) we observed no alteration of working memory in VGLUT3<sup>-/-</sup> mice. Since mice lacking VGLUT3 tend to explore less due to their anxious phenotype, we increased the test duration from 5 to 10 min to have substantial exploration levels in VGLUT3<sup>-/-</sup> mice and WT mice (>100 entries). This might explain the different findings, since poor exploration can directly affect behavioral performances. Therefore, in our hands, VGLUT3<sup>-/-</sup> mice show no deficit or facilitation of their working memory that could explain their better initial performance in fear extinction.

Some studies found VGLUT3-amacrine cells in mouse retina (Kim et al., 2015.; Lee et al., 2021, 2016) coreleasing glutamate and glycine at glycinergic synapses. How the absence of VGLUT3 could impact the function of these synapses in these animals, and therefore their ability to see properly, has yet to be determined. What seems to be accepted is the lack of VGLUT3 impacting the vision of movement (Kim et al., 2015.; Lee et al., 2016). However, based on our results, it is unlikely that the observed initial lack of cue conditioning can be due to visual impairment. First, we use a flashing light as a cue, that is a major visual information. Then, VGLUT3 deficient mice have intact performances in the spatial reference memory task in the watermaze, and in the object recognition tasks, both of which mainly rely on visual cues. Overall, we

cannot rule out differences in visual detection between controls and VGLUT3-deficient mice, but this alone
cannot explain the initial lack of cue conditioning observed.
Our findings on impaired fear-related memories in mice lacking VGLUT3 are in good agreement with the
electrophysiological reports (Fasano et al, 2017). However, this interpretation should be taken with care,
since a constitutive VGLUT3 deletion was used in the present study. Cholinergic fibers from the basal fore-
brain projecting to the basolateral amygdala are crucial in reinforcing learning and consolidating aversive
memories (Jiang et al. 2016; Crouse et al. 2020; Aitta-aho et al. 2018). Interestingly, a subset of those fibers
does express VGLUT3 (Nickerson Poulin et al. 2006). It is possible that this cholinergic pathway could also
be involved in fear-related disorders. A thorough description of the involvement of these different pathways
would require the deletion of VGLUT3 in specific subpopulation of neurons.
In conclusion, the present study suggests an important role of VGLUT3 in aversive memory processing such
as contextual generalization of fear memory which could be crucial in trauma- and stress-related disorders.

## **Table 1.** Cohorts used.

Experiment (Figure)	Sex	N of VGLUT3 <sup>+/+</sup> (WT)	N of VGLUT3 <sup>-/-</sup> (KO)
Watermaze 22°C (Fig1D-F)	F	15	11
Watermaze 19°C (Fig1G-I)	F	13	11
Object recognition (Fig2)	М	13	12
Shock sensitivity (Fig3A)	М	8	6
Fear conditioning (Fig3B-F)	М	12	12
Pattern separation (Fig4)	М	11	10
Immediate shock (Fig5)	М	14	20
Fear extinction (Fig6)	М	12	12
Y Maze (Fig7)	М	9	8
TOTAL		107	102

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751	

752	Figure Legends
753	Figure 1. Cue and Spatial Reference Memory in VGLUT3 <sup>-/-</sup> mice. (A-C) Watermaze experimental de-
754	sign: A) Cuetask, B) SRM task and C) SRM-Reversal task. (D-F) Mice were trained in 22°C water.
755	VGLUT3 <sup>-/-</sup> mice showed no deficit of learning either in the Cue- (D), or the SRM and reversal (E) tasks. (F)
756	Memory assessment was performed 10min (PT1 for SRM and SRM-R), 72h (PT2 for SRM) and 48h (PT2
757	for SRM-R) post training, and VGLUT3 <sup>-/-</sup> mice showed better performance at 72h post training, and in the
758	PT average. (G-I) Mice were trained in water at 19°C. VGLUT3 <sup>-/-</sup> mice show no deficit of learning either in
759	the Cue (G), or the SRM and reversal (H) tasks. (I) No differences were observed in memory tests done at
760	different times or on average. Data are mean $\pm$ SEM. Differences between genotypes: * p<0.05. PT: probe
761	test; R: reversal. All corresponding statistics are presented in Figure 1-1
762	
763	Figure 2. Object and Spatial Recognition in VGLUT3 <sup>-/-</sup> mice. (A) Object Recognition (OR): both groups
764	show OR memory, with VGLUT3-/ mice having higher scores. (B) Spatial Recognition (SR): both groups
765	show comparable SR memory level. (C) There is no correlation between learning sessions duration and
766	VGLUT3-/- mice performances in OR. Slopes are -0.01563 for WT and -0.001889 for VGLUT3-/- mice.
767	Data are mean ± SEM. Differences between genotypes: * p<0.05; Differences to chance level: ## p<0.01,
768	### p<0.001. All corresponding statistics are presented in Figure 2-1
769	
770	Figure 3. Contextual and Cued Fear memories of VGLUT3 <sup>-/-</sup> mice. (A) Shock sensitivity assessment, ie.
771	the intensity for which the mice express a given behaviour (movement, vocalization, running or jump). (B-F)
772	Fear memories in VGLUT3 <sup>-/-</sup> mice. (B) Freezing levels during fear conditioning consisting of 2 CS-US pai-
773	rings. (C-D) Contextual memory was tested 24h after conditioning and revealed a more stable memory in
774	VGLUT3 <sup>-/-</sup> mice. (E-F) Cued memory test revealed high level of freezing to new context for VGLUT3 <sup>-/-</sup>
775	mice. Data are mean ± SEM. post-hoc comparisons: * p<0.05. All corresponding statistics are presented in
776	Figure 3-1
777	
778	

779	Figure 4. Pattern separation of VGLUT3 <sup>-/-</sup> mice. (A) Behavioral protocol; (B) VGLUT3 <sup>+/+</sup> mice perfor-
780	mances; (C) VGLUT3 <sup>-/-</sup> mice performances. (D-G) Freezing levels on different days: (D) Day 0, before
781	conditioning, (E) Day 1, VGLUT3 <sup>-/-</sup> mice already show a higher freezing level, (F) Day7, VGLUT3 <sup>+/+</sup> mice
782	start to discriminate the different contexts, (G) On Day10, VGLUT3-/- mice still do not discriminate the dif-
783	ferent contexts. Data are mean ± SEM. * p<0.05, ** p<0.01, *** p<0.001, **** p<0.0001. All correspon-
784	ding statistics are presented in Figure 4-1
785	
786	Figure 5. Immediate shock in VGLUT3 <sup>-/-</sup> mice. Mice were subjected to a no shock (NS) or immediate
787	shock (IS) protocol to assess their levels of spontaneous freezing after experiencing an electric footshock.
788	They were either tested in the same context (IS-SC) or in a novel context (IS-NC). WT mice did not show
789	any freezing after either procedure, whereas VGLUT3 mice expressed a significant increase of freezing
790	behaviour only after the IS, in either context. Data are mean $\pm$ SEM. post-hoc comparisons: *** p<0.001.
791	All corresponding statistics are presented in Figure 5-1
792	Figure 6. Extinction Fear in VGLUT3 <sup>-/-</sup> mice. (A) Extinction learning over a 7-day period. Filled black
793	and red circles represent the freezing levels of VGLUT3+/+ mice and VGLUT3-/- mice (respectively) before
794	the presentation of the first CS. Open circles are used for the 10 subsequent CS. (B-E) Fear memory in
795	VGLUT3 <sup>-/-</sup> mice. (B-C) The Learning Index (LI) was calculated to illustrate learning efficacy over time (B)
796	and on average (C). (D-E) Original memory was recalled on Day15. (F) On D18, freezing to CS was asses-
797	sed in a new hexagonal context. Data are mean $\pm$ SEM. * p<0.05, ** p<0.01, *** p<0.001. L: light (CS). All
798	corresponding statistics are presented in Figure 6-1
799	Figure 7. Working memory in VGLUT3 <sup>-/-</sup> mice. (A) Number of entries in the Y-Maze arms for the first 5
800	min of the test (0-5min) or the total 10 min test (0-10min). (B) Percentage of alternation. Data are mean $\pm$
801	SEM. Differences between genotype: * p<0.05, *** p<0.001; Differences compared to chance level: #
802	p<0.05, ## p<0.01. All corresponding statistics are presented in Figure 7-1
803	

- 805 Extended data Legends
- 806 Extended Figure 1-1. Statistics for Watermaze experiments. 1: SRM 10min; 2: SRM 72h; 3:
- 807 SRM-R 10min; 4: SRM-R 48h; 5: PTs average
- 808 Extended Figure 2-1. Statistics for Object Recognition experiments.
- 809 Extended Figure 3-1. Statistics for Fear conditioning experiments.
- 810 Extended Figure 4-1. Statistics for Pattern separation experiment.
- 811 Extended Figure 5-1. Statistics for Immediate shock experiments. NS: no shock; IS: immediate
- 812 shock; SC: same context; NC: new context
- 813 Extended Figure 6-1. Statistics for Fear extinction experiments.
- 814 Extended Figure 7-1. Statistics for the Y-Maze experiment.

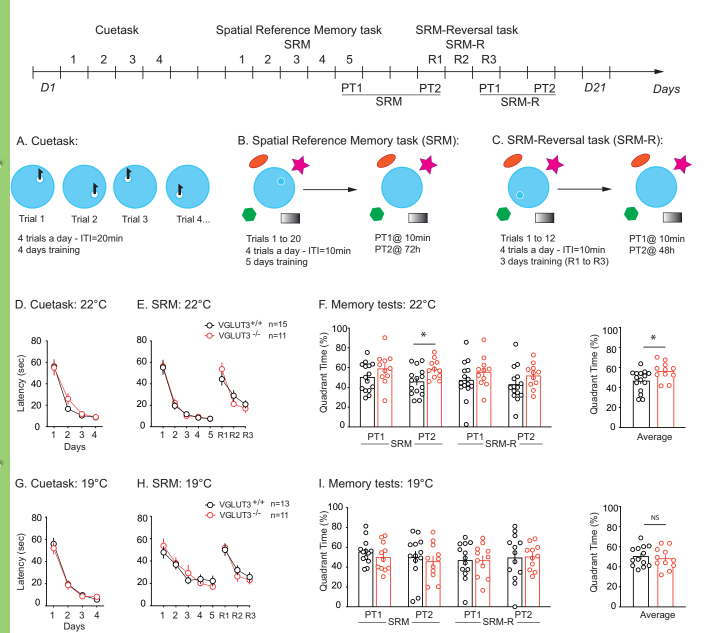
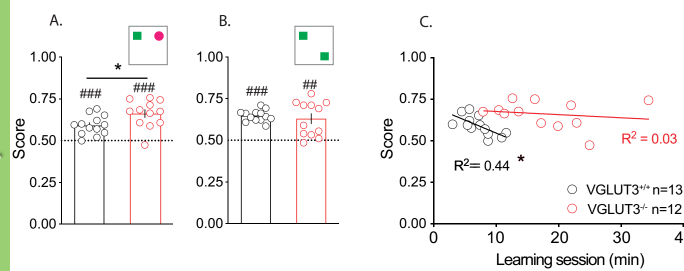


Figure 1

Figure 2



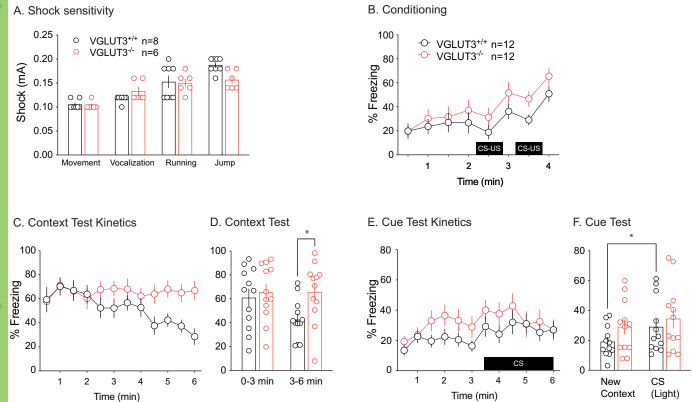


Figure 3

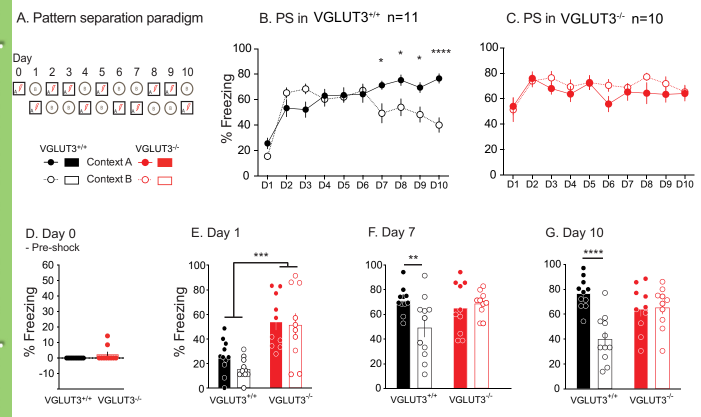
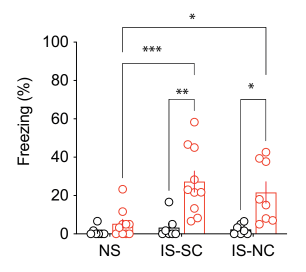


Figure 4



- O VGLUT3+/+
- VGLUT3-/-

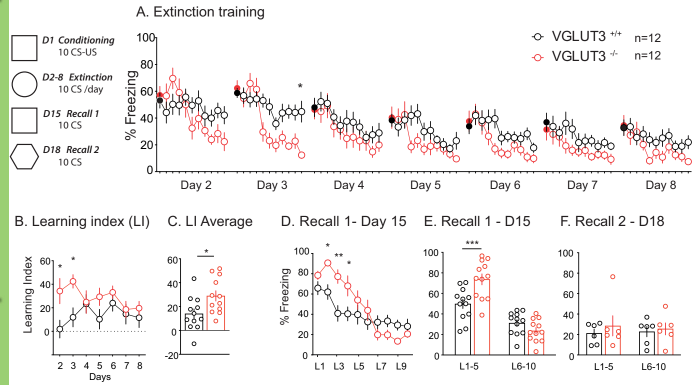


Figure 4

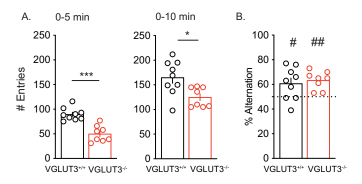


Figure 7